

# A Practical Introduction to Regression Discontinuity Designs: Volume II

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\*\* PRELIMINARY AND INCOMPLETE – COMMENTS WELCOME \*\*

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## Acknowledgments

This monograph, together with its accompanying first part ([Cattaneo, Idrobo and Titiunik, 2018a](#)), collects and expands the instructional materials we prepared for more than 30 short courses and workshops on Regression Discontinuity (RD) methodology taught over the years 2014–2017. These teaching materials were used at various institutions and programs, including the Asian Development Bank, the Philippine Institute for Development Studies, the International Food Policy Research Institute, the ICPSR’s Summer Program in Quantitative Methods of Social Research, the Abdul Latif Jameel Poverty Action Lab, the Inter-American Development Bank, the Georgetown Center for Econometric Practice, and the Universidad Católica del Uruguay’s Winter School in Methodology and Data Analysis. The materials were also employed for teaching at the undergraduate and graduate level at Brigham Young University, Cornell University, Instituto Tecnológico Autónomo de México, Pennsylvania State University, Pontificia Universidad Católica de Chile, University of Michigan, and Universidad Torcuato Di Tella. We thank all these institutions and programs, as well as their many audiences, for the support, feedback and encouragement we received over the years.

The work collected in our two-volume monograph evolved and benefited from many insightful discussions with our present and former collaborators: Sebastián Calonico, Robert Erikson, Juan Carlos Escanciano, Max Farrell, Yingjie Feng, Brigham Frandsen, Sebastián Galiani, Michael Jansson, Luke Keele, Marko Klašnja, Xinwei Ma, Kenichi Nagasawa, Brendan Nyhan, Jasjeet Sekhon, Gonzalo Vazquez-Bare, and José Zubizarreta. Their intellectual contribution to our research program on RD designs has been invaluable, and certainly made our monographs much better than they would have been otherwise. We also thank Alberto Abadie, Joshua Angrist, Ivan Canay, Richard Crump, David Drukker, Sebastian Galiani, Guido Imbens, Patrick Kline, Justin McCrary, David McKenzie, Douglas Miller, Aniceto Orbeta, Zhuan Pei, and Andres Santos for the many stimulating discussions and criticisms we received from them over the years, which also shaped the work presented here in important ways. The co-Editors Michael Alvarez and Nathaniel Beck offered useful and constructive comments on a preliminary draft of our manuscript, including the suggestion of splitting the content into two stand-alone volumes. Last but not least, we gratefully acknowledge the support of the National Science Foundation through grant [SES-1357561](#).

The goal of our two-part monograph is purposely practical and hence we focus on the empirical analysis of RD designs. We do not seek to provide a comprehensive literature review on RD designs nor discuss theoretical aspects in detail. In this second part, we employ the

data of Meyersson (2014), Lindo, Sanders and Oreopoulos (2010), Amarante, Manacorda, Miguel and Vigorito (2016), Chay, McEwan and Urquiola (2005) and Keele and Titiunik (2015) for empirical illustration of the different topics covered. We thank these authors for making their data and codes publicly available. We provide complete replication codes in both **R** and **Stata** for all the empirical work discussed throughout the monograph. In addition, we provide replication codes for another empirical illustration using the data of Cattaneo, Frandsen and Titiunik (2015), which is not discussed in the text to conserve space and because it is already analyzed in our companion software articles. The general purpose, open-source software used in this monograph, as well as other supplementary materials, can be found at:

<https://sites.google.com/site/rdpackages/>

# 1 Introduction

The Regression Discontinuity (RD) design has emerged as one of the most credible research designs in the social, behavioral, biomedical and statistical sciences for program evaluation and causal inference in the absence of experimental treatment assignment. In this manuscript we continue the discussion given in [Cattaneo, Idrobo and Titiunik \(2018a\)](#), the first part of our two-part monograph offering a practical introduction to the analysis and interpretation of RD designs. While the present manuscript is meant to be self-contained, it is advisable to consult Part I of our two-part monograph as several concepts and ideas discussed previously will naturally feature in this volume.

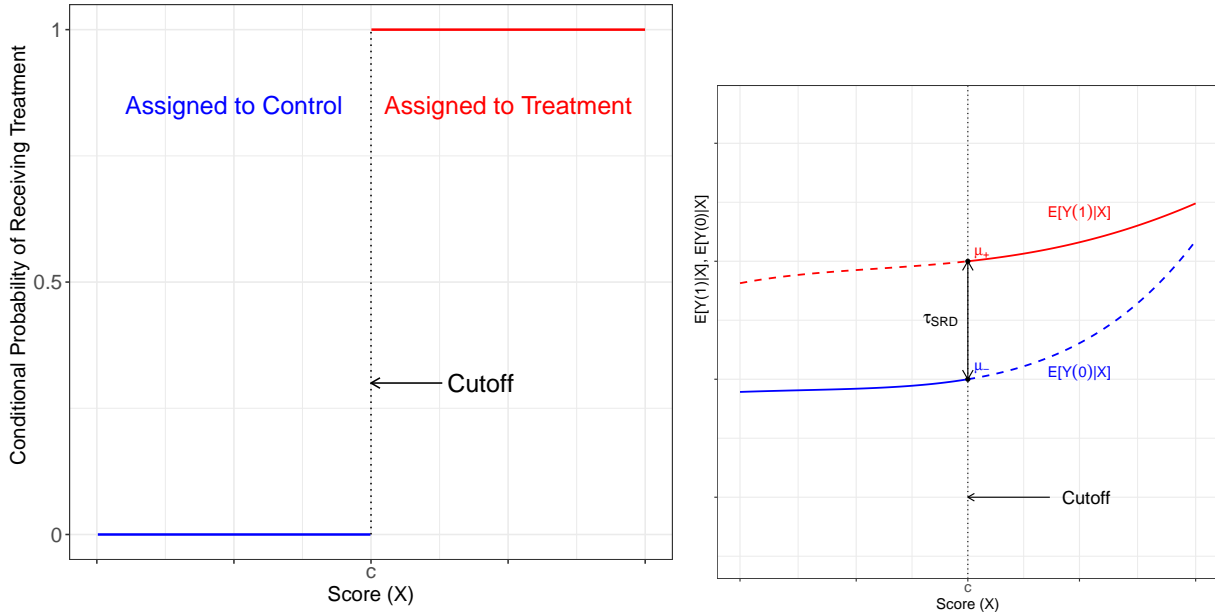
The RD design is defined by three fundamental ingredients: a score (also known as running variable, forcing variable, or index), a cutoff, and a treatment rule that assigns units to treatment or control based on a hard-thresholding rule using the score and cutoff. All units are assigned a score, and a treatment is assigned to those units whose value of the score exceeds the cutoff and not assigned to units whose value of the score is below the cutoff. This treatment assignment rule implies that the probability of treatment assignment changes abruptly at the known cutoff. If units are not able to perfectly determine or manipulate the exact value of the score that they receive, this discontinuous change in the treatment assignment probability can be used to study the effect of the treatment on outcomes of interest, at least locally, because units with scores barely below the cutoff can be used as counterfactuals for units with scores barely above it.

In the accompanying Part I of our monograph ([Cattaneo, Idrobo and Titiunik, 2018a](#)), we focused exclusively on the canonical Sharp RD design, where the running variable is continuous and univariate, there is a single cutoff determining treatment assignment and treatment compliance is perfect, and the analysis is conducted using continuity-based methods (e.g., local polynomial approximations). To be more precise, assume that there are  $n$  units, indexed by  $i = 1, 2, \dots, n$ , and each unit receives score  $X_i$ . Units with  $X_i \geq \bar{x}$  are assigned to the treatment condition, and units with  $X_i < \bar{x}$  are assigned to the untreated or control condition, where  $\bar{x}$  denotes the RD cutoff. Thus, in the canonical Sharp RD design, the univariate  $X_i$  is continuously distributed (i.e., all units received different values), and all units comply with their treatment assignment.

We denote treatment assignment by  $T_i = \mathbb{1}(X_i \geq \bar{x})$ , where  $\mathbb{1}(\cdot)$  is the indicator function. In the canonical RD treatment assignment rule is deterministic, once the score is assigned to each unit, and obeyed by all units (perfect treatment compliance). More generally, the key defining feature of any RD design is that the probability of treatment assignment given

the score changes discontinuously at the cutoff, that is, the conditional probability of being assigned to treatment given the score,  $\mathbb{P}(T_i = 1|X_i = x)$ , jumps discontinuously at the cutoff point  $x = \bar{x}$ . Figure 1.1a illustrates this graphically.

Figure 1.1: Canonical Sharp RD Design



(a) Conditional Probability of Treatment

(b) RD Treatment Effect

We adopt the potential outcomes framework to discuss causal inference and policy evaluation employing RD designs; see [Imbens and Rubin \(2015\)](#) for an introduction to potential outcomes and causality, and [Abadie and Cattaneo \(2018\)](#) for a review of program evaluation methodology. Each unit has two potential outcomes,  $Y_i(1)$  and  $Y_i(0)$ , which correspond, respectively, to the outcomes that would be observed under treatment or control. Treatment effects are therefore defined in terms of comparisons between features of (the distribution of) both potential outcomes, such as their means, variances or quantiles. If unit  $i$  receives the treatment, we observe the unit's outcome under treatment,  $Y_i(1)$ , but  $Y_i(0)$  remains unobserved, while if unit  $i$  is assigned to the control condition, we observe  $Y_i(0)$  but not  $Y_i(1)$ . This is known as the fundamental problem of causal inference. The observed outcome  $Y_i$  is therefore defined as

$$Y_i = (1 - T_i) \cdot Y_i(0) + T_i \cdot Y_i(1) = \begin{cases} Y_i(0) & \text{if } X_i < \bar{x} \\ Y_i(1) & \text{if } X_i \geq \bar{x} \end{cases}.$$

The canonical Sharp RD design, discussed in our prior monograph ([Cattaneo, Idrobo and](#)

[Titiunik, 2018a](#)), assumes that the potential outcomes  $(Y_i(1), Y_i(0))_{i=1}^n$  are random variables, and focuses on the average treatment effect at the cutoff

$$\tau_{\text{SRD}} \equiv \mathbb{E}[Y_i(1) - Y_i(0)|X_i = \bar{x}].$$

This (causal) parameter is sometimes called the Sharp RD treatment effect, and is depicted in [Figure 1.1b](#). In that figure we also plot the regression functions  $\mathbb{E}[Y_i(0)|X_i = x]$  and  $\mathbb{E}[Y_i(1)|X_i = x]$  for values of the score  $X_i = x$ , where solid and dashed lines correspond to their estimable and non-estimable portions, respectively. The continuity-based framework for RD analysis, assumes that the regression functions  $\mathbb{E}[Y_i(1)|X_i = x]$  and  $\mathbb{E}[Y_i(0)|X_i = x]$ , seen as functions of  $x$ , are continuous at  $x = \bar{x}$ , which gives

$$\tau_{\text{SRD}} = \lim_{x \downarrow \bar{x}} \mathbb{E}[Y_i|X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[Y_i|X_i = x]. \quad (1.1)$$

In words, [Equation 1.1](#) says that, if the average potential outcomes are continuous functions of the score at  $\bar{x}$ , the difference between the limits of the treated and control average observed outcomes as the score approaches the cutoff is equal to the average treatment effect at the cutoff. This identification result is due to [Hahn, Todd and van der Klaauw \(2001\)](#), and has spiked a large body of methodological work on identification, estimation, inference, graphical presentation, and falsification/validation for many RD design settings. In our first volume we focused exclusively on the canonical Sharp RD designs, and gave a practical introduction to the methods developed by [Lee \(2008\)](#), [McCrary \(2008\)](#), [Imbens and Kalyanaraman \(2012\)](#), [Calonico, Cattaneo and Titiunik \(2014b, 2015a\)](#), [Calonico, Cattaneo and Farrell \(2018b,a\)](#), and [Calonico, Cattaneo, Farrell and Titiunik \(2018c\)](#), among others.

In the present Part II of our monograph, we offer a practical discussion of several topics in RD methodology building on and extending the continuity-based analysis of the canonical Sharp RD design. Our first goal in this manuscript is to introduce readers to an alternative RD setup based on local randomization ideas that can be useful in some practical applications and complements the continuity-based approach to RD analysis. Then, employing both continuity and local randomization approaches, we extend the canonical Sharp RD design in multiple directions.

[Section 2](#) discusses the local randomization framework for RD designs, where the assignment of score values is viewed as-if randomly assigned in a small window around the cutoff, so that placement above or below the cutoff and hence treatment assignment can be interpreted to be as-if experimental. This contrasts with the continuity-based approach, where extrapolation to the cutoff plays a predominant role. Once this local randomization



assumption is invoked, the analysis can proceed by using standard tools in the analysis of experiments literature. This alternative approach, which we call the Local Randomization RD approach, requires stronger assumptions than the continuity-based approach discussed in Part I, and for this reason it is not always applicable. We discuss the main features of the local randomization approach in Section 2 below, including how to interpret the required assumptions, and how to perform estimation, inference, presentation and falsification within this alternative framework.

Next, in Section 3, we discuss RD designs where the running variable is discrete instead of continuous, and multiple units share the same value of the score. This situation is common in applications. For example, universities' Grade Point Average (GPA) are often calculated up to one or two decimal places, and collecting data on all students in a college campus would result in a dataset where hundreds or thousands of students would have the same GPA. In the RD design, the existence of such "mass points" in the score variable often requires to different methods, as the standard continuity-based methods discussed in Part I are no longer generally applicable. In Section 3, we discuss when and why continuity-based methods will be inadequate to analyze RD designs with discrete scores, and discuss how the local randomization approach can be a useful alternative framework for analysis.

We continue in Section 4 with a discussion of the so-called Fuzzy RD design, where compliance with treatment assignment is no longer perfect (in contrast to the Sharp RD case). In other words, these are RD designs where some units above the cutoff fail to take the treatment despite being assigned to take it, and/or some units below the cutoff take the treatment despite being assigned to the untreated condition. Our discussion defines several parameters of interest that can be recovered under noncompliance, and discusses how to employ both continuity-based and local randomization approaches for analysis. We also discuss the important issue of how to perform falsification analysis under noncompliance.

We devote the last two sections to generalize the assumption of a treatment assignment rule that depends on a single score and a single cutoff. In Section 5, we discuss RD designs with multiple running variables, which we refer to as Multi-Score RD designs. Such designs occur, for example, when students must obtain a grade above a cutoff in two different exams in order to receive a scholarship. In this case, the treatment rule is thus more general, requiring that both scores be above the cutoff in order to receive the treatment. Another example RD designs with multiple scores that is used frequently in applications is the Geographic RD design, where assignment to treatment of interest changes discontinuously at the border that separates two geographic areas. We discuss how to generalize the methods discussed both in Part I and in the first sections of this monograph to the multiple-score case, and

illustrate how to apply this methods with a Geographic RD example. Finally, in Section 6 we consider RD designs with multiple cutoffs, a setup where all units have a score value, but different subsets of units face different cutoff values. In our discussion, we highlight how the Multiple-Cutoff RD design can be recast as Multiple-Score RD design with two running variables.

Each section in this manuscript illustrates the methods with a different empirical application. In Section 2, we use the data provided by [Meyersson \(2014\)](#) to study the effect of Islamic parties' victory on the educational attainment of women in Turkey. This is the same empirical application employed in Part I to illustrate the analysis of the canonical continuity-based Sharp RD design. In Section 3, we re-analyze the data in [Lindo et al. \(2010\)](#), who analyze the effects of academic probation on subsequent academic achievement. In Section 4 we use the data provided by [Amarante et al. \(2016\)](#) to study the effect of a social assistance program on the birth weight of babies born to beneficiary mothers. In Section 5, we re-analyze the data in [Chay, McEwan and Urquiola \(2005\)](#), who study the effect of a school improvement program on test scores, where units in different geographic regions facing different cutoff values. Finally, in Section 5, we re-analyze the Geographic RD design in [Keele and Titiunik \(2015\)](#), where the focus is to analyze the effect of campaign ads on voter turnout. We hope that the multiple empirical applications we re-analyze across multiple disciplines will be useful to a wide range of researchers.

As in the companion Part I of our monograph [Cattaneo, Idrobo and Titiunik \(2018a\)](#), all the RD methods we discuss and illustrate are implemented using various general-purpose software packages, which are free and available for both **R** and **Stata**, two leading statistical software environments in the social sciences. Each numerical illustration we present includes an **R** command with its output, and the analogous **Stata** command that reproduces the same analysis—though we omit the **Stata** output to avoid repetition. The local polynomial methods for continuity-based RD analysis are implemented in the package **rdrobust**, which is presented and illustrated in three companion software articles: [Calonico, Cattaneo and Titiunik \(2014a\)](#), [Calonico, Cattaneo and Titiunik \(2015b\)](#) and [Calonico, Cattaneo, Farrell and Titiunik \(2017\)](#). This package has three functions specifically designed for continuity-based RD analysis: **rdbwselect** for data-driven bandwidth selection methods, **rdrobust** for local polynomial point estimation and inference, and **rdplot** for graphical RD analysis. In addition, the package **rddensity**, discussed by [Cattaneo, Jansson and Ma \(2018c\)](#), provides manipulation tests of density discontinuity based on local polynomial density estimation methods. The accompanying package **rdlocrand**, which is presented and illustrated by [Cattaneo, Titiunik and Vazquez-Bare \(2016b\)](#), implements the local randomization methods

discussed in the second part accompanying this monograph (Cattaneo, Idrobo and Titiunik, 2018b).

The full R and Stata codes that replicate all our analysis are available at <https://sites.google.com/site/rdpackages/replication>. In that website, we also provide replication codes for two other empirical applications, both following closely our discussion. One employs the data on U.S. Senate incumbency advantage originally analyzed by Cattaneo, Frandsen and Titiunik (2015), while the other uses the Head Start data originally analyzed by Ludwig and Miller (2007) and recently employed in Cattaneo, Titiunik and Vazquez-Bare (2017).

Finally, we remind the reader that our main goal is not to offer a comprehensive review of the literature on RD methodology (we do offer references to further readings after each topic is presented), but rather to provide an accessible practical guide for empirical RD analysis. For early review articles on RD designs see Imbens and Lemieux (2008) and Lee and Lemieux (2010), and for an edited volume with a contemporaneous overview of the RD literature see Cattaneo and Escanciano (2017). We are currently working on a comprehensive literature review that complements our practical two-part monograph (Cattaneo and Titiunik, 2018).

## 2 The Local Randomization Approach to RD Analysis

In the first monograph (Cattaneo et al., 2018a), we discuss in detail the continuity-based approach to RD analysis. This approach, which is based on assumptions of continuity (and further smoothness) of the regression functions  $\mathbb{E}[Y_i(1)|X_i = x]$  and  $\mathbb{E}[Y_i(0)|X_i = x]$ , is by now the standard and most widely used method to analyze RD designs in practice. In this section, we discuss a different framework for RD analysis that is based on a formalization of the idea that the RD design can be interpreted as a sort of randomized experiment near the cutoff  $\bar{x}$ . This alternative framework can be used as a complement and robustness check to the continuity-based analysis when the running variable is continuous, and is the most natural framework when the running variable is discrete and has few mass points, a case we discuss extensively in Section 3 below.

When the RD design was first introduced by Thistlethwaite and Campbell (1960), the justification for this novel research design was not based on approximation and extrapolation of smooth regression functions, but rather on the idea that the abrupt change in treatment status that occurs at the cutoff leads to a treatment assignment mechanism that, near the cutoff, resembles the assignment that we would see in a randomized experiment. Indeed, the authors described a hypothetical experiment where the treatment is randomly assigned near the cutoff as an “experiment for which the regression-discontinuity analysis may be regarded as a substitute” (Thistlethwaite and Campbell, 1960, p. 310).

The idea that the treatment assignment is “as good as” randomly assigned in a neighborhood of the cutoff is often invoked in the continuity-based framework to describe the required identification assumptions in an intuitive way, and it has also been used to develop formal results. However, within the continuity-based framework, the formal derivation of identification and estimation results always relies on continuity and differentiability of regression functions, and the idea of local randomization is used as a heuristic device only. In contrast, what we call the *local randomization approach* to RD analysis formalizes that idea that the RD design behaves like a randomized experiment near the cutoff by imposing explicit randomization-type assumptions that are stronger than the standard continuity-type conditions. In a nutshell, this approach imposes conditions so that units whose score values lie in a small window around the cutoff can be analyzed as-if they were randomly assigned to treatment or control. The local randomization approach adopts the local randomization assumption explicitly, not as a heuristic interpretation, and builds a set of statistical tools exploiting this specific assumption.

We now introduce the local randomization approach in detail, discussing how adopting

an explicit randomization assumption near the cutoff allows for the use of new methods of estimation and inference for RD analysis. We also discuss the differences between the standard continuity-based approach and the local randomization approach. When the running variable is continuous, the local randomization approach typically requires stronger assumptions than the continuity-based approach; in these cases, it is natural to use the continuity-based approach for the main RD analysis, and to use the local randomization approach as a robustness check. But in settings where the running variable is discrete (with few mass points) or other departures from the canonical RD framework occur, the local randomization approach can not only be very useful but also possibly the only valid method for estimation and inference in practice.

Recall that we are considering a RD design where the (continuous) score is  $X_i$ , the treatment assignment is  $T_i = \mathbb{1}(X_i \geq \bar{x})$ ,  $Y_i(1)$  and  $Y_i(0)$  are the potential outcomes under treatment and control, respectively, and  $Y_i$  is the observed outcome. Throughout this section, we maintain the assumption that the RD design is sharp—that is, we assume that all units with score above the cutoff receive the treatment, and no units with score below the cutoff receive it.

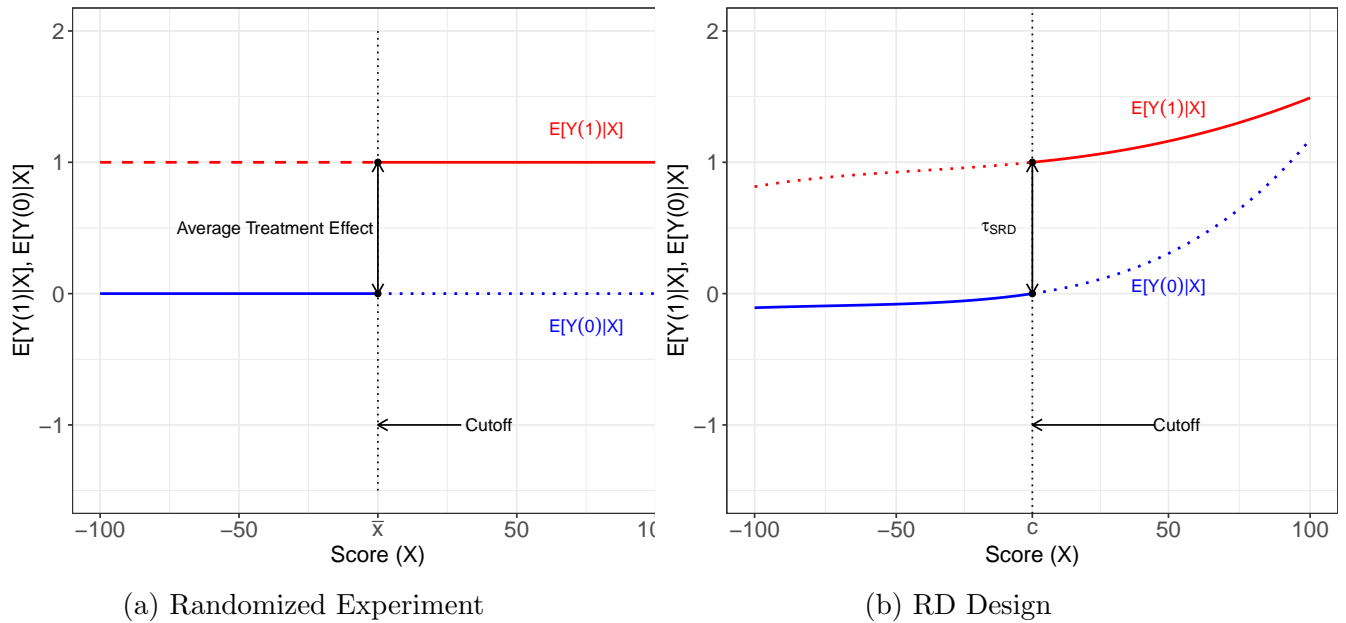
## 2.1 Local Randomization Approach: Overview

When the RD is based on a local randomization assumption, instead of assuming that the unknown regression functions  $\mathbb{E}[Y_i(1)|X_i = x]$  and  $\mathbb{E}[Y_i(0)|X_i = x]$  are continuous at the cutoff, the researcher assumes that there is a small window around the cutoff,  $W_0 = [\bar{x} - w_0, \bar{x} + w_0]$ , such that for all units whose scores fall in that window their placement above or below the cutoff is assigned as in a randomized experiment—an assumption that is sometimes called *as if random assignment*. Formalizing the assumption that the treatment is (locally) assigned as it would have been assigned in an experiment requires careful consideration of the conditions that are guaranteed to hold in an actual experimental assignment.

There are important differences between the RD design and an actual randomized experiment. To discuss such differences, we start by noting that any simple experiment can be recast as an RD design where the score is a randomly generated number, and the cutoff is chosen to ensure a certain treatment probability. For example, consider an experiment in a student population that randomly assigns a scholarship with probability 1/2. This experiment can be seen as an RD design where each student is assigned a random number with uniform distribution between 0 and 100, say, and the scholarship is given to students whose number or score is above 50. We illustrate this scenario in Figure 2.1(a).

The crucial feature of a randomized experiment recast as an RD design is that the running variable, by virtue of being a randomly generated number, is unrelated to the average potential outcomes. This is the reason why, in Figure 2.1(a), the average potential outcomes  $\mathbb{E}[Y_i(1)|X_i = x]$  and  $\mathbb{E}[Y_i(0)|X_i = x]$  are constant for all values of  $x$ . Since the regression functions are flat, the vertical distance between them can be recovered by the difference between the average observed outcomes among all units in the treatment and control groups, i.e.  $\mathbb{E}[Y_i|X_i \geq 50] - \mathbb{E}[Y_i|X_i < 50] = \mathbb{E}[Y_i(1)|X_i \geq 50] - \mathbb{E}[Y_i(0)|X_i < 50] = \mathbb{E}[Y_i(1)] - \mathbb{E}[Y_i(0)]$ , where the last equality follows from the assumption that  $X_i$  is a randomly generated number and thus is unrelated to (i.e., independent of)  $Y_i(1)$  and  $Y_i(0)$ .

Figure 2.1: Experiment versus RD Design



In contrast, in the standard continuity-based RD design there is no requirement that the potential outcomes be unrelated to the running variable over its support. Figure 2.1(b) illustrates a standard continuity-based RD design where the average treatment effect at the cutoff is the same as in the experimental setting in Figure 2.1(a),  $\tau_{\text{SRD}}$ , but where the average potential outcomes are non-constant functions of the score. This relationship between running variable and potential outcomes is characteristic of many RD designs: since the score is often related to the units' ability, resources, or performance (poverty index, vote shares, test scores), units with higher scores are often systematically different from units whose scores are lower. For example, a RD design where the score is a party's vote share in a given election and the outcome of interest is the party's vote share in the following election, the overall relationship between the score and the outcome will likely have a strongly positive slope,

as districts that strongly support the party in one election are likely to continue to support the party in the near future. As illustrated in Figure 2.1(a), a nonzero slope in the plot of  $\mathbb{E}[Y_i|X_i = x]$  against  $x$  does not occur in an actual experiment, because in an experiment  $x$  is an arbitrary random number unrelated to the potential outcomes.

The crucial difference between the scenarios in Figures 2.1(a) and 2.1(b) is our knowledge about the functional form of the regression functions. In a continuity-based approach, the RD treatment effect in 2.1(b) can be estimated by calculating the limit of the average observed outcomes as the score approaches the cutoff for the treatment and control groups,  $\lim_{x \downarrow \bar{x}} \mathbb{E}[Y_i|X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[Y_i|X_i = x]$ . The estimation of these limits requires that the researcher approximate the regression functions, and this approximation will typically contain an error that may directly affect estimation and inference. This is in stark contrast to the experiment depicted in Figure 2.1(a), where the random assignment of the score implies that the average potential outcomes are unrelated to the score and estimation does not require functional form assumptions—by construction, the regression functions are constant in the entire region where the score is randomly assigned.

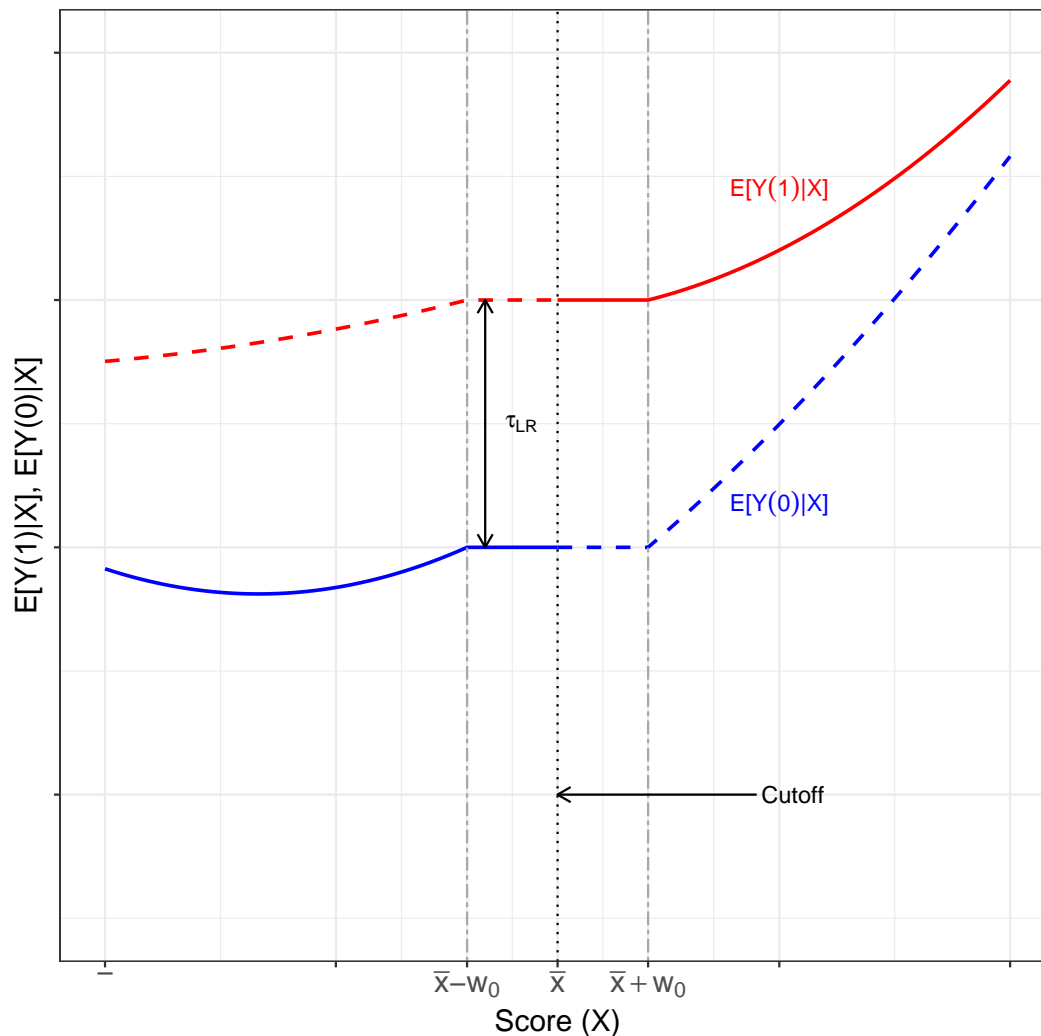
A point often overlooked is that the known functional form of the regression functions in a true experiment does not follow from the random assignment of the score per se, but rather from the score being an arbitrary computer-generated number that is unrelated to the potential outcomes. If the value of the score were randomly assigned but had a direct effect on the average outcomes, the regression functions in Figure 2.1(a) would not necessarily be flat. Thus, a local randomization approach to RD analysis must be based not only on the assumption that placement above or below the cutoff is randomly assigned within a window of the cutoff, but also on the assumption that the value of the score within this window is unrelated to the potential outcomes—a condition that is guaranteed neither by the random assignment of the score  $X_i$ , nor by the random assignment of the treatment  $T_i$ .

Formally, letting  $W_0 = [\bar{x} - w, \bar{x} + w]$ , the local randomization assumption can be stated as the two following conditions:

- (LR1) The distribution of the running variable in the window  $W_0$ ,  $F_{X_i|X_i \in W_0}(x)$ , is known, is the same for all units, and does not depend on the potential outcomes:  $F_{X_i|X_i \in W_0}(x) = F(x)$
- (LR2) Inside  $W_0$ , the potential outcomes depend on the running variable solely through the treatment indicator  $T_i = \mathbb{1}(X_i \geq \bar{x})$ , but not directly:  $Y_i(X_i, T_i) = Y_i(T_i)$  for all  $i$  such that  $X_i \in W_0$ .

Under these conditions, inside the window  $W_0$ , placement above or below the cutoff is unrelated to the potential outcomes, and the potential outcomes are unrelated to the running variable; therefore, the regression functions are flat inside  $W_0$ . This is illustrated in Figure 2.2, where  $\mathbb{E}[Y_i(1)|X_i = x]$  and  $\mathbb{E}[Y_i(0)|X_i = x]$  are constant for all values of  $x$  inside  $W_0$ , but have non-zero slopes outside of it.

Figure 2.2: Local Randomization RD



The contrast between Figures 2.1(a), 2.1(b), and 2.2 illustrates the differences between the actual experiment where the score is a randomly generated number, a continuity-based RD design, and a local randomization RD design. In the actual experiment where the score is a random number, the potential outcomes are unrelated to the score for all possible score values—i.e., in the entire support of the score. In this case, there is no uncertainty about the functional forms of  $\mathbb{E}[Y_i(1)|X_i = x]$  and  $\mathbb{E}[Y_i(0)|X_i = x]$ . In the continuity-based



RD design, the potential outcomes can be related to the score everywhere; the functions  $\mathbb{E}[Y_i(1)|X_i = x]$  and  $\mathbb{E}[Y_i(0)|X_i = x]$  are completely unknown, and estimation and inference is based on approximating them at the cutoff. Finally, in the local randomization RD design, the potential outcomes can be related to the running variable far from the cutoff, but there is a window around the cutoff where this relationship ceases. In this case, the functions  $\mathbb{E}[Y_i(1)|X_i = x]$  and  $\mathbb{E}[Y_i(0)|X_i = x]$  are unknown over the entire support of the running variable, but inside the window  $W_0$  they are assumed to be constant functions of  $x$ .

In some applications, assuming that the score has no effect on the (average) potential outcomes near the cutoff may be regarded as unrealistic or too restrictive. However, such an assumption can be taken as an approximation, at least for the very few units with scores extremely close to the RD cutoff. As we will discuss below, a key advantage of the local randomization approach is that it leads to valid and powerful finite sample inference methods, which remain valid and can be used even when only a handful of observations very close to the cutoff are included in the analysis.

Furthermore, the restriction that the score cannot directly affect the (average) potential outcomes near the cutoff can be relaxed if the researcher is willing to impose more parametric assumptions (locally to the cutoff). The local randomization assumption assumes that, inside the window where the treatment is assumed to have been randomly assigned, the potential outcomes are entirely unrelated to the running variable. This assumption, also known as the exclusion restriction, leads to the flat regression functions in Figure 2.2. It is possible to consider a slightly weaker version of this assumption where the potential outcomes are allowed to depend on the running variable, but there exists a transformation that, once applied to the potential outcomes of the units inside the window  $W_0$ , leads to transformed potential outcomes that are unrelated to the running variable.

More formally, the exclusion restriction in (LR2) requires that, for units with  $X_i \in W_0$ , the potential outcomes satisfy  $Y_i(X_i, T_i) = Y_i(T_i)$ —that is, the potential outcomes depend on the running variable only via the treatment assignment indicator and not via the particular value taken by  $X_i$ . In contrast, the weaker alternative assumption requires that, for units with  $X_i \in W_0$ , there exists a transformation  $\phi(\cdot)$  such that

$$\phi(Y_i(X_i, T_i), X_i, T_i) = \tilde{Y}_i(T_i).$$

This condition says that, although the potential outcomes are allowed to depend on the running variable  $X_i$  directly, the transformed potential outcomes  $\tilde{Y}_i(T_i)$  depend only on the treatment assignment indicator and thus satisfy the original exclusion restriction in (LR2).

For implementation, a transformation  $\phi(\cdot)$  must be assumed; for example, one can use a polynomial of order  $p$  on the unit's score, with slopes that are constant for all individuals on the same side of the cutoff. This transformation has the advantage of linking the local randomization approach to RD analysis to the continuity-based approach discussed in Part I ([Cattaneo et al., 2018a](#)).

## 2.2 Local Randomization Estimation and Inference

Adopting a local randomization approach to RD analysis implies assuming that the assignment of units above or below the cutoff was random inside the window  $W_0$  (condition LR1), and that in this window the potential outcomes are unrelated to the score (condition LR2)—or can be somehow transformed to be unrelated to the score.

Therefore, given knowledge of  $W_0$ , under a local randomization RD approach, we can analyze the data as we would analyze an experiment. If the number of observations inside  $W_0$  is large, researchers can use the full menu of standard large-sample experimental methods, all of which are based on large-sample approximations—that is, on the assumption that the number of units inside  $W_0$  is large enough to be well approximated by large sample limiting distributions. These methods may or may not involve the assumption of random sampling, and may or may not require LR2 per se (though removing LR2 will change the interpretation of the RD parameter in general). In contrast, if the number of observations inside  $W_0$  is very small, as it is often the case when local randomization methods are invoked in RD designs, estimation and inference based on large-sample approximations may be invalid; in this case, under appropriate assumptions, researchers can still employ randomization-based inference methods that are exact in finite samples and do not require large-sample approximations for their validity. These methods rely on the random assignment of treatment to construct confidence intervals and hypothesis tests. We review both types of approaches below.

The implementation of experimental methods to analyze RD designs requires knowledge or estimation of two important ingredients: (i) the window  $W_0$  where the local randomization assumption is invoked; and (ii) the randomization mechanism that is needed to approximate the assignment of units within  $W_0$  to the treatment and control conditions (i.e., to placement above or below the cutoff). In real applications,  $W_0$  is fundamentally unknown and must be selected by the research (ideally in an objective and data-driven way). Once  $W_0$  has been estimated, the choice of the randomization mechanism can be guided by the structure of the data, and it is not needed if large sample approximations are invoked. In most applications, the most natural assumption for the randomization mechanism is either complete

randomization or a Bernoulli assignment, where all units in  $W_0$  are assumed to have the same probability of being placed above or below the cutoff. We first assume that  $W_0$  is known and choose a particular random assignment mechanism inside  $W_0$ . In Section 2.3, we discuss a principled method to choose the window  $W_0$  in a data-driven way.

### 2.2.1 Finite Sample Methods

In many RD applications, a local randomization assumption will only be plausible in a very small window around the cutoff, and by implication this small window will often contain very few observations. In this case, it is natural to employ a Fisherian inference approach, which is valid in any finite sample, and thus leads to correct inferences even when the number of observations inside  $W_0$  is very small.

The Fisherian approach sees the potential outcomes as non-stochastic. This stands in contrast to the approach in the continuity-based RD framework, where the potential outcomes are random variables as a consequence of random sampling. More precisely, in Fisherian inference, the total number of units in the study,  $n$ , is seen as fixed—i.e., there is no random sampling assumption; moreover, inferences do not rely on assuming that this number is large. This setup is then combined with the so-called *sharp null hypothesis* that the treatment has no effect for any unit:

$$H_0^F : Y_i(0) = Y_i(1) \text{ for all } i.$$

The combination of non-stochastic potential outcomes and the sharp null hypothesis leads to inferences that are (type-I error) correct for any sample size because, under  $H_0^F$ , both potential outcomes— $Y_i(1)$  and  $Y_i(0)$ —can be imputed for every unit and there is no missing data. In other words, under the sharp null hypothesis, the observed outcome of each unit is equal to the unit’s two potential outcomes,  $Y_i = Y_i(1) = Y_i(0)$ . When the treatment assignment is known, observing all potential outcomes are under the null hypothesis allows us to derive the null distribution of any test statistic from the randomization distribution of the treatment assignment alone. Since the latter distribution is finite-sample exact, the Fisherian framework allows researchers to make inferences without relying on large-sample approximations.

#### *A hypothetical example*

To illustrate how Fisherian inference leads to the exact distribution of test statistics, we use a hypothetical example. We imagine that we have five units inside  $W_0$ , and we randomly assign  $n_{W_0,+} = 3$  units to treatment and  $n_{W_0,-} = n_{W_0} - n_{W_0,+} = 5 - 3 = 2$  units to control,

where  $n_{W_0}$  is the total number of units inside  $W_0$ . We choose the difference-in-means as the test-statistic,  $\bar{Y}_+ - \bar{Y}_- = \frac{1}{n_{W_0,+}} \sum_{i:i \in W_0} Y_i T_i + \frac{1}{n_{W_0,-}} \sum_{i:i \in W_0} Y_i (1 - T_i)$ . The treatment indicator continues to be  $T_i$ , and we collect in the set  $\mathcal{T}_{W_0}$  all possible  $n_{W_0}$ -dimensional treatment assignment vectors  $\mathbf{t}$  within the window.

For implementation, we must choose a particular treatment assignment mechanism. In other words, after assuming that placement above and below the cutoff was done as it would have been done in an experiment, we must choose a particular randomization distribution for the assignment. Of course, a crucial difference between an actual experiment and the RD design is that, in the RD design, the true mechanism by which units are assigned a value of the score smaller or larger than  $\bar{x}$  inside  $W_0$  is fundamentally unknown. Thus, the choice of the particular randomization mechanism is best understood as an approximation. A common choice is the assumption that, within  $W_0$ ,  $n_{W_0,+}$  units are assigned to treatment and  $n_{W_0} - n_{W_0,+}$  units are assigned to control, where each unit has probability  $\binom{n_{W_0}}{n_{W_0,+}}^{-1}$  of being assigned to the treatment (i.e. above the cutoff) group. This is commonly known as a complete randomization mechanism or a fixed margins randomization—under this mechanism, the number of treated and control units is fixed, as all treatment assignment vectors result in exactly  $n_{W_0,+}$  treated units and  $n_{W_0} - n_{W_0,+}$  control units.

In our example, under complete randomization, the number of elements in  $\mathcal{T}_{W_0}$  is  $\binom{5}{3} = 10$ —that is, there are ten different ways to assign five units to two groups of size three and two. We assume that  $Y_i(1) = 5$  and  $Y_i(0) = 2$  for all units, so that the treatment effect,  $Y_i(1) - Y_i(0)$ , is constant and equal to 3 for all units. The top panel of Table 2.1 shows the ten possible treatment assignment vectors,  $\mathbf{t}_1, \dots, \mathbf{t}_{10}$ , and the two potential outcomes for each unit.

Suppose that the observed treatment assignment inside  $W_0$  is  $\mathbf{t}_6$ , so that units 1, 4 and 5 are assigned to treatment, and units 2 and 3 are assigned to control. Given this assignment, the vector of observed outcomes is  $\mathbf{Y} = (5, 2, 2, 5, 5)$ , and the observed value of the difference-in-means statistic is  $S^{\text{obs}} = \bar{Y}_+ - \bar{Y}_- = \frac{5+5+5}{3} - \frac{2+2}{2} = 5 - 2 = 3$ . The bottom panel of Table 2.1 shows the distribution of the test statistic under the null—that is, the ten different possible values that the difference-in-means can take when  $H_0^F$  is assumed to hold. The observed difference-in-means  $S^{\text{obs}}$  is the largest of the ten, and the exact p-value is therefore  $p^F = 1/10 = 0.10$ . Thus, we can reject  $H_0^F$  with a test of level  $\alpha = 0.10$ . Note that, since the number of possible treatment assignments is ten, the smallest value that  $p^F$  can take is  $1/10$ . This p-value is finite-sample exact, because the null distribution in Table 2.1 was derived directly from the randomization distribution of the treatment assignment, and does not rely on any statistical model or large-sample approximations.

Table 2.1: Hypothetical Randomization Distribution with Five Units

All Possible Treatment Assignments												
	$Y_i(1)$	$Y_i(0)$	$\mathbf{t}_1$	$\mathbf{t}_2$	$\mathbf{t}_3$	$\mathbf{t}_4$	$\mathbf{t}_5$	$\mathbf{t}_6$	$\mathbf{t}_7$	$\mathbf{t}_8$	$\mathbf{t}_9$	$\mathbf{t}_{10}$
Unit 1	5	2	1	1	1	1	1	1	0	0	0	0
Unit 2	5	2	1	1	1	0	0	0	1	1	1	0
Unit 3	5	2	1	0	0	1	1	0	1	1	0	1
Unit 4	5	2	0	1	0	1	0	1	1	0	1	1
Unit 5	5	2	0	0	1	0	1	1	0	1	1	1
$\Pr(\mathbf{T} = \mathbf{t})$			1/10	1/10	1/10	1/10	1/10	1/10	1/10	1/10	1/10	1/10
Distribution of Difference-in-Means When $\mathbf{T} = \mathbf{t}_6$ and $\mathbf{Y} = (5, 2, 2, 5, 5)$												
			$S_{\mathbf{t}_1}^{\text{obs}}$	$S_{\mathbf{t}_2}^{\text{obs}}$	$S_{\mathbf{t}_3}^{\text{obs}}$	$S_{\mathbf{t}_4}^{\text{obs}}$	$S_{\mathbf{t}_5}^{\text{obs}}$	$S_{\mathbf{t}_6}^{\text{obs}}$	$S_{\mathbf{t}_7}^{\text{obs}}$	$S_{\mathbf{t}_8}^{\text{obs}}$	$S_{\mathbf{t}_9}^{\text{obs}}$	$S_{\mathbf{t}_{10}}^{\text{obs}}$
$\bar{Y}_+$			3	4	4	4	4	5	3	3	4	4
$\bar{Y}_-$			5	3.5	3.5	3.5	3.5	2	5	5	3.5	3.5
$\bar{Y}_+ - \bar{Y}_-$			-2	0.5	0.5	0.5	0.5	3	-2	-2	0.5	0.5
$\Pr(S = S_{\mathbf{t}_j}^{\text{obs}})$			1/10	1/10	1/10	1/10	1/10	1/10	1/10	1/10	1/10	1/10

This example illustrates that, in order to implement a local randomization RD analysis, we need to specify, in addition to the choice of  $W_0$ , the particular way in which the treatment was randomized—that is, knowledge of the distribution of the treatment assignment. In practice, the latter will not be known, but in many applications it can be approximated by assuming a complete randomization within  $W_0$ . Moreover, we need to choose a particular test statistic; the difference-in-means is a simple choice, but below we discuss other options.

### ***The General Fisherian Inference Framework***

We can generalize the above example to provide a general formula for the exact p-value associated with a test of  $H^F$ . As before, we let  $\mathbf{T}_{W_0}$  be the treatment assignment for the  $n_{W_0}$  units in  $W_0$ , and collect in the set  $\mathcal{T}_{W_0}$  all the possible treatment assignments that can occur given the assumed randomization mechanism. In a complete or fixed margins randomization,  $\mathcal{T}_{W_0}$  includes all vectors of length  $n_{W_0}$  such that each vector has  $n_{W_0,+}$  ones and  $n_{W_0,-} = n_{W_0} - n_{W_0,+}$  zeros. Similarly,  $\mathbf{Y}_{W_0}$  collects the  $n_{W_0}$  observed outcomes for units with  $X_i \in W_0$ . We also need to choose a test statistic, which we denote  $S(\mathbf{T}_{W_0}, \mathbf{Y}_{W_0})$ , that is a function of the treatment assignment  $\mathbf{T}_{W_0}$  and the vector  $\mathbf{Y}_{W_0}$  of observed outcomes for the  $n_{W_0}$  units in the experiment that is assumed to occur inside  $W_0$ .

Of all the possible values of the treatment vector  $\mathbf{T}_{W_0}$  that can occur, only one will have occurred in  $W_0$ ; we call this value the observed treatment assignment,  $\mathbf{t}_{W_0}^{\text{obs}}$ , and we denote  $S^{\text{obs}}$  the observed value of the test-statistic associated with  $\mathbf{t}_{W_0}^{\text{obs}}$ , i.e.  $S^{\text{obs}} = t(\mathbf{t}_{W_0}^{\text{obs}}, \mathbf{Y}_{W_0})$ . (In the hypothetical example discussed above, we had  $\mathbf{t}_{W_0}^{\text{obs}} = \mathbf{t}_6$ .) Then, the one-sided finite-sample exact p-value associated with a test of the sharp null hypothesis  $\mathbf{H}_0^F$  is the probability that the test-static exceeds its observed value:

$$p^F = \mathbb{P}(S(\mathbf{T}_{W_0}, \mathbf{Y}_{W_0}) \geq S^{\text{obs}}) = \sum_{\mathbf{t}_{W_0} \in \mathcal{T}_{W_0}} \mathbb{1}(S(\mathbf{t}_{W_0}, \mathbf{Y}_{W_0}) \geq S^{\text{obs}}) \cdot \mathbb{P}(\mathbf{T}_{W_0} = \mathbf{t}_{W_0}).$$

When each of the treatment assignments in  $\mathcal{T}_{W_0}$  is equally likely,  $\mathbb{P}(\mathbf{T} = \mathbf{t}) = \frac{1}{\#\{\mathcal{T}_{W_0}\}}$  with  $\#\{\mathcal{T}_{W_0}\}$  the number of elements in  $\mathcal{T}_{W_0}$ , and this expression simplifies to the number of times the test-statistic exceeds the observed value divided by the total number of test-statistics that can possibly occur,

$$p^F = \mathbb{P}(S(\mathbf{Z}_{W_0}, \mathbf{Y}_{W_0}) \geq S^{\text{obs}}) = \frac{\#\{S(\mathbf{t}_{W_0}, \mathbf{Y}_{W_0}) \geq S^{\text{obs}}\}}{\#\{\mathcal{T}_{W_0}\}}.$$

Under the sharp null hypothesis, all potential outcomes are known and can be imputed. To see this, note that under  $\mathbf{H}_0^F$  we have  $\mathbf{Y}_{W_0} = \mathbf{Y}_{W_0}(1) = \mathbf{Y}_{W_0}(0)$ , so that  $S(\mathbf{T}_{W_0}, \mathbf{Y}_{W_0}) = S(\mathbf{T}_{W_0}, \mathbf{Y}_{W_0}(0))$ . Thus, under  $\mathbf{H}_0^F$ , the only randomness in  $S(\mathbf{Z}_{W_0}, \mathbf{Y}_{W_0})$  comes from the random assignment of the treatment, which is assumed to be known.

In practice, it often occurs that the total number of different treatment vectors  $\mathbf{t}_{W_0}$  that can occur inside the window  $W_0$  is too large, and enumerating them exhaustively is unfeasible. For example, assuming a fixed-margins randomization inside  $W_0$  with 15 observations on each side of the cutoff, there are  $\binom{n_{W_0}}{n_{W_0,t}} = \binom{30}{15} = 155,117,520$  possible treatment assignments, and calculating  $p^F$  by complete enumeration is very time consuming and possibly unfeasible. When exhaustive enumeration is unfeasible, we can approximate  $p^F$  using simulations, as follows:

1. Calculate the observed test statistic,  $S^{\text{obs}} = S(\mathbf{t}_{W_0}^{\text{obs}}, \mathbf{Y}_{W_0})$ .
2. Draw a value  $\mathbf{t}_{W_0}^j$  from the treatment assignment distribution,  $\mathbb{P}(\mathbf{T}_{W_0} \leq \mathbf{t}_{W_0})$ .
3. Calculate the test statistic for the  $j^{\text{th}}$  draw  $\mathbf{t}_{W_0}^j$ ,  $S(\mathbf{t}_{W_0}^j, \mathbf{Y}_{W_0})$ .
4. Repeat steps 2 and 3  $B$  times.

5. Calculate the simulation approximation to  $p^F$  as

$$\tilde{p}^F = \frac{1}{B} \sum_{j=1}^B \mathbb{1}(S(\mathbf{t}_{W_0}^j, \mathbf{Y}_{W_0}) \geq S^{\text{obs}}).$$

Fisherian confidence intervals can be obtained by specifying sharp null hypotheses about treatment effects, and then inverting these tests. In order to apply the Fisherian framework, the null hypotheses to be inverted must be sharp—that is, under these null hypotheses, the full profile of potential outcomes must be known. This requires specifying a treatment effect model, and testing hypotheses about the specified parameters. A simple and common choice is a constant treatment effect model,  $Y_i(1) = Y_i(0) + \tau$ , which leads to the null hypothesis  $\mathbf{H}_{\tau_0}^F : \tau = \tau_0$ —note that  $\mathbf{H}_0^F$  is a special case of  $\mathbf{H}_{\tau_0}^F$  when  $\tau_0 = 0$ . Under this model, a  $1 - \alpha$  confidence interval for  $\tau$  can be obtained by collecting the set of all the values  $\tau_0$  that fail to be rejected when we test  $\mathbf{H}_{\tau_0}^F : \tau = \tau_0$  with an  $\alpha$ -level test.

To test  $\mathbf{H}_{\tau_0}^F$ , we build test statistics based on an adjustment to the potential outcomes that renders them constant under this null hypothesis. Under  $\mathbf{H}_{\tau_0}^F$ , the observed outcome is

$$\begin{aligned} Y_i &= T_i \cdot Y_i(1) + (1 - T_i) \cdot Y_i(0) \\ &= T_i \cdot (Y_i(0) + \tau_0) + (1 - T_i) \cdot Y_i(0) \\ &= T_i \cdot \tau_0 + Y_i(0). \end{aligned}$$

Thus, the adjusted outcome  $\ddot{Y}_i \equiv Y_i - T_i \tau_0 = Y_i(0)$  is constant under the null hypothesis  $\mathbf{H}_{\tau_0}^F$ . A randomization-based test of  $\mathbf{H}_{\tau_0}^F$  proceeds by first calculating the adjusted outcomes  $\ddot{Y}_i$  for all the units in the window, and then computing the test statistic using the adjusted outcomes instead of the raw outcomes, i.e. computing  $S(\mathbf{T}_{W_0}, \ddot{\mathbf{Y}}_{W_0})$ . Once the adjusted outcomes are used to calculate the test statistic, we have  $S(\mathbf{T}_{W_0}, \ddot{\mathbf{Y}}_{W_0}) = S(\mathbf{T}_{W_0}, \mathbf{Y}_{W_0}(0))$  as before, and a test of  $\mathbf{H}_{\tau_0}^F : \tau = \tau_0$  can be implemented as a test of the sharp null hypothesis  $\mathbf{H}_0^F$ , using  $S(\mathbf{Z}_{W_0}, \ddot{\mathbf{Y}}_{W_0})$  instead of  $S(\mathbf{Z}_{W_0}, \mathbf{Y}_{W_0})$ . We use  $p_{\tau_0}^F$  to refer to the p-value associated with a randomization-based test of  $\mathbf{H}_{\tau_0}^F$ .

In practice, assuming that  $\tau$  takes values in  $[\tau_{\min}, \tau_{\max}]$ , computing these confidence intervals requires building a grid  $G^{\tau_0} = \{\tau_0^1, \tau_0^2, \dots, \tau_0^G\}$ , with  $\tau_0^1 \geq \tau_{\min}$  and  $\tau_0^G \leq \tau_{\max}$ , and collecting all  $\tau_0 \in G^{\tau_0}$  that fail to be rejected with an  $\alpha$ -level test of  $\mathbf{H}_{\tau_0}^F$ . Thus, the Fisherian  $(1 - \alpha) \times 100\%$  confidence intervals is

$$\text{CI}^{\text{LRF}} = \{\tau_0 \in G^{\tau_0} : p_{\tau_0}^F \leq \alpha\}.$$

The general principle of Fisherian inference is to use the randomization-based distribution of the test statistic under the sharp null hypothesis to derive p-values and confidence intervals. In our hypothetical example, we illustrated the procedure using the difference-in-means test statistic and the fixed margins randomization mechanism. But the Fisherian approach to inference is general and works for any appropriate choice of test statistic and randomization mechanism.

Other test statistics that could be used include the Kolmogorov-Smirnov (KS) statistic and the Wilcoxon rank sum statistic. The KS statistic is defined as  $S_{\text{KS}} = \sup_y |\hat{F}_1(y) - \hat{F}_0(y)|$ , and measures the maximum absolute difference in the empirical cumulative distribution functions (CDF) of the treated and control outcomes—denoted respectively by  $\hat{F}_1(\cdot)$  and  $\hat{F}_0(\cdot)$ . Because  $S_{\text{KS}}$  is the treated-control difference in the outcome CDFs, it is well suited to detect departures from the null hypothesis that involve not only differences in means but also differences in other moments and in quantiles. Another test statistic commonly used is the Wilcoxon rank sum statistic, which is based on the ranks of the outcomes, denoted  $R_i^y$ . This statistic is  $S_{\text{WR}} = \sum_{i:T_i=1} R_i^y$ , that is, it is the sum of the ranks of the treated observations. Because  $S_{\text{WR}}$  is based on ranks, it is not affected by the particular values of the outcome, only by their ordering. Thus, unlike the difference-in-means,  $S_{\text{WR}}$  is insensitive to outliers.

In addition to different choices of test statistics, the Fisherian approach allows for different randomization mechanisms. An alternative to the complete randomization mechanism discussed above is a Bernoulli assignment, where each unit is assigned to treatment with some fixed equal probability. For implementation, researchers can set this probability equal to 1/2 or, alternatively, equal to the proportion of treated units in  $W_0$ . The disadvantage of a Bernoulli assignment is that it can result in a treated or a control group with few or no observations—a phenomenon that can never occur under complete randomization. However, in practice, complete randomization and Bernoulli randomization often lead to very similar conclusions for the same window  $W_0$ .

## 2.2.2 Large Sample Methods

Despite the conceptual elegance of finite-sample Fisherian methods, the most frequently chosen methods in the analysis of experiments are based on large-sample approximations. These alternative methods are appropriate to analyze RD designs under a local randomization assumption when the number of observations inside  $W_0$  is sufficiently large to ensure that the moment and/or distributional approximations are sufficiently similar to the finite-sample distributions of the statistics of interest.



A classic framework for experimental analysis is known as the Neyman approach. This approach relies on large-sample approximations to the randomization distribution of the treatment assignment, but still assumes that the potential outcomes are fixed or non-stochastic. In other words, the Neyman approach assumes that the sample size grows to infinity but does not assume that the data is a (random) sample from a super-population. Unlike in the Fisherian approach, in the Neyman framework point estimation is one of the main goals, and the parameter of interest is typically the finite-sample average treatment effect. Inference procedures in this framework usually focus on the null hypothesis that the average treatment effect is zero.

To be more specific, consider the *local randomization sharp RD effect*, defined as

$$\tau_{\text{SRD}}^{\text{LR}} = \bar{Y}(1) - \bar{Y}(0), \quad \bar{Y}(1) = \frac{1}{n_{W_0}} \sum_{i: X_i \in W_0} Y_i(1), \quad \bar{Y}(0) = \frac{1}{n_{W_0}} \sum_{i: X_i \in W_0} Y_i(0)$$

where  $\bar{Y}(1)$  and  $\bar{Y}(0)$  are the average potential outcomes inside the window. In this definition, we have assumed that the potential outcomes are non-stochastic.

The parameter  $\tau_{\text{SRD}}^{\text{LR}}$  is different from the more conventional continuity-based RD parameter  $\tau_{\text{SRD}}$  defined in the introduction and discussed extensively in our companion Part I monograph. While  $\tau_{\text{SRD}}^{\text{LR}}$  is an average effect inside an interval (the window  $W_0$ ),  $\tau_{\text{SRD}}$  is an average at a single point (the cutoff  $\bar{x}$ ) where, by construction, the number of observations is zero. This means that the decision to adopt a continuity-based approach versus a local randomization approach directly affects the definition of the parameter of interest. Naturally, if the window  $W_0$  is extremely small,  $\tau_{\text{SRD}}^{\text{LR}}$  and  $\tau_{\text{SRD}}$  become more conceptually similar.

Under the assumption of complete randomization inside  $W_0$ , the observed difference-in-means is an unbiased estimator of  $\tau_{\text{SRD}}^{\text{LR}}$ . Thus a natural estimator for the RD effect  $\tau_{\text{SRD}}^{\text{LR}}$  is the difference between the average observed outcomes in the treatment and control groups,

$$\hat{\tau}_{\text{SRD}}^{\text{LR}} = \bar{Y}_+ - \bar{Y}_-, \quad \bar{Y}_+ = \frac{1}{n_{W_0,+}} \sum_{i: X_i \in W_0} Y_i \mathbb{1}(X_i \geq \bar{x}), \quad \bar{Y}_- = \frac{1}{n_{W_0,-}} \sum_{i: X_i \in W_0} Y_i \mathbb{1}(X_i < \bar{x}),$$

where  $\bar{Y}_+$  and  $\bar{Y}_-$  are the average treated and control observed outcomes inside  $W_0$  and, as before,  $n_{W_0,+}$  and  $n_{W_0,-}$  are the number of treatment and control units inside  $W_0$ , respectively. In this case, a conservative estimator of the variance of  $\tau_{\text{SRD}}^{\text{LR}}$  is given by  $\hat{V} = \frac{\hat{\sigma}_+^2}{n_{W_0,+}} + \frac{\hat{\sigma}_-^2}{n_{W_0,-}}$ , where  $\hat{\sigma}_+^2$  and  $\hat{\sigma}_-^2$  denote the sample variance of the outcome for the treatment and control units within  $W_0$ , respectively. A  $100(1 - \alpha)\%$  confidence interval can be constructed in the usual way by relying on a normal large-sample approximation to the randomization distribution of the treatment assignment. For example, an approximate 95% confidence interval

is

$$\text{CI}^{\text{LRN}} = \left[ \hat{\tau}_{\text{SRD}}^{\text{LR}} \pm 1.96 \cdot \sqrt{\widehat{\mathbb{V}}} \right].$$

Hypothesis testing is based on Normal approximations. The Neyman null hypothesis is

$$\mathbf{H}_0^{\text{N}} : \bar{Y}(1) - \bar{Y}(0) = 0.$$

In contrast to Fisher’s sharp null hypothesis  $\mathbf{H}_0^{\text{F}}$ ,  $\mathbf{H}_0^{\text{N}}$  does not allow us to calculate the full profile of potential outcomes for every possible realization of the treatment assignment vector  $\mathbf{t}$ . Thus, unlike the Fisherian approach, the Neyman approach to hypothesis testing must rely on approximation and is therefore not exact. In the Neyman approach, we can construct the usual t-statistic using the point and variance estimators,  $S = \frac{\bar{Y}_+ - \bar{Y}_-}{\sqrt{\widehat{\mathbb{V}}}}$ , and then use the Normal approximation to its distribution. For example, for a one-sided test, the p-value associated with a test of  $\mathbf{H}_0^{\text{N}}$ , is  $p^{\text{N}} = 1 - \Phi(t)$ , where  $\Phi(\cdot)$  is the Normal CDF.

Finally, it is possible to consider a setup where, in addition to using large-sample approximations to the randomization mechanism as in the Neyman approach, the data  $\{Y_i, X_i\}_{i=1}^n$  is seen as random sample from a larger population—the same assumption made by the continuity-based methods discussed in Cattaneo et al. (2018a). When random sampling is assumed, the potential outcomes  $Y_i(1)$  and  $Y_i(0)$  are considered random variables, and the units inside  $W_0$  are seen as a random sample from a (large) super-population. Because in this case the potential outcomes within  $W_0$  become stochastic by virtue of the random sampling, the parameter of interest is the super-population average treatment effect,  $\mathbb{E}[Y_i(1) - Y_i(0) | X_i \in W_0]$ . Adopting this super-population perspective does not change the Neyman estimation or inference procedures discussed above, though it does affect the interpretation of the results.

### 2.2.3 The Effect of Islamic Representation on Female Educational Attainment

We illustrate the local randomization methods with the study originally conducted by Meyersson (2014)—henceforth Meyersson. This is the same example we used for illustration purposes throughout our companion Part I monograph. Meyersson employs a Sharp RD design in Turkey to study the effect of Islamic parties’ control of local governments on the educational attainment of young women. (For brevity, we refer to a mayor who belongs to one of the Islamic parties as an “Islamic mayor”, and to a mayor who belongs to a non-Islamic party as a “secular mayor”.) Naturally, municipalities where Islamist parties win elections may be systematically different from municipalities where Islamist parties are de-

feated, making the RD an appealing strategy to circumvent these methodological challenges and estimate the causal effect of local Islamic rule.

Meyersson’s study is focused exclusively on the 1994 Turkish mayoral elections. The unit of analysis is the municipality, and the score or running variable is the Islamic margin of victory—defined as the difference between the vote share obtained by the largest Islamic party, and the vote share obtained by the largest secular party opponent. Although two Islamic parties compete in the 1994 mayoral elections, *Refah* and *Büyük Birlik Partisi* (BBP), BBP won in only 11 out of the 329 municipalities where an Islamic mayor was elected; thus, the results correspond largely to the effect of a *Refah* victory.

The Islamic margin of victory can be positive or negative, and the cutoff that determines an Islamic party victory is located at zero. The treatment group is the of municipalities that elect a mayor from an Islamic party in 1994, and the control group is the municipalities that elect a mayor from a secular party. The outcome that we re-analyze is the share of the cohort of women aged 15 to 20 in 2000 who had completed high school by 2000. For brevity, we refer to it interchangeably as *female high school attainment share*, *female high school attainment*, or *high school attainment for women*.

In our analysis below, we rename the variables in the following way:

- **Y**: high school attainment for women in 2000, measured as the share of women aged 15 to 20 in 2000 who had completed high school by 2000.
- **X**: vote margin obtained by the Islamic candidate for mayor in the 1994 Turkish elections, measured as the vote percentage obtained by the Islamic candidate minus the vote percentage obtained by its strongest opponent.
- **T**: electoral victory of the Islamic candidate in 1994, equal to 1 if Islamic candidate won the election and 0 if the candidate lost.

The Meyersson dataset also contains several predetermined covariates that we use in our re-analysis: the Islamic vote share in 1994 (`vshr_islam1994`), the number of parties receiving votes in 1994 (`partycount`), the logarithm of the population in 1994 (`lpop1994`), an indicator equal to one if the municipality elected an Islamic party in the previous election in 1989 (`i89`), a district center indicator (`merkezi`), a province center indicator (`merkezp`), a sub-metro center indicator (`subbuyuk`), and a metro center indicator (`buyuk`).

Table 2.2, also presented in Part I, presents descriptive statistics for the three RD variables (Y, X, and T), and the municipality-level predetermined covariates. The outcome of

Table 2.2: Descriptive Statistics for Meyersson

Variable	Mean	Median	Std. Deviation	Min.	Max.	Obs.
Y	16.306	15.523	9.584	0.000	68.038	
X	-28.141	-31.426	22.115	-100.000	99.051	
T	0.120	0.000	0.325	0.000	1.000	
Percentage of men aged 15-20 with high school education	19.238	18.724	7.737	0.000	68.307	
Islamic percentage of votes in 1994	13.872	7.029	15.385	0.000	99.526	
Number of parties receiving votes 1994	5.541	5.000	2.192	1.000	14.000	
Log population in 1994	7.840	7.479	1.188	5.493	15.338	
Percentage of population below 19 in 2000	40.511	39.721	8.297	6.544	68.764	
Percentage of population above 60 in 2000	9.222	8.461	3.960	1.665	27.225	
Gender ratio in 2000	107.325	103.209	25.293	74.987	1033.636	
Household size in 2000	5.835	5.274	2.360	2.823	33.634	
District center	0.345	0.000	0.475	0.000	1.000	
Province center	0.023	0.000	0.149	0.000	1.000	
Sub-metro center	0.022	0.000	0.146	0.000	1.000	

interest (Y) has a minimum of 0 and a maximum of 68.04, with a mean of 16.31. Thus, on average 16.31% of women in this cohort had completed high school by the year 2000. The Islamic vote margin (X) ranges from  $-100$  (party receives zero votes) to  $100$  (party receives 100% of the vote), with a mean of  $-28.14$ , implying that the Islamic party loses by 28.14 percentage points in the average municipality. Consistent with this, the mean of the treatment variable (T) is 0.120, indicating that in 1994 an Islamic mayor was elected in only 12.0% of the municipalities.

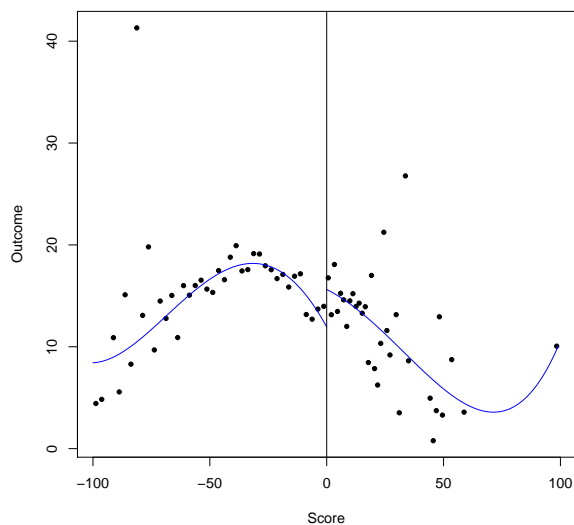


Figure 2.3: Mimicking Variance RD Plot with Evenly-Spaced Bins—Meyersson Data

Figure 2.3 presents an RD plot that illustrates the continuity-based average treatment effect at the cutoff that we estimated in our companion Part I monograph. The figure plots the

female educational attainment outcome against the Islamist margin of victory, where the solid line represents a fourth-order polynomial fit, the dots are local means computed in mimicking-variance evenly-spaced bins, and observations above the cutoff correspond to municipalities where an Islamic party won the 1994 mayoral election. Right at the cutoff, the average female educational attainment seems lower for municipalities where the Islamic party loses than for municipalities where the Islamic party barely wins. Using a continuity-based analysis, in Part I we show that the local-polynomial estimate of this effect is roughly 3 percentage points. We now reproduce these results for comparability with the local randomization analysis that we report below. We use `rdrobust` to fit a local linear polynomial within a mean-squared-error (MSE) optimal bandwidth—for further details, see Section 4 in Part I). With these specifications, the local polynomial effect of a bare Islamic victory on the female educational attainment share is 3.020, with robust p-value of 0.076.

```
> out = rdrobust(Y, X, kernel = "triangular", p = 1, bwselect = "mserd")
> summary(out)
Call: rdrobust
```

```
Number of Obs.          2629
BW type                mserd
Kernel                 Triangular
VCE method             NN

Number of Obs.          2314          315
Eff. Number of Obs.    529          266
Order est. (p)         1            1
Order bias (p)         2            2
BW est. (h)            17.239        17.239
BW bias (b)            28.575        28.575
rho (h/b)              0.603         0.603
```

```
=====
              Method      Coef. Std. Err.      z    P>|z|      [ 95% C.I. ]
=====
Conventional    3.020      1.427      2.116   0.034   [0.223 , 5.817]
Robust          -          -          1.776   0.076   [-0.309 , 6.276]
=====
```

Analogous Stata command

```
. rdrobust Y X, kernel(triangular) p(1) bwselect(mserd)
```

### 2.2.4 Estimation and Inference in Practice

We start the local randomization analysis of the Meyersson application using the function `rdrandinf`, which is part of the `rdlocrand` library/package. The main arguments of `rdrandinf` include the outcome variable `Y`, the running variable `X`, and the upper and lower limits of the window where inferences will be performed (`wr` and `wl`). We first choose the ad-hoc window  $[-2.5, 2.5]$ , postponing the discussion of automatic data-driven window selection until the next section. To make inferences in  $W = [-2.5, 2.5]$ , we set `wl` =  $-2.5$  and `wr` =  $2.5$ . Since Fisherian methods are simulation-based, we also choose the number of simulations via the argument `reps`, in this case choosing 1,000 simulations. Finally, in order to be able to replicate the Fisherian simulation-based results at a later time, we set the random seed using the `seed` argument.

```
> out = rdrandinf(Y, X, wl = -2.5, wr = 2.5, seed = 50)
```

```
Selected window = [-2.5;2.5]
```

```
Running randomization-based test...
```

```
Randomization-based test complete.
```

```
Number of obs =      2629
Order of poly =      0
Kernel type   =      uniform
Reps          =      1000
Window        =      set by user
H0:          tau =      0
Randomization =      fixed margins
```

Cutoff c = 0	Left of c	Right of c
Number of obs	2314	315
Eff. number of obs	68	62
Mean of outcome	13.972	15.044
S.d. of outcome	8.541	9.519
Window	-2.5	2.5

		Finite sample	Large sample	
Statistic	T	P> T	P> T	Power vs d =
4.27				
Diff. in means	1.072	0.488	0.501	0.765

Analogous Stata command

```
. rdrandinf Y X, wl(-2.5) wr(2.5) seed(50)
```

The output is divided in three panels. The top panel first presents the total number of observations in the entire dataset (that is, in the entire support of the running variable), the order of the polynomial used to transform the outcomes, and the kernel function that is used to weigh the observations. By default, `rdlocrand` uses a polynomial of order zero, which means the outcomes are not transformed. In order to transform the outcomes via a polynomial as explained above, users can use the option `p` in the call to `rdlocrand`. The default is also to use a uniform kernel, that is, to compute the test statistic using the unweighted observations. This default behavior can be changed with the option `kernel`. The rest of the top panel reports the number of simulations used for Fisherian inference, the method used to choose the window, and the null hypothesis that is tested (default is  $\tau_0 = 0$ , i.e. a test of  $H_0^F$  and  $H_0^N$ ). Finally, the last row of the top panel reports the chosen randomization mechanism, which by default is fixed margins (i.e. complete) randomization.

The middle panel reports the number of observations to the left and right of the cutoff in both the entire support of the running variable, and in the chosen window. Although there is a total of 2314 control observations and 315 treated observations in the entire dataset, the number of observations in the window  $[-2.5, 2.5]$  is much smaller, with only 68 municipalities below the cutoff and 62 municipalities above it. The middle panel also reports the mean and standard deviation of the outcome inside the chosen window.

The last panel reports the results. The first column reports the type of test statistic employed for testing the Fisherian sharp null hypothesis (the default is the difference-in-means), and the column labeled `T` reports its value. In this case, the difference-in-means is 1.072; given the information in the `Mean of outcome` row in the middle panel, we see that this is the difference between a female education share of 15.044 percentage points in municipalities where the Islamic party barely wins, and a female education share of 13.972 percentage points in municipalities where the Islamic party barely loses. The `Finite sample` column reports the p-value associated with a randomization-based test of the Fisherian sharp null hypothesis  $H_0^F$  (or the alternative sharp null hypothesis  $H_{\tau_0}^F$  based on a constant treatment effect model if the user sets  $\tau_0 \neq 0$  via the option `nulltau`). This p-value is 0.488, which means we fail to reject the sharp null hypothesis.

Finally, the `Large sample` columns in the bottom panel report Neyman inferences based on the large sample approximate behavior of the (distribution of the) statistic. The p-value reported in the large-sample columns is thus  $p^N$ , the p-value associated with a test of the Neyman null hypothesis  $H_0^N$  that the average treatment effect is zero. The last column in the bottom panel reports the power of the Neyman test to reject a true average treatment effect equal to `d`, where by default `d` is set to one half of the standard deviation of the outcome

variable for the control group, which in this case is 4.27 percentage points. The value of  $d$  can be modified with the options `d` or `dscale`. Like  $p^M$ , the calculation of the power versus the alternative hypothesis  $d$  is based on the Normal approximation. The large-sample p-value is 0.501, indicating that the Neyman null hypothesis also fails to be rejected at conventional levels. The power calculation indicates that the probability of rejecting the null hypothesis when the true effect is equal to half a (control) standard deviation is relatively high, at 0.765. Thus, it seems that the failure to reject the null hypothesis stems from the small size of the average treatment effect estimated in this window, which is just  $1.072/(4.27 \times 2) = 1.072/8.54 = 0.126$  standard deviations of the control outcome—a small effect.

It is also important to note the different interpretation of the difference-in-means test statistic in the Fisherian and Neyman frameworks. In Fisherian inference, the difference-in-means is simply one of the various test statistics that can be chosen to test the sharp null hypothesis, and should not be interpreted as an estimated effect—in Fisherian framework, the focus is on testing null hypotheses that are sharp, not on point estimation. In contrast, in the Neyman framework, the focus is on the sample average treatment effect; since the difference-in-means is an unbiased estimator of this parameter, it can be appropriately interpreted as an estimated effect.

To illustrate how robust Fisherian inferences can be to the choice of randomization mechanism and test statistic, we modify our call to `randinf` to use a binomial randomization mechanism, where every unit in the ad-hoc window  $[-2.5, 2.5]$  has a  $1/2$  probability of being assigned to treatment. For this, we must first create an auxiliary variable that contains the treatment assignment probability of every unit in the window; this auxiliary variable is then passed as an argument to `rdrandinf`.

```
> bern_prob = numeric(length(X))
> bern_prob[abs(X) > 2.5] = NA
> bern_prob[abs(X) <= 2.5] = 1/2
> out = rdrandinf(Y, X, wl = -2.5, wr = 2.5, seed = 50, bernoulli = bern_
  prob)
```

```
Selected window = [-2.5;2.5]
```

```
Running randomization-based test...
```

```
Randomization-based test complete.
```

```
Number of obs =      130
Order of poly =       0
Kernel type   =    uniform
Reps         =     1000
```



Window	=	set by user		
H0:	tau =	0		
Randomization	=	Bernoulli		
Cutoff c = 0		Left of c	Right of c	
Number of obs		68	62	
Eff. number of obs		68	62	
Mean of outcome		13.972	15.044	
S.d. of outcome		8.541	9.519	
Window		-2.5	2.5	
			Finite sample	Large sample
Statistic	T		P> T	Power vs d =
4.27				
Diff. in means	1.072	0.469	0.501	0.765

Analogous Stata command

```
. gen bern_prob = 1/2 if abs(X) <= 2.5
. rdrandinf Y X, wl(-2.5) wr(2.5) seed(50) bernoulli(bern_prob)
```

The last row of the top panel now says `Randomization = Bernoulli`, indicating that the Fisherian randomization-based test of the sharp null hypothesis is assuming a Bernoulli treatment assignment mechanism, where each unit has probability  $q$  of being placed above the cutoff—in this case, given our construction of the `bern_prob` variable,  $q = 1/2$  for all units. The Fisherian finite-sample p-value is now 0.469, very similar to the 0.488 p-value obtained above under the assumption of a fixed margins randomization. The conclusion of failure to reject  $H_0^F$  is therefore unchanged. This robustness of the Fisherian p-value to the choice of fixed margins versus Bernoulli randomization is typical in applications. Note also that the large-sample results are exactly the same as before—this is expected, since the choice of randomization mechanism does not affect the large-sample Neyman inferences.

We can also change the test statics used to test the Fisherian sharp null hypothesis. For example, to use the Kolmogorov-Smirnov (KS) test statistic instead of the difference-in-means, we set the option `statistic = "ksmirnov"`.

```
> out = rdrandinf(Y, X, wl = -2.5, wr = 2.5, seed = 50, statistic = "
ksmirnov")

Selected window = [-2.5;2.5]

Running randomization-based test...

Randomization-based test complete.
```

```

Number of obs =      2629
Order of poly  =       0
Kernel type   =    uniform
Reps          =     1000
Window        =    set by user
H0:          tau =       0
Randomization =    fixed margins

```

Cutoff c = 0	Left of c	Right of c
Number of obs	2314	315
Eff. number of obs	68	62
Mean of outcome	13.972	15.044
S.d. of outcome	8.541	9.519
Window	-2.5	2.5

		Finite sample	Large sample	
Statistic	T	P> T	P> T	Power vs d =
4.27				
Kolmogorov-Smirnov	0.101	0.846	0.898	NA

Analogous Stata command

```
. rdrandinf Y X, wl(-2.5) wr(2.5) seed(50) statistic(ksmirnov)
```

The bottom panel now reports the value of the KS statistic in the chosen window, which is 0.101. The randomization-based test of the Fisherian sharp null hypothesis  $H^F$  based on this statistic has p-value 0.846, considerably larger than the 0.488 p-value found in the same window (and with the same fixed-margins randomization) when the difference-in-means was chosen instead. Note that now the large-sample results report a large-sample approximation to the KS test p-value, and *not* a test of the Neyman null hypothesis  $H^N$ . Moreover, the KS statistic has no interpretation as a treatment effect in either case.

Finally, we illustrate how to obtain confidence intervals in our call to `rdrandinf`. Remember that in the Fisherian framework, confidence intervals are obtained by inverting tests of sharp null hypothesis. To implement this inversion, we must specify a grid of  $\tau$  values; `rdrandinf` will then test the null hypotheses  $H_{\tau_0}^F : Y_i(1) - Y_i(0) = \tau_0$  for all values of  $\tau_0$  in the grid, and collect in the confidence interval all the hypotheses that fail to be rejected in a randomization-based test of the desired level (default is level  $\alpha = 0.05$ ). To calculate these confidence intervals, we create the grid, and then call `rdrandinf` with the `ci` option. For this example, we choose a grid of values for  $\tau_0$  between  $-10$  and  $10$ , with  $0.25$  increments. Thus, we test  $H_{\tau_0}$  for all  $\tau_0 \in G^{\tau_0} = \{-10, -9.75, -9.50, \dots, 9.50, 9.75, 10\}$ .

```
> ci_vec = c(0.05, seq(from = -10, to = 10, by = 0.25))
> out = rdrandinf(Y, X, wl = -2.5, wr = 2.5, seed = 50, reps = 1000,
+ ci = ci_vec)
```

```
Selected window = [-2.5;2.5]
```

```
Running randomization-based test...
```

```
Randomization-based test complete.
```

```
Running sensitivity analysis...
```

```
Sensitivity analysis complete.
```

```
Number of obs =      2629
Order of poly =      0
Kernel type    =      uniform
Reps           =      1000
Window        =      set by user
H0:          tau =      0
Randomization =      fixed margins
```

Cutoff c = 0	Left of c	Right of c
Number of obs	2314	315
Eff. number of obs	68	62
Mean of outcome	13.972	15.044
S.d. of outcome	8.541	9.519
Window	-2.5	2.5

		Finite sample	Large sample	
Statistic	T	P> T	P> T	Power vs d =
4.27				
Diff. in means	1.072	0.488	0.501	0.765

```
95% confidence interval: [-2,4]
```

#### Analogous Stata command

```
. rdrandinf Y X, wl(-2.5) wr(2.5) seed(50) ci(0.05 -10(0.25)10)
```

The Fisherian 95% confidence interval is  $[-2, 4]$ . As explained, this confidence interval assumes a constant treatment effect model. The interpretation is therefore that, given the assumed randomization mechanism, all values of  $\tau$  between -2 and 4 in the constant treatment effect model  $Y_i(1) = Y_i(0) + \tau$  fail to be rejected with a randomization-based 5%-level test. In other words, in this window, and given a constant treatment effect model, the empirical evidence based on a local randomization RD framework is consistent with both negative and

positive true effects of Islamic victory on the female education share.

### 2.3 How to Choose the Window

In the previous sections, we assumed that  $W_0$  was known. However, in practice, even when a researcher is willing to assume that there *exists* a window around the cutoff where the treatment is as-if randomly assigned, the location of this window will be typically unknown. This is another fundamental difference between local randomization RD designs and actual randomized controlled experiments, since in the latter there is no ambiguity about the population of units that were subject to the random assignment of the treatment. Thus, the most important step in the implementation of the local randomization RD approach is to select the window around the cutoff where the treatment can be plausibly assumed to have been as-if randomly assigned.

One option is to choose the randomization window in an *ad-hoc* way, selecting a small neighborhood around the cutoff where the researcher is comfortable assuming local randomization. For example, a scholar may believe that elections decided by 0.5 percentage points or less are essentially decided as if by the flip of a coin, and chose the window  $[\bar{x} - 0.5, \bar{x} + 0.5]$ . The obvious disadvantage of selecting the window arbitrarily is that the resulting choice is based neither on empirical evidence nor on a systematic procedure, and thus lacks objectivity and replicability.

A preferred alternative is to choose the window using the information provided by relevant predetermined covariates—variables that reflect important characteristics of the units, and whose values are determined before the treatment is assigned and received. This approach requires assuming that there exists at least one important predetermined covariate of interest,  $Z$ , that is related to the running variable everywhere except inside the window  $W_0$ .

Figure 2.4 shows a hypothetical illustration, where the conditional expectation of  $Z$  given the score,  $\mathbb{E}(Z|X = x)$  is plotted against  $X$ . (We focus on the conditional expectation of the covariate for illustration purposes only; the approach applies more generally to any case where the distribution of the covariate is a function of the score.) Outside of  $W_0$ ,  $\mathbb{E}(Z|X)$  and  $X$  are related: a mild U-shaped relationship to the left of  $\bar{x}$ , and monotonically increasing to the right—possibly due to correlation between the score and another characteristic that also affects  $Z$ . However, inside the window  $W_0$  where local randomization holds, this relationship disappears by virtue of applying conditions LR1 and LR2 to  $Z$ , taking  $Z$  as an “outcome” variable. Moreover, because  $Z$  is a predetermined covariate, the effect of the treatment on  $Z$  is zero by construction. In combination, these assumptions imply that there is no association

between  $\mathbb{E}(Z|X)$  and  $X$  inside  $W_0$ , but these two variables are associated outside of  $W_0$ .

This suggests a data-driven method to choose  $W_0$ . We define a null hypothesis  $H_0$  stating that the treatment is unrelated to  $Z$  (or that  $Z$  is “balanced” between the groups). In theory, this hypothesis could be the Fisherian hypothesis  $H_0^F$  or the Neyman hypothesis  $H_0^N$ . However, since the procedure will typically involve some windows with very few observations, we recommend using randomization-based tests of the Fisherian hypothesis  $H_0^F$ , which takes the form  $H_0^F : Z_i(1) = Z_i(0)$ . Naturally, the effect of the treatment on  $Z$  is zero for all units inside  $W_0$  because the covariate is predetermined. However, the window selection procedure is based on the assumption that, outside  $W_0$ , the treatment and control groups *differ* systematically in  $Z$ —not because the treatment has a causal effect on  $Z$  but rather because the running variable is correlated with  $Z$  outside  $W_0$ . This assumption is important; without it, the window selector will not recover the true  $W_0$ .

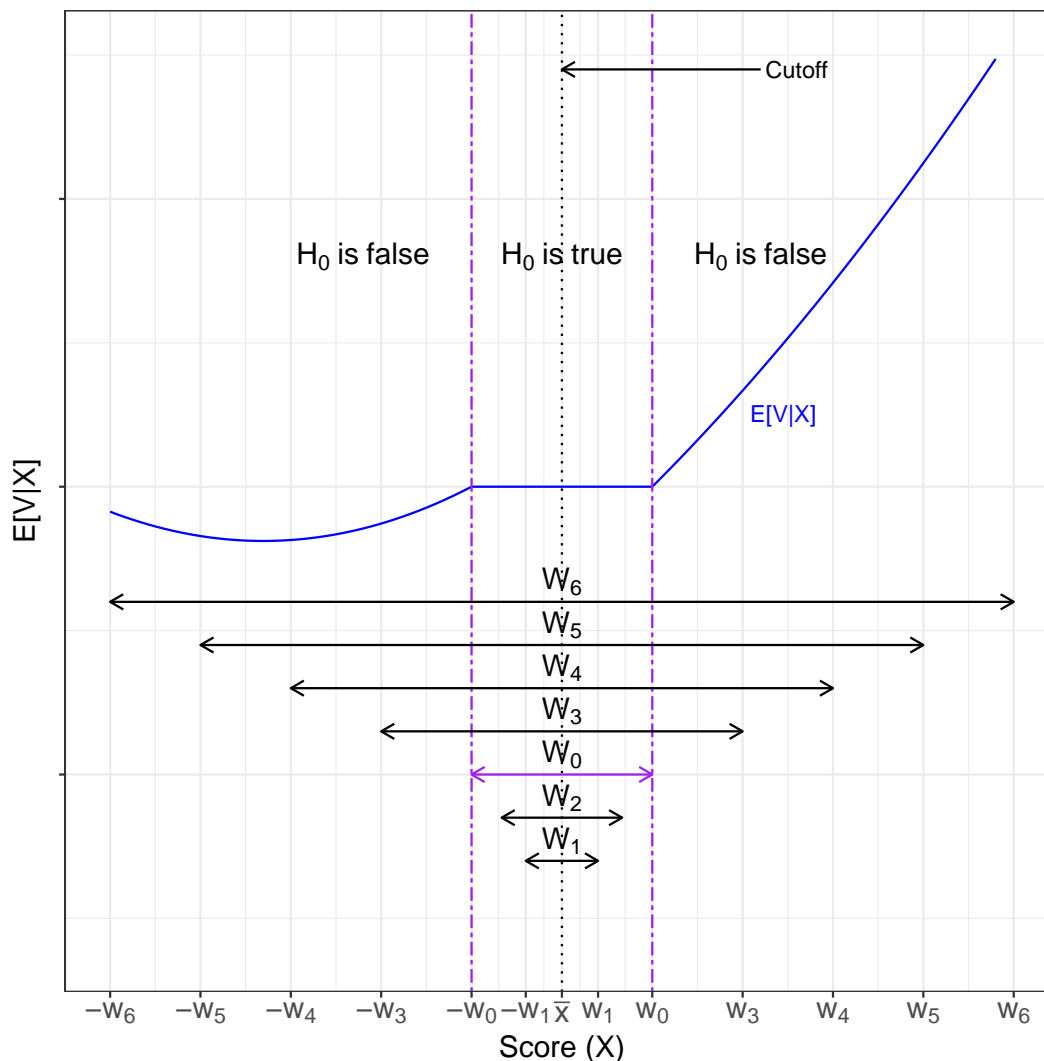
The procedure starts with the smallest possible window— $W_1$  in Figure 2.4—and tests the null hypothesis of no effect  $H_0$ . Since there is no relationship between  $\mathbb{E}(Z|X)$  and  $Z$  inside  $W_1$ ,  $H_0$  will fail to be rejected. Once  $H_0$  fails to be rejected, a smaller window  $W_2$  is selected, and the null hypothesis is tested again inside  $W_2$ . The procedure keeps increasing the length of the window and re-testing  $H_0$  in each larger window, until a window is reached where  $H_0$  is rejected at the chosen significance level  $\alpha^* \in (0, 1)$ . In the figure, assuming the test has perfect power, the null hypothesis will not be rejected in  $W_0$ , nor will it be rejected in  $W_2$  or  $W_1$ . The chosen window is the largest window such that  $H_0$  fails to be rejected inside that window, and in all windows contained in it. In Figure 2.4, the chosen window is  $W_0$ .

In spirit, this nested procedure is analogous to the use of covariate balance tests in randomized controlled experiments. In essence, the procedure chooses the largest window such that covariate balance holds in that window and all smaller windows inside it. As we show in our empirical illustration, this data-driven window selection method can be implemented with several covariates, for example, rejecting a particular window choice when  $H_0$  is rejected for at least one covariate.

As mentioned, we recommend choosing  $H_0^F$  as the null hypothesis. In addition, the practical implementation of the procedure requires several other choices:

- *Choose the relevant covariates.* Researchers must decide which covariates to use in the window selection procedure; these covariates should be related to both the outcome and the treatment assignment. If multiple covariates are chosen, the procedure can be applied using either the p-value of an omnibus test statistic, or by testing  $H_0$  for each

Figure 2.4: Window Selector Based on Covariate Balance in Locally Random RD



covariate separately and then making the decision to reject  $H_0$  based on the minimum p-value across all covariates—i.e., rejecting a particular window choice when  $H_0$  is rejected for at least one covariate.

- *Choose the test statistic.* Researchers must choose the statistic on which the randomization-based test of the Fisherian null hypothesis will be based. This can be the difference-in-means, one of the alternatives statistics discussed above, or other possibilities.
- *Choose the randomization mechanism.* Researchers must select the randomization mechanism that will be assumed inside the window to test the sharp null hypothesis  $H_0^F$  using Fisherian methods. In many applications, an appropriate choice is a complete randomization mechanism where every unit in the window has treatment assignment

probability  $1/\binom{n_{w_0}}{n_{w_0,t}}$ .

- *Choose a minimum number of observations in the smallest window.* In actual applications, if the smallest window around  $\bar{x}$  where the null hypothesis is tested is too small, it will contain too few observations and the test will not have enough power to reject the null hypothesis even if it is false. Thus, researchers must ensure that the smallest window considered contains a minimum number of observations to ensure acceptable power; we recommend that the minimum window have at least roughly ten observations on either side of the cutoff.
- *Choose  $\alpha^*$ .* The threshold significance level determines when a null hypothesis is considered rejected. Since the main concern is failing to reject a null hypothesis when it is false—in contrast to the usual concern about rejecting a true null hypothesis—the level of the test should be higher than conventional levels. When we test  $H_0$  at a higher level, we tolerate a higher probability of Type I error and a lower probability of concluding that the covariate is unrelated to the treatment assignment when in fact it is. We recommend setting  $\alpha^* \geq 0.15$  if possible, and ideally no smaller than 0.10.

Once researchers have selected the relevant covariates, the test statistic, the randomization mechanism in the window, and the threshold significance level  $\alpha^*$ , the window selection procedure can be implemented with the following algorithm:

1. Start with a symmetric window of length  $2w_j$ ,  $W_j = [\bar{x} - w_j, \bar{x} + w_j]$ .
2. For every covariate  $Z_k$ ,  $k = 1, 2, \dots, K$ , use the test statistic  $T_k$  and the chosen randomization mechanism to test  $H_0^F$  using a Fisherian framework. Use only observations with scores inside  $W_j$ . Compute the associated p-value,  $p_k$ . (If using an omnibus test, compute the single omnibus p-value.)
3. Compute the minimum p-value  $p_{\min} = \min(p_1, p_2, \dots, p_K)$ . (If using an omnibus test, set  $p_{\min}$  to the single omnibus p-value.)
  - (a) If  $p_{\min} > \alpha^*$ , do not reject the null hypothesis and increase the length of the window by  $2w_{\text{step}}$ ,  $W_{j+1} = [\bar{x} - w_j - w_{\text{step}}, \bar{x} + w_j + w_{\text{step}}]$ . Set  $W_j = W_{j+1}$  and go back to step (2).
  - (b) If  $p_{\min} \leq \alpha^*$ , reject the null hypothesis and conclude that the largest window where the local randomization assumption is plausible is  $W_j$ .

To see how the procedure works in practice, we use it to select a window in the Meyerson application using the set of predetermined covariates described above: `vshr_islam1994`, `partycount`, `lpop1994`, `i89`, `merkezi`, `merkezp`, `subbuyuk`, `buyuk`. We use the function `rdwinselect`, which is one of the functions in the `rdlocrand` library or package. The main arguments are the score variable  $X$ , the matrix of predetermined covariates, and the sequence of nested windows; for simplicity, only symmetric windows are considered. We also choose 1,000 simulations for the calculation of Fisherian p-values in each window.

There are two ways to increment the length of the windows in `rdwinselect`. One is to increment the length of the window in fixed steps, which can be implemented with the option `wstep`. For example, if the first window selected is  $[0.1, 0.1]$  and `wstep` = 0.1, the sequence is  $W_1 = [0.1, 0.1], W_2 = [0.2, 0.2], W_3 = [0.3, 0.3]$ , etc. The other is to increase the length of the window so that the number of observations increases by a minimum fixed amount on every step, which can be done via the option `wobs`. For example, by setting `wobs` = 2, every window in the sequence is the *smallest* symmetric window such that the number of added observations on each side of the cutoff relative to the prior window is at least 2. By default, `rdwinselect` starts with the smallest window that has at least 10 observations on either side, but this default behavior can be changed with the options `wmin` or `obsmin`. Finally, `rdwinselect` uses the chosen level  $\alpha^*$  to recommend the chosen window; the default is  $\alpha^* = 0.15$ , but this can be modified with the `level` option.

We start by considering a sequence of symmetric windows where we increase the length in every step by the minimum amount so that we add at least 2 observations on either side in each step. We achieve this by setting `wobs=2`.

```
> Z = cbind(data$i89, data$vshr_islam1994, data$partycount, data$lpop1994,
+ data$merkezi, data$merkezp, data$subbuyuk, data$buyuk)
> colnames(Z) = c("i89", "vshr_islam1994", "partycount", "lpop1994",
+ "merkezi", "merkezp", "subbuyuk", "buyuk")
> out = rdwinselect(X, Z, seed = 50, reps = 1000, wobs = 2)
```

Window selection for RD under local randomization

```
Number of obs =          2629
Order of poly  =           0
Kernel type    =      uniform
Reps           =         1000
Testing method =      rdrandinf
Balance test   =      diffmeans

Cutoff c = 0          Left of c          Right of c
Number of obs      2314                  315
1st percentile     24                    3
```



Window length / 2 Obs >= c	p-value	Var. name	Bin. test	Obs < c
5th percentile	115			15
10th percentile	231			31
20th percentile	463			62
0.446 10	0.451	i89	1	9
0.486 12	0.209	i89	1	11
0.536 13	0.253	i89	1	12
0.699 17	0.238	i89	0.585	13
0.856 19	0.27	i89	0.377	13
0.944 21	0.241	i89	0.627	17
1.116 25	0.048	i89	0.28	17
1.274 26	0.059	i89	0.371	19
1.343 28	0.352	merkezi	0.312	20
1.42 31	0.365	i89	0.272	22

Recommended window is [-0.944;0.944] with 38 observations (17 below, 21 above).

#### Analogous Stata command

```
. global covs "i89 vshr_islam1994 partycount lpop1994 merkezi merkezp subbuyuk buyuk"
. rdwinselect X $covs, seed(50) wobs(2)
```

The top and middle panels in the `rdwinselect` output are very similar to the corresponding panels in the `rdrandinf` output. One difference is the **Testing method**, which indicates whether randomization-based methods are used to test  $H_0^F$ , or Normal approximations methods are used to test  $H_0^N$ . The default is randomization-based methods, but this can be changed with the `approximate` option. The other difference in the output is the **Balance test**, which indicates the type of test statistic used for testing the null hypothesis—the default is `diffmeans`, the difference-in-means. The option `statistic` allows the user to select a different test statistic; the available options are the Kolmogorov-Smirnov statistic (`ksmirnov`), the Wilcoxon-Mann-Whitney studentized statistic (`ranksum`), and Hotelling’s T-squared statistic (`hotelling`).

The bottom panel shows tests of the null hypothesis for each window considered. By default, `rdwinselect` starts with the smallest symmetric window that has at least 10 observations on either side of the cutoff. Since we set `wobs=2`, we continue to consider the smallest possible (symmetric) windows so that at least 2 observations are added on each side of the cutoff in every step. For every window, the column `p-value` reports  $p_{\min}$ —the minimum of the p-values associated with the  $K$  tests of the null hypothesis of no effect performed for each of the  $K$  covariates, or the unique p-value if an omnibus test is used. The column `Var. name` reports the covariate associated with the minimum p-value—that is, the covariate  $Z_k$  such that  $p_k = p_{\min}$ .

Finally, the column `Bin. test` uses a Binomial test to calculate the probability of observing  $n_{W,+}$  successes out of  $n_W$  trials, where  $n_{W,t}$  is the number of observations within the window that are above the cutoff (reported in column `Obs>c`) and  $n_W$  is the total number of observations within the window (which can be calculated by adding the number reported in columns `Obs<c` and `Obs>c`). We postpone discussion of this test until the upcoming section on falsification of RD designs under a local randomization framework.

The output indicates that the p-values are above 0.20 in all windows between the minimum window  $[-0.446, 0.446]$  and  $[-0.944, 0.944]$ . In the window immediately after  $[-0.944, 0.944]$ , the p-value drops to 0.048, considerably below the suggested 0.15 threshold. The data-driven window is therefore  $W^* = [-0.944, 0.944]$ . After this window, the p-values start decreasing, albeit initially this decrease is not necessarily monotonic. By default, `rdwinselect` only shows the first 20 windows; in order to see the sharp decrease in p-values that occurs as larger windows are considered, we set the option `nwindows=50` to see the output of the first 50 windows. We also set the option `plot=TRUE`, to create a plot of the minimum p-values associated with the length of each window considered—the plot is shown in Figure 2.5 below.

```
> Z = cbind(data$i89, data$vshr_islam1994, data$partycount, data$lpop1994,
+ data$merkezi, data$merkezp, data$subbuyuk, data$buyuk)
> colnames(Z) = c("i89", "vshr_islam1994", "partycount", "lpop1994",
+ "merkezi", "merkezp", "subbuyuk", "buyuk")
> out = rdwinselect(X, Z, seed = 50, reps = 1000, wobs = 2, nwindows = 50,
+ plot = TRUE)
```

Window selection for RD under local randomization

```
Number of obs =      2629
Order of poly  =         0
Kernel type    =    uniform
Reps           =     1000
Testing method =    rdrandinf
Balance test   =    diffmeans
```

Cutoff $c = 0$	Left of $c$	Right of $c$
Number of obs	2314	315
1st percentile	24	3
5th percentile	115	15
10th percentile	231	31
20th percentile	463	62

Window length / 2 Obs $\geq c$	p-value	Var. name	Bin. test	Obs $< c$
0.446 10	0.451	i89	1	9
0.486 12	0.209	i89	1	11
0.536 13	0.253	i89	1	12
0.699 17	0.238	i89	0.585	13
0.856 19	0.27	i89	0.377	13
0.944 21	0.241	i89	0.627	17
1.116 25	0.048	i89	0.28	17
1.274 26	0.059	i89	0.371	19
1.343 28	0.352	merkezi	0.312	20
1.42 31	0.365	i89	0.272	22
1.49 33	0.171	merkezi	0.229	23
1.556 35	0.31	merkezi	0.245	25
1.641 37	0.188	merkezi	0.26	27
1.858 38	0.206	merkezi	0.328	29
1.914 39	0.244	merkezi	0.403	31
2.019 41	0.26	merkezi	0.416	33
2.097 42	0.211	vshr_islam1994	0.653	37
2.158 43	0.096	vshr_islam1994	0.657	38
2.319 45	0.048	vshr_islam1994	1	44
2.433 47	0.016	vshr_islam1994	0.839	50
2.583 49	0.008	vshr_islam1994	0.842	52
2.643	0.006	vshr_islam1994	0.922	53

51				
2.746	0.002	vshr_islam1994	1	54
54				
3.009	0.002	vshr_islam1994	1	56
56				
3.051	0.005	vshr_islam1994	1	58
58				
3.094	0.001	vshr_islam1994	0.928	62
60				
3.178	0	vshr_islam1994	0.929	64
62				
3.462	0	vshr_islam1994	0.491	72
63				
3.595	0	vshr_islam1994	0.444	74
64				
3.704	0	vshr_islam1994	0.356	77
65				
3.821	0	vshr_islam1994	0.251	82
67				
3.963	0	vshr_islam1994	0.258	84
69				
4.181	0	vshr_islam1994	0.153	89
70				
4.287	0	vshr_islam1994	0.138	92
72				
4.417	0	vshr_islam1994	0.192	94
76				
4.488	0.001	vshr_islam1994	0.168	95
76				
4.585	0	vshr_islam1994	0.113	99
77				
4.719	0	vshr_islam1994	0.1	101
78				
4.899	0	vshr_islam1994	0.067	106
80				
5.03	0	vshr_islam1994	0.095	107
83				
5.2	0	vshr_islam1994	0.098	109
85				
5.346	0	vshr_islam1994	0.089	112
87				
5.421	0	vshr_islam1994	0.078	114
88				
5.482	0	vshr_islam1994	0.092	114
89				
5.593	0	vshr_islam1994	0.081	116
90				
5.676	0	vshr_islam1994	0.073	119
92				
5.779	0	vshr_islam1994	0.075	120
93				
5.878	0	vshr_islam1994	0.119	121
97				
6.039	0	vshr_islam1994	0.053	128



than  $[-3, 3]$ , suggesting that there are sharp differences between municipalities where the Islamic party wins and municipalities where it loses, even when the election is decided by moderate margins. This shows that, the local randomization, if it holds at all, will hold in a small window near the cutoff.

If we want to choose the window using a sequence of symmetric windows of fixed length rather than controlling the minimum number of observations, we use the `wstep` option. Calling `rdwinselect` with `wstep=0.1` performs the covariate balance tests in a sequence of windows that starts at the minimum window and increases the length by 0.1 at each side of the cutoff.

```
> Z = cbind(data$i89, data$vshr_islam1994, data$partycount, data$lpop1994,
+ data$merkezi, data$merkezp, data$subbuyuk, data$buyuk)
> colnames(Z) = c("i89", "vshr_islam1994", "partycount", "lpop1994",
+ "merkezi", "merkezp", "subbuyuk", "buyuk")
> out = rdwinselect(X, Z, seed = 50, reps = 1000, wstep = 0.1,
+ nwindows = 25)
```

Window selection for RD under local randomization

```
Number of obs =          2629
Order of poly  =           0
Kernel type    =    uniform
Reps           =         1000
Testing method =    rdrandinf
Balance test   =    diffmeans
```

Cutoff c = 0	Left of c	Right of c
Number of obs	2314	315
1st percentile	24	3
5th percentile	115	15
10th percentile	231	31
20th percentile	463	62

Window length / 2 Obs >= c	p-value	Var. name	Bin.test	Obs < c
0.446	0.451	i89	1	9
10				
0.546	0.241	i89	1	12
13				
0.646	0.22	i89	0.851	13
15				
0.746	0.241	i89	0.473	13
18				
0.846	0.243	i89	0.377	13
19				
0.946	0.234	i89	0.627	17
21				

1.046	0.061	i89	0.349	17
24				
1.146	0.069	i89	0.28	17
25				
1.246	0.057	i89	0.451	19
25				
1.346	0.366	merkezi	0.312	20
28				
1.446	0.254	merkezi	0.22	22
32				
1.546	0.24	i89	0.298	25
34				
1.646	0.187	merkezi	0.26	27
37				
1.746	0.198	merkezi	0.215	27
38				
1.846	0.224	i89	0.268	28
38				
1.946	0.242	merkezi	0.477	32
39				
2.046	0.259	merkezi	0.567	35
41				
2.146	0.095	vshr_islam1994	0.657	38
43				
2.246	0.066	vshr_islam1994	0.914	42
44				
2.346	0.056	vshr_islam1994	1	45
45				
2.446	0.012	vshr_islam1994	0.839	50
47				
2.546	0.011	vshr_islam1994	0.841	51
48				
2.646	0.006	vshr_islam1994	0.922	53
51				
2.746	0	vshr_islam1994	1	53
54				
2.846	0.003	vshr_islam1994	1	55
55				

Recommended window is [-0.946;0.946] with 38 observations (17 below, 21 above).

#### Analogous Stata command

```
. global covs "i89 vshr_islam1994 partycount lpop1994 merkezi merkezp subbuyuk buyuk"
. rdwinselect X $covs, seed(50) wstep(0.1) nwindows(25)
```

The suggested window is  $[-0.946, 0.946]$ , very similar to the  $[-0.944, 0.944]$  window chosen above with the `wobs=2` option.

We can now use `rdrandinf` to perform a local randomization analysis in the chosen

window. For this, we use the options `wl` and `wr` to input, respectively, the lower and upper limit of the chosen window  $W^* = [-0.944, 0.944]$ . We also use the option `d = 3.020` to calculate the power of a Neyman test to reject the null hypothesis of null average treatment effect when the true average difference is 3.020. This value is the continuity-based linear polynomial point estimate reproduced above (and discussed extensively in Part I).

The difference-in-means in the chosen window  $W^* = [-0.944, 0.944]$  is 2.638, considerably similar to the continuity-based local linear point estimate of 3.020. However, using a Neyman approach, we cannot distinguish this average difference from zero, with a p-value of 0.333. Similarly, we fail to reject the Fisherian sharp null hypothesis that an electoral victory by the Islamic party has no effect on the female education share for any municipality (p-value 0.386). As shown in the last column, the large-sample power to detect a difference of around 3 percentage points is only 19.4%. Naturally, the small number of observations in the chosen window (23 and 27 below and above the cutoff, respectively) limits statistical power. In addition, the effect of 2.638, although much larger than the 1.072 estimated in the ad-hoc  $[-2.5, 2.5]$  window, is still a small effect, as it is less than a third of one standard deviation of the female education share in the control group—we see this by calculating  $2.638/8.615 = 0.306$ . In accordance with these results, the Fisherian 95% confidence interval under a constant treatment effect model is  $[-2.8, 8.5]$ , consistent with both positive and negative effects.

```
> ci_vec = c(0.05, seq(from = -10, to = 10, by = 0.1))
> out = rdrandinf(Y, X, wl = -0.944, wr = 0.944, seed = 50, reps = 1000,
+ ci = ci_vec)
```

```
Selected window = [-0.944;0.944]
```

```
Running randomization-based test...
```

```
Randomization-based test complete.
```

```
Running sensitivity analysis...
```

```
Sensitivity analysis complete.
```

```
Number of obs =      2629
Order of poly  =         0
Kernel type    =    uniform
Reps           =     1000
Window         =    set by user
H0:           tau =         0
Randomization =    fixed margins
```

```
Cutoff c = 0           Left of c           Right of c
Number of obs         2314                   315
```



Eff. number of obs	23	27		
Mean of outcome	13.579	16.217		
S.d. of outcome	8.615	10.641		
Window	-0.944	0.944		
		Finite sample	Large sample	
Statistic	T	P> T	P> T	Power vs d =
4.307				
Diff. in means	2.638	0.386	0.333	0.353
95% confidence interval: [-2.8,8.5]				

## Analogous Stata command

```
. rdrandinf Y X, wl(-0.944) wr(0.944) seed(50) ci(0.05 -10(0.1)10)
```

Finally, we mention that instead of calling `rdwinselect` first and `rdrandinf` second, we can choose the window and perform inference in one step by using the `covariates` option in `rdrandinf`.

```
> Z = cbind(data$i89, data$vshr_islam1994, data$partycount, data$lpop1994,
+ data$merkezi, data$merkezp, data$subbuyuk, data$buyuk)
> colnames(Z) = c("i89", "vshr_islam1994", "partycount", "lpop1994",
+ "merkezi", "merkezp", "subbuyuk", "buyuk")
> out = rdrandinf(Y, X, covariates = Z, seed = 50, d = 3.019522)
```

```
Running rdwinselect...
```

```
rdwinselect complete.
```

```
Selected window = [-0.917068123817444;0.917068123817444]
```

```
Running randomization-based test...
```

```
Randomization-based test complete.
```

```
Number of obs =      2629
Order of poly  =        0
Kernel type    =    uniform
Reps           =     1000
Window        =    rdwinselect
H0:           tau =        0
Randomization =    fixed margins
```

```
Cutoff c = 0          Left of c          Right of c
Number of obs      2314          315
Eff. number of obs  22           27
```

Mean of outcome	13.266	16.217		
S.d. of outcome	8.682	10.641		
Window	-0.917	0.917		
		Finite sample	Large sample	
Statistic	T	P> T	P> T	Power vs d =
3.02				
Diff. in means	2.952	0.301	0.285	0.194

#### Analogous Stata command

```
. global covs "i89 vshr_islam1994 partycount lpop1994 merkezi merkezp subbuyuk buyuk"
. rdrandinf Y X, covariates($covs) seed(50) d(3.019522)
```

However, it is usually better to first choose the window using `rdwinselect` and then use `rdrandinf`. The reason is that calling `rdwinselect` by itself will never show outcome results, and will reduce the possibility of choosing the window where the outcome results are in the “expected” direction—in other words, choosing the window without looking at the outcome results minimizes pre-testing and specification-searching issues.

## 2.4 Falsification Analysis In The Local Randomization Approach

In Part I, we discussed the importance of conducting falsification tests to provide evidence in support of the RD assumptions. Falsification and validation analyses are as important in the local randomization framework as they are in the continuity-based framework. The difference resides in their implementation. Instead of providing empirical evidence in favor of continuity assumptions as in the continuity-based approach, the main goal in a local randomization approach is to provide evidence consistent with the local randomization assumption. Thus, the methods employed to perform a falsification analysis in the local randomization approach must be different.

We now discuss four types of empirical falsification tests for a local randomization RD design, all of which we discussed in our companion Part I monograph in the context of the continuity-based approach: (i) tests of a null treatment effect on pre-treatment covariates or placebo outcomes, (ii) tests to assess the density of the score around the cutoff, (iii) treatment effect estimation at artificial cutoffs values, and (v) sensitivity to neighborhood choices. We now discuss and illustrate how to implement these tests in the local randomization approach. In Part I, we also discussed an additional falsification test based on an analysis that excludes the observations that are closest to the cutoff, an idea sometimes referred to as the “donut

hole” test. This test is most natural in a continuity-based approach, where one is interested in knowing whether the extrapolation is heavily influenced by the few observations closest to the cutoff and the analysis typically contains at least hundreds of observations. In contrast, in a local randomization approach, the chosen window is likely to be very small, and removing the observations closest to the cutoff may result in too few observations and render the analysis unfeasible. For this reason, we omit this method from our discussion.

### 2.4.1 Predetermined Covariates and Placebo Outcomes

This crucial falsification test focuses on two types of variables: predetermined covariates—variables that are determined before the treatment is assigned, and placebo outcomes—variables that are determined after the treatment is assigned but for which we have scientific reasons to believe that they are unaffected by the treatment. The idea is that, in a valid RD design, there should be no systematic differences between treated and control groups at the cutoff in terms of both placebo outcomes and predetermined covariates, because these variables could not have been affected by the treatment. For implementation, the researcher conducts a test of the hypothesis that the treatment effect is zero for each predetermined covariate and placebo outcome. If the treatment does have an effect on these variables, the plausibility of the RD assumptions is called into question.

An important principle behind this type of falsification analysis is that all predetermined covariates and placebo outcomes should be analyzed in the same way as the outcome of interest. In the local randomization approach, this means that the null hypothesis of no treatment effect should be tested within the window where the assumption of local randomization is assumed to hold, using the same inference procedures and the same treatment assignment mechanism and test statistic used for the analysis of the outcome of interest. Since in this approach  $W_0$  is the window where the treatment is assumed to have been randomly assigned, all covariates and placebo outcomes should be analyzed within this window. This illustrates a fundamental difference between the continuity-based and the randomization-based approach: in the former, estimation and inference requires approximating unknown regression functions, which requires estimating separate bandwidths for each variable analyzed; in the latter, since the treatment is assumed to be as-if-randomly assigned in  $W_0$ , all analysis occur within the same window,  $W_0$ .

In our analysis above we chose the window  $W_0 = [-0.944, 0.944]$  for the local-randomization approach. In order to test if the predetermined covariates are balanced within this window, we analyze their behavior in this window on each side of the cutoff using randomization

inference techniques. We use the difference-in-means statistic, the same statistic we used for the outcome of interest. Under the local randomization assumption, we expect the difference-in-means between treated and control groups for each covariate to be indistinguishable from zero within  $W_0$ . Naturally, we already know that the covariates that we used to choose the window are balanced in the chosen window. In this sense, the window selector procedure is itself a validation procedure. We note, however, that it is possible (and indeed common) for researchers to choose the window based on a given set of covariates, and then assess balance on a different set.

In order to test this formally, we use the `rdrandinf` function, using each covariate as the outcome of interest. For example, when we study the covariate `vshr_islam1994`, we see that the difference-in-means statistic is very small ( $0.328 - 0.319 = -0.009$ ), and the finite sample p-value is large (0.689). That is, this covariate is balanced inside  $W_0$  according to this test-statistic.

```
> out = rdrandinf(data$vshr_islam1994, X, seed = 50, wl = -0.944,
+ wr = 0.944)

Selected window = [-0.944;0.944]

Running randomization-based test...

Randomization-based test complete.
```

Number of obs =	2629			
Order of poly =	0			
Kernel type =	uniform			
Reps =	1000			
Window =	set by user			
H0: tau =	0			
Randomization =	fixed margins			
Cutoff c = 0	Left of c	Right of c		
Number of obs	2314	315		
Eff. number of obs	23	27		
Mean of outcome	32.816	31.872		
S.d. of outcome	9.41	8.141		
Window	-0.944	0.944		
			Finite sample	Large sample
Statistic	T	P> T	P> T	Power vs d =
4.705				
Diff. in means	-0.944	0.702	0.707	0.466

## Analogous Stata command

```
. rdrandinf vshr_islam1994 X, wl(-.944) wr(.944)
```

Table 2.3: Formal Local-Randomization Analysis for Covariates

Variable	Mean of Controls	Mean of Treated	Diff-in-Means Statistic	Fisherian p-value	Number of Observations
Percentage of men aged 15-20 with high school education	19.219	21.742	2.523	0.275	50
Islamic Mayor in 1989	0.000	0.150	0.150	0.222	37
Islamic percentage of votes in 1994	32.816	31.872	-0.944	0.712	50
Number of parties receiving votes 1994	6.348	6.074	-0.274	0.794	50
Log population in 1994	8.566	8.594	0.028	0.954	50
District center	0.609	0.519	-0.090	0.595	50
Province center	0.087	0.037	-0.050	0.580	50
Sub-metro center	0.087	0.074	-0.013	1.000	50
Metro center	0.043	0.037	-0.006	1.000	50

Table 2.3 contains a summary of this analysis for all covariates using randomization inference. We cannot conclude that the control and treatment means are different for any covariate, since the p-values are above 0.15 in all cases. There is no statistical evidence of imbalance (in terms of their means) inside of this window. Note that the number of observations is fixed in all cases. This happens because the window in which we analyze these covariates is not changing, is always  $W_0$ , whereas in the continuity-based approach the MSE-optimal bandwidth depends on the particular outcome being used, and therefore the number of observations is different for every covariate.

This analysis can also be carried out visually, by using `rdplot` restricted to  $W_0$ , with  $p = 0$  and a uniform kernel. If the window was chosen appropriately, then all covariates should have similar means above and below the cutoff. For example, we can construct an RD plot with these characteristics for the covariate `vshr_islam1994`.

```
> rdplot(data$vshr_islam1994[abs(X) <= 0.944], X[abs(X) <= 0.944],
+ p = 0, kernel = "uniform", x.label = "Score", y.label = "",
+ title = "", x.lim = c(-1.25, 1.25), y.lim = c(0.2, 0.45))
```

## Analogous Stata command

```
. rdplot vshr_islam1994 X if abs(X) <= .944, h(.944) p(0) kernel(uniform)
```

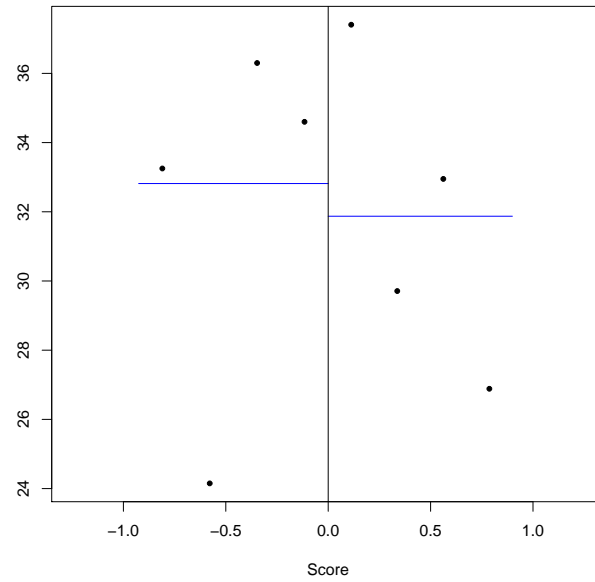
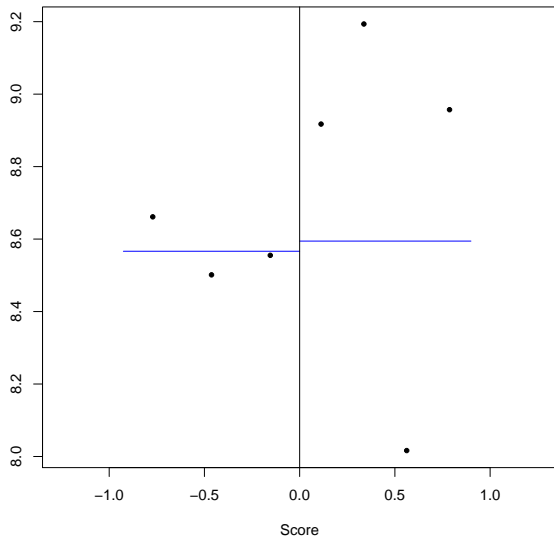
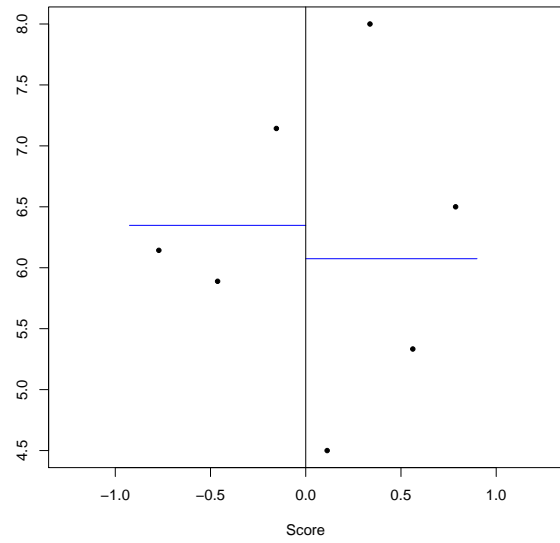


Figure 2.6 contains analogous RD plots for the six predetermined covariates analyzed above. In most cases, the visual inspection shows that the means of these covariates seem to be similar on each side of the cutoff, consistent with the results from the window selection procedure discussed in Section 2 and the formal analysis in Table 2.3.

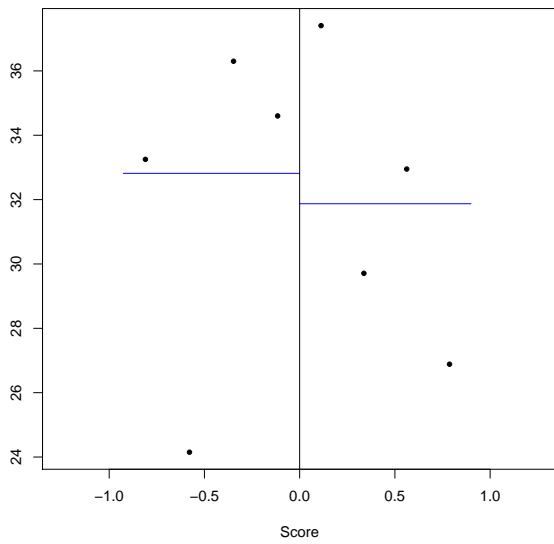
Figure 2.6: Illustration of Local Randomization RD Effects on Covariates—Meyersson Data



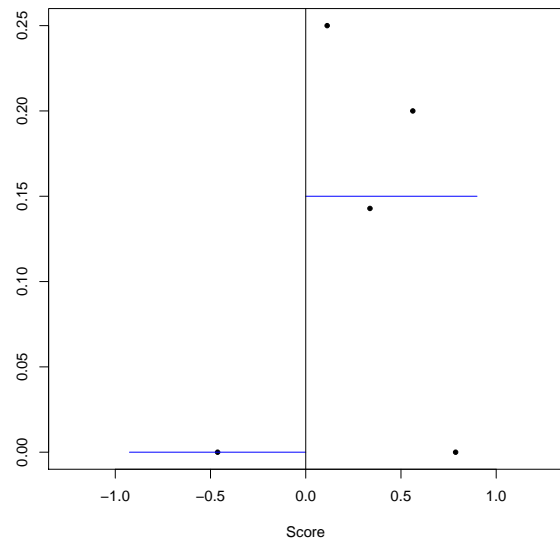
(a) Log Population in 1994



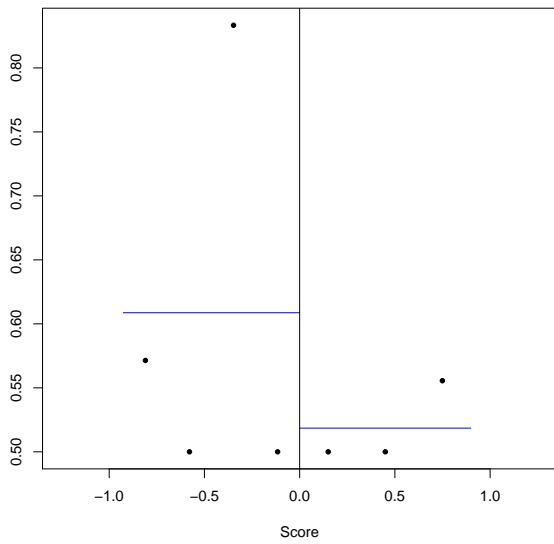
(b) Number of Parties Receiving Votes in 1994



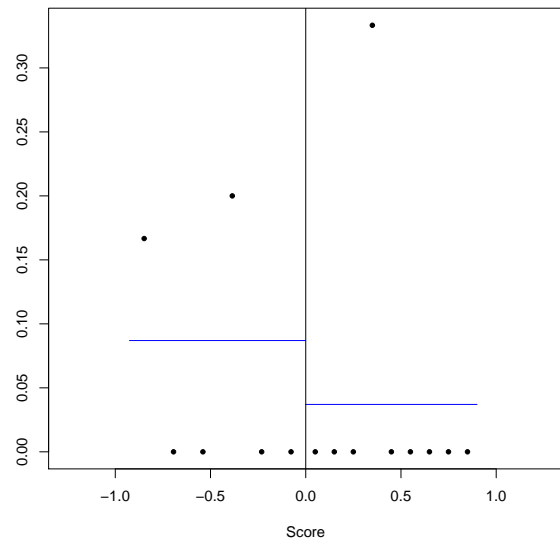
(c) Islamic Vote Share in 1994



(d) Islamic Mayor in 1989



(e) District center indicator



(f) Province center indicator

### 2.4.2 Density of Running Variable

Another important falsification test, already discussed in Part I, is a density test that analyzes whether the number of observations just above the cutoff is roughly similar to the number of observations just below the cutoff. The idea is that, if units lack the ability to control precisely the value of the score they receive, they should be as likely to receive a score value just above the cutoff as they are to receive a score value just below it. In a local randomization approach, this falsification analysis is implemented by testing the null hypothesis that, within the window  $W_0$  where the treatment is assumed to be randomly assigned, the number of treated and control observations is consistent with whatever assignment mechanism is assumed inside  $W_0$ .

For example, assuming a simple “coin flip” or Bernoulli trial with probability of success  $q$ , we would expect that the control sample size,  $n_{W_0,-}$ , and treatment sample size,  $n_{W_0,+}$ , within  $W_0$  to be compatible with the numbers generated by these  $n_{W_0,-} + n_{W_0,+} = n_{W_0}$  Bernoulli trials. In this case, the number of treated units in  $W_0$  follows a binomial distribution, and the null hypothesis of the test is that the probability of success in the  $n_{W_0}$  Bernoulli experiments is  $q$ . As discussed, the true probability of treatment is unknown. In practice, researchers can choose  $q = 1/2$  (a choice that can be justified from a large sample perspective when the score is continuous), or can set the value of  $q$  equal to the observed proportion of treated units in the window.

The binomial test is implemented in all common statistical software, and is also part of the `rdlocrand` package via the `rdwinselect` command. Using the Meyersson data, we can implement this falsification test in our selected window  $W_0 = [-0.944, 0.944]$  employing `rdwinselect` using the score as the single argument. Since we only want to see the binomial test in this window, we use the option `nwindows(1)`.

```
> out = rdwinselect(X, wmin = 0.944, nwindows = 1)
```

```
Window selection for RD under local randomization
```

```
Number of obs =          2629
Order of poly =           0
Kernel type   =      uniform
Reps          =          1000
Testing method =      rdrandinf
Balance test  =      diffmeans
```

```
Cutoff c = 0           Left of c           Right of c
Number of obs         2314                   315
1st percentile        24                     3
```



5th percentile	115		15	
10th percentile	231		31	
20th percentile	463		62	
Window length / 2 Obs >= c	p-value	Var. name	Bin. test	Obs < c
0.944 27	NA	NA	0.672	23

## Analogous Stata command

```
. rdwinselect X, wmin(0.944) nwindows(1)
```

There are 23 control observations and 27 treated observations in the window. The column `Bin. test` shows the p-value of a binomial test that uses a success probability equal to  $1/2$ . The p-value is 0.672, so we find no evidence of “sorting” around the cutoff in the window  $W_0 = [-0.944, 0.944]$ —the difference in the number of treated and control observations in this window is entirely consistent with what would be expected if municipalities were assigned to an Islamic win or loss by the flip of an unbiased coin. This is expected, as the observed share of treated observations in the window,  $\frac{27}{23+27} = 0.54$ , is very close to  $1/2$ .

We can also implement the binomial test using  $q = 1/2$ . For this, we can simply use the base distribution in R or Stata.

```
> binom.test(27, 50, 1/2)

Exact binomial test

data: 27 and 50
number of successes = 27, number of trials = 50, p-value = 0.6718
alternative hypothesis: true probability of success is not equal to 0.5
95 percent confidence interval:
 0.3932420 0.6818508
sample estimates:
probability of success
          0.54
```

## Analogous Stata command

```
. bitesti 50 27 1/2
```

As expected, the two-sided p-value is 0.6718, which is equal (after rounding) to the p-value obtained using `rdwinselect`.

### 2.4.3 Placebo Cutoffs

This falsification test chooses one or more artificial or “fake” cutoff values at which the probability of treatment assignment does not change, and analyzes the outcome of interest at these artificial cutoffs using the same methods used to conduct the analysis at the actual cutoff. The expectation is that no effect should be found at any of the artificial cutoffs. To avoid contamination from the actual treatment effect, only treated observations are included for fake cutoffs above the actual cutoff, and only control observations are included for cutoffs below the actual cutoff.

In the local-randomization approach, one possible implementation is to choose several artificial cutoff values, and then conduct a randomization-based analysis of the outcome using a symmetric window of the same length as the original window  $W_0$  around each of the fake cutoffs. Since our chosen window in the Meyersson application is  $W_0 = [-0.944, 0.944]$ , we consider windows of length  $\pm 0.944$  around each artificial cutoff. For example, for the cutoff 2, we analyze the outcome in the window  $[1.056, 2.944]$ .

### 2.4.4 Sensitivity to Window Choice

Just like in a continuity-based approach researchers are interested in the sensitivity of the results to the bandwidth choice, in a local randomization approach we are often interested in sensitivity to the window choice. To assess sensitivity to window choice, researchers can simply consider windows smaller than  $W_0$  and repeat the randomization-based analysis for the outcome of interest as conducted in the original window—that is, using the same test-statistic, same randomization mechanism, etc. This analysis should be implemented carefully, however. If the window  $W_0$  was chosen based on covariate balance as we recommend, results in windows larger than  $W_0$  will not be reliable because in such windows the treated and control groups will be imbalanced in important covariates. Thus, the sensitivity analysis should only consider windows smaller than  $W_0$ ; unfortunately, in many applications this analysis will be limited by the small number observations that is likely to occur in these windows. In the Meyersson application, our chosen window is  $W_0 = [-0.944, 0.944]$ , and has 23 and 27 observations on either side of the cutoff. We consider the smaller windows  $W_0 = [-0.85, 0.85]$ ,  $W_0 = [-0.75, 0.75]$ , and  $W_0 = [-0.75, 0.75]$  below.

## 2.5 When To Use The Local Randomization Approach

Unlike an experiment, the RD treatment assignment rule does not imply that the treatment is randomly assigned within some window. Like the continuity assumption, the local randomization assumption must be made *in addition* to the RD assignment mechanism, and is not directly testable. But the local randomization assumption is strictly stronger than the continuity assumption, in the sense that if there is a window around  $\bar{x}$  in which the regression functions are constant functions of the score, these regression functions will also be continuous functions of the score at  $\bar{x}$ —but the converse is not true. Why, then, would researchers want to impose stronger assumptions to make their inferences?

It is useful to remember that the local-polynomial approach, although based on the weaker condition of continuity, relies on extrapolation because there are no observations exactly at the cutoff. The continuity assumption does not imply a specific functional form of the regression functions near the cutoff, as it approximates these functions using non-parametric methods; however, this approximation relies on extrapolation methods and introduces an approximation error that is only negligible if the sample size is large enough. This makes the continuity-based approach more appealing if there are enough observations near the cutoff to approximate the regression functions with reasonable accuracy—but possibly inadequate when the number of observations is small. In applications with few observations, the local randomization approach has the advantage of requiring minimal extrapolation and avoiding the use of smoothing methods.

Another situation in which a local randomization approach may be preferable to a continuity-based approach is when the running variable is discrete—i.e., when multiple units share the same value of the score, as the continuity-based approach is not directly applicable in this case. We consider this issue in the next section, where we discuss how to analyze RD designs with discrete running variables.

## 2.6 Further Readings

Textbook reviews of Fisherian and Neyman estimation and inference methods in the context of analysis of experiments are given by [Rosenbaum \(2002, 2010\)](#) and [Imbens and Rubin \(2015\)](#); the latter also discusses super-population approaches and their connections to finite population inference methods. [Ernst \(2004\)](#) discusses the connection and distinctions between randomization and permutation inference methods. [Cattaneo et al. \(2015\)](#) propose Fisherian randomization-based inference to analyze RD designs based on a local random-

ization assumption, and the window selection procedure based on covariate balance tests. [Cattaneo et al. \(2017\)](#) relax the local randomization assumption to allow for a weaker exclusion restriction, and also compare RD analysis in continuity-based and randomization-based approaches. The interpretation of the RD design as a local experiment and its connection to the continuity-based framework is also discussed by [Sekhon and Titiunik \(2016, 2017\)](#).

Regarding falsification analysis, [Frandsen \(2017\)](#) discusses a manipulation test for cases where the score is discrete. The importance of falsification tests and the use of placebo outcomes is discussed in general in the literature on analysis of experiments (e.g., [Rosenbaum, 2002, 2010](#); [Imbens and Rubin, 2015](#)). [Lee \(2008\)](#) applies and further develops these ideas in the RD design context, and [Canay and Kamat \(2018\)](#) develops a permutation inference approach for RD designs. [Ganong and Jäger \(2018\)](#) propose a permutation inference approach based on the idea of placebo RD cutoffs for the Kink RD designs, Regression Kink designs, and related settings. Falsification testing based on donut hole specifications are discussed in [Bajari et al. \(2011\)](#) and [Barreca et al. \(2016\)](#).

### 3 RD Designs with Discrete Running Variables

The canonical continuity-based RD design assumes that the score that determines treatment is a continuous random variable. A random variable is continuous when the set of values that it can take contains an uncountable number of elements. For example, a share such as a party’s proportion of the vote is continuous, because it can take any value in the  $[0, 1]$  interval. In practical terms, when a variable is continuous, all the observations in the dataset have distinct values—i.e., there are no ties. In contrast, a discrete random variable such as date of birth can only take a finite number of values; as a result, a random sample of a discrete variable will contain “mass points”—that is, values that are shared by many observations.

When the RD score is not a continuous random variable, the local polynomial methods we described in Part I are not directly applicable. This is a practically relevant issue, because many real RD applications have a discrete score that can only take a finite number of values. We now consider an empirical RD example where the running variable has mass points in order to illustrate some of the strategies that can be used to analyze RD designs with a discrete running variable. We employ this empirical application to illustrate how identification, estimation, and inference can be modified when the dataset contains multiple observations per value of the running variable.

The key issue when deciding how to analyze a RD design with a discrete score is the number of distinct mass points. Local polynomial methods will behave essentially as if each mass point is a single observation. Thus, if the score is discrete but the number of mass points is sufficiently large, using local polynomial methods may still be appropriate. In contrast, if the number of mass points is very small, local polynomial methods will not be directly applicable. In this case, analyzing the RD design using the local randomization approach is a natural alternative. When the score is discrete, the local randomization approach has the advantage of that the window selection procedure is no longer needed, as the smallest window is well defined. Regardless of the estimation and inference method employed, issues of interpretability and extrapolation will naturally arise. In this section, we discuss and illustrate all these issues using an example from the education literature.

#### 3.1 The Effect of Academic Probation on Future Academic Achievement

The example we re-analyze is the study by [Lindo, Sanders and Oreopoulos \(2010, LSO hereafter\)](#), who use an RD design to investigate the impact of placing students on academic

probation on their future academic performance. Our choice of an education example is intentional. The origins of the RD design can be traced to the education literature, and RD methods continue to be used extensively in education because interventions such as scholarships or probation programs are often assigned on the basis of a test score and a fixed approving threshold. Moreover, despite being continuous in principle, it is common for test scores and grades to be discrete in practice.

LSO analyze a policy at a large Canadian university that places students on academic probation when their grade point average (GPA) falls below a certain threshold. As explained by LSO, the treatment of placing a student on academic probation involves setting a standard for the student's future academic performance: a student who is placed on probation in a given term must improve her GPA in the next term according to campus-specific standards, or face suspension. Thus, in this RD design, the unit of analysis is the student, the score is the student's GPA, the treatment of interest is placing the student on probation, and the cutoff is the GPA value that triggers probation placement. Students come from three different campuses. In campuses 1 and 2, the cutoff is 1.5. In campus 3 the cutoff is 1.6. In their original analysis, the authors normalize the score, centering each student's GPA at the appropriate cutoff, and pooling the observations from the three campuses in a single dataset. The resulting running variable is therefore the difference between the student's GPA and the cutoff; this variable ranges from -1.6 to 2.8, with negative values indicating that the student was placed on probation, and positive values indicating that the student was not placed on probation.

Table 3.1 contains basic descriptive statistics for the score, treatment, outcome and predetermined covariates that we use in our re-analysis. There are 40,582 student-level observations coming the 1996-2005 period. LSO focus on several outcomes that can be influenced by academic probation. In order to simplify our illustration, we focus on two of them: the student's decision to permanently leave the university (**Left University After 1st Evaluation**), and the GPA obtained by the student in the term immediately after he was placed on probation (**Next Term GPA**). Naturally, the second outcome is only observed for students who decide to continue at the university, and thus the effects of probation on this outcome must be interpreted with caution, as the decision to leave the university may itself be affected by the treatment. We also investigate some of the predetermined covariates included in the LSO dataset: an indicator for whether the student is male (**male**), the student's age at entry (**age**), the total number of credits for which the student enrolled in the first year (**totcredits\_year1**), an indicator for whether the student's first language is English (**english**), an indicator for whether the student was born in North America

(`bpl_north_america`), the percentile of the student’s average GPA in standard classes taken in high school (`hsgrade_pct`), and indicators for whether the student is enrolled in each of the three different campuses (`loc_campus1`, `loc_campus2`, and `loc_campus3`). As LSO, we employ these covariates to study the validity of the RD design.

Table 3.1: Descriptive Statistics for [Lindo et al. \(2010\)](#)

Variable	Mean	Median	Std. Deviation	Min.	Max.	Obs.
Next Term GPA (normalized)	1.047	1.170	0.917	-1.600	2.800	40582
Left University After 1st Evaluation	0.049	0.000	0.216	0.000	1.000	44362
Next Term GPA (not normalized)	2.571	2.700	0.910	-2.384	4.300	40582
Distance from cutoff	-0.913	-0.980	0.899	-2.800	1.600	44362
Treatment Assignment	0.161	0.000	0.368	0.000	1.000	44362
High school grade percentile	50.173	50.000	28.859	1.000	100.000	44362
Credits attempted in first year	4.573	5.000	0.511	3.000	6.500	44362
Age at entry	18.670	19.000	0.743	17.000	21.000	44362
Male	0.383	0.000	0.486	0.000	1.000	44362
Born in North America	0.871	1.000	0.335	0.000	1.000	44362
English is first language	0.714	1.000	0.452	0.000	1.000	44362
At Campus 1	0.584	1.000	0.493	0.000	1.000	44362
At Campus 2	0.173	0.000	0.379	0.000	1.000	44362
At Campus 3	0.242	0.000	0.429	0.000	1.000	44362

### 3.2 Counting the Number of Mass Points in the RD Score

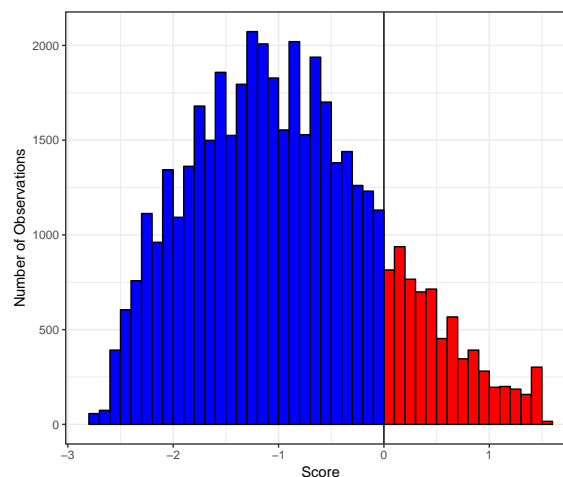
The crucial issue in the analysis of RD designs with discrete scores is the number of mass points that actually occur in the dataset. When this number is large, it will be possible to apply the tools from the continuity-based approach to RD analysis, after possibly changing the interpretation of the treatment effect of interest. When this number is either moderately small or very small, a local randomization approach will be most appropriate. In the latter situation, (local or global) polynomial fitting will be useful only as an exploratory device unless the research is willing to impose strong parametric assumptions. Therefore, the first step in the analysis of an RD design with a discrete running variable is to analyze the empirical distribution of the score and determine the total number of observations, the total number of mass points, and the total number of observations per mass point. We continue to illustrate this step of the analysis using the LSO application.

Since only those students who have GPA below a certain level are placed on probation, the treatment—the assignment to probation—is administered to students whose GPA is to the left of the cutoff. It is customary to define the RD treatment indicator as equal to one for units whose score is above (i.e., to the right of) the cutoff. To conform to this convention,

we invert the original running variable in the LSO data. We multiply the original running variable—the distance between GPA and the campus cutoff—by -1, so that, according to the transformed running variable, students placed on probation (i.e. those with GPA below the cutoff) are now above the cutoff, and students not placed on probation (i.e. those with GPA above the cutoff) are now below the cutoff.

For example, a student who has  $X_i = -0.2$  in the original score is placed on probation because her GPA is 0.2 units below the threshold. The value of the transformed running variable for this treated student is  $\tilde{X}_i = 0.2$ . Moreover, since we define the treatment as  $\mathbb{1}(\tilde{X}_i \geq 0)$ , this student will now be placed above the cutoff. The only caveat is that we must shift slightly those control students whose original GPA is exactly equal to the cutoff, since for these students the original normalized running variable is exactly zero and thus multiplying by -1 does not alter their score. In the scale of the transformed variable, we need these students to be below zero to continue to assign them to the control condition (i.e., the non-probation condition). We manually change the score of students who are exactly at zero to  $X_i = -0.000005$ . A histogram of the transformed running variable is shown in Figure 3.1.

Figure 3.1: Histogram of Transformed Running Variable—LSO Data



We first check how many total observations we have in the dataset, that is, how many observations have a non-missing value of the score.

```
> length(X)
[1] 44362
```

Analogous Stata command

```
. count if X != .
```



The total sample size in this application is large, with 44,362 observations. However, because the running variable is discrete, the crucial step is to calculate how many mass points we have.

```
> length(unique(X))
[1] 430
```

Analogous Stata command

```
. codebook X
```

The 44,362 total observations in the dataset take only 430 distinct values. This means that, on average, there are roughly 100 observations per value. To have a better idea of the density of observations near the cutoff, Table 3.2 shows the number of observations for the five mass points closest to the cutoff; this table also illustrates how the score is transformed. Since the original score ranges between -1.6 and 2.8, our transformed score ranges from -2.799 to 1.6. Both the original and the transformed running variables are discrete, because the GPA increases in increments of 0.01 units and there are many students with the same GPA value. For example, there are 76 students who are 0.02 GPA units below the cutoff. Of these 76 students,  $44 + 5 = 49$  have a GPA of 1.48 (because the cutoff in Campuses 1 and 2 is 1.5), and 27 students have a GPA of 1.58 (because the cutoff in Campus 3 is 1.6). The same phenomenon of multiple observations with the same value of the score occurs at all other values of the score; for example, there are 228 students who have a value of zero in the original score (and -0.000005 in our transformed score).

Table 3.2: Observations at Closest Mass Points

Original Score	Transformed Score	Treatment Status	Number of Observations			
			All Campuses	Campus 1	Campus 2	Campus 3
⋮	⋮	⋮	⋮	⋮	⋮	⋮
-0.02	0.02	Treated	76	44	5	27
-0.01	0.01	Treated	70	25	16	29
0	-0.000005	Control	228	106	55	67
0.01	-0.01	Control	77	30	11	36
0.02	-0.02	Control	137	58	34	45
⋮	⋮	⋮	⋮	⋮	⋮	⋮

### 3.3 Using the Continuity-Based Approach when the Number of Mass Points is Large

When the number of mass points in the discrete running variable is sufficiently large, it is possible to use the tools from the continuity-based approach to RD analysis. We discussed these tools extensively in Part I. The LSO application illustrates a case in which a continuity-based analysis might be possible, since the total number of mass points is 430, a moderate value. Because there are mass points, extrapolation between them is unavoidable; however, in practical terms, this is no different from analyzing a (finite sample) dataset with a sample of size 430.

We start with a falsification analysis, doing a continuity-based density test and a continuity-based analysis of the effect of the treatment on predetermined covariates. First, we use `rddensity` to test whether the density of the score is continuous at the cutoff.

```
> out = rddensity(X)
> summary(out)
```

RD Manipulation Test using local polynomial density estimation.

Number of obs =	44362	
Model =	unrestricted	
Kernel =	triangular	
BW method =	comb	
VCE method =	jackknife	
Cutoff c = 0	Left of c	Right of c
Number of obs	37211	7151
Eff. Number of obs	10083	4137
Order est. (p)	2	2
Order bias (q)	3	3
BW est. (h)	0.706	0.556
Method	T	P >  T
Robust	-0.4544	0.6496

Analogous Stata command

```
. rddensity X
```

The p-value is 0.6496, and we fail to reject the hypothesis that the density of the score changes discontinuously at the cutoff point.

Next, we use `rdrobust` to use local polynomial methods to estimate the RD effect of being placed on probation on several predetermined covariates. We use the default specifications

in `rdrobust`, that is, a MSE-optimal bandwidth that is equal on each side of the cutoff, a triangular kernel, a polynomial of order one, and a regularization term.

For example, we can estimate the RD effect of probation on `hsgrade_pct`, the measure of high school performance.

```
> out = rdrobust(data$hsgrade_pct, X)
> summary(out)
Call: rdrobust

Number of Obs.          44362
BW type                mserd
Kernel                 Triangular
VCE method             NN

Number of Obs.          37211      7151
Eff. Number of Obs.    6115      3665
Order est. (p)         1          1
Order bias (p)         2          2
BW est. (h)            0.465     0.465
BW bias (b)            0.759     0.759
rho (h/b)              0.612     0.612
```

```
=====
              Method      Coef. Std. Err.      z    P>|z|      [ 95% C.I. ]
=====
Conventional    1.328      1.010     1.315   0.189   [-0.652 , 3.309]
Robust          -          -       1.298   0.194   [-0.788 , 3.878]
=====
```

Analogous Stata command

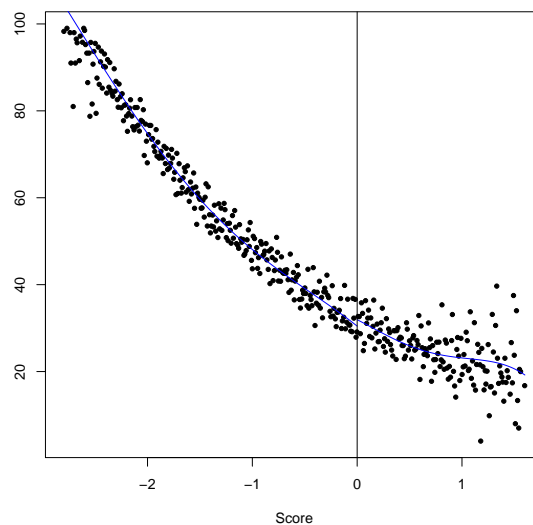
```
. rdrobust hsgrade_pct X
```

And we can also explore the RD effect graphically using `rdplot`.

```
> rdplot(data$hsgrade_pct, X, x.label = "Score", y.label = "",
+ title = "")
```

Analogous Stata command

```
. rdplot hsgrade_pct X
```

Figure 3.2: RD plot for `hsgrade_pct`—LSO data

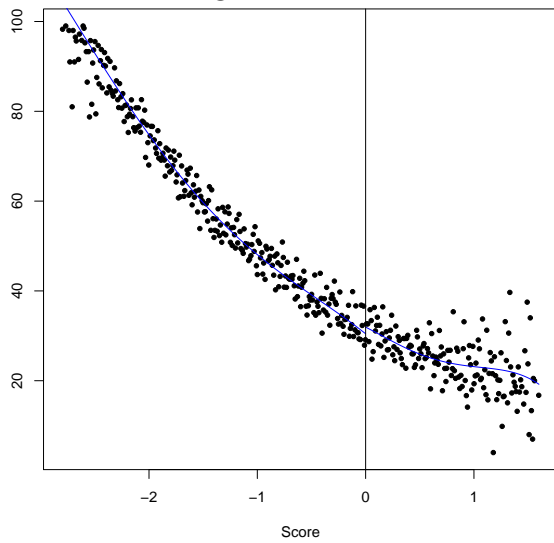
Both the formal analysis and the graphical analysis indicate that, according to this continuity-based local polynomial analysis, the students right above and below the cutoff are similar in terms of their high school performance.

We repeat this analysis for the nine predetermined covariates mentioned above. Table 3.3 presents a summary of the results, and Figure 3.3 shows the associated RD plots for six of the nine covariates.

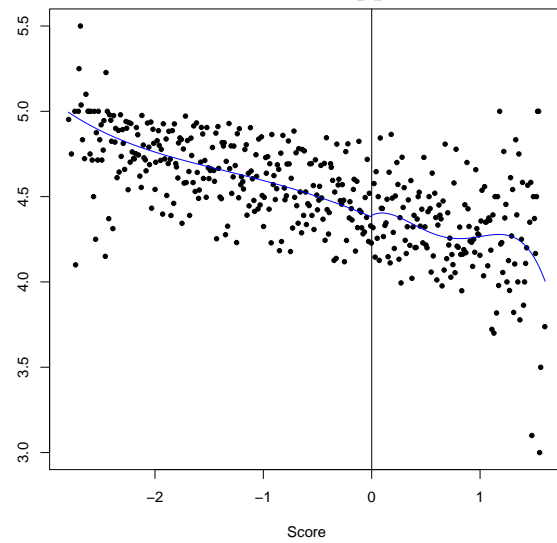
Table 3.3: RD Effects on Predetermined Covariates—LSO data, Continuity-Based Approach

Variable	MSE-Optimal	RD	Robust Inference		Number of Observations
	Bandwidth	Estimator	p-value	Conf. Int.	
High school grade percentile	0.465	1.328	0.194	[-0.788, 3.878]	9780
Credits attempted in first year	0.301	0.081	0.005	[0.027, 0.157]	6443
Age at entry	0.463	0.017	0.637	[-0.060, 0.099]	9780
Male	0.482	-0.012	0.506	[-0.067, 0.033]	10121
Born in North America	0.505	0.014	0.374	[-0.018, 0.049]	10757
English is first language	0.531	-0.035	0.085	[-0.083, 0.005]	11239
At Campus 1	0.289	-0.020	0.356	[-0.093, 0.034]	5985
At Campus 2	0.476	-0.018	0.333	[-0.063, 0.021]	9910
At Campus 3	0.271	0.036	0.139	[-0.015, 0.109]	5786

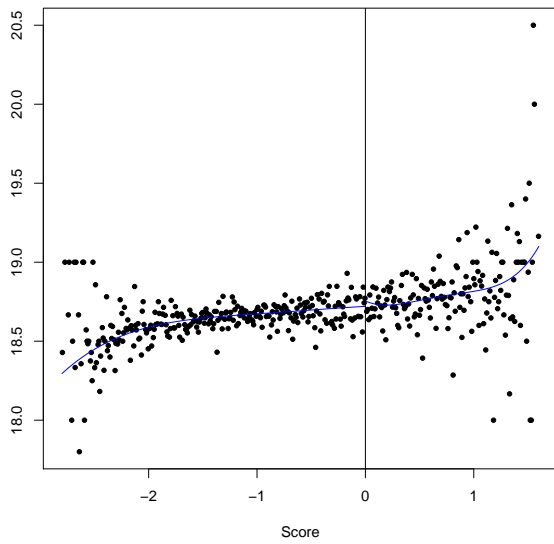
Figure 3.3: RD Plots for Predetermined Covariates—LSO Application



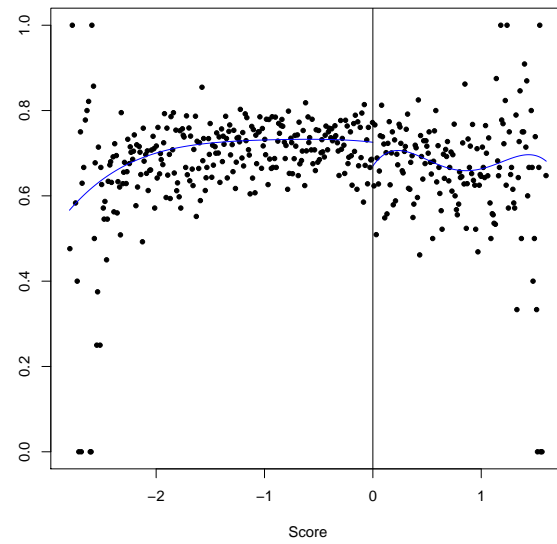
(a) High School Grade Percentile



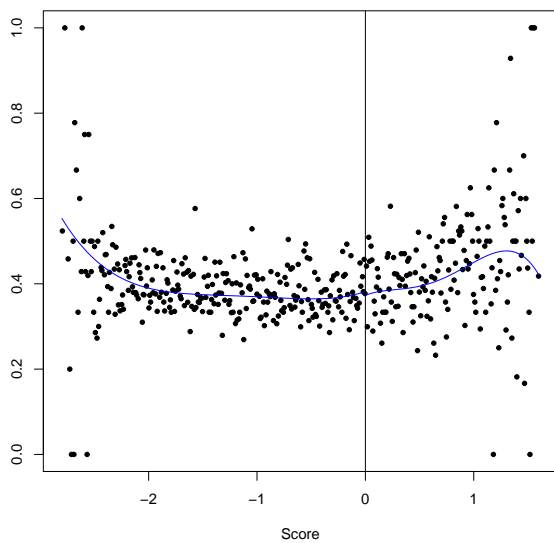
(b) Total Credits in First Year



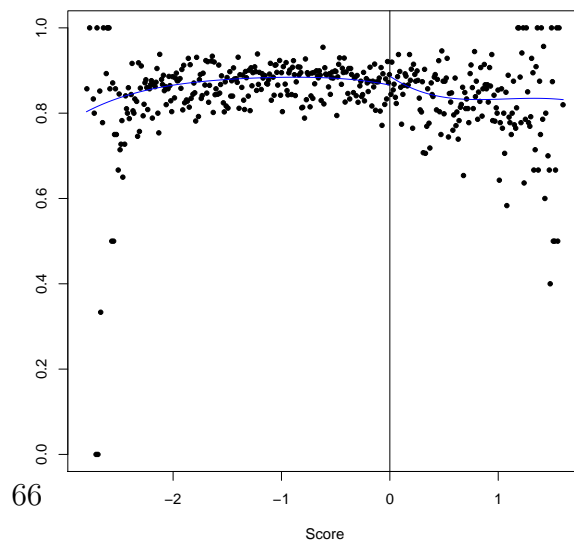
(c) Age at Entry



(d) English First Language Indicator



(e) Male Indicator



(f) Born in North America Indicator

Overall, the results indicate that the probation treatment has no effect on the pre-terminated covariates, with two exceptions. First, the effect on `totcredits_year1` has associated p-value 0.004, rejecting the hypothesis of no effect at standard levels. However, the point estimate is small: treated students take an additional 0.081 credits in the first year, but the average value of `totcredits_year1` in the overall sample is 4.43, with a standard deviation of roughly 0.5. Second, students who are placed on probation are 3.5 percentage points less likely to speak English as a first language, an effect that is significant at 10%. This difference is potentially more worrisome, and is also somewhat noticeable in the RD plot in Figure 3.3(d).

Next, we analyze the effect of being placed on probation on the outcome of interest, `nextGPA`, the GPA in the following academic term. We first use `rdplot` to visualize the effect.

```
> out = rdplot(nextGPA_nonorm, X, binselect = "esmv")
> summary(out)
Call: rdplot

Number of Obs.          40582
Kernel                  Uniform

Number of Obs.          34854          5728
Eff. Number of Obs.    34854          5728
Order poly. fit (p)     4          4
BW poly. fit (h)        2.800        1.600
Number of bins scale    1          1

Bins Selected           690          362
Average Bin Length      0.004        0.004
Median Bin Length       0.004        0.004

IMSE-optimal bins      44          13
Mimicking Variance bins 690          362

Relative to IMSE-optimal:
Implied scale           15.682        27.846
WIMSE variance weight   0.000        0.000
WIMSE bias weight       1.000        1.000
```

Analogous Stata command

```
. rdplot nextGPA_nonorm X, binselect(esmv) ///
> graph_options(graphregion(color(white))) ///
> xtitle(Score) ytitle(Outcome)
```

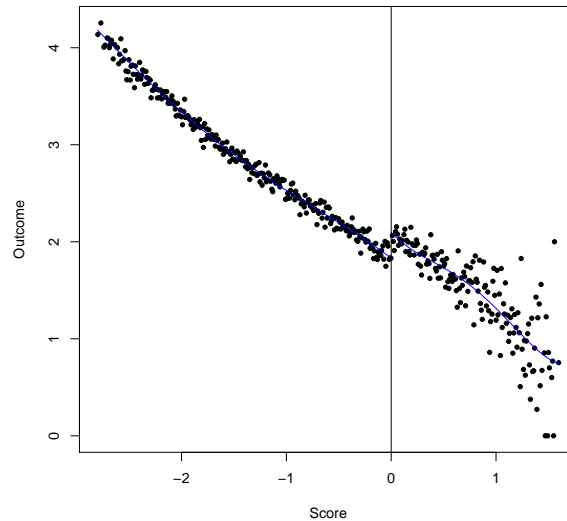


Figure 3.4: RD Plot for nextGPA—LSO Data

Overall, the plot shows a very clear negative relationship between the running variable and the outcome: students who have a low GPA in the current (and thus have a higher value of the running variable) tend to also have a low GPA in the following term. The plot also shows that students with scores just above the cutoff (who are just placed on probation) tend to have a higher GPA in the following term relative to students who are just below the cutoff and just avoided probation. These results are confirmed when we use a local linear polynomial and robust inference to provide a formal statistical analysis of the RD effect.

```
> out = rdrobust(nextGPA_nonorm, X, kernel = "triangular", p = 1,
+ bwselect = "mserd")
> summary(out)
Call: rdrobust
```

Number of Obs.	40582	
BW type	mserd	
Kernel	Triangular	
VCE method	NN	
Number of Obs.	34854	5728
Eff. Number of Obs.	5249	3038
Order est. (p)	1	1
Order bias (p)	2	2
BW est. (h)	0.437	0.437
BW bias (b)	0.717	0.717
rho (h/b)	0.610	0.610

Method	Coef.	Std. Err.	z	P> z	[ 95% C.I. ]
Conventional	0.222	0.040	5.615	0.000	[0.145 , 0.300]
Robust	-	-	4.572	0.000	[0.122 , 0.304]

## Analogous Stata command

```
. rdrobust nextGPA_nonorm X, kernel(triangular) p(1) bwselect(mserd)
```

As shown, students who are just placed on probation improve their GPA in the following term by 0.2221 additional points, relative to students who just missed probation. The robust p-value is less than 0.00005, and the robust 95% confidence interval ranges from 0.1217 to 0.3044. Thus, the evidence indicates that, conditional on not leaving the university, being placed on academic probation translates into an increase in future GPA. The point estimate of 0.2221—obtained with `rdrobust` within a MSE-optimal bandwidth of 0.4375—is very similar to the effect of 0.23 grade points found by LSO within an ad-hoc bandwidth of 0.6.

To better understand this effect, we may be interested in knowing the point estimate for the controls and treated students separately. To see this information, we explore the information returned by `rdrobust`.

```
> rdout = rdrobust(nextGPA_nonorm, X, kernel = "triangular", p = 1,
+ bwselect = "mserd")
> print(names(rdout))
[1] "Estimate" "bws" "coef" "bws" "se" "z" "pv"
"ci" "h_l" "h_r" "b_l" "b_r" "N_h_l"
"N_h_r" "beta_p_l" "beta_p_r" "V_cl" "V_rb" "N"
Nh" "Nb" "c" "p" "q" "bias" "beta_p"
[27] "kernel" "vce" "bwselect" "level" "all" "call"
> print(rdout$beta_p_r)
[,1]
[1,] 2.0671526
[2,] -0.6713546
> print(rdout$beta_p_l)
[,1]
[1,] 1.8450372
[2,] -0.6804159
```

## Analogous Stata command

```
. rdrobust nextGPA_nonorm X
. ereturn list
```



This output shows the estimated intercept and slope from the two local regressions estimated separately to the right (`beta_p_r`) and left (`beta_p_l`) of the cutoff. At the cutoff, the average GPA in the following term for control students who just avoid probation is 1.8450372, while the average future GPA for treated students who are just placed on probation is 2.0671526. The increase is the estimated RD effect reported above,  $2.0671526 - 1.8450372 = 0.2221154$ . This represents approximately a 12% GPA increase relative to the control group, a considerable effect.

An alternative to the simplest use of `rdrobust` illustrated above is to cluster the standard errors by every value of the score—this is the approach recommended by [Lee and Card \(2008\)](#), as we discuss in the Further Readings section below. We implement this using the `cluster` option in `rdrobust`.

```
> clustervar = X
> out = rdrobust(nextGPA_nonorm, X, kernel = "triangular", p = 1,
+ bwselect = "mserd", vce = "hc0", cluster = clustervar)
> summary(out)
Call: rdrobust
```

```
Number of Obs.          40582
BW type                mserd
Kernel                 Triangular
VCE method              HCO
```

```
Number of Obs.          34854      5728
Eff. Number of Obs.    4357      2709
Order est. (p)         1          1
Order bias (p)         2          2
BW est. (h)            0.377      0.377
BW bias (b)            0.627      0.627
rho (h/b)              0.602      0.602
```

```
=====
              Method      Coef. Std. Err.      z    P>|z|      [ 95% C.I. ]
=====
Conventional    0.215      0.033      6.463    0.000      [0.150 , 0.280]
Robust          -          -          5.396    0.000      [0.132 , 0.282]
=====
```

Analogous Stata command

```
. gen clustervar=X
. rdrobust nextGPA_nonorm X, kernel(triangular) p(1) bwselect(mserd) vce(cluster clustervar)
```

The conclusions remain essentially unaltered, as the 95% robust confidence interval

changes only slightly from  $[0.1217, 0.3044]$  to  $[0.1316, 0.2818]$ . Note that the point estimate moves slightly from 0.2221 to 0.2149 because the MSE-optimal bandwidth with clustering shrinks to 0.3774 from 0.4375, and the bias bandwidth also decreases.

### 3.4 Interpreting Continuity-Based RD Analysis with Mass Points

Provided that the number of mass points in the score is reasonably large, it is possible to analyze an RD design with a discrete score using the tools from the continuity-based approach. However, it is important to understand how to correctly interpret the results from such analysis. We now analyze the LSO application further, with the goal of clarifying these issues.

When there are mass points in the running variable, local polynomial methods for RD analysis behave essentially as if we had as many observations as mass points, and therefore the method implies extrapolation from the closest mass point on either side to the cutoff. In other words, when applied to a RD design with a discrete score, the effective number of observations used by continuity-based methods is the number of mass points or distinct values, not the total number of observations. Thus, in practical terms, fitting a local polynomial to the raw data with mass points is roughly equivalent to fitting a local polynomial to a “collapsed” version of the data, where we aggregate the original observations by the discrete score values, calculating the average outcome for all observations that share the same score value. Thus, the total number of observations in the collapsed dataset is equal to the number of mass points in the running variable.

To illustrate this procedure with the LSO data, we calculate the average outcome for each of the 430 mass points in the score value. The resulting dataset has 430 observations, where each observation consists of a score-outcome pair: every score value is paired with the average outcome across all students in the original dataset whose score is equal to that value. When then use `rdrobust` to estimate the RD effect with a local polynomial.

```
> data2 = data.frame(nextGPA_nonorm, X)
> dim(data2)
[1] 44362    2
> collapsed = aggregate(nextGPA_nonorm ~ X, data = data2, mean)
> dim(collapsed)
[1] 429    2
> out = rdrobust(collapsed$nextGPA_nonorm, collapsed$X)
> summary(out)
Call: rdrobust
```

```
Number of Obs.          429
```

BW type	mserd	
Kernel	Triangular	
VCE method	NN	
Number of Obs.	274	155
Eff. Number of Obs.	51	50
Order est. (p)	1	1
Order bias (p)	2	2
BW est. (h)	0.506	0.506
BW bias (b)	0.805	0.805
rho (h/b)	0.628	0.628

Method	Coef.	Std. Err.	z	P> z	[ 95% C.I. ]
Conventional	0.246	0.032	7.640	0.000	[0.183 , 0.309]
Robust	-	-	6.279	0.000	[0.166 , 0.316]

Analogous Stata command

```
. collapse (mean) nextGPA_nonorm, by(X)
. rdrobust nextGPA_nonorm X
```

The estimated effect is 0.2456, with robust p-value less than 0.00005. This is similar to the 0.2221 point estimate obtained with the raw dataset. The similarity between the two point estimates is remarkable, considering that the former was calculated using 430 observations, while the latter was calculated using 40,582 observations, more than a ninety-fold increase. Indeed, the inference conclusions from both analysis are extremely consistent, as the robust 95% confidence interval using the raw data is [0.1217 , 0.3044], while the robust confidence interval for the collapsed data is [0.1659, 0.3165], both indicating that the plausible values of the effect are in roughly the same positive range.

This analysis shows that the seemingly large number of observations in the raw dataset is effectively much smaller, and that the behavior of the continuity-based results is governed by the average behavior of the data at every mass point. Thus, a natural point of departure for researchers who wish to study a discrete RD design with many mass points is to collapse the data and estimate the effects on the aggregate results. As a second step, these aggregate results can be compared to the results using the raw data—in most cases, both sets of results should lead to the same conclusions.

While the mechanics of local polynomial fitting using a discrete running variable are clear now, the actual relevance and interpretation of the treatment effect may change. As we will

discuss below, researchers may want to change the parameter of interest altogether when the score is discrete. Alternatively, (parametric) extrapolation is unavoidable to point identification. To be more precise, because the score is discrete, it is not possible to nonparametrically point identify the vertical distance between  $\tau_{\text{SRD}} = \mathbb{E}[Y_i(1)|X_i = \bar{x}] - \mathbb{E}[Y_i(0)|X_i = \bar{x}]$ , even asymptotically, because conceptually the lack of denseness in  $X_i$  makes it impossible to appeal to large sample approximations. Put differently, if researchers insist on retaining the same parameter of interest as in the canonical RD design, then extrapolation from the closest mass point to the cutoff will be needed, no matter how large the sample size is.

Of course, there is no reason why the same RD treatment effect would be of interest when the running variable is discrete or, if it is, then any extrapolation method would be equally valid. Thus, continuity-based methods, that is, simple local linear extrapolation towards the cutoff point is natural and intuitive. When only a few mass points are present, then bandwidth selection makes little sense, and the research may just conduct linear (parametric) extrapolation globally, as this is essentially the only possibility, if the goal is to retain the same canonical treatment effect parameter.

### 3.5 Local Randomization RD Analysis with Discrete Score

A natural alternative to analyze an RD design with a discrete running variable is to use the local randomization approach, which effectively changes the parameter of interest from the RD treatment effect at the cutoff to the RD treatment effect in the neighborhood around the cutoff where local randomization is assumed to hold. A key advantage of this alternative conceptual framework is that, unlike the continuity-based approach illustrated above, it can be used even when there are very few mass points in the running variable: indeed, it can be used with as few as two mass points.

To compare the change in RD parameter of interest, consider the extreme case where the score takes five values  $-2, -1, 0, 1, 1$  and the RD cutoff is  $\bar{x} = 0$ . Then, the continuity-based parameter of interest is  $\tau_{\text{SRD}} = \mathbb{E}[Y_i(1)|X_i = 0] - \mathbb{E}[Y_i(0)|X_i = 0]$ , which is *not* nonparametrically identifiable, because the score of control observations can never get close enough to 0. However, if the true window is, say,  $W_0 = [-1, 0]$ , the local randomization parameter will be  $\tau_{\text{LR}} = \mathbb{E}[Y_i(1)|X_i = 0] - \mathbb{E}[Y_i(0)|X_i = -1]$ , which is nonparametrically identifiable under the conditions discussed in Section 2. Going from  $\tau_{\text{LR}}$  to  $\tau_{\text{SRD}}$  requires extrapolating from  $\mathbb{E}[Y_i(0)|X_i = -1]$  to  $\mathbb{E}[Y_i(0)|X_i = 0]$ , which is impossible without additional assumptions even in large samples because of the intrinsic discreteness of the running variable. In some specific applications, additional features may allow researchers to extrapolate (e.g., round-

ing), but in general extrapolation will require additional restrictions on the data generating process. Furthermore, from a conceptual point of view, it can be argued that the parameter  $\tau_{LR}$  is more interesting and policy relevant than the parameter  $\tau_{SRD}$  when the running variable is discrete.

When the score is discrete using the local randomization approach for inference does not require choosing a window in most applications. In other words, with a discrete running variable the researcher knows the exact location of the minimum window around the cutoff: this window is the interval of the running variable that contains the two mass points, one on each side of the cutoff, that are immediately consecutive to the cutoff value. Crucially, if local randomization holds, then it must hold for the *smallest* window in the absence of design failures such as manipulation of the running variable. To illustrate, as shown in Table 3.2, in the LSO application the original score has a mass point at zero where all observations are control (because they reach the minimum GPA required to avoid probation), and the mass point immediately below it occurs at -0.01, where all students are placed on probation because they fall short of the threshold to avoid probation. Thus, the smallest window around the cutoff in the scale of the original score is  $W_0 = [0.00, -0.01]$ . Analogously, in the scale of the transformed score, the minimum window is  $W_0 = [-0.000005, 0.01]$ .

Regardless of the scale used, the important point is that the minimum window around the cutoff in a local randomization analysis of an RD with a discrete score is precisely the interval between the two consecutive mass points where the treatment status changes from zero to one. Note that the particular values taken by the score are irrelevant, as the analysis will proceed to assume that the treated and control groups were assigned to treatment as-if randomly, and will typically make the exclusion restriction assumption that the particular value of the score has no direct impact on the outcome of interest. Moreover, the location of the cutoff is no longer meaningful, as any value of the cutoff between the minimum value of the score on the treated side and the maximum value of the score in the control side will produce identical treatment and control groups.

Once the researcher finds the treated and control observations located at the two mass points around the cutoff, the local randomization analysis can proceed as explained in Section 2. We first conduct a falsification analysis, to determine whether the assumption of local randomization in the window  $[-0.00005, 0.1]$  seems consistent with the empirical evidence. We conduct a density test using the `rdwinselect` function using the option `nwindows=1` to see only results for this window, to test whether the density of observations in this window is consistent with the density that would have been observed in a series of unbiased coin flips.

```
> out = rdwinselect(X, wmin = 0.01, nwindows = 1, cutoff = 5e-06)
```

Window selection for RD under local randomization

```
Number of obs = 44362
Order of poly = 0
Kernel type = uniform
Reps = 1000
Testing method = rdrandinf
Balance test = diffmeans
```

Cutoff c = 0	Left of c	Right of c
Number of obs	37211	7151
1st percentile	298	0
5th percentile	1817	269
10th percentile	3829	663
20th percentile	7588	1344

Window length / 2 Obs >= c	p-value	Var. name	Bin. test	Obs < c
0.01 77	NA	NA	0	228

Analogous Stata command

```
. rdwinselect X, wmin(0.01) nwindows(1) cutoff(0.000005)
```

As shown in the `rdwinselect` output and also showed previously in Table 3.2, there are 228 control observations immediately below the cutoff, and 77 above the cutoff. In other words, there are 228 students who get exactly the minimum GPA needed to avoid probation, and 77 students who get the maximum possible GPA that still places them on probation. The number of control observations is roughly three times higher than the number of treated observations, a ratio that is inconsistent with the assumption that the probability of treatment assignment in this window was  $1/2$ —the p-value of the Binomial test reported in column `Bin. test` is indistinguishable from zero.

We can also obtain this result by using the Binomial test commands directly.

```
> binom.test(77, 305, 1/2)
```

Exact binomial test

```
data: 77 and 305
number of successes = 77, number of trials = 305, p-value < 2.2e-16
alternative hypothesis: true probability of success is not equal to 0.5
95 percent confidence interval:
```

```

0.2046807 0.3051175
sample estimates:
probability of success
          0.252459

```

Analogous Stata command

```
. bitesti 305 77 1/2
```

Although these results alone do not imply that the local randomization RD assumptions are violated, the fact that there are many more control than treated students is consistent with what one would expect if students were actively avoiding an undesirable outcome. The results raise some concern that students may have been aware of the probation cutoff, and may have tried to appeal their final GPA in order to avoid being placed on probation.

Strictly speaking, an imbalanced number of observations would not pose any problems if the types of students in the treated and control groups were on average similar. To establish whether treated and control students at the cutoff are similar in terms of observable characteristics, we use `rdrandinf` to estimate the RD effect of probation on the predetermined covariates introduced above.

We report the full results for the covariate `hsgrade_pct`.

```
> out = rdrandinf(data$hsgrade_pct, X, wl = -0.005, wr = 0.01,
+ seed = 50)
```

```
Selected window = [-0.005;0.01]
```

```
Running randomization-based test...
```

```
Randomization-based test complete.
```

```

Number of obs =      44362
Order of poly =         0
Kernel type   =    uniform
Reps          =     1000
Window        =    set by user
H0:          tau =         0
Randomization =    fixed margins

Cutoff c = 0          Left of c          Right of c
Number of obs        37211                7151
Eff. number of obs   228                  77
Mean of outcome       29.118               32.675
S.d. of outcome       21.737               21.625
Window                -0.005               0.01

```

Statistic	T	Finite sample	Large sample	
		P> T	P> T	Power vs d =
10.868				
Diff. in means	3.557	0.219	0.213	0.968

## Analogous Stata command

```
. rdrandinf hsgrade_pct X , seed(50) wl(-.005) wr(.01)
```

We repeated this analysis for all predetermined covariates, but do not present the individual runs to conserve space. A summary of the results is reported in Table 3.4. As shown, treated and control students seem indistinguishable in terms of prior high school performance, total number of credits, age, sex, and place of birth.

On the other hand, the Fisherian sharp null hypothesis is that the treatment has no effect on the English-as-first-language indicator is rejected with p-value of 0.009. The average differences in this variable are very large: 75% of control students speak English as first language, but only 62.3% of treated students do. This difference is consistent with the local polynomial results we reported for this variable in Table 3.3, although the difference is much larger (an average difference of -3.5 percentage points in the continuity-based analysis, versus an average difference of -15 percentage points in the local randomization analysis). A similar phenomenon occurs for the Campus 2 and Campus 3 indicators, which appear imbalanced in the local randomization analysis (with Fisherian p-values of 0.075 and 0.009) but appear balanced with a continuity-based analysis.

These differences illustrate how a continuity-based analysis of a discrete RD design can mask differences that occurs in mass points closest to the cutoff. In general, when analyzing a RD design with a discrete running variable, it is advisable to perform falsification tests with the two mass points closest to the cutoff in order to detect phenomena of sorting or selection that may go unnoticed when a continuity-based approach is used.



Table 3.4: RD Effects on Predetermined Covariates—LSO data, Local Randomization Approach

Variable	Mean of Controls	Mean of Treated	Diff-in-Means Statistic	Fisherian p-value	Number of Observations
High school grade percentile	29.118	32.675	3.557	0.201	305
Credits attempted in first year	4.228	4.318	0.090	0.157	305
Age at entry	18.772	18.688	-0.084	0.421	305
Male	0.377	0.442	0.064	0.336	305
Born in North America	0.890	0.844	-0.046	0.322	305
English is first language	0.772	0.623	-0.149	0.009	305
At Campus 1	0.465	0.390	-0.075	0.259	305
At Campus 2	0.241	0.143	-0.098	0.075	305
At Campus 3	0.294	0.468	0.174	0.009	305

Finally, we also investigate the extent to which the particular window around the cutoff, including only two mass points, is driving the empirical results by repeating the analysis using different nested windows. These exercise is easily implemented using the command `rdwinselect`:

```
> Z = cbind(data$hsgrade_pct, data$totcredits_year1, data$age_at_entry,
+ data$male, data$bpl_north_america, data$english, data$loc_campus1,
+ data$loc_campus2, data$loc_campus3)
> colnames(Z) = c("hsgrade_pct", "totcredits_year1", "age_at_entry",
+ "male", "bpl_north_america", "english", "loc_campus1", "loc_campus2",
+ "loc_campus3")
> out = rdwinselect(X, Z, p = 1, seed = 50, wmin = 0.01, wstep = 0.01,
+ cutoff = 5e-06)
```

Window selection for RD under local randomization

```
Number of obs = 44362
Order of poly = 1
Kernel type = uniform
Reps = 1000
Testing method = rdrandinf
Balance test = diffmeans
```

```
Cutoff c = 0
Number of obs Left of c Right of c
1st percentile 298 0
5th percentile 1817 269
10th percentile 3829 663
20th percentile 7588 1344
```

```
Window length / 2 p-value Var. name Bin.test Obs<c
Obs>=c
```

0.01	0.008	loc_campus3	0	228
77				
0.02	0	totcredits_year	0	298
214				
0.03	0	hsgrade_pct	0	374
269				
0.04	0	loc_campus1	0	494
375				
0.05	0	totcredits_year	0	636
418				
0.06	0	hsgrade_pct	0	714
497				
0.07	0	hsgrade_pct	0	807
663				
0.08	0	totcredits_year	0	877
727				
0.09	0	totcredits_year	0	1049
815				
0.1	0	totcredits_year	0.001	1131
973				

Analogous Stata command

```
. rdwinselect X $covariates, cutoff(0.00005) wmin(0.01) wstep(0.01)
```

The empirical results continue to provide evidence of invariance in at least one pre-intervention covariate for each window considered, using randomization inference methods for the difference in means test statistic.

To complete the empirical illustration, we investigate the local randomization RD treatment effect on the main outcome of interest using `rdrandinf`.

```
> out = rdrandinf(nextGPA_nonorm, X, wl = -0.005, wr = 0.01, seed = 50)
```

```
Selected window = [-0.005;0.01]
```

```
Running randomization-based test...
```

```
Randomization-based test complete.
```

```
Number of obs =      40582
Order of poly  =         0
Kernel type   =    uniform
Reps          =      1000
Window        =    set by user
H0:          tau =         0
Randomization =    fixed margins
```

Cutoff $c = 0$	Left of $c$	Right of $c$		
Number of obs	34854	5728		
Eff. number of obs	208	67		
Mean of outcome	1.83	2.063		
S.d. of outcome	0.868	0.846		
Window	-0.005	0.01		
		Finite sample	Large sample	
Statistic	T	$P >  T $	$P >  T $	Power vs $d =$
0.434				
Diff. in means	0.234	0.063	0.051	0.952

#### Analogous Stata command

```
. rdrandinf nextGPA_nonorm X, seed(50) wl(-0.005) wr(0.01)
```

Remarkably, the difference-in-means between the 208 control students and the 67 treated students in the smallest window around the cutoff is 0.234 grade points, extremely similar to the continuity-based local polynomial effects of 0.2221 and 0.2456 that we found using the raw and aggregated data, respectively. (The discrepancy between the treated and control sample sizes of 77 and 288 reported in Table 3.2 and the sample sizes of 67 and 208 reported in the `rdrandinf` output occurs because there are missing values in the `nextGPA` outcome, as students who leave the university do not have any future GPA.) Moreover, we can reject the null hypothesis of no effect at 10% level using both the Fisherian and the Neyman inference approaches. This shows that the results for next term GPA are remarkably robust: we found very similar results using the  $208 + 67 = 275$  observations closest to the cutoff in a local randomization analysis, the total 40,582 observations using a continuity-based analysis, and the 429 aggregated observations in a continuity-based analysis.

### 3.6 Further Readings

Lee and Card (2008) proposed alternative assumptions under which the local polynomial methods in the continuity-based RD framework can be applied when the running variable is discrete. Their method requires assuming a random specification error that is orthogonal to the score, and modifying inferences by using standard errors that are clustered at each of the different values taken by the score. Similarly, Dong (2015) discusses the issue rounding in the running variable. Both approaches have in common that the score is assumed to be inherently continuous, but somehow imperfectly measured—perhaps because of rounding errors—in such a way that the dataset available to the researcher contains mass points.

[Cattaneo et al. \(2015, Section 6.2\)](#) discuss explicitly the connections between discrete scores and the local randomization approach; see also [Cattaneo et al. \(2017\)](#).

## 4 The Fuzzy RD Design

We now discuss how to modify the analysis and interpretation of the RD design when some units fail to comply with the treatment condition that is assigned to them. In all RD designs, the assignment of treatment follows the rule  $T_i = \mathbb{1}(X_i \geq \bar{x})$ , which assigns all units whose score is below the cutoff  $\bar{x}$  to the control condition, and all units whose score is above  $\bar{x}$  are to the treatment condition. In the Sharp RD design we discussed in Part I, we assumed that (i) all units assigned to the treatment condition do in fact take the treatment, and (ii) no units assigned to the control condition take the treatment. In that case, the rule  $T_i = \mathbb{1}(X_i \geq \bar{x})$  indicates not only the treatment assigned to the units, but also the treatment received by the units.

In practice, however, it is very common to encounter RD designs where some of the units with  $X_i \geq \bar{x}$  fail to receive the treatment and/or some of the units with  $X_i < \bar{x}$  receive the treatment despite being assigned to the control condition. The phenomenon of units receiving a treatment condition different from the condition that is originally assigned to them is generally known as imperfect compliance or non-compliance. The RD design with imperfect compliance is usually referred to as the Fuzzy RD design, to distinguish it from the Sharp RD design where compliance with treatment is perfect. Imperfect compliance is very common in controlled randomized experiments, and is no less common in RD designs.

The Fuzzy RD treatment assignment rule is still  $T_i = \mathbb{1}(X_i \geq \bar{x})$ . However, compliance with treatment is imperfect. As a consequence, although the probability of receiving treatment still jumps abruptly at the cutoff  $\bar{x}$ , it no longer changes from 0 to 1 as in the Sharp RD case. (Naturally, the probability of being assigned to treatment still jumps from 0 to 1 at  $\bar{x}$ .) To describe this design in generality, we need to introduce additional notation. We use the binary variable  $D_i$  to denote whether the treatment was actually received by unit  $i$ . Our notation now distinguishes between the treatment assigned,  $T_i$ , and the treatment actually received,  $D_i$ . Using this notation, we can say that the defining feature of the Fuzzy RD design is that there are some units for which  $T_i \neq D_i$ .

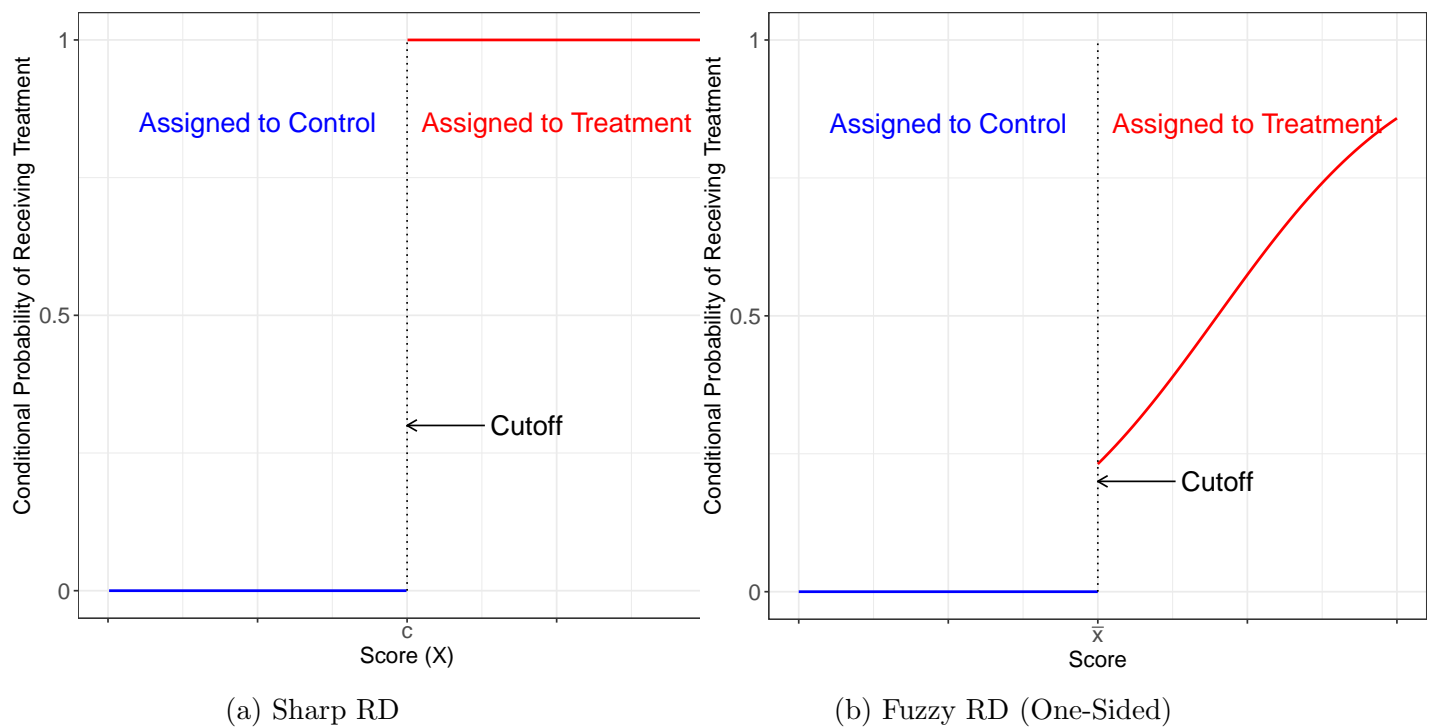
Because the treatment received is not always equal to the treatment assigned, now the treatment received  $D_i$  (also known as “treatment take-up”) has two potential values,  $D_i(1)$  and  $D_i(0)$ , corresponding, respectively, to the treatment taken when the unit is assigned to treatment condition (i.e, when  $X_i \geq \bar{x}$  and  $T_i = 1$ ) and the treatment taken when the unit is assigned to the control condition (i.e, when  $X_i < \bar{x}$  and  $T_i = 0$ ). The observed treatment taken is  $D_i = T_i \cdot D_i(1) + (1 - T_i) \cdot D_i(0)$  and, as occurred for the outcome  $Y_i$ , the fundamental problem of causal inference now means that we do not observe, for example,

whether a unit that was assigned to the treatment condition and took the treatment would have taken the treatment if it had been assigned to the control condition instead. The potential outcomes under treatment and control are still denoted, respectively,  $Y_i(1)$  and  $Y_i(0)$ , although it is important to note that 1 and the 0 in the function  $Y_i(\cdot)$  now refer to the treatment take-up status  $D_i = 1$  and  $D_i = 0$ , respectively, not to the treatment assignment status. Notice that our potential outcomes notation,  $(Y_i(1), Y_i(0))$ , also imposes the restriction that the treatment assignment  $T$  has no direct effect on  $Y$ , since  $T$  is not an argument in the potential outcome functions. The treatment assignment  $T$  affects the outcome  $Y$  only because this assignment induces a change in the actual treatment taken  $D$ , which in turns affects  $Y$ . This type of restriction is sometimes called *exclusion* restriction.

We illustrate the difference between the Sharp and Fuzzy RD designs is in Figure 4.1, where we plot the conditional probability of receiving treatment given the score,  $\mathbb{P}(D_i = 1|X_i = x)$ , for different values of the running variable  $X_i$ . As shown in Figure 4.1(a), in a Sharp RD design the probability of receiving treatment changes exactly from zero to one at the cutoff. In contrast, in a Fuzzy RD design, the change in the probability of being treated at the cutoff is always less than one. Figure 4.1(b) illustrates a particular case of a Fuzzy RD design where units whose score is below the cutoff comply perfectly with the treatment, but compliance with the treatment is imperfect for units whose score is above the cutoff. This case is sometimes called one-sided non-compliance. In general, Fuzzy RD designs can (and often will) exhibit two-sided non-compliance, where  $\mathbb{P}(D_i = 1|X_i = x)$  will be neither zero nor one for units with  $X_i$  near the cutoff  $\bar{x}$ . Below we present an empirical example with two-sided non-compliance.

In the Fuzzy RD design, researchers are typically interested in both the effect of being assigned to treatment (i.e., the effect of  $T$ ) and in the effect of actually receiving treatment (i.e., the effect of  $D$ ) on the outcome of interest. Since units lack the ability to change their treatment assignment, compliance with the assignment is always perfect. Thus, the analysis of the effect of the treatment assigned on the outcome follows the standard analysis of Sharp RD designs. In contrast, the study of the effect of the treatment itself requires modifications and additional assumptions. We devote this section to discuss such assumptions and modifications. We organize our discussion of the analysis and interpretation of Fuzzy RD designs around the same topics previously discussed for the Sharp RD case—estimation of effects, inference, falsification, graphical illustration, and interpretation. Analogously to the Sharp RD case, the analysis of Fuzzy RD designs can be based on a continuity-based approach or a local randomization approach, depending on the identification assumptions invoked. After introducing our empirical example, we discuss and illustrate the continuity-

Figure 4.1: Conditional Probability of Receiving Treatment in Sharp vs. Fuzzy RD Designs



based approach in Subsection 4.2, and the local randomization approach in Section ??.

## 4.1 Empirical Application: The Effect of Cash Transfers on Birth Weight

We now present the empirical application that we use as a running example throughout this section. We re-analyze the study by [Amarante et al. \(2016\)](#) on the effects of in utero exposure to a social assistance program. The authors study Uruguay's *Plan de Atención Nacional a la Emergencia Social* (PANES), a program that gave assistance to the poorest 10 percent of Uruguayan households in the mid 2000s. The most important component of the program was a monthly cash transfer of UY\$1,360—equivalent to roughly US\$102 in Purchasing Power Parity (PPP) terms. Beneficiary households received the transfer for the duration of the program (April 2005 through December 2007) except when their income exceeded a predetermined level. Most households received their first payment in 2005, but the time of first payment differed considerably between households due to delays in program implementation.

The program also had a smaller component, implemented in mid 2006, which consisted

of an electronic food card whose value was significantly less than the value of the monthly transfer. In contrast to most social assistance cash transfer programs implemented in developing countries, PANES was an unconditional cash transfer, as there were no enforced mandatory requirements such as health checks or school attendance to receive the transfer.

Program eligibility depended on household income. Using various household socioeconomic characteristics available in a baseline survey, program administrators computed a predicted income score, which was used to determine program eligibility according to a discontinuous rule: households with predicted income below a predetermined cutoff were eligible to receive the transfers, while households with predicted income score above the cutoff were declared ineligible. However, the eligibility rules were not enforced perfectly, and some ineligible households received the program while some eligible households failed to receive it. The combination of (i) program eligibility assigned discontinuously based on a score and a cutoff, and (ii) imperfect compliance with eligibility status, makes this application an example of a Fuzzy RD design where the running variable is the baseline predicted income score, and the treatment is receiving the cash transfer. [Amarante et al. \(2016\)](#) analyze the effect of the PANES program on the birth weight of babies born during the program period.

In the replication dataset that we analyze, the unit of observation is a mother who gave birth to one or more children during the program period, and the running variable is the income score of the mother according to the baseline survey—the mother is eligible to receive the program if her standardized income score is above zero. The mother is considered to receive the actual treatment if she receives at least one monthly transfer during her pregnancy. The outcomes we re-analyze include baby-level and mother-level variables. We follow the analysis in [Amarante et al. \(2016\)](#), and keep only the births that took place when the program was active. In our analysis of baby-level outcomes, we only include information about the first baby born to the mother during the program period, and discard information about subsequent babies—this affects only 6% of our sample, since 94% of the mothers had only one child during this period.

We present descriptive statistics for the main variables in [Table 4.1](#). There are 24,910 total observations (the differences in sample size across variables are due to missing values). The running variable  $X$  is the predicted income score for every mother, which is standardized and ranges from -0.19 to 0.946. The treatment assignment ( $T$ ) is an indicator equal to one when the running variable is above zero, and the treatment take-up ( $D$ ) is an indicator equal to one if the mother actually received the transfer, regardless of her income score value. As shown in the table, 68.9% of the mothers in this sample are eligible to receive the PANES program, and 60.1% actually receive it. As we will see, the 60.1% of mothers who do receive



the program include both mothers with income score below the eligibility cutoff, and mothers with score above the cutoff, resulting in two-sided non-compliance.

Table 4.1: Descriptive Statistics for AMMV

Variable	Mean	Median	Std. Deviation	Min.	Max.	Obs.
Age	24.067	23.000	6.924	0.000	79.000	24870
Age at Which the Baby was Born	17.948	18.000	2.613	0.000	42.000	9889
Average Pre-PANES Birthweight in Area of Residence	3193.436	3191.066	41.425	2955.000	3576.471	23912
Average Pre-PANES Birthweight in Health Center	3173.876	3210.299	76.312	2373.333	3477.500	24907
Birthweight Below 2.500gr (Y1)	0.088	0.000	0.283	0.000	1.000	24910
Has Any Job (Y2)	0.970	1.000	0.170	0.000	1.000	16519
Running Variable (X)	0.160	0.084	0.262	-0.190	0.946	24910
Treatment Assignment (T)	0.689	1.000	0.463	0.000	1.000	24910
Treatment Take-up (D)	0.601	1.000	0.490	0.000	1.000	22654
Value of Program Income Transfer During Pregnancy (Y3)	629.981	519.988	604.628	0.000	2104.469	22654

Table 4.1 also shows the main outcome of interest: an indicator equal to one if the mother’s first baby during the program period weighed less than 2,500 grams ( $Y_1$ ). We also include the value of the income transfer received ( $Y_2$ ), an outcome that is above zero only for program beneficiaries, and is useful to understand the nature of the treatment and the extent of the noncompliance. Finally, the table also shows four predetermined covariates that we use below for the falsification analysis—the mother’s age, the mother’s age when the first baby was born, the average birthweight in the mother’s area of residence in the period before PANES was active, and the average birthweight in the mother’s health center in the pre-program period.

## 4.2 Continuity-based Analysis

We start by defining the Fuzzy RD design from a continuity-based perspective—the local randomization perspective is analyzed in the next section. The non-compliance phenomenon results in important changes to the kind of effects that can be recovered from the data. When some of the units fail to comply with the treatment that has been assigned to them, it is always possible to analyze the effect of the treatment assignment  $T$  on the outcome. Since compliance with the assignment rule  $T_i = \mathbb{1}(X_i \geq \bar{x})$  is always perfect, the analysis of the effect of  $T$  on  $Y$  follows a Sharp RD design. As we discussed in Part I, a continuity-based analysis of a Sharp RD proceeds by separately (and locally) estimating the average observed outcome given the score,  $\mathbb{E}(Y_i|X_i = x)$ , for observations above and below the cutoff, and then taking the limit of those averages as  $x$  approaches the cutoff  $\bar{x}$ . In the context of a Fuzzy RD design, this estimator captures the effect of being assigned to the treatment condition, which is no longer equivalent to the effect of receiving the treatment.

Under appropriate continuity and regularity conditions, it can be shown that the Sharp RD estimator of the effect of treatment assignment  $T_i$  on the outcome  $Y_i$  consistently estimates the quantity

$$\lim_{x \downarrow \bar{x}} \mathbb{E}[Y_i | X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[Y_i | X_i = x] = \mathbb{E}[(D_i(1) - D_i(0))(Y_i(1) - Y_i(0)) | X_i = \bar{x}] \equiv \tau_{\text{ITT}} \quad (4.1)$$

The quantity on the right-hand-side of Equation 4.2,  $\tau_{\text{ITT}}$ , is usually called the average “intention-to-treat” effect, and it captures the local (i.e., at the cutoff) average effect of being assigned to the treatment. Note that this parameter is different from the Sharp RD parameter  $\tau_{\text{SRD}}$  under perfect compliance,

$$\tau_{\text{SRD}} = \mathbb{E}[Y_i(1) - Y_i(0) | X_i = \bar{x}].$$

The difference between  $\tau_{\text{SRD}}$  and  $\tau_{\text{ITT}}$  is the factor  $D_i(1) - D_i(0)$ , which captures the effect that being above the cutoff (and thus being assigned to the treatment) has on the actual treatment received. Units with  $D_i(1) = 1$  and  $D_i(0) = 0$  are called compliers, because they take the treatment when their score is above the cutoff and refuse the treatment when their score is below the cutoff. In a Fuzzy RD design, some units are non-compliers and thus have  $D_i(1) - D_i(0) \neq 1$ .

To see the relationship between the Sharp RD effect and the ITT RD effect, note that when all units are compliers,  $D_i(1) - D_i(0) = 1 - 0 = 1$  for all  $i$ , we have

$$\tau_{\text{ITT}} = \mathbb{E}[(D_i(1) - D_i(0))(Y_i(1) - Y_i(0)) | X_i = \bar{x}] = \mathbb{E}[1 \cdot (Y_i(1) - Y_i(0)) | X_i = \bar{x}] = \tau_{\text{SRD}}$$

and the effect of the treatment assignment at the cutoff is equivalent to the effect of the treatment at the cutoff. However, when some units are non-compliers, the  $\tau_{\text{ITT}}$  captures the effect of the treatment assignment, which will be in general different from the effect of actually receiving the treatment (this difference is reflected in the factor  $D_i(1) - D_i(0)$  multiplying the difference in potential outcomes).

In the presence of non-compliance, it is also of interest to define the average local (i.e. at the cutoff) effect of being assigned to treatment on receiving the treatment itself. Under standard continuity and regularity conditions, it can be shown that a Sharp RD estimator of the treatment assignment  $T_i$  on the treatment take-up  $D_i$  consistently estimates the quantity

$$\lim_{x \downarrow \bar{x}} \mathbb{E}[D_i | X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[D_i | X_i = x] = \mathbb{E}[D_i(1) - D_i(0) | X_i = \bar{x}] \equiv \tau_{\text{FS}}. \quad (4.2)$$

The parameter  $\tau_{\text{FS}}$ , usually called the *first-stage* effect, captures the average effect at the cutoff of being assigned to the treatment on receiving the treatment. In particular, since we are assuming that  $D_i$  is binary,  $\tau_{\text{FS}}$  captures the difference in the probability of actually receiving the treatment at the cutoff between units assigned to treatment vs. control.

Because both  $\tau_{\text{FS}}$  and  $\tau_{\text{ITT}}$  are Sharp RD parameters, estimation and inference can be performed following standard continuity-based Sharp RD methods—for details, see Part 1. In particular, the analysis uses  $X_i$  as the running variable, the assignment indicator  $T_i = \mathbb{1}(X_i \geq \bar{x})$  as the treatment of interest, and  $D_i$  and  $Y_i$  as outcomes.

Although the parameters  $\tau_{\text{FS}}$  and  $\tau_{\text{ITT}}$  are of genuine interest, they only capture the effect of being assigned to the treatment on the outcomes of interest. In most cases, researchers are also interested in the effect of actually receiving the treatment, not only on the effect of being assigned to receive it. Unfortunately, the average effect of receiving the treatment at the cutoff cannot be obtained in general—the reason is that the individual level “treatment effect” can be related with the decision to comply with the treatment, which makes the random variables  $Y_i(1) - Y_i(0)$  and  $D_i(1) - D_i(0)$  not independent in general.

However, the average treatment effect at the cutoff can be recovered from a Fuzzy RD design under additional assumptions. For example, assuming local independence between  $Y_i(1) - Y_i(0)$  and  $D_i(\mathbb{1}(X_i \geq \bar{x}))$  conditional on  $X_i$  being near the cutoff  $\bar{x}$ , [Hahn et al. \(2001\)](#) show that

$$\lim_{x \downarrow \bar{x}} \mathbb{E}[Y_i | X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[Y_i | X_i = x] = \mathbb{E}[D_i(1) - D_i(0) | X_i = \bar{x}] \cdot \mathbb{E}[Y_i(1) - Y_i(0) | X_i = \bar{x}],$$

which, together with Equation 4.2, implies

$$\frac{\lim_{x \downarrow \bar{x}} \mathbb{E}[Y_i | X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[Y_i | X_i = x]}{\lim_{x \downarrow \bar{x}} \mathbb{E}[D_i | X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[D_i | X_i = x]} = \mathbb{E}[Y_i(1) - Y_i(0) | X_i = \bar{x}]. \quad (4.3)$$

It is also possible to arrive at a related interpretation assuming a monotonicity condition instead of local independence. In this context, the monotonicity condition says that a unit with score  $X_i$  who refuses the treatment when the cutoff is  $\bar{x}$  must also refuse the treatment for any cutoff  $x > \bar{x}$ , and a unit who takes the treatment when the cutoff is  $\bar{x}$  must also take the treatment for any cutoff  $x < \bar{x}$ . Assuming monotonicity and continuity, it can be shown that

$$\frac{\lim_{x \downarrow \bar{x}} \mathbb{E}[Y_i | X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[Y_i | X_i = x]}{\lim_{x \downarrow \bar{x}} \mathbb{E}[D_i | X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[D_i | X_i = x]} = \mathbb{E}[Y_i(1) - Y_i(0) | X_i = \bar{x}, D_i(1) > D_i(0)], \quad (4.4)$$

that is, the ratio on the left-hand-side identifies the average effect of the treatment at the cutoff for the subset of units who are compliers—i.e., for units who do take the treatment when their score is above the cutoff and refuse the treatment when their score is below the cutoff. Following the instrumental variables literature, this is sometimes called the Local Average Treatment Effect (LATE). See [Imbens and Lemieux \(2008\)](#) and [Cattaneo et al. \(2016a\)](#) for further discussion.

Given the above results, it is very common for researchers who analyze a Fuzzy RD design to focus not only on the first stage and intention-to-treat effects  $\tau_{\text{FS}}$  and  $\tau_{\text{ITT}}$ , but also on the parameter

$$\tau_{\text{FRD}} = \frac{\lim_{x \downarrow \bar{x}} \mathbb{E}[Y_i | X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[Y_i | X_i = x]}{\lim_{x \downarrow \bar{x}} \mathbb{E}[D_i | X_i = x] - \lim_{x \uparrow \bar{x}} \mathbb{E}[D_i | X_i = x]},$$

which is the ratio between the average intention-to-treat effect  $\tau_{\text{ITT}}$  and the average effect of the treatment assignment on the treatment take-up,  $\tau_{\text{FS}}$ , both at the cutoff.

## 5 The Multi-Cutoff RD Design

TO BE ADDED.

### 5.1 Empirical Application: To be added

TO BE ADDED.

### 5.2 Taxonomy of Multiple Cutoffs

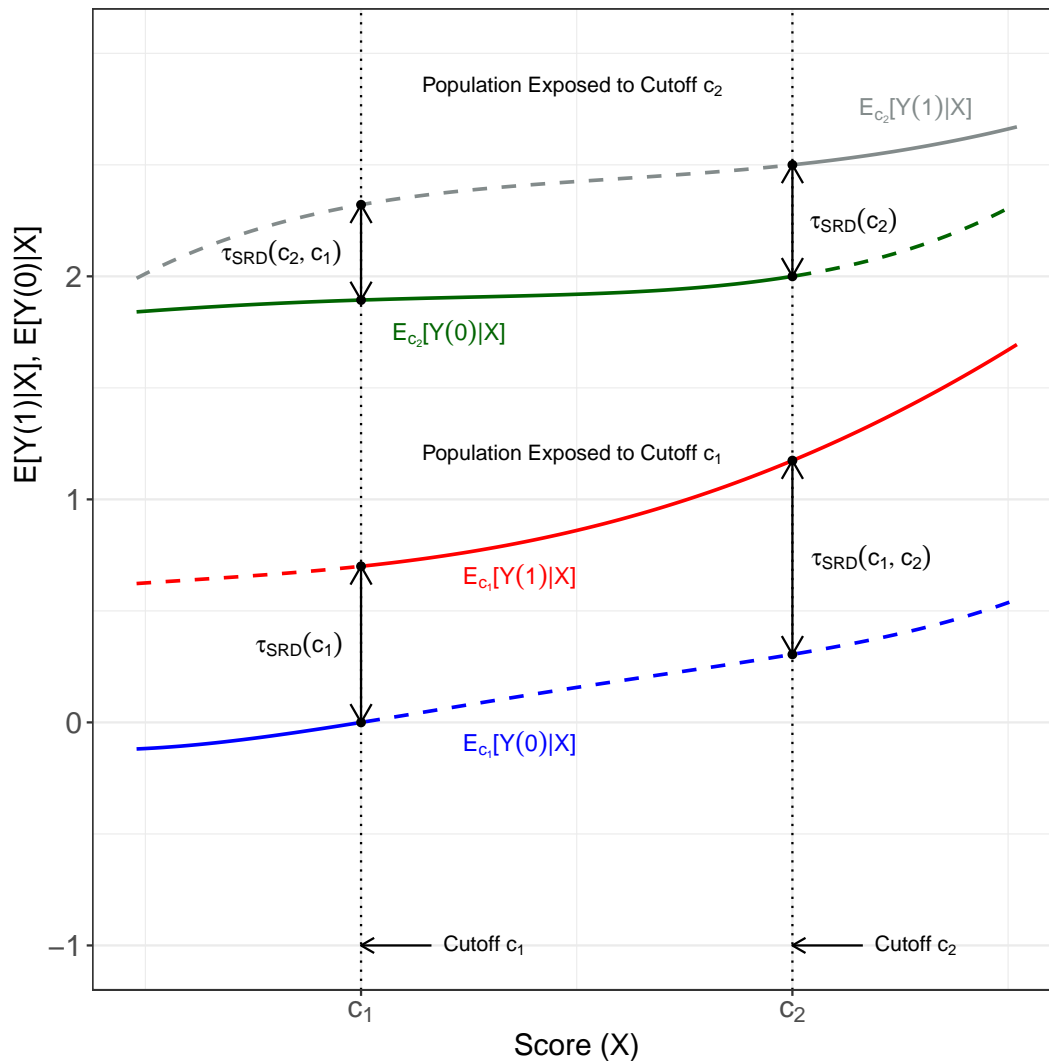
Another generalization of the RD design that is commonly seen in practice occurs when the treatment is assigned using different cutoff values for different subgroups of units. In the standard RD design, all units face the same cutoff value  $\bar{x}$ ; as a consequence, the treatment assignment rule is  $T_i = \mathbb{1}(X_i \geq \bar{x})$  for all units. In contrast, in the Multi-Cutoff RD design different units face different cutoff values.

An example occurs in RD designs where the running variable  $X_i$  is a political party's vote share in a district, and the treatment is winning that district's election. When there are only two parties contesting the election, the cutoff for the party of interest to win the election is always 50%, because the party's strongest opponent always obtains  $(100 - X_i)\%$  of the vote. However, when there are three or more parties contesting the race and the winner is the party with the highest vote share, the party can win the election barely in many different ways. For example, if there are three parties, the party of interest could barely win with 50% of the vote against two opponents who get, respectively, 49% and 1% of the vote; but it could also barely win with 39% of the vote against two opponents who got 38% and 23%. Indeed, in this context, there is an infinite number of ways in which one party can barely win the election—the party just needs to obtain a barely higher vote share than the vote share obtained by its strongest opponent, whatever value the latter takes.

Another common example occurs when a federal program is administered by sub-national units, and each of the units chooses a different cutoff value to determine program eligibility. For example, in order to target households that were most in need in a given locality, the Mexican conditional cash transfer program Progresa determined program eligibility based on a household-level poverty index. In rural areas, the cutoff that determined program eligibility varied geographically, with seven distinct cutoffs used depending on the geographic location of each locality. This type of situation arises in many other contexts where the cutoff for eligibility varies among the units in the analysis.

Cattaneo et al. (2016a) introduced an RD framework based on potential outcomes and continuity conditions to analyzing Multi-Cutoff RD designs, and established a connection with the most common practice of normalizing-and-pooling the information for empirical implementation. Suppose that the cutoff is a random variable  $C_i$ , instead of a known constant, taking on  $J$  distinct values  $\mathcal{C} = \{c_1, c_2, \dots, c_J\}$ . The continuous case is discussed below, though in practice it is often hard to implement RD designs with more than a few cutoff points due to data limitations. In a multi-cutoff RD setting, the treatment assignment is generalized to  $T_i = \mathbb{1}(X_i \geq C_i)$ , where  $C_i$  is a random variable with support  $\mathcal{C}$ . Of course, the single cutoff RD design is contained in this generalization when  $\mathcal{C} = \{\bar{x}\}$  and thus  $\mathbb{P}[C_i = \bar{x}] = 1$ , though more generally  $\mathbb{P}[C_i = c] \in (0, 1)$  for each  $c \in \mathcal{C}$ .

Figure 5.1: RD Design with Multiple Cutoffs



In multi-cutoff RD settings, one approach commonly used in practice is to normalize the

running variable so that all units face the same common cutoff value at zero, and then apply the standard RD design machinery to the normalized score and the common cutoff. To do this, researchers define the normalized score  $\tilde{X}_i = X_i - C_i$ , and pool all observations using the same cutoff of zero for all observations in a standard RD design, with the normalized score used in place of the original score. In this normalizing-and-pooling approach, the treatment assignment indicator is therefore  $T_i = \mathbb{1}(X_i - C_i \geq 0) = \mathbb{1}(\tilde{X}_i \geq 0)$  for all units. In the case of the single-cutoff RD design discussed so far, this normalization is achieved without loss of generality as the interpretation of the estimands remain unchanged; only the score ( $X_i \mapsto \tilde{X}_i$ ) and cutoff ( $\bar{x} \mapsto 0$ ) change.

More generally, the normalize-and-pool strategy employing the score variable  $\tilde{X}_i$ , usually called the normalized (or centered) running variable, changes the parameters already discussed above in an intuitive way: they become weighted averages of RD treatment effects for each cutoff value underlying the original score variable. For example, the Sharp RD treatment effect now is

$$\bar{\tau}_{\text{SRD}} = \mathbb{E}[Y_i(1) - Y_i(0) | \tilde{X}_i = 0] = \sum_{c \in \mathcal{C}} \tau_{\text{SRD}}(c) \omega(c), \quad \omega(c) = \frac{f_{X|C}(c|c) \mathbb{P}[C_i = c]}{\sum_{c \in \mathcal{C}} f_{X|C}(c|c) \mathbb{P}[C_i = c]}$$

with  $\bar{\tau}_{\text{SRD}}$  denoting the normalized-and-pooled sharp RD treatment effect,  $\tau_{\text{SRD}}(c)$  denoting the cutoff-specific sharp RD treatment effect, and  $f_{X|C}(x|c)$  denoting the conditional density of  $X_i|C_i$ . See [Cattaneo et al. \(2016a\)](#) for more details on notation and interpretation, and for analogous results for Fuzzy and Kink RD designs.

From a practical perspective, Multi-Cutoff RD designs can be analyzed as a single-cutoff RD design by either normalizing-and-pooling or by considering each cutoff separately. For example, the first approach maps the Multi-Cutoff RD design to a single sharp/fuzzy/kink RD Design, and thus the discussion in this monograph and/or its extension to fuzzy and kink designs applies directly: under standard assumptions on the normalized score, we have the analogous identification result to the standard Sharp RD design, given by

$$\bar{\tau}_{\text{SRD}} = \lim_{x \downarrow 0} \mathbb{E}[Y_i | \tilde{X}_i = x] - \lim_{x \uparrow 0} \mathbb{E}[Y_i | \tilde{X}_i = x],$$

which implies that estimation and inference for  $\bar{\tau}_{\text{SRD}}$  can proceed in the same way as in the standard Sharp RD design with a single cutoff. Alternatively, by considering each subsample  $C_i = c$  with  $c \in \mathcal{C}$ , the methods discuss in this monograph can be applied directly to each cutoff point, and then collected for further analysis and interpretation under additional regularity conditions. Either way, as mentioned before, we focus exclusively on the practical

aspects of implementing estimation, inference and falsification for the single-cutoff Sharp RD design to conserve space and avoid side discussions.

### 5.3 Local Polynomial Analysis

TO BE ADDED.

### 5.4 Local Randomization Analysis

TO BE ADDED.

### 5.5 Further Readings

TO BE ADDED.

## 6 The Multi-Score RD Design

### 6.1 The General Setup

Yet another generalization of canonical RD designs occurs when two or more running variables are determining a treatment assignment, which by construction induces a multi-cutoff RD design with infinite many cutoffs. For example, a grant or scholarship may be given to students who score above a given cutoff in both a mathematics and a language exam. This leads to two running variables—the student’s score in the mathematics exam and her score in the language exam—and two (possibly different) cutoffs. Another popular example is related to geographic boundaries inducing discontinuous treatment assignments. This type of designs have been studied in [Papay et al. \(2011\)](#), [Reardon and Robinson \(2012\)](#), and [Wong et al. \(2013\)](#) for generic multi-score RD settings, and in [Keele and Titiunik \(2015\)](#) for geographic RD settings.

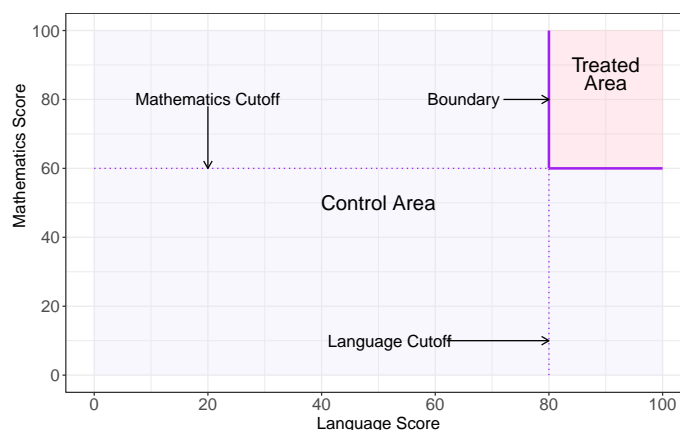
To allow for multiple running variables, we assume each unit’s score is a vector (instead of a scalar as before) denoted by  $\mathbf{X}_i$ . When there are two running variables, the score for unit  $i$  is  $\mathbf{X}_i = (X_{1i}, X_{2i})$ , and the treatment assignment is, for example,  $T_i = \mathbb{1}(X_{1i} > b_1) \cdot \mathbb{1}(X_{2i} > b_2)$  where  $b_1$  and  $b_2$  denote the cutoff points along each of the two dimensions. For simplicity, we assume the potential outcome functions are  $Y_i(1)$  and  $Y_i(0)$ , which implicitly imposes



additional assumptions (e.g., no spill-overs in a geographic setting). See [Cattaneo et al. \(2016a\)](#) for more discussion on this type of restrictions on potential outcomes.

The parameter of interest changes, as discussed before in the context of Multi-Cutoff RD designs, because there is no longer a single cutoff at which the probability of treatment assignment changes discontinuously. Instead, there is a set of values at which the treatment changes discontinuously. To continue our education example, assume that the scholarship is given to all students who score above 60 in the language exam and above 80 in the mathematics exam, letting  $X_{1i}$  denote the language score and  $X_{2i}$  the math score, and  $b_1 = 80$  and  $b_2 = 60$  be the respective cutoffs. According to this hypothetical treatment assignment rule, a student with score  $\mathbf{x}_i = (80, 59.9)$  is assigned to the control condition, since  $\mathbb{1}(80 \geq 80) \cdot \mathbb{1}(59.9 \geq 60) = 1 \cdot 0 = 0$ , and misses the treatment only barely—had she scored an additional 1/10 of a point in the mathematics exam, she would have received the scholarship. Without a doubt, this student is very close to the cutoff criteria for receiving the treatment. However, scoring very close to both cutoffs is not the only way for a student to be barely assigned to treatment or control. A student with a perfect 100 score in language would still be barely assigned to control if he scored 59.9 in the mathematics exam, and a student with a perfect math score would be barely assigned to control if she got 79.9 points the language exam. Thus, with multiple running variables, there is no longer a single cutoff value at which the treatment status of units changes from control to treated. Instead, the discontinuity in the treatment assignment occurs along a boundary of points. This is illustrated graphically in [Figure 6.1](#).

Figure 6.1: Example of RD Design With Multiple Scores: Treated and Control Areas



Consider once again for simplicity a sharp RD design (or an intention-to-treat situation). The parameter of interest in the Multi-Score RD design is therefore a generalization of the standard Sharp RD design parameter, where the average treatment effect is calculated at

all (or, more empirically relevant, at some) points along the boundary between the treated and control areas, that is, at points where the treatment assignment changes discontinuously from zero to one:

$$\mathbb{E}[Y_i(1) - Y_i(0)|\mathbf{X}_i = \mathbf{b}], \quad \mathbf{b} \in \mathcal{B},$$

where  $\mathcal{B}$  denotes the boundary determining the control and treatment areas. For example, in the hypothetical education example in Figure 6.1,  $\mathcal{B} = \{(x_1, x_2) : x_1 = 80 \text{ and } x_2 = 60\}$ .

Although notationally more complicated, conceptually a Multi-Score RD design is very easy to analyze. For example, in the sharp example we are discussing, the identification result is completely analogous to single score case:

$$\tau_{\text{SRD}}(\mathbf{b}) = \lim_{\mathbf{x} \rightarrow \mathbf{b}; \mathbf{x} \in \mathcal{B}_t} \mathbb{E}[Y_i|\mathbf{X}_i = \mathbf{x}] - \lim_{\mathbf{x} \rightarrow \mathbf{b}; \mathbf{x} \in \mathcal{B}_c} \mathbb{E}[Y_i|\mathbf{X}_i = \mathbf{x}], \quad \mathbf{b} \in \mathcal{B},$$

where  $\mathcal{B}_t$  and  $\mathcal{B}_c$  denote the treatment and control areas, respectively. In other words, for each cutoff point along the boundary, the treatment effect at that point is identifiable by the observed bivariate regression functions for each treatment group, just like in the single-score case. The only conceptually important distinction is that Multi-Score RD designs generate a  $\tau_{\text{SRD}}(\mathbf{b})$  family or curve of treatment effects, one for each boundary point  $\mathbf{b} \in \mathcal{B}$ . For example, two potentially distinct sharp RD treatment effects are  $\tau_{\text{SRD}}(80, 70)$  and  $\tau_{\text{SRD}}(90, 60)$ .

## 6.2 The Geographic RD Design

An important special case of the RD design with multiple running variables is the Geographic RD design, where the boundary  $\mathcal{B}$  at which the treatment assignment changes discontinuously is a geographic boundary that separates a geographic treated area from a geographic control area. A typical Geographic RD design is one where the treated and control areas are adjacent administrative units such as counties, districts, municipalities, states, etc., with opposite treatment status. In this case, the boundary at which the treatment status changes discontinuously is the border that separates the adjacent administrative units. For example, some counties in Colorado have all-mail elections where voting can only be conducted by mail and in-person voting is not allowed, while other counties have traditional in-person voting. Where the two types of counties are adjacent, the administrative border between the counties induces a discontinuous treatment assignment between in-person and all-mail voting, and a Geographic RD design can be used to estimate the effect of adopting all-mail elections on voter turnout. This RD design can be formalized as a RD design with two running variables, where the score  $\mathbf{X}_i = (X_{1i}, X_{2i})$  contains two coordinates such as latitude

and longitude that determine the exact geographic location of unit  $i$ . In practice, the score  $\mathbf{X}_i = (X_{1i}, X_{2i})$ —that is, the geographic location of each unit in the study—is obtained using Geographic Information Systems (GIS) software, which allows researchers to locate each unit on a map as well as to locate the entire treated and control areas, and all points on the boundary between them.

For implementation, in both the geographic and non-geographic cases, there are two main approaches mirroring the discussion for the case of Multi-Cutoff RD designs. One approach is the equivalent of normalizing-and-pooling, while the other approach estimates many RD treatment effects along the boundary. For example, consider first the latter approach in a sharp RD context: the RD effect at a given boundary point  $\mathbf{b} = (b_1, b_2) \in \mathcal{B}$  may be obtained by calculating each unit’s distance to  $\mathbf{b}$ , and using this one-dimensional distance as the unit’s running variable, giving negative values to control units and positive values to treated units. Letting the distance between a unit’s score  $\mathbf{X}_i$  and a point  $\mathbf{x}$  be  $d_i(\mathbf{x})$ , we can re-write the above identification result as

$$\tau_{\text{SRD}}(\mathbf{b}) = \lim_{d \uparrow 0} \mathbb{E}[Y_i | d_i(\mathbf{b}) = d] - \lim_{d \downarrow 0} \mathbb{E}[Y_i | d_i(\mathbf{b}) = d], \quad \mathbf{b} \in \mathcal{B}.$$

The choice of distance metric  $d_i(\cdot)$  depends on the particular application. A typical choice is the Euclidean distance  $d_i(\mathbf{b}) = \sqrt{(X_{1i} - b_1)^2 + (X_{2i} - b_2)^2}$ . In practice, this approach is implemented for a finite collection of evaluation points along the boundary, and all the methods and discussion presented in this monograph can be apply to this case directly, one cutoff at the time. The normalizing-and-pooling approach is also straightforward in the case of Multi-Score RD designs, as the approach simply pools together all the units closed to boundary and conducts inference as in a single-cutoff RD design.

As in the previous cases, we do not elaborate on practical issues for this specific setting to conserve space and because all the main methodological recommendations, codes and discussions apply directly. However, to conclude our discussion, we do highlighting an important connection between RD designs with multiple running variables and RD designs with multiple cutoffs. In the Multi-Cutoff RD design, our discussion was based on a discrete set of cutoff points, which would be the natural setting in Multi-Score RD designs applications. In such case, we can map each cutoff point on the boundary to one of the cutoff points in  $\mathcal{C}$  and each observation can be assigned a running variable relative to each cutoff point via the distance function. With these two simple modifications, any Multi-Score RD design can be analyzed as a Multi-Cutoff RD design over finitely many cutoff points on the boundary. In particular, this implies that all the conclusions and references given in the previous section apply to this case as well. See the supplemental appendix of [Cattaneo et al. \(2016a\)](#) for more

discussion on this idea and further generalizations.

### **6.3 Further Readings**

TO BE ADDED.

## **7 Final Remarks**

TO BE ADDED.

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