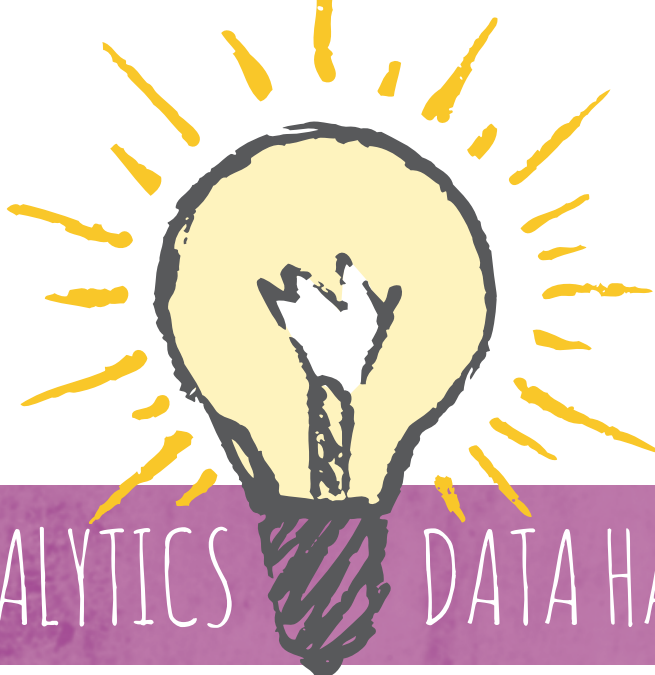


DEVELOPMENT RESEARCH IN PRACTICE



DIME ANALYTICS DATA HANDBOOK



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DEVELOPMENT
RESEARCH
IN PRACTICE:
THE DIME ANALYTICS
DATA HANDBOOK

DIME ANALYTICS

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This version of the book has been revised since its release in June 2019 with feedback from readers and experts. It contains most of the content we plan for the finished version. This book is a living product that is written and maintained publicly. The code and edit history are at: <https://github.com/worldbank/dime-data-handbook>. You can get a PDF copy at: <https://worldbank.github.com/dime-data-handbook>. The website includes updated instructions for providing feedback, as well as notes on updates to the content.

Whether you work with DIME, the World Bank, or another organization or university, we ask that you read the contents of this book critically. We welcome feedback and corrections to improve the book. Please visit <https://worldbank.github.com/dime-data-handbook/feedback> to provide feedback. You can also email us at dimeanalytics@worldbank.org, and we will be very thankful. We hope you enjoy *Development Research in Practice!*

Abbreviations

- 2SLS** – Two-Stage Least Squares
AEA – American Economic Association
CAPI – Computer-Assisted Personal Interviewing
DD or DiD – Differences-in-Differences
DIME – Development Impact Evaluations
DOI – Digital object identifier
eGAP – Evidence in Governance and Politics
EU – European Union
GDPR – Global Data Protection Regulation
HFC – High-Frequency Checks
IPA – Innovations for Poverty Action
IRB – Institutional Review Board
IV – Instrumental Variables
JPAL – The Abdul Lateef Jameel Poverty Action Lab
MDE – Minimum Detectable Effect
NGO – Non-Governmental Organization
ODK – Open Data Kit
OLS – Ordinary Least Squares
OSF – Open Science Foundation
PI – Principal Investigator
PII – Personally-Identifying Information
RA – Research Assistant
RD – Regression Discontinuity
RCT – Randomized Control Trial
SSC – Statistical Software Components

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*Dedicated to all the research assistants
who have wrangled data without being
taught how, hustled to get projects done
on time, wondered if they really should get
their PhD after all, and in doing so made
this knowledge necessary and possible.*

Introduction: Development Research in Practice

Welcome to *Development Research in Practice: The DIME Analytics Data Handbook*. This book is intended to teach all users of development data how to handle data effectively, efficiently, and ethically. An empirical revolution has changed the face of development research rapidly over the last decade. Increasingly, researchers are working not just with complex data, but with *original* data: datasets collected by the research team themselves or acquired through a unique agreement with a project partner. Research teams must carefully document how original data is created, handled, and analyzed. These tasks now contribute as much weight to the quality of the evidence as the research design and the statistical approaches do. At the same time, the scope and scale of empirical research projects is expanding: more people are working on the same data over longer timeframes. For that reason, the central premise of this book is that data work is a “social process”. This means that the many different people on a team need to have the same ideas about what is to be done, and when and where and by whom, so that they can collaborate effectively on a large, long-term research project.

Despite the growing importance of managing data work, little practical guidance is available for practitioners. There are few guides to the conventions, standards, and best practices that are fast becoming a necessity for empirical research. *Development Research in Practice* aims to fill that gap. It covers the full data workflow for a complex research project using original data. In this book, we share the lessons, tools, and processes developed based on DIME’s diverse research experience, and compile them into a single narrative of best practices for data work. This book is not sector-specific; it will not teach you econometrics, or how to design an impact evaluation. There are many excellent existing resources on those topics. Instead, it will teach you how to think about all aspects of your research from a data perspective, how to structure research projects to ensure data quality, and how to institute transparent and reproducible workflows.

The team behind this book

DIME is the Development Impact Evaluations department of the World Bank.¹ Its mission is to generate high-quality and operationally relevant data and research to transform development policy, help reduce extreme poverty, and secure shared prosperity.² DIME develops customized data and evidence ecosystems to produce actionable information and recommend specific policy pathways to maximize impact. The department conduct research in 60 countries with 200 agencies, leveraging a US\$180 million research budget to shape the design and implementation of US\$18 billion in development finance. DIME also provides advisory services to 30 multilat-

¹ <https://www.worldbank.org/en/research/dime>

² Legovini et al. (2015)

eral and bilateral development agencies.³ DIME research is organized into four primary topic pillars: Economic Transformation and Growth; Gender, Economic Opportunity, and Fragility; Governance and Institution Building; and Infrastructure and Climate Change. Over the years, DIME has employed dozens of research economists, and hundreds of full-time research assistants, field coordinators, and other staff. The team has conducted over 325 impact evaluations. *Development Research in Practice* exists to take advantage of that concentration and scale of research, to synthesize many resources for data collection and research, and to make DIME tools available to the larger community of development researchers.

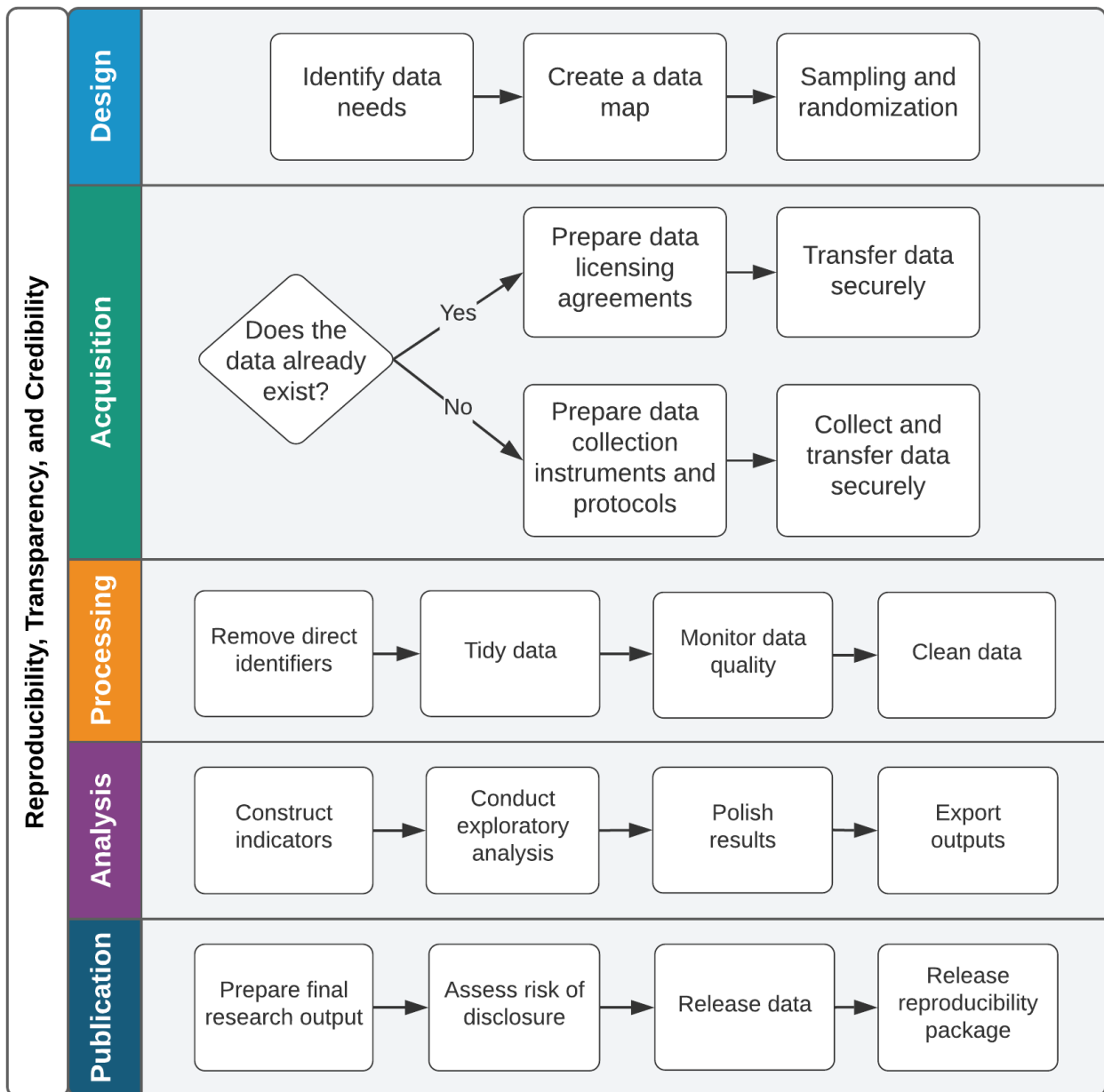
As part of its broader mission, DIME invests in public goods to improve the quality and reproducibility of development research around the world. One key early innovation at DIME was the creation of DIME Analytics, the team responsible for writing and maintaining this book.⁴ DIME Analytics is a centralized unit that develops and ensures adoption of high quality research practices across the department's portfolio. This is done through an intensive, collaborative innovation cycle: DIME Analytics onboards and supports research assistants and field coordinators, provides standard tools and workflows to all teams, delivers hands-on support when new tasks or challenges arise, and then develops and integrates lessons from those engagements to bring to the full team. Resources developed and tested in DIME are converted into public goods for the global research community, through open-access trainings and open-source tools. The DIME Analytics Resource Directory appendix provides an introduction to public materials.

DIME Analytics has invested many hours over the past years learning from data work across DIME's portfolio, identifying inefficiencies and barriers to success, developing tools and trainings, and standardizing best-practice workflows adopted in DIME projects. It has also invested significant energy in the language and materials used to teach these workflows to new team members, and, in many cases, in software tools that support these workflows explicitly. DIME team members often work on diverse portfolios of projects with a wide range of teammates, and we have found that standardizing core processes across all projects results in higher-quality work with fewer opportunities to make mistakes. In that way, the Analytics team is DIME's method of institutionalizing tools and practices, developed and refined over time, that give the department a common base of knowledge and practice. In 2018, for example, DIME adopted universal reproducibility checks conducted by the Analytics team; the lessons from this practice helped move the DIME team from where 50% of submitted papers in 2018 required significant revision to pass

³ Legovini et al. (2019)

⁴ <https://www.worldbank.org/en/research/dime-data-and-analytics>

to where 64% of papers passed in 2019 without revision required.



How to read this book

This book aims to be a highly practical resource so the reader can immediately begin to collaborate more effectively on large, long-term research projects that use the methods and tools discussed.

This introduction outlines the basic philosophies that motivate this book and our approach to research data. We want all readers to understand at the outset our mindset that research data work is primarily about communicating effectively within a team and that standardization and simplification of data tasks is a major enabler of effective collaboration. The main chapters of this book will walk you through the data workflow at each stage of an empirical research project, from design to publication. The figure in this introduction visualizes the data workflow. Chapters 1 and 2 contextualize the workflow, and set the stage for the hands-on data tasks which are described in detail in Chapters 3 to 7.

Chapter 1 outlines a set of practices and ideals to ensure that research consumers can be confident in the conclusions reached, and that the reliability of research work can be verified. It begins with ethical principles to guide empirical research, focusing on research reproducibility, transparency, and credibility. It then introduces tools to document the aims and methods of a research project, ensuring that meta-information about your research is available and that you approach data work with an eye towards the future.

Chapter 2 teaches you to structure your data work for collaborative research, while ensuring the privacy and security of research participants. It discusses the importance of planning datawork and associated tools in advance, long before any data is acquired. It also describes ethical concerns common to development data, common pitfalls in legal and practical management of data, and how to respect the rights of research participants at all stages of data work.

Chapter 3 turns to the measurement framework, and how to translate research design to a data work plan. It details DIME's data map template, a set of tools to communicate the project's data requirements both across the team and across time. It also discusses how to implement random sampling and random assignment in a reproducible and credible manner.

Chapter 4 covers data acquisition. It starts with the legal and institutional frameworks for data ownership and licensing, to ensure that you are aware of the rights and responsibilities of using data collected by the research team or by others. It provides a deep dive on collecting high-quality primary electronic survey data, including developing and deploying survey instruments. Finally, it discusses secure data handling during transfer, sharing, and storage, which is essential in protecting the privacy of respondents in any data.

Chapter 5 describes data processing tasks. It details how to construct "tidy" data at the appropriate units of analysis, how to ensure uniquely identified datasets, and how to routinely incorporate data quality checks into the workflow. It also provides guidance on de-

identification and cleaning of personally-identified data, focusing on how to understand and structure data so that it is ready for indicator construction and analytical work.

Chapter 6 discusses data analysis tasks. It begins with data construction, or the creation of new variables from the raw data acquired or collected in the field. It introduces core principles for writing analytical code and creating, exporting, and storing research outputs such as figures and tables reproducibly using dynamic documents.

Chapter 7 outlines the publication of research outputs, including manuscripts, code, and data. This chapter discusses how to effectively collaborate on technical writing using dynamic documents. It also covers how and why to publish datasets in an accessible, citable, and safe fashion. Finally, it provides guidelines for preparing functional and informative reproducibility packages that contain all the code, data, and meta-information needed for others to evaluate and reproduce your work.

After reading each chapter, you should understand what tasks will be performed at every stage of the workflow, and how to implement them according to best practices. You should also understand how the various stages of the workflow tie together, and what inputs and outputs are required and produced from each. The references and links contained in each chapter will lead you to detailed descriptions of individual ideas, tools, and processes to refer to when you need to implement the tasks yourself. Specific implementation details and links to further practical resources will often be found on the linked pages in the **DIME Wiki**.⁵

Handling original data is a core research task

In the past, data work was often treated as a “black box” in research. A published manuscript might exhaustively detail research designs, estimation strategies, and theoretical frameworks, but typically reserved very little space for detailed descriptions of how data was actually collected and handled. It is almost impossible to assess the quality of the data in such a paper, and whether the results could be reproduced. In the past decade, this has started to change,⁶ in part due to increasing requirements by publishers and funders to release code and data. Data handling and documentation is a key skill for researchers and research staff. Standard processes and documentation practices are important throughout the research process to accurately convey and implement the intended research design.⁷ When data work is done in an ad-hoc manner, it is very difficult for others to understand what is being done – a reader has to simply trust that the researchers did these things right. Most importantly, if any part of

⁵ The DIME Wiki is a free online collection of impact evaluation resources and best practices. <https://dimewiki.worldbank.org>

⁶ Swanson et al. (2020)

⁷ Vilhuber (2020)

the data pipeline breaks down, research results become unreliable⁸ and cannot be faithfully interpreted as being an accurate picture of the intended research design.⁹ Because we almost never have “laboratory” settings¹⁰ in this type of research, such a failure has a very high cost: we will have wasted the investments that were made into knowledge generation, and the research opportunity itself, where we intended to conduct the study.¹¹

Accurate and reproducible data management and analysis is essential to the success and credibility of modern research. Standardizing and documenting data handling processes is essential to be able to evaluate and understand the data work alongside any final research outputs. An important component of this is **process standardization**.¹² Process standardization means that there is little ambiguity about how something ought to be done, and therefore the tools to do it can be set in advance. Standard processes help other people understand your work, and they also make your work easier to document. Process standardization and documentation should allow readers of your code to: (1) quickly understand what a particular process or output is supposed to be doing; (2) evaluate whether or not it does that thing correctly; and (3) modify it either to test alternative hypotheses or to adapt into their own work. This book will discuss specific standards recommended by DIME Analytics, but we are more interested in convincing the reader to discuss the adoption of *a* standard within research teams than to necessarily use *the* particular standards that we recommend.

Documenting data work with standardized code

Modern quantitative research relies heavily on standardized statistical software tools, written with various coding languages, to standardize analytical work. Outputs like regression tables and data visualizations are created using code in statistical software for two primary reasons. The first is that using a standard command or package ensures that the work is done right, and the second is that it ensures the same procedure can be confirmed or checked at a later date or using different data. Keeping a clear, human-readable record of these code and data structures is critical. While it is often *possible* to perform nearly all the relevant tasks through an interactive user interface or even through software such as Excel, this practice is strongly advised against. In the context of statistical analysis, the practice of writing all work using standard code is widely accepted. To support this practice, DIME now maintains portfolio-wide standards about how analytical code should be maintained and made accessible before, during, and after release or publication.

⁸ McCullough et al. (2008)

⁹ <https://blogs.worldbank.org/impactevaluations/more-replication-economics>

¹⁰ See Baldwin and Mvukiyehe (2015) for an example.

¹¹ Camerer et al. (2016)

¹² **Process standardization:** Agreement within a research team about how all tasks of a specific type will be approached.

Over the last few years, DIME has extended the same principles to preparing data for analysis, which often comprises just as much (or more) of the manipulation done to the data over the life cycle of a research project. A major aim of this book is to encourage research teams to think of the tools and processes they use for designing, collecting, and handling data just as they do for analytical tasks. Correspondingly, a major contribution of DIME Analytics has been tools and standard practices for implementing these tasks using statistical software.

While we assume that you are going to do nearly all data work using code, many development researchers come from economics and statistics backgrounds and often understand code to be a means to an end rather than an output itself. We believe that this must change somewhat: in particular, we think that development practitioners must think about their code and programming workflows just as methodologically as they think about their research workflows, and think of code and data as research outputs, just as manuscripts and briefs are.

This approach arises because we see the code as the “recipe” for the analysis. The code tells others exactly what was done, how they can do it again in the future, and provides a roadmap and knowledge base for further original work.¹³ Performing every task through written code creates a record of every task you performed.¹⁴ It also prevents direct interaction with the data files that could lead to non-reproducible processes.¹⁵ Finally, DIME Analytics has invested a lot of time in developing code as a learning tool: the examples we have written and the commands we provide are designed to provide a framework for common practice across the entire DIME team, so that everyone is able to read, review, and provide feedback on the work of others starting from the same basic ideas about how various tasks are done.

Most specific code tools have a learning and adaptation process, meaning you will become most comfortable with each tool only by using it in real-world work. To support your process of learning reproducible tools and workflows, we reference free and open-source tools wherever possible, and point to more detailed instructions when relevant. **Stata**,¹⁶ as a proprietary software, is the notable exception here due to its persistent popularity in development economics and econometrics.¹⁷ This book also includes, in an appendix, the **DIME Analytics Coding Guide** which includes instructions for how to write good code, instructions on how to use the code examples in this book, as well as our Stata Style Guide. DIME projects are strongly encouraged to explicitly adopt and follow coding style guides in their work. Style guides harmonize code within and across

¹³ Hamermesh (2007)

¹⁴ Ozier (2019)

¹⁵ Chang and Li (2015)

¹⁶ StataCorp, LLC (2019)

¹⁷ https://dimewiki.worldbank.org/Software_Tools

teams making it easier to understand and reuse code, which ultimately helps teams to build on each other's best practices. Some of the programming languages used at DIME already have well-established and commonly used style guides, such as the Tidyverse style guide for R and PEP-8 for Python.¹⁸. Stata has relatively few resources of this type available, which is why we have created and included one here that we hope will be an asset to all Stata users.

¹⁸ See DIME Analytics Coding Standards: <https://github.com/worldbank/dime-standards>

Looking forward

While adopting the workflows and mindsets described in this book requires an up-front cost, it will save you (and your collaborators) a lot of time and hassle very quickly. In part this is because you will learn how to implement essential practices directly; in part because you will find tools for the more advanced practices; and most importantly because you will acquire the mindset of doing research with a high-quality data focus. We hope you will find this book helpful for accomplishing all of the above, and that mastery of data helps you make an impact. We hope that by the end of the book, you will have learned how to handle data more efficiently, effectively and ethically at all stages of the research process.

Chapter 1: Reproducibility, transparency, and credibility

Policy decisions are made every day using the results of development research, and these have wide-reaching effects on the lives of millions. As the emphasis on evidence-based policy grows, so too does the scrutiny under which research methods and results are placed. Three major components make up this scrutiny: credibility, transparency, and reproducibility. These three components contribute to one simple idea: research should be high quality and well-documented. Research consumers, including the policy makers who will use the evidence to make decisions, should be able to easily examine and recreate such evidence. In this framework, it is useful to think of research as a public service that requires researchers as a group to be accountable for their methods. This means acting to collectively protect the credibility of development research by following modern practices for research planning and documentation. Across the social sciences, the open science movement has been fueled by concerns about the proliferation of low-quality research practices, data and code that are inaccessible to the public, analytical errors in major research papers, and in some cases even outright fraud. While the development research community has not yet experienced major scandals, it has become clear that there are necessary improvements to make in the way that code and data are handled as part of research. Moreover, having common standards and practices for creating and sharing materials, code, and data with others will improve the value of the work we do.

In this chapter, we outline principles and practices that help to ensure research consumers can be confident in the conclusions reached. We discuss each of the three components – credibility, transparency, and reproducibility – in turn. The first section covers research credibility. It presents three popular methods to commit to particular research questions or methods, and avoid potential criticisms of cherry-picking results: registration, pre-analysis plans, and registered reports. The second section discusses how to apply principles of transparency to all research processes, which allows research teams to be more efficient, and research consumers to fully understand and evaluate research quality. The final section provides guidance on how to make your research fully reproducible, and explains why publishing replication materials is an important research contribution in its own right.

Developing a credible research project

The evidentiary value of research is traditionally a function of design choices,¹ such as powered through sampling and randomization, and robustness to alternative specifications and definitions. One frequent target for critics of such research² is the fact that most researchers have a lot of leeway in selecting their projects, results, or outcomes *after* already having had the experience of implementing a project or

¹ Angrist and Pischke (2010), Ioannidis (2005)

² Ioannidis et al. (2017)

collecting data in the field, which increases the likelihood of finding “false positive” results that are not true outside carefully-selected data.³ Credible research design methods are key to maintaining credibility in these choices and avoiding serious errors. This is especially relevant for research that relies on original data sources, from innovative big data sources to unique surveys. Development researchers should take these concerns seriously. Such flexibility can be a significant issue for the quality of evidence overall, particularly if researchers believe that certain types of results are substantially better for their careers or their publication chances.

This section presents three popular methods for researchers to commit to particular research questions or methods, and to avoid potential criticisms of cherry-picking results for publication: registration, pre-analysis plans, and registered reports. Each of these methods involves documenting specific research design components, ideally before carrying out the analytical component or extensively exploring the data. Study registration provides formal notice that a study is being attempted and creates a hub for materials and updates about the study results. Pre-analysis plans are a more formal commitment to use specific methods on particular questions. Writing and releasing a pre-analysis plan in advance of working with data is used to protect the credibility of approaches that have a high likelihood of producing false results.⁴ Finally, registered reports allow researchers to approach research planning itself as a process at the level of a full peer review. Registered reports enable close scrutiny of a research design, a feedback and improvement process, and a commitment from a publisher to publish the study based on the credibility of the design, rather than the specific results.

Registering research studies

Registration of research studies is an increasingly common practice, and more journals are beginning to require the registration of studies they publish.⁵ Study registration intended to ensure that a complete record of research inquiry is easily available.⁶ Registering research studies ensures that future scholars can quickly find out what work has been carried out on a given question, even if some or all of the work done never results in formal publication. Registration of studies is increasingly required by publishers and can be done before, during, or after the study with essential information about the study purpose. Some currently popular registries are operated by the AEA,⁷ 3ie,⁸ eGAP,⁹ and OSF.¹⁰ They all have different target audiences and features, so select one that is appropriate to your work.

Pre-registering studies before they begin is an extension of this

³ Simmons et al. (2011)

⁴ Wicherts et al. (2016)

⁵ Vilhuber et al. (2020)

⁶ https://dimewiki.worldbank.org/Study_Registration

⁷ <https://www.socialscienceregistry.org>

⁸ <https://ridie.3ieimpact.org>

⁹ <https://egap.org/content/registration>

¹⁰ <https://osf.io/registries>

principle.¹¹ Registration of a study before it goes to implementation or data acquisition, particularly when specific hypotheses are included in the registration, provides a simple and low-effort way for researchers to conclusively demonstrate that a particular line of inquiry was not generated by the process of data collection or analysis itself.¹² Pre-registrations need not provide exhaustive details about how a particular hypothesis will be approached; only that it will be. This can be highly valuable for the credibility of the research and requires only minor time investment or administrative effort. For this reason, the DIME team requires pre-registration of all studies in a public database with at least some primary hypotheses prespecified, prior to providing funding for impact evaluation research.

¹¹ Nosek et al. (2018)

¹² <https://dimewiki.worldbank.org/Pre-Registration>

Writing pre-analysis plans

If a research team has a large amount of flexibility to define how they approach a particular hypothesis, study registration may not be sufficient to avoid the criticism of “hypothesizing after the results are known”, or HARKing.¹³ Examples of such flexibility include a broad range of concrete measures that could be argued to measure to an abstract concept; future choices about sample inclusion or exclusion; or decisions about how to construct derived indicators. When the researcher is collecting a large amount of information and has leverage over even a moderate number of these options, it is almost guaranteed that they can come up with any result they like.¹⁴

Pre-analysis plans (PAPs) can be used to assuage these concerns by specifying some set of analyses the researchers intend to conduct.¹⁵ The pre-analysis plan should be written up in detail for areas that are known to provide a large amount of leeway for researchers to make later decisions, particularly for things like interaction effects or subgroup analysis.¹⁶ Pre-analysis plans should not, however, be viewed as binding the researcher’s hands.¹⁷ Depending on what is known about the study at the time of writing, pre-analysis plans can vary widely in the amount of detail they should include.¹⁸ The core function of a PAP is to carefully and explicitly describe one or more specific data-driven inquiries, as specific formulations are often very hard to justify in retrospect with data or projects that potentially provide many avenues to approach a single theoretical question.¹⁹ Anything outside the original plan is just as interesting and valuable as it would have been if the the plan was never published; but having pre-committed to the details of a particular inquiry makes its results immune to a wide range of criticisms of specification searching or multiple testing.²⁰

¹³ Kerr (1998)

¹⁴ Gelman and Loken (2013)

¹⁵ https://dimewiki.worldbank.org/Pre-Analysis_Plan

¹⁶ See Cusolito et al. (2018) for an example.

¹⁷ Olken (2015)

¹⁸ <https://blogs.worldbank.org/impactevaluations/pre-analysis-plans-and-registered-reports-what-new-opinion-piece-does-and-doesnt>

¹⁹ See Bedoya et al. (2019) for an example.

²⁰ Duflo et al. (2020)

Publishing registered reports

Registered reports take the process of pre-specifying a complex research design to the level of a formal publication.²¹ In a registered report, a journal or other publisher will peer review and conditionally accept a specific study for publication, typically then guaranteeing the acceptance of a later publication that carries out the analysis described in the registered report. While far stricter and more complex to carry out than ordinary study registration or pre-analysis planning, the registered report has the added benefit of peer review and expert feedback on the design and structure of the proposed study.²²

This process is in part meant to combat the “file-drawer problem”²³ and ensure that researchers are transparent in the sense that all promised results obtained from registered-report studies are actually published. This approach has the advantage of pre-specifying in great detail a complete research and analytical design, and securing a commitment for publication regardless of the outcome. This may be of special interest for researchers studying events or programs where either there is a substantial risk that they would either not be able to publish a null or negative result,²⁴ or where they may wish to avoid any pressure toward finding a particular result, for example when the program or event is the subject of substantial social or political pressures. As with pre-registration and pre-analysis, nothing in a registered report should be understood to prevent a researcher from pursuing additional avenues of inquiry once the study is complete, either in the same or separate research outputs.

²¹ https://dimewiki.worldbank.org/Registered_Reports

²² <https://blogs.worldbank.org/impactevaluations/registered-reports-piloting-pre-results-review-process-journal-development-economics>

²³ Simonsohn et al. (2014)

²⁴ See Coville et al. (2019) for an example.

Conducting research transparently

Transparent research exposes not only the code, but all research processes involved in developing the analytical approach. This means that readers are able to judge for themselves whether the research was done well and the decision-making process was sound. If the research is well-structured, and all of the relevant documentation²⁵ is shared, it is easy for the reader to understand the analysis fully. Researchers that expect process transparency also have an incentive to make better decisions, be skeptical and thorough about their assumptions, and save themselves time, because transparent research methods are labor-saving over the complete course of a project.

Clearly documenting research work is necessary to allow others to evaluate exactly what data was acquired and how it was used to obtain a particular result. Many development research projects are purpose-built to address specific questions, and often use unique data, novel methods, or small samples. These approaches can yield

²⁵ https://dimewiki.worldbank.org/Research_Documentation

new insights into essential academic questions, but need to be transparently documented so they can be reviewed or replicated by others in the future.²⁶ Unlike disciplines where data is more standardized or where research is more oriented around secondary data, the exact data used in a development project has often not been observed by anyone else in the past and may not be able to be re-collected by others in the future. Regardless of the novelty of study data, transparent documentation methods help ensure that data was collected and handled appropriately and that studies and interventions were implemented correctly. As with study registrations, project and data documentation should be released on external **archival repositories**²⁷ so they can always be accessed and verified.

Documenting data acquisition and analysis

Documenting a project in detail greatly increases transparency. Many disciplines have a tradition of keeping a “lab notebook”, and adapting and expanding this process to create a lab-style workflow in the development field is a critical step towards more transparent practices. This means explicitly noting decisions as they are made, and explaining the process behind the decision-making. Careful documentation will also save the research team a lot of time during a project, as it prevents you from having the same discussion twice (or more!), since you have a record of why something was done in a particular way. There are a number of available tools that will contribute to producing documentation, but project documentation should always be an active and ongoing process, not a one-time requirement or retrospective task. New decisions are always being made as the plan begins contact with reality, and there is nothing wrong with sensible adaptation so long as it is recorded and disclosed.

Email, however, is *not* a documentation service, because communications are rarely well-ordered, can be easily deleted, and are not available for future team members. There are various software solutions for building proper documentation over time. Some work better for field records such as implementation decisions, research design, and survey development; others work better for recording data work and code development. The **Open Science Framework**²⁸ provides one such solution, with integrated file storage, version histories, and collaborative wiki pages. **GitHub**²⁹ provides a transparent documentation system through commit messages, issues, README files, and pull requests,³⁰ in addition to version histories and wiki pages. Such services offer multiple different ways to record the decision process leading to changes and additions, track and register discussions, and manage tasks. These are flexible tools that can be adapted to different

²⁶ Duvendack et al. (2017)

²⁷ **Archival repository:** A third-party service for information storage that guarantees the permanent availability of current and prior versions of materials.

²⁸ <https://osf.io>

²⁹ <https://github.com>

³⁰ https://dimewiki.worldbank.org/Getting_started_with_GitHub

team and project dynamics. Services that log your research process can show things like modifications made in response to referee comments, by having tagged version histories at each major revision. They also allow you to use issue trackers to document the research paths and questions you may have tried to answer as a resource to others who have similar questions. Each project has specific requirements for data, code, and documentation management, and the exact transparency tools to use will depend on the team's needs, but they should be agreed upon prior to project launch. This way, you can start building a project's documentation as soon as you start making decisions.

Cataloging and archiving data

Data and data collection methods should be fully cataloged, archived, and documented, whether you are collecting data yourself or receiving it from an outside partner. In some cases this is as simple as uploading a survey instrument or an index of datasets and a code-book to an archive. In other cases this will be more complex. Proper documentation of data collection will often require a detailed description of the overall sampling procedure.³¹ For example, settings with many overlapping strata, treatment arms, excluded observations, or resampling protocols might require extensive additional field work documentation. This documentation should be continuously updated and kept with the other study materials; it is often necessary to collate these materials for an appendix for publication in any case.

When data is received from partners or collected in the field, the **original data** (including corrections)³² should be immediately placed in a secure permanent storage system. Before analytical work begins, you should create a “for-publication” copy of the original dataset by removing potentially identifying information. This will become the raw data, and must be placed in an archival repository where it can be cited.³³ This can initially be done under embargo or with limited release, in order to protect your data and future work. This type of data depositing or archiving precedes publishing or releasing any data: data at this stage may still need to be embargoed or have other, potentially permanent, access restrictions, so you can instruct the archive to formally release the data later.

Some project funders provide specific repositories in which they require the deposit of data they funded,³⁴ and you should take advantage of these when possible. If this is not provided, you must be aware of privacy issues with directly identifying data and questions of data ownership before uploading raw data to any third-party server, whether public or not; this is a legal question for your home

³¹ See Yishay et al. (2016) for an example.

³² **Original data:** A new dataset, as obtained and corrected, that becomes the functional basis for research work.

³³ Vilhuber et al. (2020)

³⁴ For example, <https://data.usaid.gov>

organization. Regardless of these considerations, all data repositories, such as DIME's standard, the World Bank Microdata Library³⁵ and the World Bank Data Catalog,³⁶ should create a record of the data's existence and provide instructions on how access might be obtained by another researcher. For more on the steps required to prepare and publish a de-identified dataset, you can refer to Chapter 6 and Chapter 7 of this book. Data publication should create a data citation and a **digital object identifier (DOI)**,³⁷ or some other persistent index that you can use in your future work to unambiguously indicate the location of your data. This data publication should also include the methodological documentation as well as complete human-readable codebooks for all the variables there.

³⁵ <https://microdata.worldbank.org>

³⁶ <https://datacatalog.worldbank.org>

³⁷ **Digital object identifier (DOI):** A permanent reference for electronic information that persistently updates to a new URL or other locations if the information is relocated.

Analyzing data reproducibly

Reproducible research makes it easy for others to apply your techniques to new data or to implement a similar research design in a different context. Development research is rapidly moving in the direction of requiring adherence to specific reproducibility guidelines.³⁸ Major publishers and funders, most notably the American Economic Association, have taken steps to require that code and data are accurately reported, cited, and preserved as research outputs that can be accessed and verified by others. Making research reproducible in this way is a public good.³⁹ It enables other researchers to re-use code and processes to do their own work more easily and effectively in the future. Regardless of what is formally required, your code should be written neatly with clear instructions. It should be easy to read and understand. The corresponding analysis data should also be made accessible to the greatest legal and ethical extent that it can be.⁴⁰

Common research standards from journals and funders feature both regulation and verification policies.⁴¹ Regulation policies require that authors provide reproducibility packages before publication which are then reviewed by the journal for completeness. Verification policies require that authors make certain materials available to the public, but their completeness is not a precondition for publication. Other journals have adopted guidance that offer checklists for reporting on whether and how various practices were implemented, without specifically requiring any.⁴² If you are personally or professionally motivated by citations, producing these kinds of resources can lead to that as well.

³⁸ Christensen and Miguel (2018)

³⁹ https://dimewiki.worldbank.org/Reproducible_Research

⁴⁰ https://dimewiki.worldbank.org/Publishing_Data

⁴¹ Stodden et al. (2013)

⁴² Nosek et al. (2015)

Preparing a reproducibility package

At DIME, all research outputs are required to satisfy **computational reproducibility**,⁴³ which is an increasingly common requirement for publication.⁴⁴ Before releasing a working paper, the research team submits a **reproducibility package** with de-identified data, and DIME Analytics verifies that the package produces exactly the same results that appear in the paper.⁴⁵ The team also comments on whether the package includes sufficient documentation. The Analytics team organizes frequent peer code review for works in progress, and our general recommendation is to ensure that projects are *always* externally reproducible instead of waiting until the final stages to prepare this material. Once the computational reproducibility check is complete, the team receives a completed reproducibility certificate that also lists any publicly available materials to accompany the package, for use as an appendix to the publication. The team also organizes regular peer code review for works in progress, and our general recommendation is to ensure that projects are *always* externally reproducible instead of waiting until the final stages to prepare this material. In this way, code is continuously maintained with clear documentation, and should be easy to read and understand in terms of structure, style, and syntax.

For research to be reproducible, all code files for data cleaning, construction and analysis should be public, unless they contain confidential information. Nobody should have to guess what exactly comprises a given index, or what controls are included in your main regression, or whether or not you clustered standard errors correctly. That is, as a purely technical matter, nobody should have to “just trust you”, nor should they have to bother you to find out what would happen if any or all of these things were to be done slightly differently.⁴⁶ Letting people play around with your data and code is a great way to have new questions asked and answered based on the valuable work you have already done.⁴⁷

A reproducibility package should include the complete materials needed to exactly re-create your final analysis, and be accessible and well-documented so that others can identify and adjust potential decision points that they are interested in. They should be able to easily identify: what data was used and how that data can be accessed; what code generates each table, figure and in-text number; how key outcomes are constructed; and how all project results can be reproduced. A well-organized reproducibility package usually takes the form of a complete directory structure, including documentation and a master script, that leads the reader through the process and rationale for the code behind each of the outputs when considered in

⁴³ **Computational reproducibility:** The ability of another individual to reuse the same code and data and obtain the exact same results as yours.

⁴⁴ https://dimewiki.worldbank.org/Reproducible_Research

⁴⁵ <https://blogs.worldbank.org/impactevaluations/what-development-economists-talk-about-when-they-talk-about-reproducibility>

⁴⁶ Simonsohn et al. (2015)

⁴⁷ <https://blogs.worldbank.org/opendata/making-analytics-reusable>

combination with the corresponding publication.

Looking ahead

With the ongoing rise of empirical research and increased public scrutiny of scientific evidence, making analysis code and data available is necessary but not sufficient to guarantee that findings will be credible. Even if your methods are highly precise, your evidence is only as good as your data – and there are plenty of mistakes that can be made between establishing a design and generating final results that would compromise its conclusions. That is why transparency is key for research credibility. It allows other researchers, and research consumers, to verify the steps to a conclusion by themselves, and decide whether their standards for accepting a finding as evidence are met. Every investment you make in documentation and transparency up front protects your project down the line, particularly as these standards continue to tighten. With these principles in mind, the approach we take to the development, structure, and documentation of data work provides a system to implementing these ideas in everyday work. In the next chapter, we will discuss the workspace you need in order to work reproducibly in an efficient, organized, and secure manner.

Chapter 2: Setting the stage for collaboration

In order to do effective data work in a team environment, you need to structure your workflow in advance. Preparation for collaborative data work begins long before you acquire any data, and involves planning both software tools and collaboration platforms for your team. This means knowing what types of data you'll acquire, whether the data will require special handling due to size or privacy considerations, which datasets and outputs you will need at the end of the process, and how all data files and versions will stay organized throughout. It's important to plan data workflows in advance because changing software or protocols halfway through a project is costly and time-consuming. Seemingly small decisions such as file-sharing services, folder structures, and filenames can be extremely painful to alter down the line in any project.

This chapter will guide you in setting up an effective environment for collaborative data work, structuring your data work to be well-organized and clearly documented, and setting up processes to handle confidential data securely. The first section outlines hows to set up your working environment to effectively collaborate on technical tasks with others, and how to document tasks and decisions. The second section discusses how to organize your code and data so that others will be able to understand and interact with it easily. The third section provides guidelines for ensuring privacy and security when working with confidential data.

Preparing a collaborative work environment

This section introduces core concepts and tools for organizing data work in an efficient, collaborative and reproducible manner. Some of these skills may seem elementary, but thinking about simple things from a workflow perspective can help you make marginal improvements every day you work; those add up to substantial gains over the course of multiple years and projects. Together, these processes form a collaborative workflow that will greatly accelerate your team's ability to get tasks done on all your projects.

Teams often develop workflows in an ad hoc fashion, solving new challenges as they arise. Adaptation is good, of course. But it is important to recognize that there are a number of tasks that exist in common for every project, and it is more efficient to agree on the corresponding workflows in advance. For example, every project requires research documentation, organized file naming, directory organization, coding standards, version control, and code review.

These tasks are common to almost every project, and their solutions translate well between projects. Therefore, there are large efficiency gains to thinking in advance about the best way to do these tasks, instead of throwing together a solution when the task arises. This section outlines the main points to discuss within the team, and suggests best practice solutions for these tasks.

Setting up your computer

First things first: almost all your data work will be done on your computer, so make sure it's set up for success. The operating system should be fully updated, it should be in good working order, and you should have a **password-protected** login. Make sure your computer is backed up to prevent information loss. Follow the **3-2-1 rule**: maintain 3 copies of all original or irreplaceable data, on at least 2 different hardware devices you have access to, with 1 offsite storage method.¹ Chapter 4 provides a protocol for implementing this.

Ensure you know how to get the **absolute file path** for any given file. Using the absolute file path, starting from the filesystem root, means that the computer will never accidentally load the wrong file. On MacOS this will be something like /users/username/git/project/..., and on Windows, C:/users/username/git/project/.... Use forward slashes (/) in file paths for folders, and whenever possible use only the 26 English characters, numbers, dashes (-), and underscores (_) in folder names and filenames.² For emphasis: *always* use forward slashes (/) in file paths in code, just like in internet addresses. Do this even if you are using a Windows machine where both forward and backward slashes are allowed, as your code will otherwise break if anyone tries to run it on a Mac or Linux machine. Making the structure of your directories a core part of your workflow is very important, since otherwise you will not be able to reliably transfer the instructions for replicating or carrying out your analytical work.

When you are working with others, you will most likely be using some kind of **file sharing** software.³ The exact services you use will depend on your tasks, but in general, there are several approaches to file sharing, and the three discussed here are the most common. **File syncing** is the most familiar method, and is implemented by software like Dropbox. Sync forces everyone to have the same version of every file at the same time, which makes simultaneous editing difficult but other tasks easier. **Distributed version control** is another method, commonly implemented through systems like GitHub, GitLab, and Bitbucket that interact with Git.⁴ Distributed version control allows everyone to access different versions of files at the same time. It

¹ [https://dimewiki.worldbank.org/
Data_Storage](https://dimewiki.worldbank.org/Data_Storage)

² [https://dimewiki.worldbank.org/
Naming_Conventions](https://dimewiki.worldbank.org/Naming_Conventions)

³ [https://dimewiki.worldbank.org/
Collaboration_Tools](https://dimewiki.worldbank.org/Collaboration_Tools)

⁴ **Git:** A distributed version control system for collaborating on and tracking changes to code as it is written.

is only optimized for specific types of files (for example, any type of code files). Finally, **server storage** is the least-common method, because there is only one version of the materials, and simultaneous access must be carefully regulated. Server storage ensures that everyone has access to exactly the same files and environment, and it also enables high-powered computing processes for large and complex data.

All three file sharing methods are used for collaborative workflows, and you should review the types of data work that you will be doing, and plan which types of files will live in which types of sharing services. It is important to note that they are, in general, not interoperable, meaning you should not have version-controlled files inside a syncing service, or vice versa, without setting up complex workarounds, and you cannot shift files between them without losing historical information. Therefore, choosing the correct sharing service for each of your team's needs at the outset is essential. At DIME we typically use file syncing for all project administrative files and data, version control in git for code, and server storage for backup and/or large scale computations when needed.

Documenting decisions and tasks

Once your technical and sharing workspace is set up, you need to decide how you are going to communicate with your team. The first habit that many teams need to break is using instant communication for management and documentation. Email is, simply put, not a system. It is not a system for anything. Neither is instant messaging apps like WhatsApp. Instant messaging tools are developed for communicating "now" and that is what they do well. They are not structured to manage group membership or to present the same information across a group of people, or to remind you when old information becomes relevant. They are not structured to allow people to collaborate over a long time or to review old discussions. It is therefore easy to miss or lose communications from the past when they have relevance in the present. Everything with future relevance that is communicated over email or any other instant medium – such as, for example, decisions about research design – should immediately be recorded in a system that is designed to keep permanent records. We call these systems collaboration tools, and there are several that are very useful.⁵

Good collaboration tools are workflow-oriented systems that allow the team to create and assign tasks, carry out discussions related to single tasks, track task progress across time, and quickly see the overall project status. They are web-based so that everyone on your team

⁵ https://dimewiki.worldbank.org/Collaboration_Tools

can access them simultaneously and have ongoing discussions about tasks and processes. Such systems link communications to specific tasks so that related decisions are permanently recorded and easy to find in the future when questions about that task come up. Choosing the right tool for your team's needs is essential to designing an effective workflow. What is important is that your team chooses a system and commits to using it, so that decisions, discussions, and tasks are easily reviewable long after they are completed.

Some popular and free collaboration tools that meet these criteria are GitHub and Dropbox Paper. Any specific list of software will quickly be outdated; we mention these as examples that have worked for our team. Different collaboration tools can be used different types of tasks. Our team, for example, uses GitHub for code-related tasks, and Dropbox Paper for more managerial tasks. GitHub creates incentives for writing down why changes were made in response to specific discussions as they are completed, creating naturally documented code. It is useful also because tasks in GitHub Issues can clearly be tied to file versions. On the other hand, Dropbox Paper provides a clean interface with task notifications, assignments, and deadlines, and is very intuitive for people with non-technical backgrounds. Therefore, it is a useful tool for managing non-code-related tasks.

Setting up a productive code environment

Taking time early in your project to choose a programming language to work in and setting up a productive code environment for that language will make your work significantly easier. Setting up a productive code environment means to make sure that the programming language and all other software your code requires will run smoothly on all the hardware you need it to run on. It also means that you have a productive way of interacting with code, and that the code has a seamless method to access your data.

It is difficult and costly to switch programming languages halfway through a project, so think ahead about the various software your team will use. Take into account the technical abilities of team members, what type of internet access the software will need, the type of data you will need to access, and the level of security required. Big datasets require additional infrastructure and may overburden the tools commonly used for small datasets. Also consider the cost of licenses, the time to learn new tools, and the stability of the tools. There are few strictly right or wrong choices for software, but what is important is that you plan in advance and understand how the chosen tools will interact with your workflows.

One option is to hold your code environment constant over the lifecycle of a single project. While this means you will inevitably have different projects with different code environments, each successive project will be better than the last, and you will avoid the costly process of migrating an ongoing project into a new code environment. Code environments should be documented as precisely as possible. The specific version number of the programming languages and the individual packages you use should be referenced or maintained so that they can be reproduced going forward, even if different releases contain changes that would break your code or change your results. DIME Analytics developed the command `ieboilststart` in the `ietoolkit` package to support version and settings stability in Stata.⁶ If your project requires more than one programming language, for example if you analyze your data in one language but visualize your results in another, then make sure to make an as clear division between the two as possible. This means that you first complete all tasks done in one language, before the completing the rest of the tasks in the other language. Frequently swapping back and forth between languages is a reproducibility nightmare.

Next, think about how and where you write and execute code. This book is intended to be agnostic to the size or origin of your data, but we are going to broadly assume that you are using one of the two most popular statistical software packages: R or Stata. (If you are using another language, like Python, many of the same principles apply but the specifics will be different.) Most of your code work will be done in a code editor. If you are working in R, **RStudio** is the typical choice.⁷ For Stata, the built-in do-file editor is the most widely adopted code editor. You might also consider using an external editor for your R or Stata code.⁸ These editors offer great accessibility and quality features. For example, they can access an entire directory – rather than a single file – which gives you directory-level views and file management actions, such as folder management, Git integration, and simultaneous work with other types of files, without leaving the editor. Using an external editor can also be preferable since your editor will not crash if the execution of your code causes your statistical software to crash.

⁶ <https://dimewiki.worldbank.org/ieboilststart>

⁷ <https://www.rstudio.com>

⁸ https://dimewiki.worldbank.org/Code_Editors

Organizing code and data

We assume you are going to do your analytical work through code, and that you want all your processes to be documented and replicable. Though it is possible to interact with some statistical software through the user interface without writing any code, we strongly advise against it. Writing code creates a record of every task you

performed. It also prevents direct interaction with the data files that could lead to non-reproducible steps. You may do some exploratory tasks by point-and-click or typing directly into the console, but anything that is included in a research output must be coded in an organized fashion so that you can release the exact code that produces your final results – up to and including individual statistics in text. Still, organizing code and data into files and folders is not a trivial task. What is intuitive to one person rarely comes naturally to another, and searching for files and folders is everybody's least favorite task. As often as not, you come up with the wrong one, and then it becomes very easy to create problems that require complex resolutions later. This section provides basic tips on managing the folder that stores your project's data work.

Maintaining an organized file structure for data work is the best way to ensure that you, your teammates, and others are able to easily edit and replicate your work in the future. It also ensures that automated processes from code and scripting tools are able to interact well with your work, whether they are yours or those of others. File organization makes data work easier as well as more transparent, and facilitates integration with tools like version control systems that aim to cut down on the amount of repeated tasks you have to perform. It is worth thinking in advance about how to store, name, and organize the different types of files you will be working with, so that there is no confusion down the line and everyone has the same expectations.

Organizing files and folder structures

Once you start a research project, the number of scripts, datasets, and outputs that you have to manage will grow very quickly. This can get out of hand just as quickly, so it's important to organize your data work and follow best practices from the beginning. You should agree with your team on a specific directory structure, and set it up at the beginning of the project. You should also agree on a file naming convention.⁹ This will help you to easily find project files and ensure that all team members can easily run the same code.

To support consistent folder organization at DIME, DIME Analytics created `iefolder`¹⁰ as a part of our `ietoolkit` package. This Stata command sets up a pre-standardized folder structure for what we call the `DataWork` folder.¹¹ The `DataWork` folder includes folders for all the steps of a typical project. Since each project will always have unique needs, we have tried to make the structure easy to adapt. Having a universally standardized folder structure across the entire portfolio of projects means that everyone can easily move between

⁹ [https://dimewiki.worldbank.org/
Naming_Conventions](https://dimewiki.worldbank.org/Naming_Conventions)

¹⁰ [https://dimewiki.worldbank.org/
iefolder](https://dimewiki.worldbank.org/iefolder)

¹¹ [https://dimewiki.worldbank.org/
DataWork_Folder](https://dimewiki.worldbank.org/DataWork_Folder)

projects without having to reorient on file and folder organization.

If you do not already have a standard file structure across projects, `iefolder` is an easy template to start from. The command creates a `DataWork` folder at the project level, and within that folder, creates standardized directory structures for each data source or survey round. Within each subdirectory, `iefolder` creates folders for raw encrypted data, raw deidentified data, cleaned data, final data, outputs, and documentation. It creates a parallel folder structure for the code files that move the data through this progression, and for the final analytical outputs. The `ietoolkit` package also includes the `iegitaddmd` command, which can place `README.md` placeholder files in your folders so that your folder structure can be shared using Git.¹² Since these placeholder files are written in a plaintext language called **Markdown**, they also provide an easy way to document the contents of every folder in the structure.

The `DataWork` folder may be created either inside an existing project folder, or it may be created as a separate root folder. We advise keeping project management materials (such as contracts, terms of reference, briefs and other administrative or management work) separate from the `DataWork` folder structure. It is useful to maintain project management materials in a sync system like Dropbox, whereas the code folder should be maintained in a version-control system like Git. However, a version-controlled folder should *never* be stored in a synced folder that is shared with other people. Combining these two types of collaboration tools almost always creates undesired functionalities.

¹² Git only tracks files, so empty folders – which most folders are in the beginning of a project – are ignored if placeholder files are not used, leading to only parts of the folder structure being shared across the team.

Connecting tasks through files

Each task in the research workflow has specific inputs and outputs which feed into one another. It is common, particularly when different tasks are performed by different people inside a team, for incompatibilities to be created. For example, if the Principal Investigators are writing a paper using `LATEX`, exporting tables from statistical software into a `.csv` format will break the workflow. Therefore, it's important to agree with your team on what tools will be used for what tasks, and where inputs and outputs will be stored, before you start creating them. Take into account ease of use for different team members, and keep in mind that learning how to use a new tool may require some time investment upfront that will be paid off as your project advances.

Knowing how code outputs will be used will help you decide the best format to export them. You can typically use the same software to save figures into various formats, such as `.eps`, `.png`, `.pdf` or `.jpg`.

However, the decision between using Office Suite software such as Word and PowerPoint versus L^AT_EX and other plain text formats may influence how you write your code, as this choice often implicates in the use of a particular format. This decision will also affect the version control systems that your team can use.

Setting up a version control system

We recommend using a **version control system** to maintain control of file history and functionality. A good version control system tracks who edited each file and when, allows you to revert to previous versions, and provides a protocol for ensuring that conflicting versions are avoided. This is important, for example, for your team to be able to find the version of a presentation that you delivered to a donor, or to understand why the significance level of your estimates has changed. Everyone who has ever encountered a file named something like `final_report_v5_LJK_KLE_jun15.docx` can appreciate how useful such a system can be.

Most syncing services offer some kind of rudimentary version control; these are usually enough to manage changes to binary files (such as office documents) without needing to rely on dreaded filename-based versioning conventions. For code files, however, a more detailed version control system is usually desirable. We recommend using Git for version-control of all data work. Git documents changes to all **plaintext** files. Plaintext files include all code files, most raw outputs, and written outputs that use code languages, such as L^AT_EX files and many dynamic documents. Git tracks all the changes you make to each plaintext file, and allows you to go back to previous versions without losing the information on changes made. It also makes it possible to work on multiple parallel versions of a file, so you don't risk breaking code for other team members as you try something new.

Documenting and organizing code

Good code is written in a way that is easily understood and run by others. Below we discuss a few crucial steps to code organization. They all come from the principle that code is an output by itself, not just a means to an end, and should be written thinking of how easy it will be for someone to read it later. At the end of this section, we include a template for a master script do-file in Stata, to provide a concrete example of the required elements and structure. Throughout this section, we refer to lines of this example do-file to give concrete examples of the required code elements, organization and structure.

To be readable, code must be well-documented. Start by adding

a code header to every file. A code header is a long **comment**¹³ that details the functionality of the entire script; refer to lines 5-10 in the example do-file. This should include simple things such as the purpose of the script and the name of the person who wrote it. If you are using a version control software, the last time a modification was made and the person who made it will be recorded by that software. Otherwise, you should include it in the header. You should always track the inputs and outputs of the script, as well as the uniquely identifying variable; refer to lines 52-54 in the example do-file. When you are trying to track down which code creates which dataset, this will be very helpful. While there are other ways to document decisions related to creating code, the information that is relevant to understand the code should always be written in the code file.

Two types of comments should be included in the script itself. The first type of comment describes what is being done. This might be easy to understand from the code itself if you know the language well enough and the code is clear, but often it is still a great deal of work to reverse-engineer the code's intent. Describing the task in plain English (or whichever language you use to communicate with your team) will make it easier for everyone to read and understand the code's purpose. It can also help you organize your own work and ensure you are following logical steps. The second type of comment explains why the code is performing a task in a particular way. As you are writing code, you are making a series of decisions that (hopefully) make perfect sense to you at the time. These are often highly specialized and may exploit a functionality that is not obvious or has not been seen by others before. Even you will probably not remember the exact choices that were made in a couple of weeks. Therefore, you must document your precise processes in your code.

Code files should be stored in an easy-to-find location and named in a meaningful way. Breaking your code into independently readable “chunks” is good practice for code organization. You should write each functional element as a chunk that can run completely on its own. This ensures that each code component is independent; it does not rely on a complex program state created by other code chunks that are not obvious from the immediate context. One way to do this is by creating sections in your script to identify where a specific task is completed. For example, if you want to find the line in your code where the directory is set, you can go straight to PART 2: Prepare folder paths and define programs, instead of reading line by line through the entire code.

RStudio makes it very easy to create sections, and it compiles them into an interactive script index for you. In Stata, you can use comments to create section headers (see line 27 of the example do-file),

¹³ **Comments:** Code components that have no function to the computer, but describe in plain language for humans to read what the code is supposed to do.

though they're just there to make the reading easier and don't have functionality. Since an index is not automated, create this manually in the code header by copying and pasting section titles (see lines 8-10 in the example do-file). You can then add and navigate through them using the `find` functionality. Since Stata code is harder to navigate, as you will need to scroll through the document, it's particularly important to avoid writing very long scripts. Therefore, in Stata at least, we recommend breaking code tasks down into separate do-files, since there is no limit on how many you can have, how detailed their names can be, and no advantage to writing longer files. One reasonable rule of thumb is to not write do-files that have more than 200 lines. This is an arbitrary limit, just like the common practice of limiting code lines to 80 characters: it seems to be "enough but not too much" for most purposes.

Working with a master script

To bring all these smaller code files together, you must maintain a master script.¹⁴ A master script is the map of all your project's data work which serves as a table of contents for the instructions that you code. Anyone should be able to follow and reproduce all your work from raw data to all outputs by simply running this single script. By follow, we mean someone external to the project who has the master script and all the input data can (i) run all the code and recreate all outputs, (ii) have a general understanding of what is being done at every step, and (iii) see how codes and outputs are related. The master script is also where all the settings are established, such as versions, folder paths, functions, and constants used throughout the project.

Try to create the habit of running your code from the master script. Creating "section switches" using macros or objects to run only the codes related to a certain task should always be preferred to manually open different scripts to run them in a certain order (see the `if (0)` switches in Part 3 of `stata-master-dofile.do` for one way of how to do this). Furthermore, running all scripts related to a particular task through the master whenever one of them is edited helps you identify unintended consequences of the changes you made. Say, for example, that you changed the name of a variable created in one script. This may break another script that refers to this variable. But unless you run both of them when the change is made, it may take time for that to happen, and when it does, it may take time for you to understand what's causing an error. The same applies to changes in datasets and results.

To link code, data and outputs, the master script reflects the struc-

¹⁴ https://dimewiki.worldbank.org/Master_Do-files

ture of the DataWork folder in code through globals (in Stata) or string scalars (in R); refer to lines 38-43 of the example do-file. These coding shortcuts can refer to subfolders, so that those folders can be referenced without repeatedly writing out their absolute file paths. Because the DataWork folder is shared by the whole team, its structure is the same in each team member's computer. The only difference between machines should be the path to the project root folders, i.e. the highest-level shared folder. Depending on your software environment you may have multiple root folders. In a typical DIME project we have one Git root folder for our code, one sync software root folder for our de-identified data, and a third for our encrypted data. This is reflected in the master script in such a way that the only change necessary to run the entire code for a new team member is to change the path to the project root folders to reflect the file system and username; refer to lines 30-35 of the example do-file. The code in `stata-master-dofile.do` shows how folder structure is reflected in a master do-file. Because writing and maintaining a master script can be challenging as a project grows, an important feature of the `iefolder` is to write sub-master do-files and add to them whenever new subfolders are created in the DataWork folder.

In order to maintain well-documented and organized code, you should agree with your team on a plan to review code as it is written. Reading other people's code is the best way to improve your coding skills. And having another set of eyes on your code will make you more comfortable with the results you find. It's normal (and common) to make mistakes as you write your code. Reading it again to organize and comment it as you prepare it to be reviewed will help you identify them. Try to have a code review scheduled frequently, every time you finish writing a piece of code, or complete a small task. If you wait for a long time to have your code reviewed, and it gets too complex, preparation and code review will require more time and work, and that is usually the reason why this step is skipped. One other important advantage of code review is that making sure that the code is running properly on other machines, and that other people can read and understand the code easily, is the easiest way to be prepared in advance for a smooth project handover or for release of the code to the general public.

stata-master-dofile.do

```

1  ****
2  *          TEMPLATE MASTER DO-FILE
3  ****
4  *
5  * PURPOSE:   Reproduce all data work, map inputs and outputs,
6  *             facilitate collaboration
7  *
8  * OUTLINE:   PART 1: Set standard settings and install packages
9  *             PART 2: Prepare folder paths and define programs
10 *            PART 3: Run do-files
11 *
12 ****
13 PART 1: Install user-written packages and harmonize settings
14 ****
15
16 local user_commands ietoolkit iefieldkit //Add required user-written commands
17 foreach command of local user_commands {
18     cap which `command'
19     if _rc == 111 ssc install `command'
20 }
21
22     *Harmonize settings accross users as much as possible
23 ieboilstart, v(13.1)
24 `r(version)'
25
26 ****
27 PART 2: Prepare folder paths and define programs
28 ****
29
30 * Research Assistant folder paths
31 if "`c(username)'" == "ResearchAssistant" {
32     global github      "C:/Users/RA/Documents/GitHub/d4di/DataWork"
33     global dropbox     "C:/Users/RA/Dropbox/d4di/DataWork"
34     global encrypted   "M:/DataWork/EncryptedData"
35 }
36
37 * Baseline folder globals
38 global bl_encrypt    "${encrypted}/Round Baseline Encrypted"
39 global bl_dt         "${dropbox}/Baseline/DataSets"
40 global bl_doc        "${dropbox}/Baseline/Documentation"
41 global bl_do         "${github}/Baseline/Dofiles"
42 global bl_out        "${github}/Baseline/Output"
43
44 ****
45 PART 3: Run do-files
46 ****
47
48 /*-----
49 PART 3.1: De-identify baseline data
50 REQUIRES:  ${bl_encrypt}/Raw Identified Data/D4DI_baseline_raw_identified.dta
51 CREATES:   ${bl_dt}/Raw Deidentified/D4DI_baseline_raw_deidentified.dta
52 IDS VAR:   hhid
53 ----- */
54 *Change the 0 to 1 to run the baseline de-identification dofile
55 if (0) do "${bl_do}/Cleaning/deidentify.do"
56
57 /*-----
58 PART 3.2: Clean baseline data
59 REQUIRES:  ${bl_dt}/Raw Deidentified/D4DI_baseline_raw_deidentified.dta
60 CREATES:   ${bl_dt}/Final/D4DI_baseline_clean.dta
61           ${bl_doc}/Codebook baseline.xlsx
62 IDS VAR:   hhid
63 ----- */
64 *Change the 0 to 1 to run the baseline cleaning dofile
65 if (0) do "${bl_do}/Cleaning/cleaning.do"
66
67 /*-----
68 PART 3.3: Construct income indicators
69 REQUIRES:  ${bl_dt}/Final/D4DI_baseline_clean.dta
70 CREATES:   ${bl_out}/Raw/D4DI_baseline_income_distribution.png
71           ${bl_dt}/Intermediate/D4DI_baseline_constructed_income.dta
72 IDS VAR:   hhid
73 ----- */
74 *Change the 0 to 1 to run the baseline variable construction dofile
75 if (0) do "${bl_do}/Construct/construct_income.do"

```

Preparing to handle confidential data

Anytime you are working with original data in a development research project, you are almost certainly handling data that include **personally-identifying information (PII)**.¹⁵ PII is information which can, without any transformation or linkage, be used to identify individual people, households, firms, (or other units) in your data. Some examples of PII variables include names, addresses, and geolocations, email addresses, phone numbers, and bank accounts or other financial details. If you are working in a context or population that is either small, specific, or has extensive linkable data sources available to others, information like someone's age and gender may be sufficient to disclose their identity, even though those variables would not be considered PII in general.

In a collaborative project, you will sometimes need to transfer and work with PII information, and sometimes you will prefer to remove or mask PII before transferring or working with data. There is no one-size-fits-all solution to determine what is PII, research teams have to use careful judgment in each case to avoid statistical disclosure.¹⁶ It is important to keep in mind that data privacy principles apply not only for the respondent giving you the information but also for their household members or other individuals who are included in the data.

In all cases where confidential information is involved, you must make sure that you adhere to several core principles. These include ethical approval, participant consent, data security, and participant privacy.¹⁷ If you are a US-based researcher, you will become familiar with a set of governance standards known as "The Common Rule".¹⁸ If you interact with European institutions or persons, you will also become familiar with the General Data Protection Regulation (GDPR),¹⁹ a set of regulations governing **data ownership** and privacy standards. No matter who you are or what exact legal requirements you face, the core principles and practices you need to consider will always be similar.

Obtaining ethical approval and consent

Most of the field research done in development involves human subjects.²⁰ As a researcher, you are asking people to trust you with personal information about themselves: where they live, how rich they are, whether they have committed or been victims of crimes, their names, their national identity numbers, and all sorts of other data. PII data carries strict expectations about data storage and handling, and it is the responsibility of the research team to satisfy

¹⁵ **Personally-identifying information:** any piece or set of information that can be used to identify an individual research subject. https://dimewiki.worldbank.org/Protecting_Human_Research_Subjects

¹⁶ https://dimewiki.worldbank.org/Data_Release

¹⁷ See Baldwin et al. (2017) for an example.

¹⁸ Bierer et al. (2017)

¹⁹ <https://gdpr-info.eu>

²⁰ https://dimewiki.worldbank.org/Human_Subjects_Approval

these expectations.²¹ Your donor or employer will most likely require you to hold a certification from a respected source.²²

For almost all such data collection and research activities, you will be required to complete some form of **Institutional Review Board (IRB)** process.²³ Most commonly this consists of a formal application for approval of a specific protocol for consent, data collection, and data handling.²⁴ Which IRB has authority over your project is not always apparent, particularly if some institutions do not have their own. It is customary to obtain an approval from a university IRB where at least one PI is affiliated, and if work is being done in an international setting, approval is often also required from an appropriate local institution subject to the laws of the country where data originates.

One primary consideration of IRBs is the protection of the people about whom information is being collected and whose lives may be affected by the research design. Some jurisdictions (especially those who governed by EU law) view all personal data as intrinsically owned by the persons who they describe. This means that those persons have the right to refuse to participate in data collection before it happens, as it is happening, or after it has already happened. It also means that they must explicitly and affirmatively consent to the collection, storage, and use of their information for any purpose.

The development of appropriate consent processes is of primary importance. All survey instruments must include a module in which the sampled respondent grants informed consent to participate. Research participants must be informed of the purpose of the research, what their participation will entail in terms of duration and any procedures, any foreseeable benefits or risks, and how their identity will be protected.²⁵ There are special additional protections in place for vulnerable populations, such as minors, prisoners, and people with disabilities, and these should be confirmed with relevant authorities if your research includes them.

IRB approval should be obtained well before any data is acquired. IRBs may have infrequent meeting schedules or require several rounds of review for an application to be approved. If there are any deviations from an approved plan or expected adjustments, report these as early as possible so that you can update or revise the protocol. IRBs have the authority to retroactively deny the right to use data which was not acquired in accordance with an approved plan. This is extremely rare, but shows the seriousness of these considerations since the institution itself may face legal penalties if its IRB is unable to enforce them. As always, as long as you work in good faith, you should not have any issues complying with these regulations.

²¹ https://dimewiki.worldbank.org/Research_Ethics

²² https://dimewiki.worldbank.org/Protecting_Human_Research_Subjects

²³ **Institutional Review Board (IRB):** An institution formally responsible for ensuring that research meets ethical standards.

²⁴ https://dimewiki.worldbank.org/IRB_Approval

²⁵ https://dimewiki.worldbank.org/Informed_Consent

Ensuring privacy and security in research data

In order to safeguard PII data and protect respondent privacy, you must set up a secure data system from the outset of a project. Secure data storage and transfer are ultimately your personal responsibility.²⁶ Later chapters will discuss how to properly protect your data depending on which method you are using to acquire, store or share data. This section will only cover the computer setup you will need for any project. There are several components to this.

First, you need a system for managing strong and unique passwords for all accounts – including personal accounts like computer logins and email. This means that all your passwords should be long, not use common words and should not be reused for multiple accounts. The only way to make that practically feasible is to use a password manager. Most password managers also allows you to securely share passwords for shared accounts with your colleagues.

Second, all machines that stores confidential data should have **hard disk encryption** enabled, if available in the operative system used.²⁷ Most hard disk encryption use your computer login password to prevent your files from ever being read by anyone but you. It is important to note that password-protection alone is not sufficient. Password protection makes it more difficult for someone else to gain access to your files, but only encryption properly protects your data if someone manages to access your files anyway. Hard disk encryption ensures that all your local files are protected, but it may only be supported in enterprise versions of operating systems, most notably Microsoft Windows. To properly secure your data you should have the habit of signing out from your computer when you leave it, no matter how briefly.

Third, you must make sure that data including confidential information remains encrypted at all times during data collection, storage, and transfer. When files are properly encrypted, the information they contain will be completely unreadable and unusable even if they were to be intercepted by a malicious “intruder” (an information-security term for any unauthorized information access) or accidentally made public. We will discuss implementations of encryption specific to different stages of data work in Chapter 4, but when setting up a software environment for you and your team, make sure that you have a solution for creating encrypted folders on your computers. DIME uses Veracrypt for encrypted folders, our protocols are available as part of the DIME Research Standards.²⁸

Handling confidential data properly will always add to your workload. The easiest way to reduce that work load is to handle it as rarely as possible. Whenever our research design allows us to,

²⁶ [https://dimewiki.worldbank.org/
Data_Security](https://dimewiki.worldbank.org/Data_Security)

²⁷ **Encryption:** Methods which ensure that files are unreadable even if laptops are stolen, databases are hacked, or any other type of unauthorized access is obtained. <https://dimewiki.worldbank.org/Encryption>

²⁸ [https://github.com/worldbank/
dime-standards](https://github.com/worldbank/dime-standards)

we should not work with confidential data at all, and not collect it or ask for it in the first place. Even when we do need confidential data for some aspect of our research, it is almost never the case that we need it for *all* aspects of our data. It is often very simple to conduct planning and analytical work using a subset of the data that does not include this type of information. Therefore we recommend that all projects plan a workflow with a version of the data where confidential data has been removed and always use that dataset when possible.

Note that it is in practice impossible to **anonymize** data. There is always some statistical chance that an individual's identity will be re-linked to the data collected about them – even if that data has had all directly identifying information removed – by using some other data that becomes identifying when analyzed together. For this reason, we recommend de-identification in two stages. The **initial de-identification** process strips the data of direct identifiers as early in the process as possible, to create a working de-identified dataset that can be shared *within the research team* without the need for encryption. This data set should always be used when possible. The **final de-identification** process involves making a decision about the trade-off between risk of disclosure and utility of the data before publicly releasing a dataset.²⁹

Finally, it is essential to have an end-of-life plan for data even before it is acquired.³⁰ This includes plans for how to transfer access and control to a new person joining the team, and how to revoke that access when someone is leaving the team. It should also include a plan for how the confidential data should be deleted. Every project should have a clear data retention and destruction plan. After a project is completed and its de-identified data has been made available as a part of data publication, research teams should not retain confidential data indefinitely.

Looking ahead

With your code environment established, you will have a firm idea about how you are going to handle the data and code that you receive and create throughout the research process. This structure should prepare you to work collaboratively, to share code and data across machines and among team members, and to document your work as a group. With an organization plan and plans to to version-control and back up files, you are ready to handle materials ethically and securely. You should also have secured the approvals needed for any planned work. You are now ready to translate your project's research design into a measurement framework to answer your re-

²⁹ <https://dimewiki.worldbank.org/De-identification>

³⁰ https://dimewiki.worldbank.org/Data_Storage

search questions. In the next chapter, we will outline how to prepare the essential elements of research data. You will learn how to map out your project's data needs according to both the research design and the planned creation and use of data across the project timeline.

Chapter 3: Establishing a measurement framework

The first step in the data workflow is to establish a measurement framework. This requires understanding your project's data requirements and how to structure the required data to answer the research questions. Setting up the measurement framework involves more than simply listing the key outcome variables. You also need to understand how to structure original data, understand how various types of data connect together, and create tools to share this understanding across your team.

In this chapter we will show how to save time and increase the quality of your research by planning this framework in advance. The first section of this chapter discusses how to determine your project's data needs, and introduces the *DIME Data Map Template*. The template includes a data linkage table, master datasets, and data flowcharts. These tools are used to communicate the project's data requirements both across the team and across time. This section also discusses how to assess data needs based on your project's research design, and how to document those in the data map. We will discuss two types of variables: variables that tie your research design to the observations in the data which we call **research variables**; and variables that correspond to observations of the real world, which we call **measurement variables**. The second section of the chapter covers how to reproducibly generate research variables, specifically through random sampling and random assignment. The final section discusses power calculations and randomization inference, which are tools for assessing the measurement ability of the framework and any randomization you undertake.

Creating a data map

In most projects, more than one data source is needed to answer a research question. These could be data from multiple survey rounds, data acquired from different partners (such as administrative data, implementation data, sensor data), technological tools like satellite imagery or web scraping, or complex combinations of these and other sources.¹ However your study data is structured, you need to know how to link data from all sources and analyze the relationships between the units that appear in them to answer your research questions. You might think that you are able to keep all the relevant details in your head, but your whole research team is unlikely to have the same understanding, at all times, of all the datasets required. The only way to make sure that the full team shares the same understanding is to create a **data map**.² DIME's data map template has three components: one *data linkage table*, one or several *master datasets*

¹ See Kondylis and Stein (2018) for an example.

² https://dimewiki.worldbank.org/Data_Map

and one or several *data flowcharts*.

Each component of the data map that you create for your project will contribute to understanding what the data is about and how to use it. First, you will create a **data linkage table**³ listing all the raw datasets that will be used in the project, what data sources they are created from, and how they relate to each other. Second, for each **unit of observation**⁴ in the data linkage table, as well for each unit of analysis you plan to use, you will maintain a **master dataset**⁵ listing all observations of that unit relevant to the project. These datasets will be the authoritative sources for all research data including unique identifiers, sample statuses, and treatment assignments. Finally, using these two resources you will continuously create **data flowcharts**,⁶ describing how the raw datasets and master datasets are to be combined and manipulated to create analysis datasets.

The purpose of the data map is to make sure that at the time of data analysis as described in Chapter 6, you have all data that you need to answer the research questions in your research design. The process of drafting the data map is also a useful opportunity for the principal investigators to communicate their vision of the data structure and requirements, and for research assistants to communicate their understanding of that vision. The data map should be drafted at the outset of a project, before any data is acquired. However, it is not a static document; each component will need to be updated and maintained as the project evolves and new data sources, observations, and research questions arise.

Data linkage table

To create a data map according to DIME’s template, the first step is to create a **data linkage table** by listing all the data sources you know you will use in a spreadsheet, and the raw datasets that will be created from them. If one source of data will result in two different raw datasets, then list each dataset on its own row. For each dataset, list the unit of observation and the name of the **project ID**⁷ variable for that unit of observation. It is important to include both plain-language terminology as well as technical file and variable names here. For example, the `hh_baseline2020_listmap.csv` dataset may be called the “Baseline Household Listing” data; it may be identified by the `hh_id` variable and said to be identified at the “Household” level. Having the plain-language terminology here early in the project allows you to use these titles unambiguously in communication.

The data linkage table will therefore help you plan out how you will identify each unit of observation in your data. When you list a dataset in the data linkage table – which should be done before that

³ https://dimewiki.worldbank.org/Data_Linkage_Table

⁴ https://dimewiki.worldbank.org/Unit_of_Observation

⁵ https://dimewiki.worldbank.org/Master_Data_Set

⁶ https://dimewiki.worldbank.org/Data_Flow_Chart

⁷ **Project ID:** The main ID used in your project to identify observations. You should never have multiple project IDs for the same unit of observation. The project ID must uniquely and fully identify all observations in the project. See https://dimewiki.worldbank.org/Project_ID for more details.

data source is acquired – you should make sure that the data will be fully and uniquely identified by the project ID, or make a plan for how the new dataset will be linked to the project ID. It is very labor-intensive to work with a dataset that do not have an unambiguous way to link to the project ID, and it is a major source of error.⁸

The data linkage table should indicate whether datasets can be merged one-to-one (for example, merging baseline and endline datasets that use the same unit of observation), or whether two datasets need to be merged many-to-one (for example, school administrative data merged with student data). Your data map must indicate which ID variables can be used – and how – when merging datasets. The data linkage table is also a great place to list other metadata, such as the source of your data, its backup locations, the nature of the data license, and so on.

⁸ See Fernandes et al. (2017) for an example.

Master datasets

The second step in creating a data map is to create one **master dataset** for each unit of observation that will be used in any research activity. Examples of such activities are data collection, data analysis, random sampling, and treatment assignment. The master dataset should be the authoritative source of all **research variables**⁹ but not include any **measurement variables**.¹⁰ Research variables and measurement variables often come from the same source, but should not be stored in the same way. For example, if you acquire administrative data that both includes information on eligibility for the study (research variable) and data on the topic of your study (measurement variable) you should first decide which variables are research variables, and store them in the master dataset, while storing the measurement variables in a cleaned dataset as described in Chapter 5. It is common that you will have to update your master datasets throughout your project.

The most important function of each master dataset is to be the authoritative source for how all observations at that level are identified. This means that the master datasets should include identifying information such as names, contact information, but also your project ID. The project ID is the ID variable used in the data linkage table, and is therefore how observations are linked across datasets. Your master dataset may list alternative IDs that are used, for example, by a partner organization. However, you must not use such an ID as your project ID, as you would then not be in control over who can re-identify data that you publish.

The project ID must be created by the project team, and the linkage to direct identifiers should only be known to people listed on

⁹ **Research variables:** Research data that identifies observations and maps research design information to those observations. Research variables are time-invariant and often, but not always, controlled by the research team. Examples include ID variables, sampling status, treatment status, and treatment uptake.

¹⁰ **Measurement variables:** Data that measures attributes or records responses of research subjects. Measurement variables are not controlled by the research team and often vary over time. Examples include characteristics of the research subject, outcome variables, and control variables among many others.

the IRB. If you receive a dataset with an alternative ID, you should immediately replace it with your project ID, and the alternative ID should be dropped as a part of your de-identification (see Chapter 5 and Chapter 7). Your master dataset serves as the linkage between all other identifying information and your project ID. Since your master dataset is full of identifying information, it must always be encrypted.

The starting point for the master dataset is typically a sampling frame (more on sampling frames later in this chapter). However, you should continuously update the master dataset with all observations ever encountered in your project, even if those observations are not eligible for the study. Examples include new observations listed during monitoring activities or observations that are connected to respondents in the study, for example in a social network module. This is useful because, if you ever need to perform a record linkage such as a fuzzy match on string variables like proper names, the more information you have the fewer errors you are likely to make. If you ever need to do a fuzzy match, you should always do that between the master dataset and the dataset without an unambiguous identifier.¹¹ You should not do anything with that dataset until you have successfully merged the project IDs from the master dataset. When adding new observations to the master datasets, always convince yourself beyond reasonable doubt that the new observation is indeed a new observation. Sometimes you will encounter different spellings in identifiers and identifiers such as addresses may become outdated.

Since the master dataset is the authoritative source of the project ID and all research variables for the corresponding unit of observation, it serves as an unambiguous method of mapping the observations in your study to your research design.

¹¹ See Benhassine et al. (2018) for an example.

Data flowcharts

The third and final step in creating the data map is to create **data flowcharts**. Each analysis dataset (see Chapter 6 for discussion on why you likely need multiple analysis datasets) should have a data flowchart showing how it was created. The flowchart is a diagram where each starting point is either a master dataset or a dataset listed in the data linkage table. The data flowchart should include instructions on how the datasets can be combined to create the analysis dataset. The operations used to combine the data could include: appending, one-to-one merging, many-to-one or one-to-many merging, collapsing, or a broad variety of others.

You must list which variable or set of variables should be used in each operation, and note whether the operation creates a new

variable or combination of variables to identify the newly linked data. These variables should be project IDs when possible. Examples of exception are time variables in longitudinal data, and sub-units like farm plots that belong to farmers with project IDs. Once you have acquired the datasets listed in the flowchart, you can add to the data flowcharts the number of observations that the starting point dataset has and the number of observation each resulting datasets should have after each operation. This is a great method to track attrition and to make sure that the operations used to combine datasets did not create unwanted duplicates or incorrectly drop any observations.

A data flowchart can be created in a flowchart drawing tool (there are many free alternatives online) or by using shapes in Microsoft PowerPoint. You can also do this simply by drawing on a piece of paper and taking a photo, but we recommend a digital tool so that flowcharts can easily be updated over time. Similarly to the data linkage table, you should include here both technical information as well as plain-language interpretations of the operations that are done and the data that is created. These will help readers understand the complex data combinations that often result from merges and appends, such as panel datasets like “person-year” structures and multi-level data like “district-school-teacher-student” structures.

Expressing research design in data maps

After you have set up your data map, you need to carefully think about your research design and which research variables you will need in the data analysis to infer the relation between differences in measurement variables and your research design. We assume you have a working familiarity with the research designs mentioned here. You should reference our **Research Design Appendix**, where you will find more details and specific references for common impact evaluation designs.

Defining research variables related to your research design

As DIME primarily works on impact evaluations, we focus our discussion here on research designs that compare a group that received some kind of **treatment**¹² group against a **counterfactual**.¹³ The key assumption is that each person, facility, or village (or whatever the unit of treatment is) had two possible states: their outcome if they did receive the treatment and their outcome if they did not receive that treatment. The average impact of the treatment,¹⁴ is defined as the difference between these two states averaged over all units.

However, we can never observe the same unit in both the treated

¹² **Treatment:** The general word for the evaluated intervention or event. This includes things like being offered a training, a cash transfer from a program, or experiencing a natural disaster, among many others.

¹³ **Counterfactual:** A statistical description of what would have happened to specific individuals in an alternative scenario, for example, a different treatment assignment outcome.

¹⁴ **Average treatment effect (ATE):** The expected average change in outcome that untreated units would have experienced had they been treated.

and untreated state simultaneously, so we cannot calculate these differences directly. Instead, the treatment group is compared to a control group that is statistically indistinguishable, which is often referred to as achieving **balance** between two or more groups. DIME Analytics maintains a Stata command to standardize and automate the creation of well-formatted balance tables: `iebaltab`.¹⁵ Each research design has a different method for identifying and balancing the counterfactual group. The rest of this section covers how research data requirements differ between those different methods. What does not differ, however, is that these data requirements are all research variables. And that the research variables discussed below should always be included in the master dataset. You will often have to merge the research variables to other datasets, but that is an easy task if you created a data linkage table.

In **experimental research designs**, such as **randomized control trials (RCTs)**,¹⁶ the research team determines which members of the studied population will receive the treatment. This is typically done by a randomized process in which a subset of the eligible population is randomly assigned to receive the treatment (see later in this chapter for how to implement this). The intuition is that if everyone in the eligible population is assigned at random to either the treatment or control group, then the two groups will, on average, be statistically indistinguishable. The treatment will therefore not be correlated with anything but the impact of that treatment.¹⁷ The randomized assignment should be done using data from the master dataset, and the result should be saved back to the master dataset, before being merged to other datasets.

Quasi-experimental research designs,¹⁸ by contrast, are based on events not controlled by the research team. Instead, they rely on “experiments of nature”, in which natural variation in exposure to treatment can be argued to approximate deliberate randomization. You must have a way to measure this natural variation, and how the variation is categorized as outcomes of a naturally randomized assignment must be documented in your master dataset. Unlike carefully planned experimental designs, quasi-experimental designs typically require the luck of having access to data collected at the right times and places to exploit events that occurred in the past. Therefore, these methods often use either secondary data, including administrative data or other classes of routinely-collected information, and it is important that your data linkage table documents how this data can be linked to the rest of the data in your project.

No matter the design type, you should be very clear about which data points you observe or collect are research variables. For example, **regression discontinuity (RD)**¹⁹ designs exploit sharp breaks

¹⁵ <https://dimewiki.worldbank.org/iebaltab>

¹⁶ https://dimewiki.worldbank.org/Randomized_Control_Trials

¹⁷ Duflo et al. (2007)

¹⁸ https://dimewiki.worldbank.org/Quasi-Experimental_Methods

¹⁹ https://dimewiki.worldbank.org/Regression_Discontinuity

or limits in policy designs.²⁰ The cutoff determinant, or running variable, should be saved in your master dataset. In **instrumental variables (IV)**²¹ designs, the **instruments** influence the *probability* of treatment.²² These research variables should be collected and stored in the master dataset. Both the running variable in RD designs and the instruments in IV designs, are among the rare examples of research variables that may vary over time. In such cases your research design should ex-ante clearly indicate what point of time they will be recorded, and this should be clearly documented in your master dataset.

In **matching** designs, observations are often grouped by a strata, grouping, index, or propensity score.²³ Like all research variables, the matching results should be stored in the master dataset. This is best done by assigning a matching ID to each matched pair or group, and create a variable in the master dataset with the matching ID each unit belongs to.²⁴ In all these designs, fidelity to the design is important to record as well.²⁵ A program intended for students that scored under 50% on a test might have some cases where the program is offered to someone that scored 51% at the test, or someone that scored 49% at the might decline to participate in the program. Differences between assignments and realizations should also be recorded in the master datasets.

Time periods in data maps

Your data map should also take into consideration whether you are using data from one time period or several. A study that observes data in only one time period is called a **cross-sectional study**. Observations over multiple time periods, referred to as **longitudinal data**, can consist of either **repeated cross-sections** or **panel data**. In repeated cross-sections, each successive round of data collection uses a new random sample of observations from the treatment and control groups, but in a panel data study the same observations are tracked and included each round.²⁶ If each round of data collection is a separate activity, then they should be treated as separate sources of data and get their own row in the data linkage table.

If the data is continuously collected, or at frequent intervals, then it can be treated as a single data source. The data linkage table must document how the different rounds will be merged or appended when panel data is collected in separate activities. You must keep track of the *attrition rate* in panel data, which is the share of observations not observed in follow-up data. It is common that the observations not possible to track can be correlated with the outcome you study.²⁷ For example, poorer households may live in more informal

²⁰ See Alix-Garcia et al. (2019) for an example.

²¹ https://dimewiki.worldbank.org/Instrumental_Variables

²² See Calderon et al. (2017) for an example.

²³ <https://dimewiki.worldbank.org/Matching>

²⁴ See Prennushi and Gupta (2014) for an example.

²⁵ See Pradhan et al. (2013) for an example.

²⁶ See Kondylis et al. (2016) for an example.

²⁷ See Bertrand et al. (2017) for an example.

dwellings, patients with worse health conditions might not survive to follow-up, and so on. If this is the case, then your results might only be an effect of your remaining sample being a subset of the original sample that were better or worse off from the beginning. You should have a variable in your master dataset that indicates attrition. A balance check using the attrition variable can provide insights as to whether the lost observations were systematically different compared to the rest of the sample.

Monitoring data

For any study with an ex-ante design, **monitoring data**²⁸ is very important for understanding if the research design corresponds to reality. The most typical example is to make sure that, in an experimental design, the treatment was implemented according to your treatment assignment. Treatment implementation is often carried out by partners, and field realities may be more complex than foreseen during research design. Furthermore, the field staff of your partner organization, might not be aware that their actions are the implementation of a research design.

Therefore, you must acquire monitoring data that tells you how well the treatment assignment in the field corresponds to your intended treatment assignment, for nearly all experimental research designs. While it is always better to monitor all activities, it might be too costly. In those cases you can sample a smaller number of critical activities and monitor them. This will not be detailed enough to be used as a control in your analysis, but it will still give a way to estimate the validity of your research design assumptions.

Monitoring data is particularly prone to errors when linking it with the rest of the data in your project. Often monitoring activities are done by sending a team to simply record the name of all people attending a training, or by a partner organization sharing their administrative data, which is rarely maintained in the same format or structure as your research data.²⁹ In both those cases it can be difficult to make sure that the project ID or any other unambiguous identifiers in our master datasets is used to record who is who. Planning ahead for this when the monitoring activity is added to the data linkage table is the best protection from ending up with poor correlation between treatment uptake and treatment assignment, without a way to tell if the poor correlation is just a result of a fuzzy link between monitoring data and the rest of your data.

²⁸ https://dimewiki.worldbank.org/Monitoring_Data

²⁹ See Goldstein et al. (2015) for an example.

Creating research variables by randomization

Random sampling and random treatment assignment are two core research design activities that generate new research variables. Random sampling and random treatment assignment directly determine the set of individuals who are going to be observed and what their status will be for the purpose of effect estimation. **Randomization**³⁰ is used to ensure that a sample is representative and that treatment groups are statistically indistinguishable.

Randomization in statistical software is non-trivial and its mechanics are unintuitive for the human brain. The principles of randomization we will outline apply not just to random sampling and random assignment, but to all statistical computing processes that have random components such as simulations and bootstrapping. Furthermore, all random processes introduce statistical noise or uncertainty into estimates of effect sizes. Choosing one random sample from all the possibilities produces some probability of choosing a group of units that are not, in fact, representative. Similarly, choosing one random assignment produces some probability of creating groups that are not good counterfactuals. *Power calculation* and *randomization inference* are the main methods by which these probabilities of error are assessed. These analyses are particularly important in the initial phases of development research – typically conducted before any data acquisition or field work occurs – and have implications for feasibility, planning, and budgeting.

Randomizing sampling and treatment assignment

Sampling is the process of randomly selecting observations from a list of individuals to create a representative or statistically similar sub-sample. This process can be used, for example, to select a subset from all eligible units to be included in data collection when the cost of collecting data on everyone is prohibitive.³¹ But it can also be used to select a sub-sample of your observations to test a computationally heavy process before running it on the full data. **Randomized treatment assignment** is the process of assigning observations to different treatment arms. This process is central to experimental research design. Most of the code processes used for randomized assignment are the same as those used for sampling, since it is also a process of randomly splitting a list of observations into groups. Where sampling determines whether a particular individual will be observed at all in the course of data collection, randomized assignment determines if each individual will be observed as a treatment observation or used as a counterfactual.

³⁰ Randomization is often used interchangeably to mean random treatment assignment. In this book however, *randomization* will only be used to describe the process of generating a sequence of unrelated numbers, i.e. a random process. *Randomization* will never be used to mean the process of assigning units in treatment and control groups, that will always be called *random treatment assignment*, or a derivative thereof.

³¹ https://dimewiki.worldbank.org/Sample_Size_and_Power_Calculations

The list of units to sample or assign from may be called a **sampling universe**, a **listing frame**, or something similar. This list should always be your **master dataset** when possible, and the result should always be saved in the master dataset before merged to any other data. One example of the rare exceptions when master datasets cannot be used is when sampling must be done in real time – for example, randomly sampling patients as they arrive at a health facility. In those cases, it is important that you collect enough data during the real time sampling, such that you can add these individuals, and the result of the sampling, to your master dataset afterwards.

The simplest form of sampling is **uniform-probability random sampling**. This means that every eligible observation in the master dataset has an equal probability of being selected. The most explicit method of implementing this process is to assign random numbers to all your potential observations, order them by the number they are assigned, and mark as “sampled” those with the lowest numbers, up to the desired proportion. There are a number of shortcuts to doing this process, but they all use this method as the starting point, so you should become familiar with exactly how it works. The do-file below provides an example of how to implement uniform-probability sampling in practice. This code uses a Stata built-in dataset and is fully reproducible (more on reproducible randomization in next section), so anyone that runs this code in any version of Stata later than 13.1 (the version set in this code) will get the exact same, but still random, results.

```

----- simple-uniform-probability-sampling.do -----
1 * Set up reproducible randomization - VERSIONING, SORTING and SEEDING
2   ieboilstart , v(13.1)      // Version
3   `r(version)'              // Version
4   sysuse bpwide.dta, clear   // Load data
5   isid patient, sort        // Sort
6   set seed 215597           // Seed - drawn using https://bit.ly/stata-random
7
8 * Generate a random number and use it to sort the observations.
9 * Then the order the observations are sorted in is random.
10  gen sample_rand = rnormal() // Generate a random number
11  sort sample_rand          // Sort based on the random number
12
13 * Use the sort order to sample 20% (0.20) of the observations.
14 *_N in Stata is the number of observations in the active dataset,
15 * and _n is the row number for each observation. The bpwide.dta has 120
16 * observations and 120*0.20 = 24, so (_n <= _N * 0.20) is 1 for observations
17 * with a row number equal to or less than 24, and 0 for all other
18 * observations. Since the sort order is randomized, this means that we
19 * have randomly sampled 20% of the observations.
20  gen sample = (_n <= _N * 0.20)
21
22 * Restore the original sort order
23  isid patient, sort
24
25 * Check your result
26  tab sample

```

Sampling typically has only two possible outcomes: observed and unobserved. Similarly, a simple randomized assignment has two outcomes: treatment and control, and the logic in the code would be identical to the sampling code example. However, randomized assignment often involves multiple treatment arms which each represent different varieties of treatments to be delivered;³² in some cases, multiple treatment arms are intended to overlap in the same sample. Complexity can therefore grow very quickly in randomized assignment and it is doubly important to fully understand the conceptual process that is described in the experimental design, and fill in any gaps before implementing it in code. The do-file below provides an example of how to implement a randomized assignment with multiple treatment arms.

³² See De Andrade et al. (2013) for an example.

```

----- simple-multi-arm-randomization.do -----
1 * Set up reproducible randomization - VERSIONING, SORTING and SEEDING
2 ieboilstart , v(13.1)      // Version
3 `r(version)'              // Version
4 sysuse bpwide.dta, clear // Load data
5 isid patient, sort        // Sort
6 set seed 654697           // Seed - drawn using https://bit.ly/stata-random
7
8 * Generate a random number and use it to sort the observation.
9 * Then the order the observations are sorted in is random.
10 gen treatment_rand = rnormal() // Generate a random number
11 sort treatment_rand        // Sort based on the random number
12
13 * See simple-sample.do example for an explanation of "(_n <= _N * X)".
14 * The code below randomly selects one third of the observations into group 0,
15 * one third into group 1 and one third into group 2.
16 * Typically 0 represents the control group
17 * and 1 and 2 represents the two treatment arms
18 generate treatment = 0          // Set all observations to 0 (control)
19 replace treatment = 1 if (_n <= _N * (2/3)) // Set only the first two thirds to 1
20 replace treatment = 2 if (_n <= _N * (1/3)) // Set only the first third to 2
21
22 * Restore the original sort order
23 isid patient, sort
24
25 * Check your result
26 tab treatment

```

Programming reproducible random processes

For statistical programming to be reproducible, you must be able to re-obtain its exact outputs in the future.³³ We will focus on what you need to do to produce truly random results for your project, to ensure you can get those results again. This takes a combination of strict rules, solid understanding, and careful programming. This section introduces strict rules: these are non-negotiable (but thankfully simple). Stata, like most statistical software, uses a **pseudo-random number generator**³⁴ which, in ordinary research use, produces sequences of numbers that are as good as random.³⁵ However, for *reproducible* randomization, we need two additional properties: we need to be able to fix the sequence of numbers generated and we need to ensure that the first number is independently randomized. In Stata, this is accomplished through three concepts: **versioning**, **sorting**, and **seeding**. We again use Stata in our examples, but the same principles translate to all other programming languages.

Versioning means using the same version of the software each time you run the random process. If anything is different, the underlying list of random numbers may have changed, and it may be impossible to recover the original result. In Stata, the `version` command ensures that the list of random numbers is fixed.³⁶ The `ieboilstart` command in `ietoolkit` provides functionality to sup-

³³ Orozco et al. (2018)

³⁴ **Pseudo-random number generator:** An algorithm that creates a long, fixed sequence of numbers which exhibits no statistical relationships between the position or value of any set of those numbers.

³⁵ https://dimewiki.worldbank.org/Randomization_in_Stata

³⁶ At the time of writing, we recommend using version 13.1 for backward compatibility; the algorithm used to create this list of random numbers was changed after Stata 14 but the improvements do not matter in practice.

port this requirement.³⁷ We recommend you use `ieboilstart` at the beginning of your master do-file.³⁸ However, testing your do-files without running them via the master do-file may produce different results, since Stata's `version` setting expires after each time you run your do-files.

Sorting means that the actual data that the random process is run on is fixed. Because random numbers are assigned to each observation row-by-row starting from the top row, changing their order will change the result of the process. In Stata, the only way to guarantee a unique sorting order is to use `isid [id_variable], sort`. (The `sort, stable` command is insufficient.) Since the exact order must be unchanged, the underlying data itself must be unchanged as well between runs. This means that if you expect the number of observations to change (for example to increase during ongoing data collection), your randomization will not be reproducible unless you split your data up into smaller fixed datasets where the number of observations does not change. You can combine all those smaller datasets after your randomization.

Seeding means manually setting the start point in the list of pseudo-random numbers. A seed is just a single number that specifies one of the possible start points. It should be at least six digits long and you should use exactly one unique, different, and randomly created seed per randomization process.³⁹ In Stata, `set seed [seed]` will set the generator to the start point identified by the seed. In R, the `set.seed` function does the same. To be clear: you should not set a single seed once in the master do-file, but instead you should set a new seed in code right before each random process. The most important thing is that each of these seeds is truly random, so do not use shortcuts such as the current date or a seed you have used before. You should also describe in your code how the seed was selected.

Other commands may induce randomness in the data, change the sorting order, or alter the place of the random generator without you realizing it, so carefully confirm exactly how your code runs before finalizing it. To confirm that a randomization has worked correctly before finalizing its results, save the outputs of the process in a temporary location, re-run the code, and use `cf` or `datasignature` to ensure nothing has changed. It is also advisable to let someone else reproduce your randomization results on their machine to remove any doubt that your results are reproducible. Once the result of a randomization is used in the field, there is no way to correct any mistakes.

³⁷ <https://dimewiki.worldbank.org/ieboilstart>

³⁸ https://dimewiki.worldbank.org/Master_Do-files

³⁹ You can draw a uniformly distributed six-digit seed randomly by visiting <https://bit.ly/stata-random>. (This link is a just shortcut to request such a random number on <https://www.random.org>.) There are many more seeds possible but this is a large enough set for most purposes.

reproducible-randomization.do

```

1 * VERSIONING - Set the version
2   `eboilstart , v(13.1)
3   `r(version)'
4
5 * Load the auto dataset (auto.dta is a test dataset included in all Stata installations)
6   sysuse auto.dta , clear
7
8 * SORTING - sort on the uniquely identifying variable "make"
9   isid make, sort
10
11 * SEEDING - Seed picked using https://bit.ly/stata-random
12   set seed 287608
13
14 * Demonstrate stability after VERSIONING, SORTING and SEEDING
15   gen check1 = rnormal() // Create random number
16   gen check2 = rnormal() // Create a second random number without resetting seed
17
18   set seed 287608      // Reset the seed
19   gen check3 = rnormal() // Create a third random number after resetting seed
20
21 * Visualize randomization results. See how check1 and check3 are identical,
22 * but check2 is random relative to check1 and check3
23   graph matrix check1 check2 check3 , half

```

Some types of experimental designs require that randomized assignment results be revealed in the field. It is possible to do this using survey software or live events, such as a live lottery. These methods typically do not leave a record of the randomization, and as such are never reproducible. However, you can often run your randomization in advance even when you do not have list of eligible units in advance. Let's say you want to, at various health facilities, randomly select a sub-sample of patients as they arrive. You can then have a pre-generated list with a random order of "in sample" and "not in sample". Your field staff would then go through this list in order and cross off one randomized result as it is used for a patient.

This is especially beneficial if you are implementing a more complex randomization. For example, a hypothetical research design may call for enumerators to: (1) sample 10% of people observed in a particular location; (2) show a video to 50% of the sample; (3) administer a short questionnaire to 80% of all those sampled and (4) administer a longer questionnaire to the remaining 20%, with the questionnaire mix equal between those who were shown the video and those that were not. In real-time, such a complex randomization is much more likely to be implemented correctly if your field staff simply can follow a list with the randomized categories where you are in control of the pre-determined proportions and the random order. This way, you can also control with precision how these categories are evenly distributed across all the locations where you plan to conduct this research.

Finally, if this real-time randomization implementation is done using survey software, then the pre-generated list of randomized categories can be preloaded into the questionnaire. Then the field team can follow a list of respondent IDs that are randomized into the appropriate categories, and the survey software can show a video and control which version of the questionnaire is asked. This way, you reduce the risk of errors in field randomization.

Clustering or stratifying a random sample or assignment

For a variety of reasons, random sampling and random treatment assignment are rarely as straightforward as a uniform-probability draw. The most common variants are **clustering** and **stratification**.⁴⁰

Clustering occurs when your unit of analysis is different than the unit of sampling or treatment assignment.⁴¹ For example, a policy may be implemented at the village level, or you may only be able to send enumerators to a limited number of villages, or the outcomes of interest for the study are measured at the household level.⁴² The groups in which units are assigned to treatment are called clusters.⁴³

Stratification breaks the full set of observations into subgroups before performing randomized assignment within each subgroup.⁴⁴ The subgroups are called **strata**. This ensures that members of every subgroup are included in all groups of the randomized assignment process, or that members of all groups are observed in the sample. Without stratification, it is possible that the randomization would put all the members of a subgroup into just one of the treatment arms, or fail to select any of them into the sample. For both clustering and stratification, implementation is nearly identical in both random sampling and random assignment.

Clustering is procedurally straightforward in Stata, although it typically needs to be performed manually. To cluster a sampling or randomized assignment, randomize on the master dataset for the unit of observation of the cluster, and then merge the results to the master dataset for the unit of analysis. This is a many-to-one merge and your data map should document how those datasets can be merged correctly. If you do not have a master dataset yet for the unit of observation of the cluster, then you first have to create that and update your data map accordingly. When sampling or randomized assignment is conducted using clusters, the clustering variable should be clearly identified in the master dataset for the unit of analysis since it will need to be used in subsequent statistical analysis. Namely, standard errors for these types of designs must be clustered at the level at which the randomization was clustered.⁴⁵ This accounts for the design covariance within the cluster – the

⁴⁰ Athey and Imbens (2017a)

⁴¹ https://dimewiki.worldbank.org/Unit_of_Observation

⁴² [https://dimewiki.worldbank.org/Multi-stage_\(Cluster\)_Sampling](https://dimewiki.worldbank.org/Multi-stage_(Cluster)_Sampling)

⁴³ See Keating et al. (2011) for an example.

⁴⁴ https://dimewiki.worldbank.org/Stratified_Random_Sample

⁴⁵ <https://blogs.worldbank.org/impactevaluations/when-should-you-cluster-standard-errors-new-wisdom-econometrics-oracle>

information that if one individual was observed or treated there, the other members of the clustering group were as well.

By contrast, implementing stratified designs in statistical software is prone to error. Even for a simple multi-armed design, the basic method of randomly ordering the observations will often create very skewed assignments.⁴⁶ The user-written `randtreat` command properly implements stratification.⁴⁷ The options and outputs (including messages) from the command should be carefully reviewed so that you understand exactly what has been implemented. Notably, it is extremely hard to target precise numbers of observations in stratified designs, because exact allocations are rarely round fractions and the process of assigning the leftover “misfit” observations imposes an additional layer of randomization above the specified division.

Doing power calculations for research design

Random sampling and treatment assignment are noisy processes: it is impossible to predict the result in advance. By design, we know that the exact choice of sample or treatment will be uncorrelated with our key outcomes, but this lack of correlation is only true “in expectation”—that is, across a large number of randomizations. In any *particular* randomization, the correlation between the sampling or randomized assignments and the outcome variable is guaranteed to be *nonzero*: this is called the **in-sample** or **finite-sample correlation**.

Since we know that the true correlation (over the “population” of potential samples or randomized assignments) is zero, we think of the observed correlation as an **error**. In sampling, we call this the **sampling error**, and it is defined as the difference between a true population parameter and the observed mean due to chance selection of units.⁴⁸ In randomized assignment, we call this the **randomization noise**, and define it as the difference between a true treatment effect and the estimated effect due to placing units in groups. The intuition for both measures is that from any group, you can find some possible subsets that have higher-than-average values of some measure; similarly, you can find some that have lower-than-average values. Your random sample or treatment assignment will fall in one of these categories, and we need to assess the likelihood and magnitude of this occurrence. *Power calculation* and *randomization inference* are the two key tools to doing so.⁴⁹

Power calculations report the likelihood that your experimental design will be able to detect the treatment effects you are interested in given these sources of noise. This measure of **power** can be described in various different ways, each of which has different practical uses. The purpose of power calculations is to identify where the

⁴⁶ <https://blogs.worldbank.org/impactevaluations/tools-of-the-trade-doing-stratified-randomization-with-uneven-numbers-in-some-strata>

⁴⁷ Carril (2017)

⁴⁸ <https://dimewiki.worldbank.org/Sampling>

⁴⁹ https://dimewiki.worldbank.org/Sample_Size_and_Power_Calculations

strengths and weaknesses of your design are located, so you know the relative tradeoffs you will face by changing your randomization scheme for the final design.

The **minimum detectable effect (MDE)**⁵⁰ is the smallest true effect that a given research design can reliably detect. This is useful as a check on whether a study is worthwhile. If, in your field, a “large” effect is just a few percentage points or a small fraction of a standard deviation, then it is nonsensical to run a study whose MDE is much larger than that. This is because, given the sample size and variation in the population, the effect needs to be much larger to possibly be statistically detected, so such a study would never be able to say anything about the effect size that is practically relevant. Conversely, the **minimum sample size** pre-specifies expected effect sizes and tells you how large a study’s sample would need to be to detect that effect, which can tell you what resources you would need to implement a useful study.

Randomization inference⁵¹ is used to analyze the likelihood that the randomized assignment process, by chance, would have created a false treatment effect as large as the one you estimated. Randomization inference is a generalization of placebo tests, because it considers what the estimated results would have been from a randomized assignment that did not in fact happen in reality. Randomization inference is particularly important in quasi-experimental designs and in small samples, where the number of possible *randomizations* is itself small. Randomization inference can therefore be used proactively during experimental design, to examine the potential spurious treatment effects your exact design is able to produce. If there is significant heaping at particular result levels, or if results seem to depend dramatically on the outcomes for a small number of individuals, randomization inference will flag those issues before the experiment is fielded and allow adjustments to the design to be made.

⁵⁰ https://dimewiki.worldbank.org/Minimum_Detectable_Effect

⁵¹ https://dimewiki.worldbank.org/Randomization_Inference

Looking ahead

This chapter provided a framework for you to develop a data map that documents your data acquisition plan and how each component relates to the design of your study. The data map contains research variables and the instructions for using them in combination with measurement variables, which together will form the dataset for your analytical work. It also provided ways for you to use this planning data to inform and execute research design tasks, such as randomized sampling and assignment, and to produce concrete measures of whether the project design is sufficient to answer the research

questions you pose. In the next chapter, we will turn to the first step in actually answering those questions – data acquisition. The next chapter will detail the processes of obtaining original data, whether you are collecting it yourself or receiving it from another entity.

Chapter 4: Acquiring development data

Many research questions require the team to acquire original data, because no source of publicly available data addresses the inputs or outcomes of interest for the relevant population. Data acquisition can take many forms, including: primary data generated through surveys; private sector partnerships granting access to new data sources, such as administrative and sensor data; digitization of paper records, including administrative data; web scraping; primary data capture by unmanned aerial vehicles or other types of remote sensing; or novel integration of various types of datasets, such as combining survey and sensor data. Much of the recent push toward credibility in the social sciences has focused on analytical practices. However, credible development research depends, first and foremost, on the quality of the acquired data. Clear and careful documentation of the data acquisition process is essential for research reproducibility.

This chapter covers reproducible data acquisition, special considerations for generating high-quality survey data, and protocols for safely and securely handling confidential data. The first section discusses reproducible data acquisition: how to establish and document your right to use the data. This applies to all original data, whether collected for the first time through surveys or sensors or acquired through a unique partnership. The second section goes into detail on data acquisition through surveys, as this process is typically more involved than acquisition of secondary data, and has more in-built opportunities for quality control. It provides detailed guidance on the electronic survey workflow, from questionnaire design to programming and monitoring data quality. We conclude with a discussion of safe data handling, providing guidance on how to receive, transfer, store, and share confidential data. Secure file management is a basic requirement to comply with the legal and ethical agreements that allow access to personal information for research purposes.

Acquiring data from research partners

Clearly establishing and documenting data access is critical for reproducible research. This section provides guidelines for receiving data from development partners, establishing data ownership, and documenting the research team's right to use the data. It is the researchers' responsibility to respect the rights of people who own the data and the people who are described by it; but also to make sure that information is as available and accessible as possible. These twin responsibilities can and do come into tension, so it is important to be fully informed about what other researchers are doing and to fully inform other researchers of what you are doing. Writing down and agreeing to specific details is a good way of doing that.

Receiving data from development partners

Research teams granted access to existing data may receive that data in a number of different ways. You may receive access to an existing server, or physical access to extract certain information, or a one-time data transfer. In all cases, you must take action to ensure that data is transferred through secure channels so that confidentiality is not compromised. See the section *Handling data securely* later in this chapter for how to do that. Keep in mind that compliance with ethical research standards may in some cases require a stricter level of security than initially proposed by the partner agency.

At this stage, it is very important to assess documentation and cataloging of the data and associated metadata. It is not always clear what pieces of information will jointly constitute a research dataset, and many of the datasets you receive will not be organized for research. You should always retain the original data exactly as received alongside a copy of the corresponding ownership agreement or license. You should make a simple README document noting the date of receipt, the source and recipient of the data, and a brief description of each file received. All too often data will be provided as vaguely-named spreadsheets, or digital files with non-specific titles, and documentation will be critical for future access and reproducibility.

Eventually, you will want to make sure that you are creating a set of documents that can be properly submitted to a data catalog and given a reference and citation. The metadata – documentation about the data – is critical for future use of the data. Metadata should include documentation of how the data was created, what they measure, and how they are to be used. In the case of survey data, this includes the survey instrument and associated manuals; the sampling protocols and field adherence to those protocols, and any sampling weights; what variable(s) uniquely identify the dataset(s), and how different datasets can be linked; and a description of field procedures and quality controls. DIME uses as a standard the Data Documentation Initiative (DDI), which is supported by the World Bank's Microdata Catalog.¹

As soon as the desired pieces of information are stored together, think about which ones are the components of what you would call a dataset. Often, when you are receiving data from a partner, even highly-structured materials such as registers or records are not, as received, equivalent to a research dataset, and require initial cleaning, restructuring, or recombination to reach the stage we would consider a raw research dataset. This is as much an art than a science: you want to keep information together that is best contextualized together, but you also want to information granular as much as pos-

¹ <https://microdata.worldbank.org>

sible, particularly when there are varying units of observation. There usually won't be a single correct way to answer this question, and the research team will need to decide how the materials received. Soon, you will begin to build research datasets from this set of information, and these will become your original clean data, which will be the material published, released, and cited as the starting point of your data. (If funders or publishers request that "raw" data be published or cataloged, for example, this is the dataset that you should provide.) These first datasets created from the received materials are the objects you need to catalog, release, and license. Now is a good time to begin assessing disclosure risk and/or seek publication licenses in collaboration with data providers, while you are in close contact with them.

Data ownership

Except in the case of primary surveys funded by the research team, the data is typically not owned by the research team. Rather, they must enter into data licensing agreements to access the data and publish research outputs based on it. These agreements should make clear from the outset whether the research team can make the original data, any portion thereof, or derivatives² of the data, public.

Data ownership³ can sometimes be challenging to establish, as various jurisdictions have differing laws regarding data and information, and the research team may have their own information regulations. In some, data is implicitly owned by the people who it is about. In others, it is owned by the people who collected it. In still more, it is highly unclear and there are varying norms. The best approach is always to consult with a local partner, and enter into specific legal agreements establishing ownership, access, and publication rights. This is particularly critical where confidential data is involved – that is, when people are disclosing information to you that you could not obtain simply by observation or through public records.

For original data generated by the research team, such as survey data, it is important to clarify up front who owns the data, and who will have access to it. These details need to be shared with respondents when they are offered the opportunity to consent to participate in the study. If the research team is not collecting the data directly – for example, if a government, private company, or research partner is doing the data collection – make sure that you have an explicit agreement about who owns the resulting data.

The contract for data collection should include specific terms as to the rights and responsibilities of each stakeholder. It must clearly stipulate which party owns the data produced, and that the research

² **Derivatives** of data can be indicators, aggregates, visualizations etc. created from the original data.

³ https://dimewiki.worldbank.org/Data_Ownership

team maintains full intellectual property rights. The contract should also explicitly indicate that the contracted firm is responsible for protecting respondent privacy, that the data collection will not be delegated to any third parties, and that the data will not be used by the firm or subcontractors for any purpose not expressly stated in the contract, before, during or after the assignment. The contract should also stipulate that the vendor is required to comply with ethical standards for social science research, and adhere to the specific terms of agreement with the relevant Institutional Review Board (IRB)⁴ or applicable local authority. Finally, it should include policies on reuse, storage, and retention or destruction of data.

Research teams that acquire original data must also consider data ownership downstream, through the terms they will use to release that data to other researchers or to the general public. The team should consider whether they can publish the data in full after removing personal identifiers. For example, the team must consider whether it would be acceptable for their data to be copied and stored on servers anywhere in the world, whether they would prefer to manage permissions on a case-by-case basis, and whether they expect that data users would cite or credit them. Similarly, the team can require users in turn to release their derivative datasets or publications under similar licenses, or offer use without restriction. There are simple license templates for offering many of these permissions, but, at the planning stage, the team should make sure that all licensing agreements, data collection contracts, and informed consent processes used to acquire the data specifically detail those future uses.

Data licensing

Data licensing⁵ is the formal act of the dataset owner giving some data rights to a specific user, while retaining ownership of the dataset. If you are not the owner of the dataset you want to analyze, you must enter into a licensing agreement to access it for research purposes. Similarly, when you own a dataset, you must consider whether you will make the dataset accessible to other researchers, and what terms-of-use you require.

If the research team requires access to existing data for novel research, terms of use should be agreed on with the data owner, typically through a data licensing agreement. Keep in mind that the data owner is likely not highly familiar with the research process, and therefore may be surprised at some of the things you want to do if you are not clear up front. You will typically want intellectual property rights to all research outputs developed using the data, a license for all uses of derivative works, including public distribution

⁴ https://dimewiki.worldbank.org/IRB_Approval

⁵ https://dimewiki.worldbank.org/Data_License_Agreement

(unless ethical considerations contraindicate this). This is important to allow the research team to store, catalog, and publish, in whole or in part, either the original licensed dataset or datasets derived from the original. Make sure that the license you obtain from the data owner allows these uses, and that you consult with the owner if you foresee exceptions with specific portions of the data.

The World Bank has a template Data License Agreement which DIME follows. The template specifies the specific objectives of the data sharing, and whether the data can be used only for the established purpose or for other objectives after it is obtained. It classifies the data into one of four access categories, depending on who can access the data by default and whether case-by-case authorization for access is needed. The data provider may impose similar restrictions to sharing derivative data and any or all of the associated metadata. The template also specifies the required citation for the data. While you do not need to use the World Bank's template or its access categories if you do not work on a World Bank project, we still think it is important that you use this information in two ways. First, make sure to base your Data License Agreement on some template. Handwritten agreements can leave many legal ambiguities or gaps where the permissions given to the research team are unclear or incomplete. Second, we strongly recommend you to categorize data using some variation of this system. Then you should have different standard procedures for each category, so that the intended processes for handling the data are clear.

Collecting original data using electronic surveys

In this section, we detail specific considerations for acquiring high-quality data through direct surveys of study subjects. There are many survey software options available to researchers, and the market is rapidly evolving. There are also many excellent resources on questionnaire design and field supervision, but few covering the particular challenges and opportunities presented by electronic surveys. Therefore, we focus on specific workflow considerations for digitally-collected data, and on basic concepts rather than software-specific tools. If you are collecting data directly from the research subjects yourself, you are most likely designing and fielding an electronic survey. These types of data collection technologies have greatly accelerated our ability to bring in high-quality data using purpose-built survey instruments, and therefore improved the precision of research. At the same time, electronic surveys create new pitfalls to avoid. Programming surveys efficiently requires a very different mindset than simply designing them in word processing software, and ensuring

that they flow correctly and produce data that can be used in statistical software requires careful organization. This section will outline the major steps and technical considerations you will need to follow whenever you field a custom survey instrument, no matter the scale.

Developing a data collection instrument

A well-designed questionnaire results from careful planning, consideration of analysis and indicators, close review of existing questionnaires, survey pilots, and research team and stakeholder review. There are many excellent resources on questionnaire design, such as from the World Bank's Living Standards Measurement Survey.⁶ The focus of this section is the design of electronic field surveys, often referred to as Computer Assisted Personal Interviews (CAPI).⁷ Although most surveys are now collected electronically, by tablet, mobile phone or web browser, **questionnaire design**⁸ (content development) and **questionnaire programming**⁹ (functionality development) should be seen as two strictly separate tasks. Therefore, the research team should agree on all questionnaire content and design a version of the survey on paper before beginning to program the electronic version. This facilitates a focus on content during the design process and ensures teams have a readable, printable version of their questionnaire. Most importantly, it means the research, not the technology, drives the questionnaire design.

We recommend this approach because an easy-to-read paper questionnaire is very useful for training data collection staff, because it focuses on the survey content and structure before diving into the technical components. It is much easier for enumerators to understand the range of possible participant responses and how to handle them correctly on a paper survey than on a tablet, and it is easier to translate that logic to digital functionality later. Finalizing this version of the questionnaire before beginning any programming avoids version control concerns that arise from concurrent work on paper and electronic survey instruments. Finally, a readable paper questionnaire is a necessary component of data documentation, since it is difficult to work backwards from the survey program to the intended concepts.

The workflow for designing a questionnaire will feel much like writing an essay: begin from broad concepts and slowly flesh out the specifics.¹⁰ It is essential to start with a clear understanding of the **theory of change**¹¹ and **research design**¹² for your project. The first step of questionnaire design is to list key outcomes of interest, as well as the main covariates to control for and any variables needed for the specific research design. The ideal starting point for this is a

⁶ Glewwe and Grosh (2000)

⁷ [https://dimewiki.worldbank.org/Computer-Assisted_Personal_Interviews_\(CAPI\)](https://dimewiki.worldbank.org/Computer-Assisted_Personal_Interviews_(CAPI))

⁸ https://dimewiki.worldbank.org/Questionnaire_Design

⁹ https://dimewiki.worldbank.org/Questionnaire_Programming

¹⁰ https://dimewiki.worldbank.org/Questionnaire_Design

¹¹ https://dimewiki.worldbank.org/Theory_of_Change

¹² https://dimewiki.worldbank.org/Research_Design

pre-analysis plan.¹³

Use the list of key outcomes to create an outline of questionnaire *modules*. Do not number the modules; instead use a short prefix as numbers quickly get outdated when modules are reordered. For each module, determine if the module is applicable to the full sample, or only to specific respondents, and whether or how often the module should be repeated. A few examples: a module on maternal health only applies to households with a woman who has children, a household income module should be answered by the person responsible for household finances, and a module on agricultural production might be repeated for each crop the household cultivated. Each module should then be expanded into specific indicators to observe in the field.¹⁴ Questionnaires for impact evaluation must also include ways to document the reasons for **attrition** and treatment **contamination**. These are essential data components for completing CONSORT records, a standardized system for reporting enrollment, intervention allocation, follow-up, and data analysis through the phases of a randomized trial.¹⁵

At this point, it is useful to do a **survey pilot**¹⁶ of the questionnaire. A comprehensive survey pilot is a critical stage in survey design.¹⁷ The pilot must be done out-of-sample, but in a context as similar as possible to the study sample. The survey pilot includes three steps: a **pre-pilot**, a **content-focused pilot**, and a **data-focused pilot**.¹⁸ The first step is a **pre-pilot**. The pre-pilot is a qualitative exercise, done early in the questionnaire design process. The objective is to answer broad questions about how to measure key outcome variables, and gather qualitative information relevant to any of the planned survey modules. A pre-pilot is particularly important when designing new survey instruments.

The second step is a **content-focused pilot**¹⁹. The objectives at this stage are to improve the structure and length of the questionnaire, refine the phrasing and translation of specific questions, and confirm coded response options are exhaustive.²⁰ In addition, it is an opportunity to test and refine all survey protocols,²¹ such as how units will be sampled or pre-selected units identified. The content-focused pilot is best done on pen and paper, before the questionnaire is programmed, because changes at this point may be deep and structural, which are hard to adjust in code. After testing the content, proceed to program a draft version of the electronic survey instrument. The final stage is a **data-focused pilot**. The objective at this stage is to refine the questionnaire programming; this is discussed in more detail in the following section.²²

¹³ https://dimewiki.worldbank.org/Pre-Analysis_Plan

¹⁴ https://dimewiki.worldbank.org/Literature_Review_for_Questionnaire

¹⁵ Begg et al. (1996)

¹⁶ https://dimewiki.worldbank.org/Piloting_Survey_Content

¹⁷ https://dimewiki.worldbank.org/Survey_Pilot

¹⁸ https://dimewiki.worldbank.org/Structuring_a_Survey_Pilot

¹⁹ https://dimewiki.worldbank.org/Piloting_Survey_Content

²⁰ [https://dimewiki.worldbank.org/index.php?title=Checklist:_Refine_the_Questionnaire_\(Content\)](https://dimewiki.worldbank.org/index.php?title=Checklist:_Refine_the_Questionnaire_(Content))

²¹ https://dimewiki.worldbank.org/Piloting_Survey_Protocols

²² https://dimewiki.worldbank.org/Timeline_of_Survey_Pilot

Designing surveys for electronic deployment

Once the team is satisfied with the content and structure of the survey, it is time to move on to implementing it electronically. Electronic data collection has great potential to simplify survey implementation and improve data quality. Electronic questionnaires are typically developed in a spreadsheet (usually using Excel or Google Sheets) or a software-specific form builder, all of which are accessible even to novice users.²³ We will not address software-specific form design in this book; rather, we focus on coding and design conventions that are important to follow for electronic surveys regardless of software choice.²⁴ Survey software tools provide a wide range of features designed to make implementing even highly complex surveys easy, scalable, and secure. However, these are not fully automatic: you need to actively design and manage the survey. Here, we discuss specific practices that you need to follow to take advantage of electronic survey features and ensure that the exported data is compatible with your statistical software.

From a data perspective, questions with **coded response options**²⁵ are always preferable to **open-ended questions**.²⁶ The content-based pilot is an excellent time to ask open-ended questions and refine fixed responses for the final version of the questionnaire – do not count on coding up lots of free text after a full survey. Coding responses helps to ensure that the data will be useful for quantitative analysis. Two examples help illustrate the point. First, instead of asking “How do you feel about the proposed policy change?”, use techniques like **Likert scales**.²⁷ Second, if collecting data on things like medication use or food supplies, you could collect: the brand name of the product; the generic name of the product; a coded identifier for the item; or the broad category to which each product belongs (“antibiotics” or “staple foods”, for example).²⁸ All four may be useful for different reasons, but the latter two are likely to be the most useful for rapid data analysis. The coded identifiers require providing a translation dictionary to field staff, but enables automated rapid recoding for analysis with no loss of information. The broad category requires agreement on the groups of interest, but allows for much more comprehensible top-line statistics and data quality checks. Rigorous field testing is required to ensure that answer categories are comprehensive; however, it is best practice to include an *other, specify* option. Keep track of those responses in the first few weeks of field work. Adding an answer category for a response frequently showing up as *other* can save time, as it avoids extensive post-coding.

It is essential to name the fields in your questionnaire in a way that will also work in your data analysis software. Most survey

²³ https://dimewiki.worldbank.org/Questionnaire_Programming

²⁴ https://dimewiki.worldbank.org/SurveyCTO_Coding_Practices

²⁵ **Coded response options:** Responses to questions which require respondents to select from a list of choices, corresponding to underlying numerical response codes.

²⁶ **Open-ended questions:** Responses to questions which do not impose any structure on the response, typically recorded as free-flowing text.

²⁷ **Likert scale:** an ordered selection of choices indicating the respondent’s level of agreement or disagreement with a proposed statement.

²⁸ See Wafula et al. (2017) for an example.

programs will not enforce this by default, since limits vary across statistical software, and survey software will encourage you to use long sentences as question labels and detailed descriptions as choice options. This is what you want for the enumerator-respondent interaction, but you should already have analysis-compatible labels programmed in the background so the resulting data can be rapidly imported in analytical software. There is some debate over how exactly individual questions should be identified: formats like hq_1 are hard to remember and unpleasant to reorder, but formats like hq_asked_about_loans quickly become cumbersome. We recommend using descriptive names with clear prefixes so that variables within a module stay together when sorted alphabetically.²⁹ Variable names should never include spaces or mixed cases (we prefer all-lowercase naming). Take special care with the length: very long names will be cut off in some softwares, which could result in a loss of uniqueness and lots of manual work to restore compatibility. We further discourage explicit question numbering, at least at first, as it discourages re-ordering questions, which is a common recommended change after the pilot. In the case of follow-up surveys, numbering can quickly become convoluted, too often resulting in uninformative variables names like ag_15a, ag_15_new, ag_15_fup2, and so on.

²⁹ https://dimewiki.worldbank.org/Variable_Names

Using electronic survey features

Electronic surveys are more than simply a paper questionnaire displayed on a mobile device or web browser. All common survey software allows you to automate survey logic and include hard or soft constraints on survey responses. These features make enumerators' work easier, and they create the opportunity to identify and resolve data issues in real time, simplifying data cleaning and improving response quality. Well-programmed questionnaires should include most or all of the following features:

- **Localization:** the survey instrument should display full-text questions and responses in all potential survey languages, and it should also have English and code-compatible versions of all text and labels.
- **Survey logic:** built-in tests should be included for all logic connections between questions, so that only relevant questions appear, rather than relying on enumerators to follow complex survey logic. This covers simple skip codes, as well as more complex interdependencies (e.g., a child health module is only asked to households that report the presence of a child under 5).

- **Range checks:** add range checks for all numeric variables to catch data entry mistakes (e.g. age must be less than 120).
- **Confirmation of key variables:** require double entry of essential information (such as a contact phone number in a survey with planned phone follow-ups), with automatic validation that the two entries match and rejection and re-entry otherwise.
- **Multimedia:** electronic questionnaires facilitate collection of images, video, and geolocation data directly during the survey, using the camera and GPS built into the tablet or phone.
- **Preloaded data:** data from previous rounds or related surveys can be used to prepopulate certain sections of the questionnaire, and validated during the survey.
- **Filtered response options:** filters reduce the number of response options dynamically (e.g. filtering a “cities” choice list based on the state selected).
- **Location checks:** enumerators submit their actual location using in-built GPS, to confirm they are in the right place for the interview.
- **Consistency checks:** check that answers to related questions align, and trigger a warning if not so that enumerators can probe further (e.g., if a household reports producing 800 kg of maize, but selling 900 kg of maize from their own production).
- **Calculations:** make the electronic survey instrument do all math, rather than relying on the enumerator or asking them to carry a calculator.

All established survey software include debugging and test options to correct syntax errors and make sure that the survey instruments will successfully compile. This is not sufficient, however, to ensure that the resulting dataset will load without errors in your data analysis software of choice. DIME Analytics developed the `ietestform` command,³⁰ part of the Stata package `iefieldkit`, to implement a form-checking routine for SurveyCTO, a proprietary implementation of the open source **Open Data Kit (ODK)** software. Intended for use during questionnaire programming and before field work, `ietestform` tests for best practices in coding, naming and labeling, and choice lists. Although `ietestform` is software-specific, many of the tests it runs are general and important to consider regardless of software choice. To give a few examples, `ietestform` tests that no variable names exceed 32 characters, the limit in Stata (variable

³⁰ <https://dimewiki.worldbank.org/ietestform>

names that exceed that limit will be truncated, and as a result may no longer be unique). It checks whether ranges are included for numeric variables. `ietestform` also removes all leading and trailing blanks from response lists, which could be handled inconsistently across software.

The final stage of survey piloting, the data-focused pilot, should be done at this stage (after the questionnaire is programmed). The objective of this **data-focused pilot**³¹ is to validate the programming and export a sample dataset. Significant desk-testing of the instrument is required to debug the programming as fully as possible before going to the field. It is important to plan for multiple days of piloting, so that any debugging or other revisions to the electronic survey instrument can be made at the end of each day and tested the following, until no further field errors arise. The data-focused pilot should be done in advance of enumerator training.

Handling data securely

All confidential data must be handled in such a way that only people specifically approved by an Institutional Review Board (IRB) are able to access the data. Data can be confidential for multiple reasons; two very common reasons are that the data contains personally-identifying information (PII)³² or that the data owner has specified restricted access.

Basic principles of data encryption

Data encryption is a group of tools and methods to ensure that confidential files are unreadable and unusable even if laptops are stolen, servers are hacked, or unauthorized access to the data is obtained in any other way.³³ Proper encryption is central to secure data handling, and can rarely be condensed into a single tool or method, as the data will travel through many servers, devices, and computers from the source of the data to the final analysis. Encryption should be seen as a system that is only as secure as its weakest link. This section recommends a streamlined encryption workflow, so that it is easy as possible to make sure the entire chain is easy to manage and is sufficiently secure.³⁴

All encryption relies on a password or encryption key for both encrypting and decrypting information. Encryption makes data files completely unusable to anyone who obtains them if they do not have the specific decryption key. This is a higher level of security than most password-protection, because password-protected information is often readable if the password is bypassed or the storage server is

³¹ [https://dimewiki.worldbank.org/index.php?title=Checklist:_Refine_the_Questionnaire_\(Data\)](https://dimewiki.worldbank.org/index.php?title=Checklist:_Refine_the_Questionnaire_(Data))

³² [https://dimewiki.worldbank.org/Personally_Identifiable_Information_\(PII\)](https://dimewiki.worldbank.org/Personally_Identifiable_Information_(PII))

³³ <https://dimewiki.worldbank.org/Encryption>

³⁴ For more details on the topics that follow, including specific instructions for our recommended software, see Pillar 4 of the DIME Standards repository: <https://github.com/worldbank/dime-standards/>

compromised. You will need to share and store these keys carefully; if you lose them or cannot match them to encrypted information, the information is permanently lost. Therefore, you should treat access to encryption keys as equivalent to access to the confidential information. It is never secure to share these passwords or keys by email, WhatsApp or other common modes of communication; instead, use a secure password manager built for this purpose.

There are two contexts for encryption you should be aware of. **Encryption-in-transit**³⁵ protects your data when it is sent over the internet. This is a standard, passive protection that almost all internet-based services use; you only need to worry about it if you are creating a custom transfer solution. **Encryption-at-rest**³⁶. protects your data when it is stored on a server, computer, or drive.

There are two main types of encryption algorithms and they are called **symmetric encryption**³⁷ and **asymmetric encryption**³⁸. In symmetric encryption, the same key is used to both encrypt and decrypt the data. In asymmetric encryption, one key is used to encrypt data, and another key from the same “pair” is used to decrypt it. You, as a user, need to keep track of these keys. While encryption keys for asymmetric encryption are often automatically provided to the devices recording or inputting information, only people listed on your IRB should have access to decryption keys or symmetric-encryption keys.

Typically, unless you have access to an approved enterprise version of data sharing software, you will need to set up encryption at rest for your data in two locations – server or web storage during data acquisition and local storage afterwards. You should never trust that this is automatically implemented unless a cybersecurity expert within your organization has specified that a specific service is appropriate to your use case. In all other cases you should follow the steps laid out in this section, where you set up your own encryption where you are in full control of who has access to the key.

Collecting and storing data securely

Most data collection software will automatically encrypt all data in transit (i.e., upload from field or download from server). However, it is necessary to ensure that confidential data are protected when stored on a server owned by the data collection software provider or which can be accessed by people not on your research team (including your local IT or system administrators). In most data collection platforms, encryption-at-rest needs to be explicitly enabled and operated by the user. When collecting data, the tablets or the browsers used for data collection should encrypt all information be-

³⁵ https://dimewiki.worldbank.org/Encryption#Encryption_in_Transit

³⁶ https://dimewiki.worldbank.org/Encryption#Encryption_at_Rest

³⁷ https://dimewiki.worldbank.org/Encryption#Symmetric_Encryption

³⁸ https://dimewiki.worldbank.org/Encryption#Asymmetric_Encryption

fore submitting it to the server, and you should decrypt it only after downloading to your local machine.

This is a perfect use case for **asymmetric encryption** where there are two keys, forming a “public/private key pair”. The public key can safely be sent to all tablets or browsers so it can be used for encrypting the data before it is submitted to the server. Only the private key in the same key pair can then be used to decrypt that data so it can be accessed after it has been received. The private key should be kept secret and should not be shared with anyone not listed on the IRB. Again, you must store the key pair in a secure location, such as a secure note in a password manager, as there is no way to access your data if the private key is lost. If your data collection service allows you to browse data in the browser, then the encryption is only implemented correctly if you are asked for the key each time.

The data security standards that apply when receiving confidential data from the field also apply when transferring confidential data to the field, such as sampling or enumeration lists containing PII. In some survey software, you can use the same encryption that allows you to receive data securely from the field, to also send confidential data, such as an identifying list of respondents, to the field. Otherwise, you will need to create a securely stored file, transfer it to the field team using an insecure tool, and have them decrypt the information locally using a key that is transferred using a secure password manager. This process will be more similar to that for securely *storing* data, which we discuss next.

The first thing you need to do before planning how to securely send or receive data, is to plan how to securely store data after it has been transferred. Typically, you want to store your data so that you can decrypt and access it, interact with it and then encrypt it again. (Usually, you will not want to *edit* this data, but only extract non-sensitive pieces of it to an insecure location.) That is a perfect use case for **symmetric encryption** where the same key is used to both encrypt and decrypt your files. Think of this type of encryption similarly to a physical safe, where you have one key which is used to both add and access contents.

The equivalent to the safe in secure data storage is an encrypted folder, which you can set up using, for example, VeraCrypt. You can interact with files in an encrypted folder, and modify them like any unencrypted file, if and only if you have the key. This is an implementation of encryption-at-rest. There is absolutely no way to restore the data if you lose your key, so we cannot stress enough the importance of using a password manager, or equally secure solution, to store these encryption keys.

It is becoming more and more common that development research is done on data set that is too big to store on a regular computer, and instead the data is stored and processed in a cloud environment. There are many available cloud storage solutions and you need to understand how the data is encrypted and how the keys are handled. This is likely another case where a regular research team will have to ask a cybersecurity expert. After someone have helped you to set up a secure cloud storage, if you were to download a sample of the data – for example to develop your code on – then you need to remember to encrypt the data when stored on your computer.

Backing up original data

In addition to encrypting your data, you must protect it from being accidentally overwritten or deleted. This is done through a backup protocol, which creates additional copies of your original data, exactly as received and finalized, in secure locations that will remain permanently available but are not intended for frequent access. Here is an example of such a protocol:

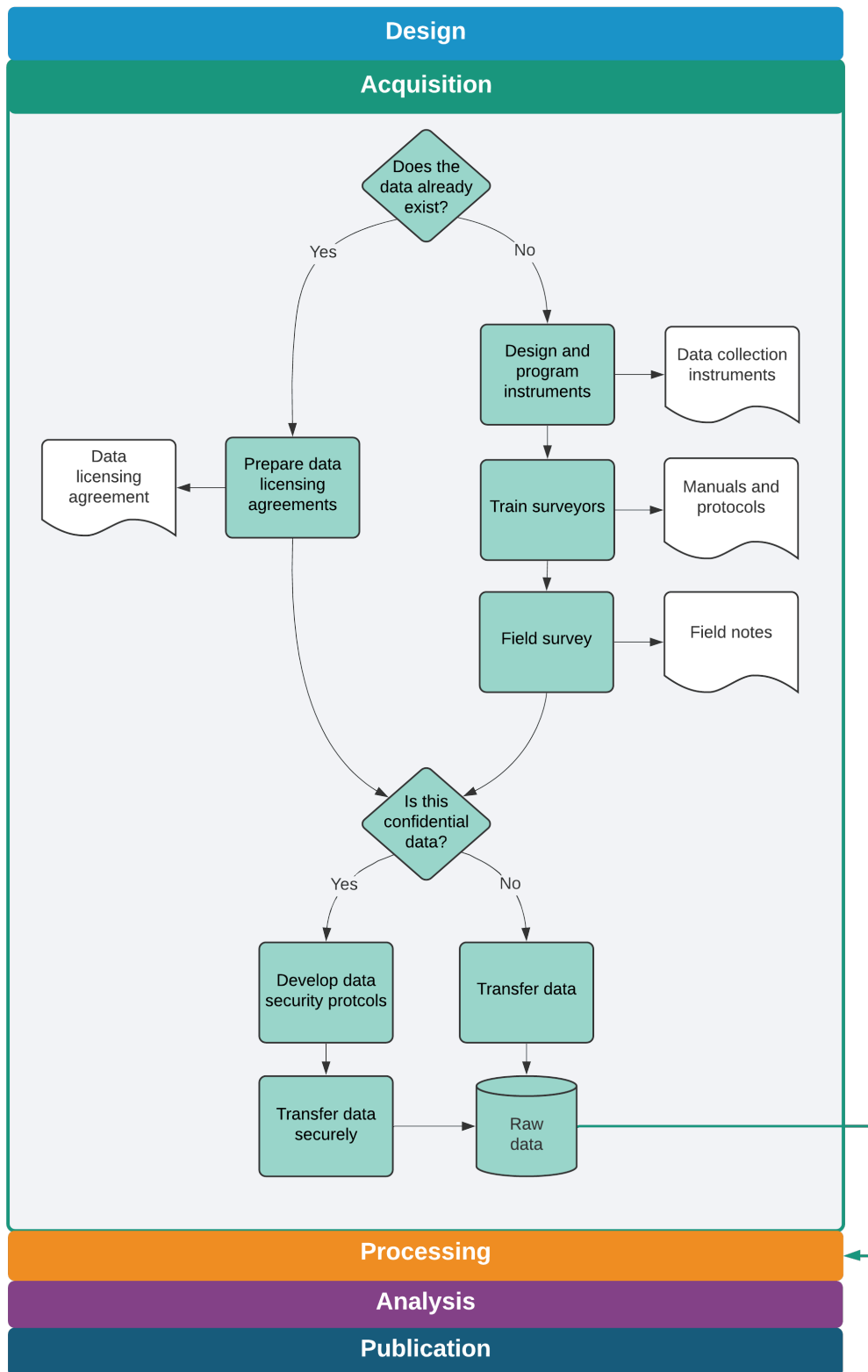
1. Create an encrypted folder in your project folder. This should be on your computer, and could be in a shared folder.
2. Download your original data from your data source to that encrypted folder. If your data source is a survey and the data was encrypted during data collection, then you will need *both* the private key used during data collection to be able to download the data, *and* the key used when you created the encrypted folder to save it there. This your first copy of your raw data, and the copy you will use for cleaning and analysis.
3. Create a second encrypted folder on an external drive that you can keep in a secure location. Copy the data you just downloaded to this second encrypted folder. This is the “master” backup copy of the raw data. You should never work with this data on a day-to-day basis. You should not use the same encrypted folder or the same key as above, because if you use the same key and lose the key, then you will have lost access to both encrypted folders. If you have an physical safe where you can securely store the external drive, then you do not need to encrypt the data and thereby do not risk losing access by losing an encryption key.
4. Finally, create a third encrypted folder. Either you can create this on your computer and upload it to a long-term cloud storage service (not a sync software), or you can create it on another external hard drive or computer that you then store in a second

location, for example, at another office of your organization. This is the “golden master” backup copy of the raw data. You should never store the “golden master” copy in a synced folder, as it would be deleted in the cloud storage if it is deleted on your computer. You should also never work with this data; it exists only for recovery purposes.

This handling satisfies the **3-2-1 rule**: there are two on-site copies of the data and one off-site copy, so the data can never be lost in case of hardware failure. If you remain lucky, you will never have to access your “master” or “golden master” copies – you just want to know it is there, safe, if you one day end up needing it.

Looking ahead

This chapter provided a road map to the acquisition of original data. It outlined guidelines for ensuring that you have effective ownership or licensing of data that you obtain or collect, and how to make those materials available in the future. It also provided an extensive guide for one of the most common – and challenging – methods of data collection: primary electronic surveys. Finally, it emphasized the secure handling of this potentially sensitive information, giving you a range of tools and a complete workflow for transferring and storing data securely at all times. Once your original data is completely transferred, securely stored, and backed up, the data acquisition stage is complete, as summarized in the figure accompanying this chapter. This is when the heavy lifting with statistical software starts. Before you can proceed to analyze your data and answer your project’s research questions, you first need to check the quality of the data acquired, make sure you have the right information in the right format, and prepare it for analysis. This process, which we call data cleaning and processing, is described in the next chapter.



Chapter 5: Cleaning and processing research data

Original data comes in a variety of formats, most of which are not immediately suited for analysis. The process of preparing data for analysis has many different names: data cleaning, data munging, data wrangling. But they all mean the same thing – transforming raw data into a convenient format for your intended use. This is the most time-consuming step of a project’s data work, particularly when primary data is involved; it is also essential for data quality. A structured workflow for preparing newly-acquired data for analysis is essential for efficient, transparent, and reproducible data work. One key point of this chapter is that no changes are made to the contents of data at this point. We consider creating new variables, imputing values and correcting outliers to be research decisions, and will discuss those in the next chapter. Therefore, the clean dataset, which is the main output from the workflow discussed in this chapter, contains the same information as the raw data, but in a format that is ready for use with statistical software.

This chapter describes the various tasks involved in making newly-acquired data ready for analysis. The first section teaches you how to make your data *tidy*. This means adjusting how the dataset is organized until the relationship between rows and columns is well-defined. The second section describes quality assurance checks, which are necessary to verify data accuracy. The third section covers de-identification, as removing direct identifiers early in the data handling process helps to ensure privacy. The final section discusses how to examine each variable in your dataset and make sure that it is as well documented and as easy to use as possible. Each of these tasks is implemented through code, and resulting datasets can be reproduced exactly by running this code. The raw data files are kept exactly as they were acquired, and no changes are made directly to them.

Making data “tidy”

The very first step in creating an analysis-friendly dataset is understanding the data acquired, and using this understanding to translate the data into an intuitive format. This section discusses what steps may be needed to make sure that each row in your **data tables**¹ represents one observation. Getting to such a format may be harder than expected, and the **unit of observation**² may be ambiguous in many raw datasets. This section will present what we call a *tidy* data format, which is, in our experience, the ideal format to handle tabular data. Although this may at first seem out of place, as it comes before quality assurance, we believe that this is the appropriate first step in data cleaning, that it should be done continuously as data is received,

¹ **Data table:** data that is structured into rows and columns. Also called *tabular datasets* or *rectangular data*. Examples of non-rectangular data are written text, NoSQL and graph databases, or files such as images.

² **Unit of observation:** the unit described by the data. In datasets, it is ideally what each row represents.
https://dimewiki.worldbank.org/Unit_of_Observation

and that quality assurance is best done using tidied data.

Unique identifiers

An important step before starting to tidy a dataset is to understand the **unit of observation**³ and find out which variable or set of variables is the **unique identifier**⁴ for each observation. As discussed in Chapter 3, the unique identifier will be used to link observations in this dataset to data in other data sources according to the **data linkage table**,⁵ and the unique identifier for all observations must be listed in the **master dataset**.⁶ Ensuring that observations are uniquely and fully identified is arguably the most important step in data cleaning. It may be the case that the variables expected to uniquely identify the raw data contain either missing or duplicate values.⁷

It is also possible for a raw dataset to not include an unique identifier, or that the identifier is not a suitable **project ID**.⁸ Suitable project IDs should, for example, not involve long strings that are difficult to work with, such as a name, or be an ID that is known outside the research team. In such cases, cleaning begins by adding a project ID to the raw data. If a project ID already exists, for this unit of observation, then you should carefully merge it from the master dataset to the raw data using other identifying information.⁹ If a project ID does not exist, then you need to generate one, add it to the master dataset, and then merge it back into the raw data. Note that while digital survey tools create unique identifiers for each data submission, that is not the same as having a unique ID variable for each observation in the sample, as there can be multiple submissions for the same observation.

DIME Analytics created an automated workflow to identify, correct and document occurrences of duplicated entries in the unique identifier using `ieduplicates` and `iecompdup`, two Stata commands included in the `iefieldkit` package.¹⁰ One advantage of using `ieduplicates`¹¹ to correct duplicated entries is that it creates *duplicates reports* which records each corrections made and documents the reason for it. Even if you are not using this command, it is important to keep a record of all cases of duplicated IDs encountered and how they were resolved.

Tidying raw data

Though raw data can be acquired in all shapes and sizes, it is most commonly received as one or multiple data tables. These data tables can organize information in multiple ways, and not all of them result in easy-to-handle datasets. Fortunately, a vast literature of database management has identified the format that makes interacting with

³ **Unit of observation:** the unit described by the data. In a data table, it is what each row represents. https://dimewiki.worldbank.org/Unit_of_Observation

⁴ https://dimewiki.worldbank.org/ID_Variable_Properties

⁵ https://dimewiki.worldbank.org/Data_Linkage_Table

⁶ https://dimewiki.worldbank.org/Master_Data_Set

⁷ We use the expression **raw data** to refer to the “data in the state it was originally received by the research team”. In other sources, you will also see it used to refer to the “corrected and compiled dataset created from received information, reflecting only that information”, which we call **clean data**. This applies to data acquired from partners as well as original data collected by the research team.

⁸ https://dimewiki.worldbank.org/Project_ID

⁹ Such operations are commonly called “merges” in Stata, and “joins” in R’s *tidyverse* dialect. We will use the term *merge* in this book.

¹⁰ <https://dimewiki.worldbank.org/iefieldkit>

¹¹ <https://dimewiki.worldbank.org/ieduplicates>

the data as easy as it can be. We call data in such format **tidy**. A data table is tidy when each column represents one **variable**,¹² each row represents one observation, and all variables in it have the same unit of observation. Every other format is *untidy*. This may seem trivial, but raw data, and raw survey data in particular, is rarely received in a tidy format.

The most common case of untidy raw data encountered in development research is a dataset with multiple units of observations stored in the same data table. Take, for example, a household survey that includes household-level questions, as well as a household member roster. Such raw datasets usually consists of a single data table where questions from the household member roster are saved in different columns, one for each member, with a corresponding member suffix, and household-level questions are represented by one column each. When your rows include multiple nested observational units, then the identifying variable does not identify all observations on that row, as there is more than one unit of observation on the same row.

Survey data containing nested units of observation is typically imported from survey platforms in **wide format**.¹³ Wide format data could have, for instance, one column for a household-level variable (for example `ownsfridge`), and a few columns for household member-level variables (for example `sex_1`, `sex_2`). Raw data is often saved in this format because it's the most efficient way to transfer it: adding different levels of observation into the same data table allows for data to be transferred in a single file. However, this leads to the widespread practice of interacting with data in wide format, although doing so is often inefficient and error-prone.

To understand how dealing with wide data can be complicated, imagine you need to calculate the share of women in each household using the household level data described above. In a wide data table you will either have to first create variables counting the number of women and the total number of household members, and then calculate the share, or you will have to transform the data to a different format. In a tidy data table, however, where each row is a household member, you can easily aggregate the share of women by household, without additional steps, and then merge the result to the household-level data tables. Tidy data tables are also easier to clean, as each attribute only needs to be checked once, and each column corresponds directly to one question in the questionnaire. Finally, as you will see in Chapter 6, summary statistics and distributions are much simpler to generate from tidy data tables.

As mentioned earlier, there are unlimited ways for data to be untidy; wide format is only one of those ways. Another example

¹² **Variable:** the collection of all data points that measure the same attribute for each observation.

¹³ **Wide data:** a data table where a single variable is divided into multiple columns, for example one for each individual in a household.

is a data table containing both information on transactions and on the firms involved in each transaction. In this case, the firm-level information will be repeated for all transactions a given firm was involved in. Analyzing data in this format would give more weight to firms that conducted more transactions, which may not be consistent with the research design.

The basic process behind tidying a data table is simple: first, identify all the variables that were measured at the same level of observation; second, create separate data tables for each level of observation; and third, reshape¹⁴ the data and remove duplicated rows until each data table is uniquely and fully identified by the identifying variable that corresponds to its unit of observation. Reshaping data tables is the most intricate task in data cleaning; you should be very familiar with commands such as `reshape` in Stata and `pivot` in R. You must be sure that identifying variables are consistent across data tables, so they can always be linked. Reshaping is the type of transformation we referred to in the example of how you calculate the share of women in a wide data set. The important difference is that in a tidy workflow, instead of transforming the data for each operation, this transformation is done once for all data during cleaning, making all subsequent operations much easier.

In the earlier household survey example, household-level variables will be stored in one tidy data table, and household-member variables are reshaped and stored in a separate, member-level, tidy data table, which also contains the household ID for each individual. The household ID is intentionally duplicated in the household members data table to allow one or several household members to be linked to the same household data. The unique identifier for the household member-level data data will be either a single household member ID or a combination of household ID and household member ID. In the transaction data example, the result of the tidying process would be one transaction-level data table, containing variables indicating the ID of all firms involved; and one firm-level data table with a single entry for each firm. Then, firm-level analysis is easily done by calculating appropriate statistics in the transactions data table (in Stata, often through `collapse`) and then merging or joining those results to the firms data table.

In a tidy workflow, your clean dataset is a set of one or more tidy data tables. In both examples above, your clean dataset is made up of two tidy data tables. There must be a clear way to connect each tidy data table to a master dataset, and thereby also to all other datasets. To implement this, you need to decide which data table is the main data table; that data table's unit of observation will be the main unit of observation of your dataset. The main unit of observation must

¹⁴ **Reshape:** transform a data table in such a way that the unit of observation represented by a row changes.

directly correspond to a master dataset, and be listed in the data linkage table. All other data tables in your dataset must have an unambiguous way to merge with the main data table. This way, it will be possible to link all data points in all your project's datasets to each other. We recommend that you save your datasets as a folder, in which the main data table shares the same name as the folder, and the name of all other data tables start with the same name, but are suffixed with the unit of observation for that data table.

In the household dataset example, the household-level data table would be the main table. This means that there must be a master dataset for households. (You may have a master dataset for household members as well if you think it is important for your research, but it is not strictly required.) The household data set would then be stored in a folder called, for example, `baseline-hh-survey/`. In that folder you would save both the household-level data table with the same name as the folder, for example `baseline-hh-survey.csv`, and the household member-level data named in the same format but with a suffix, for example `baseline-hh-survey-hhmember.csv`.

The tidying process gets more complex as the number of nested groups increases. That means the steps of identifying the unit of observation of each variable and reshaping the separated data tables need to be repeated multiple times. However, the more nested groups a dataset includes, the more efficient it is to deal with tidy data as compared to untidy. Cleaning and analyzing wide datasets, in particular, is a repetitive and error-prone process.

The next step of data cleaning, data quality monitoring, may involve comparing different units of observation. Aggregating sub-units to compare to a higher unit is much easier with tidy data, which is why we suggest tidying data as the first step in the cleaning workflow. If you are conducting primary data collection, you can start preparing or coding the data tidying even before the data is acquired, since you will know in advance the exact format in which the data will be received. In the case of survey data, tidying datasets will guarantee a one-to-one correspondence between questions in the questionnaire and columns in the data. Preparing the data for analysis, the last task in this chapter, is much simpler when that is the case.

Data quality assurance

Whether you are acquiring data from a partner or collecting it directly, it is important to make sure that data faithfully reflects ground realities. You should carefully examine and clean any data you are about to use. When reviewing raw data, you will inevitably en-

counter data entry mistakes, such as typos and inconsistent values. Data quality assurance requires a combination of real-time data checks and back-checks or validation audits, which often means tracking down the people whose information is included in the dataset.

Implementing high-frequency quality checks

A key advantage of continuous electronic data ingestion methods, as compared to traditional paper surveys and one-time data dumps, is the ability to access and analyze the data acquisition process is ongoing. Data issues can then be identified and resolved in real-time. Designing systematic data checks and running them routinely throughout data intake simplifies monitoring and improves data quality. To prepare for data acquisition, the research team should develop a **data quality assurance plan**.¹⁵ While data is being acquired, a research assistant should work closely with the field team or data licensing partner to ensure that the data is flowing as expected, and to set up and perform **high-frequency checks (HFCs)** with the incoming data.

High-frequency checks should carefully inspect key treatment and outcome variables to ensure that the data quality of core study variables is uniformly high, and that additional effort is centered where it is most important. Data quality checks should be run every time data is received to flag irregularities in the acquisition progress, in sample completeness, or in response quality. The faster issues are identified, the more likely they are to be solved. `ipacheck`¹⁶ is a very useful Stata command that automates some of these tasks, regardless of the data source.

It is important to check continuously that the observations received match the intended sample. In surveys, electronic survey software often provides case management features through which sampled units are directly assigned to individual enumerators. For data received from partners, such as administrative data, this may be harder to validate. In these cases, cross-referencing with other data sources can help to ensure completeness. It is often the case that raw data includes duplicate or missing entries, which may occur due to typos, failed submissions to data servers, or other mistakes.¹⁷ Issues with data transmission often result in missing observations, particularly when large datasets are being transferred, or when data is being collected in locations with limited internet connection. Keeping a record of what data was submitted, and comparing it to the data received as soon as transmission is complete reduces the risk of noticing that data is missing when it is no longer possible to recover

¹⁵ https://dimewiki.worldbank.org/Data_Quality_Assurance_Plan

¹⁶ <https://github.com/PovertyAction/high-frequency-checks>

¹⁷ https://dimewiki.worldbank.org/Duplicates_and_Survey_Logs

it.

Once data completeness is confirmed, observed units must be validated against the expected sample: this is as straightforward as merging the sample list with the data received and checking for mismatches. Reporting errors and duplicate observations in real time allows for efficient corrections. `ieduplicates`¹⁸ provides a workflow for resolving duplicate entries with the data provider. For surveys, it is also important to track data collection progress to monitor attrition, so that it is clear early on if a change in protocols or additional tracking will be needed.¹⁹ Remember to also check survey completion rates and sample compliance by surveyors and survey teams, and compare data missingness across administrative regions, to identify any clusters that may be providing data of suspect quality.

High frequency checks should also include checks of response quality and consistency. For example, whether the values for each variable fall within the expected range and the answers provided by the same household are consistent across all survey modules.²⁰ Electronic survey and data entry software often incorporate many quality control features, such as range restrictions and logical flows. Data received from systems that do not include such controls should be checked more carefully. Response consistency should be checked across all datasets, as this is much harder to automate. Consistency checks are project specific, so it is difficult to provide general guidance. A detailed knowledge of the variables in the dataset and a careful examination of the analysis plan is the best way to prepare. Analysis of metadata and paradata can also be useful in assessing data quality. For example, electronic survey software generates automatically collected timestamps and trace histories, showing when data was submitted, how long enumerators spent on each question, and how many times answers were changed before or after the data was submitted.

High-frequency checks will only improve data quality if the issues they catch are communicated to the data provider and corrections are documented and applied to the data. There are lots of ways to do this; what's most important is to find a way to create actionable information for your team. `ipacheck`, for example, generates a spreadsheet with flagged errors; these can be sent directly to the data collection teams. Many teams choose other formats to display results, such as online dashboards created by custom scripts. It is also possible to automate communication of errors to the field team by adding scripts to link the HFCs with a messaging platform. Any of these solutions are possible: what works best for your team will depend on such factors as cellular networks in field work areas, whether field supervisors have access to laptops, internet speed, and coding skills

¹⁸ <https://dimewiki.worldbank.org/ieduplicates>

¹⁹ See Özler et al. (2016) for an example.

²⁰ https://dimewiki.worldbank.org/Monitoring_Data_Quality

of the team preparing the HFC workflows.

Conducting back-checks and data validation

Careful validation of data is essential for high-quality original data. While we cannot control natural measurement error,²¹ which comes from variation in the realization of key outcomes, there is often an opportunity to reduce error arising from inaccuracies in the data generation process. **Back-checks**²² and other validation audits help ensure that data is not falsified, incomplete, or otherwise suspect. For survey data, this can also be a means to ensure that all field protocols are followed. For back-checks and validation audits, a random subset of observations is selected, and a subset of information from the full survey is verified through a brief targeted survey with the original respondent or a cross-referenced dataset from another source (if the original data is not a field survey). Design of the back-checks or validations follows the same survey design principles discussed above: you should use the analysis plan or list of key outcomes to establish which subset of variables to prioritize, and similarly focus on errors that would be major flags for poor quality data.

Real-time access to the data massively increases the potential utility of validation, and both simplifies and improves the rigor of the associated workflows. You can use the raw data to draw the back-check or validation sample; this ensures that the validation is correctly apportioned across observations. As soon as high frequency checks are complete, the validation data can be tested against the original data to identify areas of concern in real-time. The `bcstats` command is a useful tool for analyzing back-check data in Stata.²³ Some electronic surveys surveys also provide a unique opportunity to do audits through audio recordings of the interview, typically short recordings triggered at random throughout the questionnaire. **Audio audits**²⁴ are a useful means to assess whether enumerators are conducting interviews as expected. Do note, however, that audio audits must be included in the informed consent for the respondents, and the recordings will need to be assessed by specially trained staff.

De-identifying research data

When implementing the steps discussed up to this point, you are likely to be handling confidential data. Effective data quality monitoring frequently requires you to identify the individual observations in your dataset, and the people or other entities who provided the information. Using identified data allows you to quickly follow up on and resolve identified issues. Handling confidential data such as

²¹ See Kondylis et al. (2015) for an example.

²² https://dimewiki.worldbank.org/Back_Checks

²³ White (2016)

²⁴ https://dimewiki.worldbank.org/Random_Audio_Audits

personally-identifying information requires a secure environment and, typically, decryption. De-identifying the data will allow you to simplify that workflow, and will also reduce the risk of harmful leaks. This section describes how to de-identify data in order to share it with a wider audience.

Protecting privacy as a researcher

Most development data involves human subjects.²⁵ As a researcher, you may have access to personal information about your subjects: where they live, how much income they have, whether they have committed or been victims of crimes, their names, their national identity numbers, and other sensitive data.²⁶ There are strict requirements for safely storing and handling personally-identifying data, and it is the responsibility of the research team to satisfy these requirements.²⁷ Everyone working with human subjects research should have completed an ethics certification course.²⁸ A plan for secure data handling is typically also required for IRB approval.

The best way to avoid risk is to minimize interactions with PII as much as possible. First, only collect personally-identifying information that is strictly necessary for the research. Second, avoid the proliferation of copies of identified data. There should never be more than one copy of the raw identified dataset in the working project folder, and it must always be encrypted. Third, de-identify the data as early as possible in the workflow. Even within the research team, access to the identified data should be limited to team members who require it for their specific tasks. Data analysis that requires identifying information is rare and in most cases can be avoided by properly linking masked identifiers to research information such as treatment statuses and weights, then removing unmasked identifiers.

Once data is acquired and the data quality checks described above are completed, the next task is typically to **de-identify** the data, by removing or masking all personally-identifying variables.²⁹ Note that it is in practice impossible to **anonymize** data. There is always some statistical chance that an individual's identity will be re-linked to the stored data – even if that data has had all directly identifying information removed – by using some other data that becomes identifying when integrated. For this reason, we typically recommend de-identification in two stages. The **initial de-identification** process, performed as soon as data is acquired, strips the data of direct identifiers, to create a working de-identified dataset that can be shared *within the research team* without the need for encryption. The **final de-identification** process, performed before data is publicly released, involves careful consideration of the trade-offs between risk

²⁵ https://dimewiki.worldbank.org/Human_Subjects_Approval

²⁶ See Banerjee et al. (2019) for an example.

²⁷ https://dimewiki.worldbank.org/Research_Ethics

²⁸ Protecting Human Research Participants (<https://phrptraining.com>) and the CITI Program (<https:////citiprogram.org>) are common options.

²⁹ <https://dimewiki.worldbank.org/De-identification>

of identifying individuals and the utility of the data, and typically requires the removal of a further level of indirect identifiers. The rest of this section describes how to implement both the initial and the final de-identification processes.

De-identification in practice

Initial de-identification reduces risk and simplifies workflows. Once you create a de-identified version of the dataset, you no longer need to interact directly with the encrypted data. Note that if the data tidying resulted in multiple raw data tables, each will need to be de-identified separately, but the workflow will be the same for all of them.

During the initial round of de-identification, datasets must be stripped of personally identifying information.³⁰ To do so, you will need to identify all variables that contain such information. For data collection, where the research team designs the survey instrument, flagging all potentially identifying variables at questionnaire design stage simplifies the initial de-identification process. If you did not do that, or you received original data by another means, there are a few tools to help flag variables with personally-identifying data. JPAL's PII-scan and IPA's PII_detection, scan variable names and labels for common string patterns associated with identifying information.³¹ The World Bank's sdcMicro lists variables that uniquely identify observations, but its more refined method and higher processing capacity requirement makes it better suited for final de-identification.³² The iefieldkit command iecodebook lists all variables in a dataset and exports an Excel sheet where you can easily select which variables to keep or drop.³³

Once you have a list of variables that contain confidential information, assess them against the analysis plan and first ask yourself for each variable: *will this variable be needed for the analysis?* If not, the variable should be dropped. Don't be afraid to drop too many variables the first time, as you can always go back and extract additional variables from the raw data, but you cannot go back in time and drop a PII variable that was leaked.

For each confidential variable that is needed in the analysis, ask yourself: *can I encode or otherwise construct a variable that masks the confidential component, and then drop this variable?* For example, it is easy to encode identifiers for small localities like villages and only provide a meaningless numerical indicator showing which observations are in the same village without revealing which villages are included in the data. This is typically the case for most identifying information. If the answer to either of the two questions above is yes, all you need

³⁰ <https://dimewiki.worldbank.org/De-identification>

³¹ <https://github.com/J-PAL/PII-Scan> and https://github.com/PovertyAction/PII_detection

³² Benschop et al. (2019)

³³ <https://dimewiki.worldbank.org/iecodebook>

to do is write a script to drop the variables that are not required for analysis, encode or otherwise mask those that are required, and save a working version of the data. For example: after constructing measures of distance or area, drop the specific geolocations in the data; after constructing and verifying numeric identifiers in a social network module, drop all names. If confidential information is strictly required for the analysis and cannot be masked or encoded, it will be necessary to keep at least a subset of the data encrypted through the data analysis process.

After initial de-identification is complete, your dataset will consist of one or multiple tidy, de-identified data tables. This is the dataset that you will interact with during the remaining tasks described in this chapter. Initial de-identification should not affect the usability of the data. Note that access to the initially de-identified data should still be restricted to the research team only, as indirect identifiers may still present a high risk if disclosure. It is common, and even desirable, for teams to make data publicly available once the tasks discussed in this chapter are concluded. This will allow other researchers to conduct additional analysis and to reproduce your finding. Before that can be done, however, you should further consider whether your data can be re-identified, in a process we call **final de-identification**, which will be discussed in more detail in Chapter 7.

Cleaning and preparing data for analysis

The last step in the data cleaning process involves making the dataset easy to use and understand, and carefully examining each variable to document distributions and identify patterns that may bias the analysis. The resulting dataset will contain only the variables collected in the field, and no modifications to data points will be made, except for corrections of mistaken entries. You may have more data tables in your dataset now than originally received, and they may have a different *format*, but the information contained is still the same. Apart from the **cleaned dataset** (or datasets) itself, cleaning will also yield extensive documentation describing it.

Data cleaning yields in-depth understanding of the contents and structure of your data. This knowledge will be key to correctly constructing and analyzing final indicators, which we cover in the next chapter. Do not rush through this step! It is common for data cleaning to be the most time-consuming task in a project. In this section, we introduce some concepts and tools to make it more efficient and productive. The section is separated into three subtopics: exploring the data, making corrections, and recoding and annotating.

They are separated here because they are different in nature, and should be kept separated in your code. In practice, however, they may all done at the same point in time.

Exploring the data

The first time you interact with the data contents is during quality checks. However, these checks are usually time-sensitive, and there may not be time to explore the data at length. During data cleaning, on the other hand, you will need to inspect each variable closely. Use tabulations, summary statistics, histograms and density plots to understand the structure of data, and look for patterns or irregularities. Think critically about what you see. You should ensure that the numerical values that appear are consistent with the information the variable represents. You should ensure that statistical distributions look realistic and are not highly clumped or skewed. You should confirm that related variables are consistent with each other. You should check for outliers and missing values. Then, you should assess if unusual or unexpected distributional patterns of any of these characteristics could be caused by data entry errors.

At this point, it is more important to document your findings than to directly address any irregularities found. There is a very limited set of changes that should be made to the raw data during cleaning. They are described in the next two sections, and are usually applied to each variable as you examine it. Most of the transformations that result in new variables will be done during **data construction**,³⁴ a process discussed in the next chapter. For now, focus on creating a record of what you observe, and extensively documenting the data being explored. You will use this documentation when discussing with your team how to address irregularities once you get to the construction stage. This material will also be valuable during exploratory data analysis.

³⁴ **Data construction:** The process of creating complex or abstract measures from raw information that is directly observed or collected.

Correcting data points

As mentioned earlier, corrections to issues identified during data quality monitoring are the only changes done to individual data points during the data cleaning stage. However, there is a lot of discussion about whether one should modify such data points at all. Some argue that follow-ups to the issues identified are costly and add limited value. Since it is not possible to check each and every possible data entry error, doing so can create a false sense of security from issues identified on a few main variables. Additionally, manually-inspected data may suffer from considerable inspector variability. In many cases, the main purpose of data quality checks is

to detect fraud and identify problems with data collection protocols. On the other hand, there is also an argument to be made against keeping clear typing errors or not correcting missing values. We recommend correcting any entries that are clearly identified as errors. However, there is some subjectivity involved in deciding which cases fall into this category. A common rule of thumb is to include the set of corrections which are based on information that you have privileged access to and other research teams would not be able to make, and no more. Making this such decisions involve deep knowledge of the data and the particular circumstances of each research project.

Whether you decide to modify your data or not, you must keep a careful record of all issues that you identify. If no data points are modified, it may still be helpful to add flags to observations containing potentially problematic values, so you can verify how they affect results during analysis. If your team decides to follow up on and correct these issues, the follow-up process must also be thoroughly documented. Be very careful not to include confidential information in documentation that is not securely stored, or that you intend to release as part of a replication package or data publication. Finally, remember not to make changes directly to the raw data. Instead, any corrections must be done as part of data cleaning, applied through code, and saved to a new intermediate dataset.

Recoding and annotating data

The cleaned dataset is the starting point of data analysis. It will be extensively manipulated to construct analysis indicators, so it is important for it to be easily processed by statistical software. To make the analysis process smoother, anyone opening this dataset for the first time should have all the information needed to interact with it, even if they were not involved in the acquisition or cleaning process. This will save them time going back and forth between the dataset and its accompanying documentation.

Often times, datasets are not imported into statistical software in the most efficient format. The most common example is string (text) variables: categorical variables and open-ended responses are often read as strings. However, variables in this format cannot be used for quantitative analysis. Therefore, categorical variables must be transformed into other formats, such as `factors` in R and `labeled integers35` in Stata. Additionally, open-ended responses stored as strings usually have a high risk of including identifying information, so cleaning them requires extra attention. The choice names in categorical variables (called *value labels* in Stata and *levels*

³⁵ https://dimewiki.worldbank.org/Data_Cleaning#Value_Labels

in R) should be accurate, concise, and directly linked to the data collection instrument. Adding choice names to categorical variables makes it easier to understand your data as you explore it, and thus reduces the risk of small errors making their way through into the analysis stage.

In survey data, it is common for non-responses such as “Don’t know” and “Declined to answer” to be represented by arbitrary survey codes. The presence of these values could bias your analysis, since they don’t represent actual observations of an attribute. They need to be turned into *missing values*. However, the fact that a respondent didn’t know how to answer a question is also useful information that would be lost by simply omitting all information. In Stata, this information can be elegantly conserved using extended missing values.³⁶

We recommend that the cleaned dataset be kept as similar to the raw data as possible. This is particularly important regarding variable names: keeping them consistent with the raw data makes data processing and construction more transparent. Unfortunately, not all variable names are informative. In such cases, one important piece of documentation makes the data easier to handle: the variable dictionary. When a data collection instrument (for example a questionnaire) is available, it is often the best dictionary one could ask for. But even in these cases, going back and forth between files can be inefficient, so annotating variables in a dataset is extremely useful. In Stata, *variable labels*³⁷ must always be present in a cleaned dataset. They should include a short and clear description of the variable. A lengthier description, that may include, for example, the exact wording of a question, may be added through *variable notes*. In R, it is less common to use variable labels, and a separate dataset with a variable dictionary is often preferred, but `data frame` attributes can be used for the same purpose.

Finally, any information that is not relevant for analysis may be removed from the dataset. In primary data, it is common to collect information for quality monitoring purposes, such as notes, duration fields and surveyor IDs. Once you are past the quality monitoring phase, these variables may be removed from your dataset. In fact, to make the data easier to handle, you may choose to start from a minimal set of variables, and add new ones as you clean them. To ensure the cleaned dataset file doesn’t get too big to be handled, use commands such as `compress` in Stata so the data is always stored in the most efficient format.

Although all these tasks are key to making the data easy to use, implementing them can be quite repetitive and create convoluted scripts. The `iecodebook` command suite, part of the `iefieldkit` Stata

³⁶ https://dimewiki.worldbank.org/Data_Cleaning#Survey_Codes_and_Missing_Values

³⁷ https://dimewiki.worldbank.org/Data_Cleaning#Variable_Labels

package, is designed to make some of the most tedious components of this process more efficient.³⁸ It also creates a self-documenting workflow, so your data cleaning documentation is created alongside that code, with no extra steps. As far as we know, currently there are no similar resources in R. However, the `tidyverse`³⁹ packages compose a consistent and useful grammar to perform the same tasks.

Documenting data cleaning

Throughout the data cleaning process, you will often need extensive inputs from the people responsible for data collection. Sometimes this is your research team, but often it will be someone else. It could be a survey team, the government ministry responsible for administrative data systems,⁴⁰ the technology firm that generated remote sensing data, etc. Regardless of who originally collected the data, you should acquire and organize all documentation of how the data was generated.⁴¹ What type of documentation that is available depends on how the data was collected. For original data collection, this should include field protocols, data collection manuals, survey instruments, supervisor notes, and data quality monitoring reports. For secondary data, you should try to get the same type of information, but that is often not possible unless the data source is a well managed data publication. Independently of its exact composition, the data documentation should be stored alongside your data dictionary and codebooks. You will probably need these files during analysis, and they should be published along with the data, so other researchers may use them for their analysis as well.

Looking ahead

This chapter introduced a workflow of formatting, cleaning, and quality assurance for the data that you obtained from the field or from partners, illustrated in the figure that follows. These tasks create your first research outputs using original data: an analysis-ready dataset. This dataset is well-structured to describe your units of analysis (it is “tidy”), it faithfully represents the measurements it was intended to collect, and it does not expose the identities of the people described by it. You have also now taken the time to fully understand the patterns and structures in your data, and annotated and labeled them for your use and use by others. Combined with the data map, this dataset will be the fundamental starting point for all analysis work. In the next chapter, we will walk through the steps needed to run the analyses originally specified in your analysis plan and answer your research question – or perhaps to find out you now have even

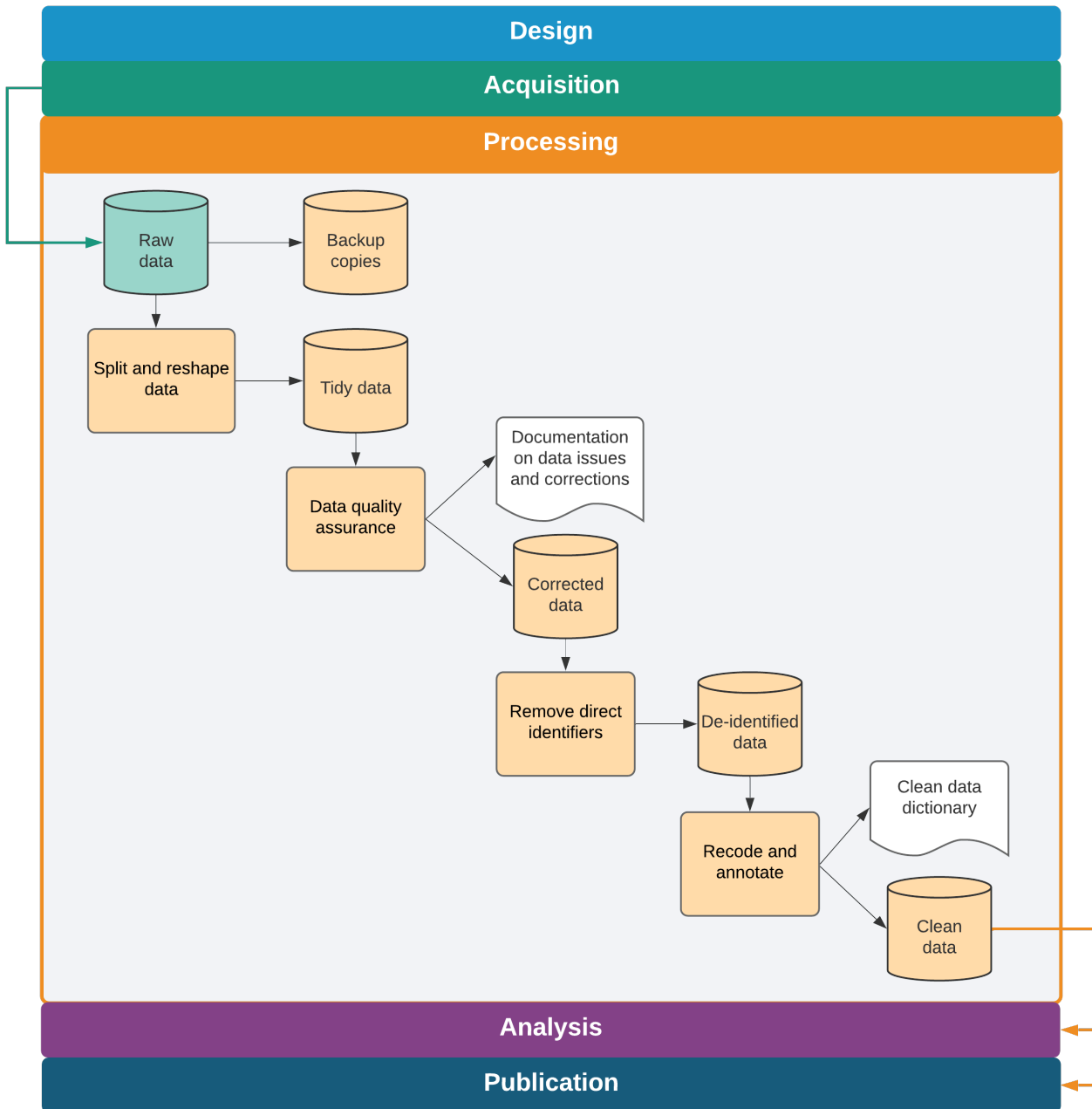
³⁸ <https://dimewiki.worldbank.org/icodebook>

³⁹ <https://www.tidyverse.org/>

⁴⁰ See Fernandes et al. (2015) for an example.

⁴¹ https://dimewiki.worldbank.org/Data_Documentation

more questions.



Chapter 6: Constructing and analyzing research data

The process of data analysis is typically a back-and-forth discussion between people with differing skill sets. To effectively do this in a team environment, data, code and outputs must be well-organized, with a clear system for version control, analysis scripts that can be run by all team members, and creation of research outputs fully automated. Putting in time upfront to structure the data analysis workflow in this way pays substantial dividends throughout the process. Transparently documenting research decisions made during data analysis is essential for reproducible research.

In this chapter, we discuss the process of transforming cleaned raw data into the final analysis dataset(s). The suggested workflow starts from clean raw data (the output of data cleaning, discussed in the previous chapter). The first section covers variable construction: transforming the raw data into economically meaningful indicators. The second section discusses the analysis itself. We do not offer instructions on how to conduct specific analyses, as that is determined by research design, and there are many excellent existing guides. Rather, we discuss how to structure analysis code so that it is easy to follow and understand, for both the full research team members and a public audience. The final section discusses ways to automate common outputs so that your work is fully reproducible.

Creating analysis datasets

For this chapter, we assume you are starting from one or multiple well-documented tidy¹ datasets. We also assume that these datasets have gone through thorough quality checks and incorporate any corrections needed. The next step is to **construct**² the variables that you will use for analysis; that is, to transform the cleaned data into analysis-ready data. It's possible the acquired data is ready for analysis, but in most cases it needs to be prepared by integrating different datasets and creating derived variables (dummies, indices, and interactions, to name a few³). The derived indicators you will construct should be planned during research design; the pre-analysis plan should serve as your guide at this stage. During construction, data will typically be reshaped, merged, and aggregated to change the level of the data points from the **unit of observation** in the raw data to the **unit of analysis**.⁴

Each analysis dataset is built to answer an analysis question. If the sub-samples and units of observation vary for different pieces of the analysis, you may need to create many purpose-built analysis

¹ Wickham and Grolemund (2017)

² **Data construction:** The process of transforming cleaned data into analysis data by creating the derived indicators that will be analyzed.

³ See Adjognon et al. (2019) for an example.

⁴ https://dimewiki.worldbank.org/Unit_of_Observation

datasets. In such cases, it is not good practice to try to create a single “one-size-fits-all” analysis dataset. For a concrete example of what this means, think of an agricultural intervention that was randomized across villages and only affected certain plots within each village. The research team may want to run household-level regressions on income, test for plot-level productivity gains, and check if village characteristics are balanced. Having three separate datasets for each of these three pieces of analysis will result in cleaner, more efficient, and less error-prone analytical code than if you started from a single analysis dataset and repeatedly transformed it.

Fitting construction into the data workflow

Construction follows data cleaning and should be treated as a separate task for two reasons. First, this helps to clearly differentiate error corrections (necessary for all data uses) from creation of analysis indicators (necessary only for specific analyses). Second, it helps to ensure that variable definitions are consistent across datasets. Construction can create many outputs combining many inputs, and, unlike in data cleaning, resulting datasets should be concise rather than exhaustive. For example, take a project that has a baseline and an endline survey. Unless the two data collection instruments are exactly the same, which is preferable but often not the case, the data cleaning for each of these rounds will require different steps, and therefore will be done separately. However, the analysis indicators must be constructed in the exact same way, so they are comparable. To do this, you will require at least two separate cleaning scripts, and a unified construction script. Maintaining one construction script guarantees that you will not accidentally make changes to an indicator from one round while forgetting to update the other.

When we visualize the research workflow, variable construction precedes data analysis, as derivative variables need to be created before they are analyzed. In practice, however, as you analyze the data, it is often useful to revisit construction, and explore different subsets of the same and other transformations of the data. Even if construction and analysis are done concurrently, you should always do the two in separate scripts. If every script that creates a table starts by loading a dataset, subsetting it, and manipulating variables, any edits to construction need to be replicated in all scripts. This increases the chances that at least one of them will have a different sample or variable definition. Doing all variable construction and data transformation in a unified script, separate from the analysis code, helps avoid this and ensures consistency across different outputs.

Integrating different data sources

To create the analysis dataset, it is typically necessary to combine information from different data sources.⁵ As discussed in Chapter 3, this process should be documented using **data flowcharts**,⁶ and different data sources should only be combined in accordance with the data linkage table.⁷ For example, you may merge administrative data with survey data in order to include demographic information in your analysis, or you may want to integrate geographic information in order to include location-specific controls. To understand how to perform such operations, you will need to consider the unit of observation for each dataset, and their respective identifying variables. Merges are frequent and complex operations, which makes them a common source of error. Whichever statistical software you are using, take the time to read through the help file of merge commands and make sure you understand their options and outputs.

When writing the code to implement merge operations, a few steps can help avoid mistakes. First, before writing code to combine the datasets, write pseudocode to understand which observations you expect to be matched or not, and why. When possible, determine exactly which and how many matched and unmatched observations should result from the merge. The best tool you have to understand this is the three components of the data map discussed in chapter 3. Second, think carefully about whether you want to keep matched and unmatched observations, or only specific matching outcomes (e.g. to create a balanced panel), and add that to the pseudocode as well. Finally, run the code to merge the datasets, and compare outcome to your expectations. Add comments to explain any exceptions, and make it so the code will return an error in case unexpected results show up in future runs.

To avoid unintentional changes to your data, pay close attention to merge results. Two issues to pay extra attention to are missing values and dropped observations. Make sure to read about how each command treats missing observations: are unmatched observations dropped, or are they kept with missing values? Whenever possible, add automated checks in the script that throw an error message if the result is different than what you expect, or you may not changes in the outcome running large portions of code. Document changes in the number of observations in your comments, and explain why they are happening. If your are subsetting your data by keeping only matched observations, write down the reason why the observation differ across datasets, as well as why you are only interested in those that matched. The same applies when you are adding new observations from the merged dataset.

⁵ [https://dimewiki.worldbank.org/
Data_Integration](https://dimewiki.worldbank.org/Data_Integration)

⁶ [https://dimewiki.worldbank.org/
Data_Flow_Charts](https://dimewiki.worldbank.org/Data_Flow_Charts)

⁷ [https://dimewiki.worldbank.org/
wiki/Data_Linkage_Table](https://dimewiki.worldbank.org/wiki/Data_Linkage_Table)

Some merges of data with different units of observation are more conceptually complex. Examples include: overlaying road location data with household data, using a spatial match; combining school administrative data, such as attendance records and test scores, with student demographic characteristics from a survey; or linking a dataset of infrastructure access points, such as water pumps or schools, with a dataset of household locations. In these cases, a key part of the research contribution is figuring out a useful way to combine the datasets. Since the conceptual constructs that link observations from the two data sources are important and can take many possible forms, it is especially important for the data integration to not be treated mechanically, and to be extensively documented, separately from other data construction tasks.

Creating analysis variables

Once you have assembled variables from different sources into a single working dataset with the right raw information and observations, it's time to create the derived indicators of interest for analysis. Before constructing new indicators, you must check and double-check units, scales, and value assignments of each variable that will be used. This is when you will use the knowledge of the data and the documentation developed during cleaning the most. First, check that all categorical variables have the same value assignment, such that labels and levels have the same correspondence across variables that use the same options. For example, it's possible that in one question 0 means "no" and 1 means "yes", while in another one the same answers were coded as 1 and 2. (We recommend coding binary questions as either 1 and 0 or TRUE and FALSE, so they can be used numerically as frequencies in means and as dummies in regressions. This often implies re-expressing categorical variables like gender as binary variables like woman.) Second, make sure that any numeric variables you are comparing are converted to the same scale or unit of measure: you cannot add one hectare and two acres and get a meaningful number. New variables should be assigned functional names, and the dataset ordered such that related variables are together. Adding notes to each variable will make your dataset more user-friendly.

At this point, you will also need to decide how to handle any outliers or unusual values identified during data cleaning. How to treat outliers is a research question.⁸ There are multiple possible approaches, and the best choice for a particular case will depend on the objectives of the analysis. Whatever your team decides, make sure to explicitly note what the decision was and how it was made. Results can be sensitive to the treatment of outliers, so keeping the

⁸ https://dimewiki.worldbank.org/Variable_Construction#Dealing_with_outliers

original variable in the dataset will allow you to test how much it affects your outputs. All these points also apply to imputation of missing values and other distributional patterns. As a general rule, never overwrite or delete original data during the construction process. Always create derived indicators with new names.

Finally, creating a panel with survey data involves additional timing complexities. It is common to construct indicators soon after receiving data from a new survey round. However, creating indicators for each round separately increases the risk of using different definitions each time. Having a well-established definition for each constructed variable helps prevent that mistake, but the best way to guarantee it won't happen is to create the indicators for all rounds in the same script. Say you constructed some analysis variables after baseline, and are now receiving midline data. Then the first thing you should do is create a cleaned panel dataset, ignoring the previous constructed version of the baseline data. Our team created `iefieldkit's iecodebook append` subcommand to help you reconcile and append data from cleaned survey rounds or similar data collected from different contexts. This is done by completing an Excel sheet to indicate what changes in names, value assignments, and value labels should be made so the data is consistent across rounds or settings.⁹ By doing so, you are also creating helpful documentation about your data work. Once data tables are consistently appended, adapt your construction script so it can be used on the complete panel dataset. In addition to preventing inconsistencies and documenting your work, this process will also save you time and give you an opportunity to review your original code.

⁹ See Daniels et al. (2017) for an example.

Documenting variable construction

Because data construction involves translating concrete observed data points to measures of abstract concepts, it is important to document exactly how each variable is derived or calculated. Careful documentation is closely linked to the research principles discussed in the first chapter. It makes research decisions transparent, as anyone can read about how you defined each variable in your analysis, and what was the reasoning behind these decisions. By reading the documentation, someone unfamiliar with the project should be able to understand the contents of the analysis datasets, the steps taken to create them, and the decision-making process through your documentation. Ideally, they should also be able reproduce your steps and recreate the constructed variables. Therefore, documentation is an output of construction as relevant as the code and data, and it is good practice for papers to have an accompanying data appendix listing analysis

variables and their definitions.

The development of construction documentation is a good opportunity to have a wider discussion with your team about creating protocols for variable definition, which will guarantee that indicators are defined consistently across projects. You must have a detailed account of how variables are created. This will be implemented in your code, but you should still add comments explaining in human language what you are doing and why. This is a crucial step both to prevent mistakes and to guarantee transparency. To make sure that these comments can be more easily navigated, it is wise to start writing a variable dictionary as soon as you begin making changes to the data.¹⁰ Carefully record how specific variables have been combined, recoded, and scaled, and refer to those records in the code.

The `iecodebook export` subcommand is a good way to ensure you have easy-to-read documentation. When all your final indicators have been created, you can use the `it` to list all variables in the dataset in an Excel sheet. You can then add the variable definitions to that file to create a concise metadata document. Take this opportunity to review your notes and make sure that your code is implementing exactly what is described in the documentation.

Writing data analysis code

After data is cleaned and indicators are constructed, you are ready to start analyzing the data. There are many existing resources for data analysis and statistical methods, such as *R for Data Science*,¹¹ *A Practical Introduction to Stata*,¹² *Mostly Harmless Econometrics*,¹³ and *Causal Inference: The Mixtape*.¹⁴ We focus on how to structure data analysis code and files, rather than how to conduct specific analyses.

¹⁰ See Jones et al. (2019) for an example.

¹¹ Wickham and Grolemund (2017)

¹² McGovern (2012a)

¹³ Angrist and Pischke (2008)

¹⁴ Cunningham (2018)

Organizing analysis code

The analysis stage usually starts with a process we call **exploratory data analysis**. This is when you are first looking for patterns in your data, creating descriptive graphs and tables, and trying different tests to understand your results. It progresses into **final analysis** when your team starts to decide which are the “main results”, or those that will make it into a research output. The way you deal with code and code outputs for exploratory and final analysis is different. During the exploratory stage, you will be tempted to write lots of analysis into one big, impressive, start-to-finish script. While this is fine when you are writing your research stream of consciousness into code, it leads to poor practices in the final code such as not clearing the workspace and not loading a fresh dataset before each analysis task.

To avoid mistakes, it's important to take the time to organize the code that you want to keep, that is, the final analysis code, in an organized manner. The result is a curated set of polished scripts that will be part of a reproducibility package. A well-organized analysis script starts with a completely fresh workspace and, for each output it creates, explicitly loads data before analyzing it. This setup encourages data manipulation to be done earlier in the workflow (that is, in separate cleaning and construction scripts). This also prevents the common problem of having analysis scripts that depend on other analysis scripts being run before them. Such dependencies tend to require manual instructions for all necessary chunks of code to be run in the right order. We encourage you to code each task so it is completely independent of all other code, except for the analysis master script. You can go as far as coding every output in a separate script.

There is nothing wrong with code files being short and simple. In fact, analysis scripts should be as simple as possible, so whoever is reading them can focus on the concepts, not the coding. Research questions and statistical decisions should be incorporated explicitly in the code through comments, and their implementation should be easy to detect from the way the code is written. This includes clustering, sampling, and controlling for different variables, to name a few. If you have multiple analysis datasets, each of them should have a descriptive name about its sample and unit of observation. As your team comes to a decision about model specification, you can create functions and globals (or objects) in the master script to use across scripts. This is a good way to make sure specifications are consistent throughout the analysis. It also makes your code more dynamic, as it's easy to update specifications and results through a master file without changing every script.

To accomplish this, you will need to make sure that you have an effective data management system, including file naming, organization, and version control. Just as for the analysis datasets, you should name each of the individual analysis files descriptively. Code files such as `spatial-diff-in-diff.do`, `matching-villages.R`, and `summary-statistics.py` are clear indicators of what each file is doing, and allow you to find code quickly. If you intend to numerically order the script files to correspond to exhibits as they appear in a paper or report, leave this to near publication time, as you will constantly re-order them during data analysis.

Visualizing data

Data visualization is increasingly popular, and is becoming a field in its own right.¹⁵ We attribute some of the difficulty of creating good data visualization to the difficulty of writing code to create them. Making a visually compelling graph would already be hard enough if you didn't have to go through many rounds of searching and reading help files to understand a command's graphical options syntax. The trickiest part of using plotting commands is getting the data into the right format. Though both Stata and R have plotting functions that graph summary statistics, a good rule of thumb is to ensure that each observation in your dataset corresponds to one data point in your desired visualization. This may seem simple, but often requires the aggregation and reshaping operations discussed earlier in this chapter.

Based on DIME's accumulated experience creating visualizations for impact evaluations, our team has developed a few resources to facilitate this workflow. First of all, we maintain easily-searchable data visualization libraries for both Stata¹⁶, and R¹⁷. These libraries feature curated data visualization examples, along with source code and example datasets, so you get a good sense of what your data should look like before you can start writing code to create a visualization. The `ietoolkit` package also contains two commands to automate common impact evaluation graphs: `iegraph` plots the values of coefficients for treatment dummies, and `iekdensity` displays the distribution of an outcome variable across groups and adds the treatment effect as a note. The DIME Wiki provides a list of other recommended resources and tools for creating good data visualizations.¹⁸

¹⁵ Healy (2018); Wilke (2019)

¹⁶ <https://worldbank.github.io/Stata-IE-Visual-Library>

¹⁷ <https://worldbank.github.io/r-econ-visual-library/>

¹⁸ https://dimewiki.worldbank.org/Data_visualization

Creating analysis outputs

A great number of outputs will be created during the course of a project. These will include both raw outputs such as tables and graphs and final products such as presentations, papers and reports. During exploratory analysis, your team will consider different approaches to answer research questions and present answers. Though it is best to be transparent about different specifications tried and tests performed, only a few will ultimately be considered “main results”. These will be **exported**¹⁹ from the statistical software. That is, they will be saved as tables and figures in format that is easier to interact with. For example, saving graphs as images will allow your team to quickly see them, as well as to add them as exhibits to other documents. When the first of these code outputs are being created,

¹⁹ **Exporting results:** The creation of publication-ready representations of results.

agree on where to store them, what software and formats to use, and how to keep track of them. This discussion will save you time and efforts on two fronts: you will spend less time formatting and polishing tables and graphs that will not make their way into final research products; and you will remember the different paths your team has already taken, so you don't do the same thing twice. This section will take you through key elements to keep in mind when making workflow decisions and outputting results.

Managing outputs

Decisions about storage of outputs are limited by technical constraints, and dependent on file format. Plain text formats like .tex and .csv and should be managed through version control systems like Git, as discussed in Chapter 2. Binary outputs like Excel files, PDFs, PowerPoints, or Word documents, on the other hand, should be kept in a synced folder. Exporting all raw outputs as plain text files, which can be done through all statistical software, facilitates the identification of changes in results. When you re-run your code from the master script, the outputs will be overwritten, and any changes (for example, in coefficients or number of observations) will be automatically flagged for you or a reviewer to check. Tracking changes to binary files is more cumbersome. They they use more space, which may cause slow down the cloud syncing. There may be exceptions to this general rule depending on the Git client you are using. GitHub Desktop, for example, displays changes in common binary image formats such as PNG files in an accessible manner.

You will need to update your outputs frequently. And if you have tried to recreate a result after a few months, you probably know that it can be hard to remember where the code that created it was saved. File naming conventions and code organization, including easily searchable file names and comments, play a key role in not re-writing scripts again and again. We recommend maintaining one “final” analysis folder and one folder with draft code or exploratory analysis. The latter contains pieces of code that are stored for reference, but not cleaned up to be included in any final outputs. Once an output presents a result in the clearest manner possible, it should be renamed and moved to the “final analysis” folder. It's typically desirable to have the names of outputs and scripts linked – so, for example, `factor-analysis.do` creates `factor-analysis.eps` and so on. Document output creation in the master script that runs your code, so that before the line that runs a particular analysis script there are a few lines of comments listing datasets and functions that are necessary for it to run, as well as all outputs created by that script.

Knowing how your code outputs will be used will help you decide the best format to export them. You can often save figures into different formats, such as .eps, .png, .pdf or .jpg. However, the decision between using Office Suite software such as Word and Power Point versus L^AT_EX and other plain text formats may influence how you write your code, as this choice often implicates in the use of a particular command. We strongly recommend that you chose software to create final products that can be linked to raw outputs in such a way that they are updated in the paper or presentation every time changes are made to them. We broadly call files that have this feature **dynamic documents**, and they will be discussed in more detail in the final section of this chapter.

Exporting analysis outputs

As briefly discussed in the previous section, you do not necessarily have to export each and every table and graph created during exploratory analysis. Most statistical software allow you to review results interactively, and this is often preferred at this stage. Final analysis scripts, on the other hand, must export outputs that are ready to be included in a paper or report. No manual edits, including formatting, should be necessary after exporting final outputs. Manual edits are difficult to reproduce; the less you make, the more reproducible your output is. You may think that it's not worth coding a small formatting adjustment, but you will inevitably need to make changes to the output, and automating them will save you time by the end of the process. (However, don't spend much time formatting tables and graphs until you have come to a decision about which will be used for your final product.²⁰) Polishing final outputs can be a time-consuming process, and you want to do it as few times as possible.

We cannot stress this enough: do not set up a workflow that requires copying and pasting results. Copying results from Excel to Word is error-prone and inefficient. Copying results from a software console is risk-prone, even more inefficient, and totally unnecessary. The amount of work needed in a copy-paste workflow increases rapidly with the number of tables and figures included in a research output, and so do the chances of having the wrong version of a result in a paper or report.

There are numerous commands to export outputs from both R and Stata. Some examples are `estout`,²¹ `outreg2`,²² and `outwrite`²³ in Stata, and `stargazer`²⁴ and `ggplot2`'s `ggsave`²⁵ in R. They allow for a wide variety of output formats. We recommend using formats that are accessible and, whenever possible, lightweight. Accessible

²⁰ For a more detailed discussion on this, including different ways to export tables from Stata, see <https://blogs.worldbank.org/impactevaluations/nice-and-fast-tables-stata>

²¹ Jann (2005), Jann (2007)

²² Wada (2014)

²³ Daniels (2019)

²⁴ Hlavac (2015)

²⁵ Wickham (2016)

means that it's easy for other people to open them. In Stata, that would mean always using `graph export` to save images as `.jpg`, `.png`, `.pdf`, etc., instead of `graph save`, which creates a `.gph` file that can only be opened by Stata. Some publications require "lossless" TIFF or EPS files, which are created by specifying the desired extension. Whichever format you decide to use, remember to always specify the file extension explicitly. For tables, there are fewer file format options. Given our recommendation to use **dynamic documents**, which will be discussed in more detail both in the next section and in Chapter 7, exporting tables to `.tex` is preferred. Excel `.xlsx` and `.csv` are also commonly used, but often require the extra step of copying the tables into the final output. The `ietoolkit` package includes two commands to export formatted tables, automating the creation of common outputs and saving time for research. `iebaltab`²⁶ creates and exports balance tables to Excel or L^AT_EX. `ieddtab`²⁷ does the same for difference-in-differences regressions.

If you need to create a table with a very specific format that is not automated by any command you know, consider writing it manually (Stata's `filewrite` and R's `cat()`, for example, allow you to do that). This will allow you to write a cleaner script that focuses on the econometrics, and not on complicated commands to create and append intermediate matrices. Keep in mind that final outputs should be self-standing. This means it should be easy to read and understand them with only the information they contain. Make sure labels and notes cover all relevant information included in your code and comments that are not otherwise visible in the output. Examples of information that should be included in labels and notes include sample, unit of observation, unit of measurement, and variable definition.²⁸

Preparing dynamic documents

Dynamic documents²⁹ are a broad class of tools that enable a streamlined, reproducible workflow. The term "dynamic" can refer to any document-creation technology that allows the inclusion of explicitly encoded linkages to raw output files. This means that, whenever outputs are updated, the next time the document is loaded or compiled, it will automatically include all changes made to all outputs without any additional intervention from the user. This is not possible in tools like Microsoft Office, although there are tools and add-ons that produce similar functionality. In Word, by default, you have to copy and paste each object individually whenever tables, graphs, or other inputs have to be updated. This workflow becomes more complex as the number of inputs grows, increasing the likelihood of making

²⁶ <https://dimewiki.worldbank.org/iebaltab>

²⁷ <https://dimewiki.worldbank.org/ieddtab>

²⁸ https://dimewiki.worldbank.org/Checklist:_Submit_Table

²⁹ **Dynamic documents:** File types that include direct references to exported materials and update them in the output automatically.

mistakes or missing updates. Dynamic documents prevent this from happening by managing document compilation and inclusion of inputs in a single integrated process, so you can skip the copying and pasting altogether.

Dynamic exploratory analysis

If all team members working on a dynamic document are comfortable using the same statistical software, built-in dynamic document engines are a good option for exploratory analysis. With these tools, you can write both text (often in Markdown³⁰) and code in the script, and the result will usually be a PDF or HTML file including code, text, and outputs. In our experience, many researchers find the entry cost to learning how to use these tools to be high. These types of dynamic document tools are typically best used by the team members working most closely with code, and can be great for creating exploratory analysis reports as you work on them, or paper appendices including large chunks of code and dynamically created graphs and tables. RMarkdown³¹ is the most widely adopted solution in R. Stata offers a built-in package for dynamic documents, `dyndoc`,³² and user-written commands such as `markstat`,³³ `markdoc`,³⁴ `webdoc`,³⁵ and `texdoc`.³⁶ The advantage of these tools in comparison with LaTeX is that they create full documents from within your scripts, so running the code and compiling the document is reduced to a single step.

Documents called “notebooks” (such as Jupyter Notebook³⁷) work similarly, as they also use the underlying code that create the document. These tools are usually appropriate for short or informal documents because it tends to be difficult for those who are not familiar with them to edit the content, and they often don’t offer as extensive formatting options as, for example, Word. There are also other simple tools for dynamic documents that do not require direct operation of the underlying code or software, simply access to the updated outputs. An example of this is Dropbox Paper, a free online writing tool that can be linked to files in Dropbox which are automatically updated anytime the file is replaced. These have limited functionality in terms of version control and formatting, and may never include any references to confidential data, but they do offer extensive collaboration features, and can be useful for working on informal outputs. Markdown files on GitHub can also provide similar functionality through the browser, and are version controlled. However, as with other Markdown options, the need to learn a new syntax may discourage take up among team members who don’t work with GitHub more extensively.

³⁰ <https://www.markdownguide.org/>

³¹ <https://rmarkdown.rstudio.com>

³² <https://www.stata.com/manuals/rptdynodoc.pdf>

³³ Rodriguez (2017)

³⁴ Haghish (2016)

³⁵ Jann (2017)

³⁶ Jann (2016)

³⁷ <https://jupyter.org>

Using L^AT_EX for dynamic research outputs

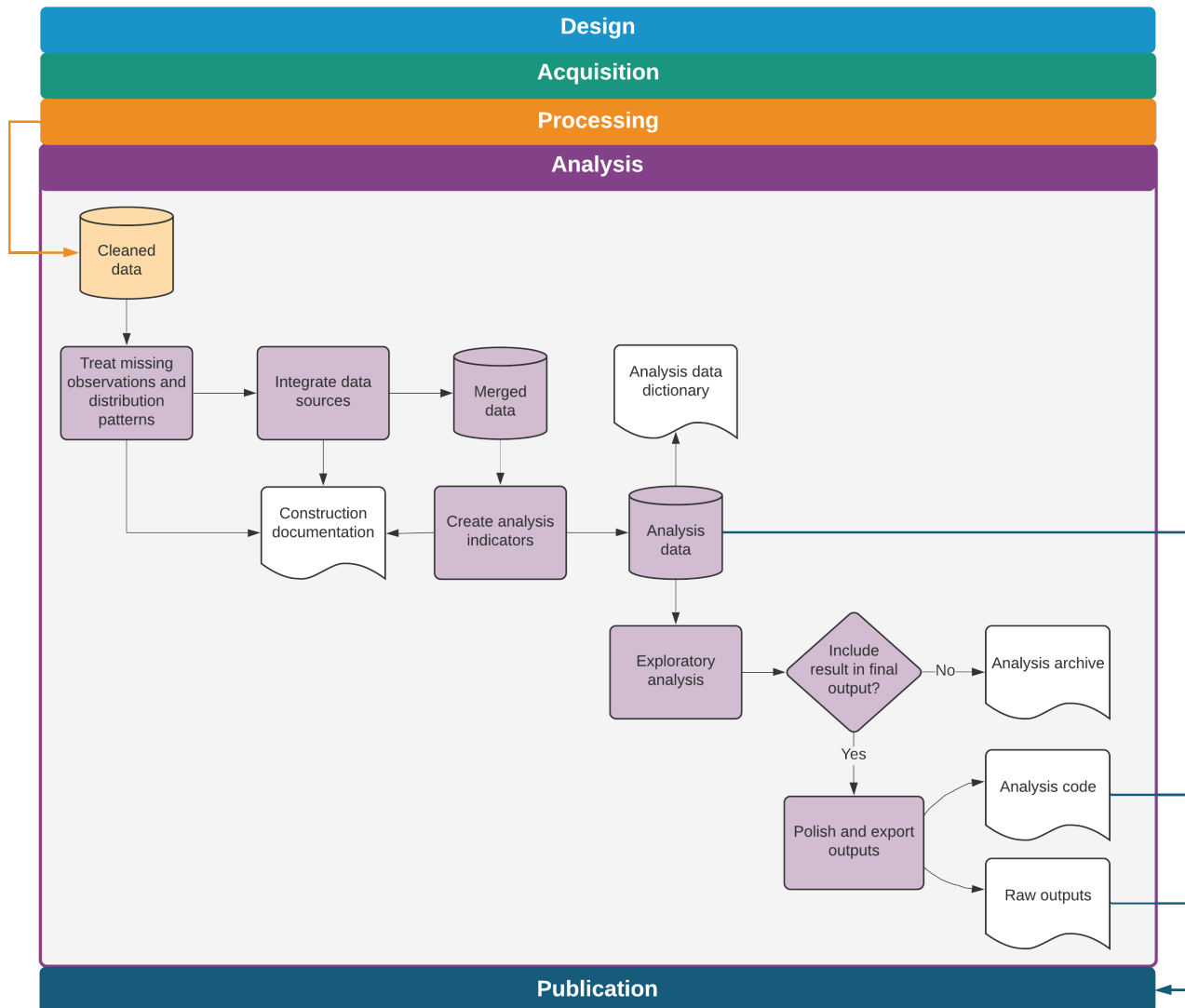
Though formatted text software such as Word and PowerPoint are still prevalent, researchers are increasingly choosing to prepare final outputs like documents and presentations using L^AT_EX. L^AT_EX is a document preparation and typesetting system with a unique code syntax.³⁸ While L^AT_EX has a significant learning curve, its enormous flexibility in terms of operation, collaboration, output formatting, and styling make it our preferred choice for most large technical outputs. In fact, L^AT_EX operates behind-the-scenes in many other dynamic document tools (discussed below). Therefore, we recommend that you learn L^AT_EX as soon as you are able to.

The main advantage of using L^AT_EX is that it updates outputs every time the document is compiled, while still allowing for text to be added and extensively formatted to publication-quality standards. Additionally, because of its popularity in the academic community, social scientists are more familiar with it than other dynamic documents tools, so the cost of entry for a team is often relatively low. Because .tex files are plain text, they can be version-controlled using Git. Creating documents in L^AT_EX using an integrated writing environment such as TeXstudio, TeXmaker or LyX is great for outputs that focus mainly on text and include figures and tables that may be updated. It is good for adding small chunks of code into an output. Finally, some journals make L^AT_EX templates available, so papers can be more easily be formatted into their specific layout.

³⁸ See <https://github.com/worldbank/DIME-LaTeX-Templates>.

Looking ahead

This chapter discussed the steps needed to create analysis datasets and outputs from original data. By combining the observed variables of interest for the analysis (measurement variables) with the information in the data map describing the study design (research variables), you created original datasets ready for analysis, as shown in the figure below. Doing so is difficult, creative work, and it is impossible to be reproduced by another person without access to detailed records and explanations of how you interpreted and modified the data. You learned that code must be well-organized and well-documented to allow others to understand how research outputs were created and how those were used to answer the research questions. The final chapter of this book will provide a guide to assembling the raw findings into publishable work, as well as describing methods for making your data, code, documentation, and other research outputs accessible and reusable alongside your primary outputs.



Chapter 7: Publishing research outputs

Publication typically involves many iterations of data, code, and code output files, with inputs from multiple collaborators. This process can quickly become unwieldy. It is in nobody's interest for a skilled and busy researcher to spend days re-numbering figures, tables, or references (and it can take days) when a reasonable amount of up-front effort can automate the task. Similarly, simultaneous collaboration should not involve the repetitive and error-prone task of manually resolving sets of tracked-changes documents with conflicting edits. Furthermore, for most development data projects, completing a research output is not the end of the task. Academic journals increasingly require reproducibility packages containing the data, code, and supporting materials needed to recreate the results. DIME requires reproducibility packages for all published outputs. Replication materials represent an intellectual contribution in their own right, because they enable others to learn from your process and better understand the results you have obtained. If you have organized your analysis process according to the general principles outlined in earlier chapters, publication will not require substantial reorganization of the work you have already done. Hence, publication is the conclusion of the system of transparent, reproducible, and credible research we introduced from the very first chapter of this book.

In this chapter, we suggest tools and workflows for efficiently managing collaboration on research and policy outputs and ensuring reproducible results. The first section discusses how to use dynamic documents to collaborate on writing. The second section covers how to prepare and publish original data, an important research contribution in its own right. The third section provides guidelines for preparing functional and informative reproducibility packages. In all cases, we note that technology is rapidly evolving and that specific tools noted here may not remain cutting-edge, but the core principles involved in publication and transparency will endure.

Preparing written documents for publication

Development research is increasingly a collaborative effort. This reflects changes in the economics discipline overall: the number of sole-authored research outputs is decreasing, and the majority of recent papers in top journals have three or more authors.¹ As a consequence, documents typically pass back and forth between several writers before they are ready for publication or release. As in all other stages of the research process, effective collaboration requires the adoption of tools and practices that enable version control and simultaneous contributions. This book, for example, was written in L^AT_EX and managed on GitHub.² As we outlined in the

¹ Kuld and O'Hagan (2017)

² <https://github.com/worldbank/dime-data-handbook>

previous chapter, **dynamic documents** are a way to simplify writing workflows: updates to code outputs that appear in these documents, such as tables and figures, can be passed in to the final research output in a single click, rather than copy-and-pasted or otherwise handled individually. Managing the writing process in this way improves organization and reduces error, such that there is no risk of materials being compiled with out-of-date results, or of completed work being lost or redundant.

Preparing formal outputs as L^AT_EX documents

As we discussed in Chapter 6, the most widely used software for dynamically managing formal manuscripts and policy outputs is L^AT_EX. It is also becoming more popular for shorter documents, such as policy briefs, with the proliferation of skills and templates for these kinds of products. L^AT_EX uses explicit references to the file path of each input (such as tables and figures), which are reloaded from these locations every time the final document is compiled. This is not possible by default in, for example, Microsoft Word. There, you have to copy and paste each object whenever tables, graphs, or other inputs are updated. As time goes on, it becomes increasingly likely that a mistake will be made or something will be missed. In L^AT_EX, instead of writing in a “what-you-see-is-what-you-get” mode as you do in Word, you write plain text in a .tex file, interlaced with coded instructions formatting the document and linking to exhibits (similar to HTML). L^AT_EX manages tables and figures dynamically and includes commands for simple markup like font styles, paragraph formatting, section headers and the like. It includes special controls for footnotes and endnotes, mathematical notation, and bibliography preparation. It also allows publishers to apply global styles and templates to already-written material, reformatting entire documents in house styles with only a few keystrokes.

While L^AT_EX *can* produce complex formatting, this is rarely needed for academic publishing, as academic manuscripts will usually be reformatted based on the style of the publisher. (By contrast, policy and other self-produced documents may desire extensive typesetting and investments in custom templates and formatting.) In academia at least, it’s rarely worth the investment to go beyond basic L^AT_EX tools: the title page, sections and subsections, figures and tables, mathematical equations, bolding and italics, footnotes and endnotes, and, last but not least, references and citations.

One of the most important tools available in L^AT_EX is the BibTeX citation and bibliography manager.³ BibTeX keeps all the references you might use in a .bib file, then references them using a simple

³ Kopka and Daly (1995)

command typed directly in the document. Specifically, L^AT_EX inserts references in text using the `cite` command. Once this is written, L^AT_EX automatically pulls all the citations into text and creates a complete bibliography based on the citations you used whenever you compile the document. The system allows you to specify exactly how references should be displayed in text (such as superscripts, inline references, etc.) as well as how the bibliography should be styled and in what order (such as Chicago, MLA, Harvard, or other common styles). The same principles that apply to figures and tables are therefore applied here: You can make changes to the references in one place (the `.bib` file), and then everywhere they are used they are updated correctly with one process. BibTeX is so widely used that it is natively integrated in Google Scholar. Since different publishers have different requirements, it is quite useful to be able to adapt this and other formatting very quickly, including through using publisher-supplied templates where available.

Because it follows a standard code format, L^AT_EX has one more useful trick: you can convert the raw document into Word or a number of other formats using utilities such as pandoc.⁴ Even though conversion to Word is required for a number of academic publishers and can even be preferred for some policy outputs, we still recommend using L^AT_EX to prepare these when possible. You should export to Word only at the final stage, when submitting materials. You can also use a CSL (Citation Styles Library) file⁵ for nearly any journal and have it applied automatically in this process. Therefore, even in the case where you are required to provide `.docx` versions of materials to others, or tracked-changes versions, you can create them effortlessly from a L^AT_EX document, then use external tools like Word's compare feature to generate integrated track-changes versions when needed.

Getting started with L^AT_EX as a team

Getting used to L^AT_EX can be challenging, but the control it offers over the writing process is invaluable. Because it is written in a plain text file format, `.tex` can be version-controlled using Git. This makes it possible to manage contributions and version histories using the same system we recommend for data work. DIME Analytics has created a variety of templates and resources that you can adapt for your own team.⁶ Integrated editing and compiling tools like TeXStudio⁷ and atom-latex⁸ offer the most flexibility to work with L^AT_EX in teams.

Although ultimately worth it, setting up L^AT_EX environments locally is not always simple, particularly if you are new to working with plain text code and file management. This is because L^AT_EX

⁴ <https://pandoc.org>

⁵ <https://github.com/citation-style-language/styles>

⁶ <https://github.com/worldbank/DIME-LaTeX-Templates>

⁷ <https://www.texstudio.org>

⁸ <https://atom.io/packages/atom-latex>

requires that all formatting be done in its special code language, and it is not always informative when you do something wrong. This can be off-putting very quickly for people who simply want to get to writing, and staff not used to programming may not easily acquire the necessary knowledge.

Cloud-based implementations of L^AT_EX can make it easier for your team to use L^AT_EX without all members having to invest in new skills or set up matching software environments, and can be particularly useful for first forays into L^AT_EX writing. One example of this is Overleaf.⁹ Most such sites offer a subscription feature with useful extensions and various sharing permissions, and some offer free-to-use versions with basic tools that are sufficient for a broad variety of applications, up to and including writing a complete academic paper with coauthors.

Cloud-based implementations of L^AT_EX have several advantageous features for teams compared to classic desktop installations. First, since they are completely hosted online, they avoid the inevitable troubleshooting required to set up a L^AT_EX installation on various personal computers run by the different members of a team. Second, they typically maintain a single, continuously synced, master copy of the document so that different writers do not create conflicted or out-of-sync copies, or need to deal with Git themselves to maintain that sync. Third, they typically allow collaborators to edit documents simultaneously, though different services vary the number of collaborators and documents allowed at each tier. Fourth, some implementations provide a “rich text” editor that behaves pretty similarly to familiar tools like Word, so that collaborators can write text directly into the document without worrying too much about the underlying L^AT_EX coding. Cloud services also usually offer a convenient selection of templates so it is easy to start up a project and see results right away without needing to know a lot of the code that controls document formatting.

Cloud-based implementations of L^AT_EX also have disadvantages. There is still some up-front learning required, unless you’re using the rich text editor. Continuous access to the internet is necessary, and updating figures and tables requires a bulk file upload that is tough to automate. They vary dramatically in their ability to integrate with file systems where you store your code and code outputs, and so you will need to practice an integrated workflow depending what is available to you. Some teams adopt cloud-based tools as a permanent solution, though our recommendation is to eventually shift to local editing and compiling using tools such as TexStudio and code editors like Atom.

⁹ <https://www.overleaf.com/>

Preparing and publishing data

While we have focused so far on written materials, you must also consider how you will publish the data used in your research. The open science community at large sees data publication both as a citable output and as a necessary transparency measure. Fortunately, it is a conceptually simple task to produce and catalog the required materials. You should be prepared to catalog two separate collections. First, you should catalog the clean data with all variables corresponding directly to fields in the original dataset or data collection instrument (this may not be necessary if you are working with secondary data that is published elsewhere in this format). If you follow the steps outlined in Chapter 5, when you get to the publication stage you will have a cleaned data set and supporting documentation ready.

Second, you should separately catalog the analysis dataset used for the research output you are publishing. This is typically included in the replication package for the research output, and should assume that another researcher is starting work with only the published clean data in hand. The package should also include the data construction scripts that create transformed and derived indicators, project-specific information such as treatment assignment and other indicators generated directly by the research team (another example is constructed record linkages). If you followed the workflow recommended in Chapter 6, by the time you reach publication stage you will already have all necessary files and documentation at hand.

De-identifying data for publication

Before publishing data, you should carefully perform a **final de-identification**. The objective of de-identification is to reduce the risk of disclosing confidential information in the published dataset. If you are following the workflow outlined in this book, you should have already removed direct identifiers as a first step after acquiring the data (see the discussion of initial de-identification in Chapter 5). For the final de-identification, you should additionally remove indirect identifiers, and assess the statistical disclosure risk of your data.¹⁰ Unlike direct identifiers, for which a link (or lack thereof) to public information is verifiable, indirect identifiers require an assessment of the likelihood that an individual can be singled out in the data and then linked to public information using combinations of available data. For example, seemingly innocuous variables such as US zip code, gender, and date of birth uniquely identify approximately 87% of the US population.¹¹ In development data, information such as the size of a household, the ages and marital statuses of the household

¹⁰ Disclosure risk: the likelihood that a released data record can be associated with an individual or organization.

¹¹ Sweeney (2000)

members, and the types of work or schooling they engage in may be more than enough to identify a person or family from a sufficiently small group.

A number of tools have been developed to help researchers de-identify data. At this stage, the World Bank's `sdcMicro` tool,¹² has a useful feature that allows you to assess the uniqueness of the records in your data. It produces simple measures of the identifiability of records from the combination of potentially indirectly identifying variables, and allows you to apply common information masking algorithms, such as binning, top-coding, and jittering data prior to release. You should determine how sensitive your results are to these transformations; it may be the case that masked data cannot be used for your reproducibility package.

There will almost always be a trade-off between accuracy and privacy. For publicly disclosed data, you should favor privacy. Stripping identifying variables from a dataset may not be sufficient to protect respondent privacy, due to the risk of re-identification. One solution is to add noise to data, as the U.S. Census Bureau has proposed.¹³ This makes the trade-off between data accuracy and privacy explicit. But there are not, as of yet, established norms for such "differential privacy" approaches: most approaches fundamentally rely on judging "how harmful" information disclosure would be. The fact remains that there is always a balance between information release (and therefore transparency) and privacy protection, and that you should engage with it actively and explicitly. The best thing you can do is make a complete record of the steps that have been taken so that the process can be reviewed, revised, and updated as necessary.

Removing variables results in loss of information, so the de-identification process requires careful assessment of the potential risk to the individual that could be caused by disclosure of their identity or personal information. This will vary widely depending on the types of information you are collecting and the overall vulnerability of the population. In extreme cases, where the population is highly vulnerable and combinations of information are highly specific, you may not be able to publicly release any data at all. You will still be expected to catalog and cite your data, even if you cannot release it publicly. In practice, this may mean publishing only a catalog entry providing information about the contents of the datasets and how future users might request permission to access them (even if you are not the person to grant that permission). In some cases, it may be possible to release the dataset but embargoing specific variables that are required for the analysis but cannot be released publicly. It may be necessary to grant access to the embargoed data for specific purposes, such as a computational reproducibility check required

¹² Benschop et al. (2019)

¹³ Abowd (2018)

for publication, if done under careful data security protocols and approved by an IRB.

Publishing clean and constructed datasets

Publicly documenting all original data acquired as part of a research project is an important contribution in its own right. Cataloging and/or archiving original datasets is a significant contribution in addition to any publication of analysis results.¹⁴ Publicly releasing data allows other researchers to validate the mechanical construction of your results, investigate what other results might be obtained from the same population, and test alternative approaches or answer other questions. This fosters collaboration and may enable researchers to explore variables and questions that you do not have time to focus on otherwise.

The first step toward data publication is choosing the platform where you will publish your data. A variety of options exist; it is important to choose one that allows you to obtain a digital object identifier (DOI) for the location of your data (even if its URL changes), and a formal citation for your data, so you can reference it in other research outputs.¹⁵ Two common platforms for development data are the World Bank's Development Data Hub and Harvard University's Dataverse. The World Bank's Development Data Hub¹⁶ includes a Microdata Catalog¹⁷ and a Geospatial Catalog, where researchers can publish data and documentation for their projects.¹⁸ The Harvard Dataverse publishes both data and code, and its Datahub for Field Experiments in Economics and Public Policy¹⁹ is especially relevant for impact evaluations. Both the World Bank Microdata Catalog and the Harvard Dataverse create data citations for deposited entries. DIME has its own collection of datasets in the Microdata Catalog, where data from our projects is published.²⁰

Once you have chosen a platform, you need to determine exactly what data you will publish. As mentioned earlier, there are typically two different types of data releases for a research project: complete (de-identified) original datasets and derivative datasets used for specific research outputs. Whether you can publish the original dataset depends on data ownership and licensing agreements. If the data was acquired through a survey that was contracted by the research team, the data most likely belongs to the research team, and therefore the team has publication rights for both the original and derivative data. If data was acquired from a partner through a licensing agreement, the terms of the license will determine publication rights. These datasets should match the survey instrument or source documentation as closely as possible, and should not include indicators

¹⁴ https://dimewiki.worldbank.org/Publishing_Data

¹⁵ <https://www.doi.org>

¹⁶ <https://datacatalog.worldbank.org>

¹⁷ <https://microdata.worldbank.org>

¹⁸ https://dimewiki.worldbank.org/Microdata_Catalog

¹⁹ <https://dataverse.harvard.edu> and <https://dataverse.harvard.edu/dataverse/DFEEP>

²⁰ <https://microdata.worldbank.org/catalog/dime>

constructed by the research team. Even if you do not have rights to publish the original data, you can typically publish derivative datasets prepared by the research team. These datasets usually contain only the constructed indicators and associated documentation,²¹ and should also be included in the replication package.

When publishing data, you will decide how the data may be used and what license you will assign to it.²² Make sure you understand the rights associated with any data release and communicate them to its future users. Material without a license may never be reused. You should prefer to offer a license that is explicit and details whether and how specific individuals may access the data. Terms of use available in the World Bank Microdata Catalog include, in order of increasing restrictiveness: *open access*, *direct access*, and *licensed access*.²³ *Open access* data is freely available to anyone, and simply requires attribution. *Direct access* data is available to registered users who agree to use the data for statistical and scientific research purposes only, to cite the data appropriately, and not to attempt to identify respondents or data providers or link the data to other datasets that could allow for re-identification. *Licensed access* data is restricted to users who submit a documented application detailing how they will use the data and then sign a formal agreement governing data use. The user must be acting on behalf of an organization, which will be held responsible in the case of any misconduct.

Published data should be released in a widely recognized format. While software-specific datasets are acceptable accompaniments to the code (since those precise materials are probably necessary), you should also consider releasing datasets in plain text formats such as CSV files with accompanying codebooks, since these can be used by any researcher. Additionally, you should also release PDF or code versions of the data collection instrument or survey questionnaire so that readers can understand which data components are collected directly in the field and which are derived. With your analysis dataset, you should also release the code that constructs any derived measures from the clean dataset, so that others can learn from your work and adapt it as they like.

Preparing a complete reproducibility package

Major journals now often require that you provide both the data and code required to recreate your results. Some even require being able to reproduce the results themselves before they will approve a paper for publication.²⁴ This set of materials, taken together, is often referred to as a **reproducibility package**. If you have followed the workflows described in this book, preparing the replication package

²¹ Dupriez, O. and Greenwell, G. (2007). Quick reference guide for data archivists. *IHSN paper*. <http://www.ihsn.org/home/download.php>

²² https://dimewiki.worldbank.org/Publishing_Data

²³ <https://microdata.worldbank.org/index.php/terms-of-use>

²⁴ Vilhuber et al. (2020)

will only require a small amount of extra work. If not, creating this package may take some time. When the replication package is completed, whoever downloads it should be able to understand how your code produces results from your data and be able to reproduce them exactly by executing the included master script.

Organizing code for reproducibility

Before releasing your code, you should edit it for content and clarity just as if it were written material. The purpose of releasing code is to allow others to understand exactly what you have done in order to obtain your results, and enable them to apply similar methods in future projects. Other researchers should be able to reproduce individual portions of your analysis by making only small adjustments to your code. In either a scripts folder or in the root directory, you should include a master script that allows someone else to run the entire project and re-create all raw code outputs by changing only a single line of code: the one setting the directory path. The code should both be functional and readable, through the use of a clear structure and extensive commenting. Code is often not written this way when it is first prepared, so it is important for you to review the content and organization so that a new reader can figure out what your code should do and how it does it. Making code clean and readable is often where you need to invest time prior to releasing your reproducibility package.

DIME requires all academic outputs to successfully pass a computational reproducibility check before being submitted for publication. We have adopted several practices and requirements to support the production of high-quality reproducibility packages. The materials for these practices are publicly available, so you can use them to check the reproducibility of your own work. This reproducibility check is initiated by submitting the Reproducibility Package Checklist.²⁵ DIME projects are required to organize code with a master script, to facilitate handovers across team members and make the computational reproducibility check a one-click exercise. Compliance with these and other coding standards at DIME is monitored through quarterly peer code review rounds, which allows research assistants to improve their code and documentation as it is written, rather than revisiting it in a rush near publication time. DIME projects are also expected to use Git and GitHub to document project work and collaboration, and to keep the main branch up-to-date as a working edition.

Before publicly releasing a reproducibility package, it is essential to make sure that the code runs identically on your individual setup

²⁵ <https://github.com/worldbank/dime-standards>

compared to a fresh installation of your software. To ensure that your code will run completely on a new computer, you must install any required user-written commands in the master script (for example, in Stata using `ssc install` or `net install` and in R adding code that gives users the option to install packages, including selecting a specific version of the package if necessary). In many cases you can even directly provide the underlying code for any user-installed packages that are needed to ensure forward-compatibility. Make sure system settings like software version and memory settings are defined. The `ieboilstart` command in `ietoolkit` defines and applies these settings for a chosen Stata version.²⁶

Finally, make sure that code inputs and outputs are clearly identified. A new user should, for example, be able to easily find and quickly recreate any files generated by the code. Readers should also be able to easily map all the outputs of the code to where they appear in the associated published material, so you must ensure that the raw components of figures or tables are clearly identified. Documentation in the master script is often used to indicate this information. For example, code outputs should clearly correspond by name to an exhibit in the paper, and vice versa. (Supplying a compiling L^AT_EX document can support this.) Code and code outputs which are not used in the final paper should be removed from the final replication package, but still archived for transparency.

Releasing a reproducibility package

Once your replication package is prepared for public release, you need to find a place to publish your materials.²⁷ At the time of writing, there is no consensus on the best solution for publishing code, and there are a variety of archives and storage providers that cater to different needs. The technologies available are likely to change dramatically over the next few years; the specific solutions we mention here highlight some current approaches as well as their strengths and weaknesses.

Features to look for in a platform to release reproducibility packages include: the possibility to store data and documentation as well as code, the creation of a static copy of its content, that cannot be changed or removed, and the assignment of a permanent digital object identifier (DOI) link. Unlike data, code usually has few external constraints to publication. The research team owns the code in almost all cases, and code is unlikely to contain identifying information (though you must verify that it does not). Publishing code also requires assigning a license to it; most code publishers offer permissive licensing options. If you do not provide a license, no one can reuse

²⁶ <https://dimewiki.worldbank.org/ieboilstart>

²⁷ https://dimewiki.worldbank.org/Reproducible_Research

your code. It is common to only require attribution and citation for code reuse, without putting any barriers or restrictions to accessing the code.

One option for creating and releasing a reproducibility package is GitHub. Making a public GitHub repository is completely free. It can hold any file types, provide a structured, compressed download of your whole project, and allow others to look at alternate versions or histories easily. It is straightforward to simply upload a fixed directory to GitHub, apply a sharing license, and obtain a URL for the whole package. There is a strict size restriction of 100MB per file and a restriction of 100GB on the size of the repository as a whole, so larger projects will need alternative solutions. However, GitHub is not the ideal platform on which to *publish* reproducibility packages. It is built to version control code, and to facilitate collaboration on it. It is not an archive, meaning that it does not guarantee the permanence of uploaded materials or the access URL, and it does not manage citations or non-code licenses by default. One suggestion is to combine GitHub with the Open Science Framework,²⁸ as OSF can easily link to and import material from GitHub and apply a permanent URL, DOI, formal citation, general license, and archival services to it.

Other options include the Harvard Dataverse²⁹ and Research-Gate.³⁰ Any of these archival services is acceptable – the main requirement is that the system can handle the structured directory that you are submitting, and that it can provide a stable URL for your project and report exactly what, if any, modifications you have made since initial publication. You can even combine more than one tool if you prefer, as long as they clearly reference each other. For example, one could publish code and the corresponding license on GitHub and point to data published on the World Bank Microdata Catalog. Emerging technologies such as the “containerization” approach of CodeOcean³¹ offer to store both code and data in one repository, and also provide an online workspace in which others can execute and modify your code without having to download your tools and match your local environment. This is particularly useful over time, as packages and other underlying software may have changed since publication.

In addition to code and data, you may also want to release an author’s copy or preprint of the article itself along with these materials. Check with your publisher before doing so; not all journals will accept material that has been publicly released before its formal publication date, although, in most development research fields, the release of working papers is a fairly common practice. This can be done on a number of preprint websites, many of which are topic-

²⁸ <https://osf.io>

²⁹ <https://dataverse.harvard.edu>

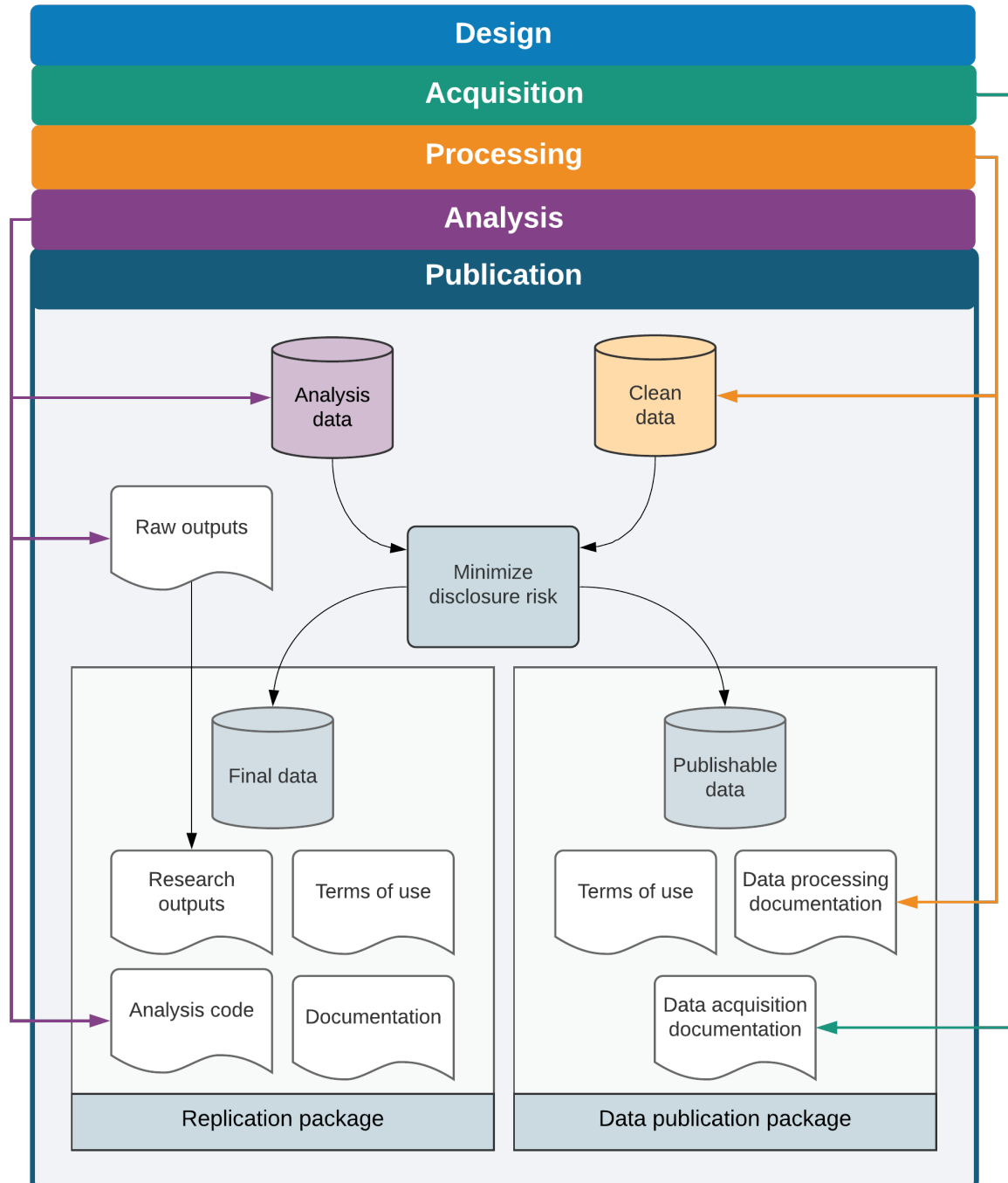
³⁰ <https://www.researchgate.net>

³¹ <https://codeocean.com>

specific. You can also use GitHub or OSF and link to the PDF file directly through your personal website or whatever medium you are sharing the preprint. We recommend against using file sharing services such as Dropbox or Google Drive for this purpose, as their access is more restrictive, and organizations limit access to platforms other than the one officially adopted.

Looking ahead

This chapter described the culmination of all the efforts in data acquisition, cleaning, processing, and analyzing – the production of materials to share with the world that answer a scientific or policy question in a way that has never been done before. The figure included in this chapter provides an overview of all the outputs created by this stage. Making sure that everything you discovered and created is as broadly available and as easy to use as possible is the last step in producing scientific evidence from original data. This is the purpose of all the rigor, organization, and documentation we encourage and detail at every step of the process: since all your research materials are continuously organized, shareable, secure, documented, and readable, they are both valuable to you and others and easy to provide access to.



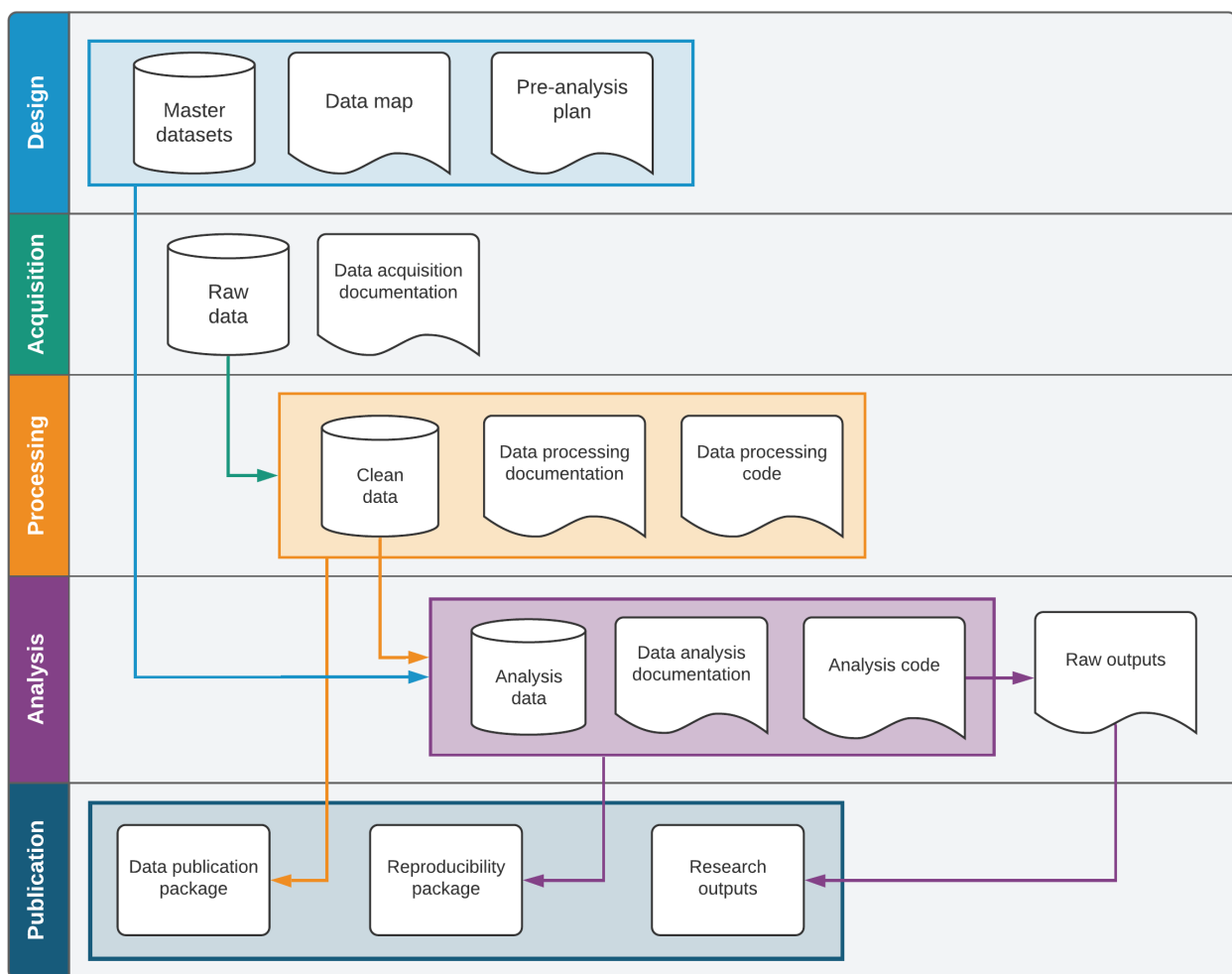
Bringing it all together

We hope you have enjoyed *Development Research in Practice: The DIME Analytics Data Handbook*. Our aim was to teach you to handle data more efficiently, effectively, and ethically. We laid out a complete vision of the tasks of a modern researcher, from planning a project's data governance to publishing code and data to accompany a research product. We have tried to set the text up as a resource guide so that you will always be able to return to it as your work requires you to become progressively more familiar with each of the topics included in the guide.

We started the book with a discussion of credibility, transparency, and reproducibility in research: an overarching idea that your work should always be accessible and available to others, both within and outside your team. We then discussed the current research work environment, which necessitates cooperation with a diverse group of collaborators using modern approaches to computing technology. We outlined methods for planning your data work and creating basic documentation for data, so that multiple data sources can be linked without error and so that you can map the structure of your data to your research design. We discussed how to implement reproducible routines for sampling and randomization, and to analyze statistical power and use randomization inference.

We detailed modern data acquisition methods and frameworks, including data licensing, data ownership, electronic data collection, and data security. We provided a detailed workflow for data cleaning and processing, emphasizing tidy data, quality control, privacy protection, and documentation. We discuss the creation of derived indicators and analysis outputs, and give an overview of the workflow required to move these results to publication – no matter the format of your work – as well as tools and practices for making this work publicly accessible. Throughout, we emphasized that data work is a “social process”, involving multiple team members with different roles and technical abilities. This mindset and workflow, from top to bottom, outline the tasks and responsibilities that are fundamental to doing credible research.

However, as you probably noticed, the text itself provides just enough detail to get you started: an understanding of the purpose and function of each of the core research steps. The references and resources get into the details of how you will realistically implement these tasks: from DIME Wiki pages detail specific code conventions and field procedures that our team considers best practices, to the theoretical papers that will help you figure out how to handle the unique cases you will undoubtedly encounter. We hope you will keep the book on your desk (or the PDF on your desktop) and come back to it anytime you need more information. We wish you all the best in your work and will love to hear any input you have on ours! You can share your comments and suggestion on this book through <https://worldbank.github.io/dime-data-handbook>.



Appendix: The DIME Analytics Coding Guide

Most academic programs that prepare students for a career in the type of work discussed in this book spend a disproportionately small amount of time teaching their students coding skills in relation to the share of their professional time they will spend writing code in their first few years after they graduate. Recent masters-level graduates joining our team have shown very good theoretical knowledge while requiring a lot of training in practical skills. To us, this is like an architecture graduate having learned how to sketch, describe, and discuss the concepts and requirements of a new building very well – but without the technical ability to contribute to a blueprint following professional standards that can be used and understood by others. The reasons for this are a topic for another book, but in today’s data-driven world, people working in quantitative development research must be proficient collaborative programmers, and that includes more than being able to compute correct numbers.

This appendix begins with a section on some general and language-agnostic principles on how to write “good” code. Good code is code that both generates correct results *and* is easily read and adapted by other professionals. The second section in this appendix contains instructions on how to access and use the code examples in this book. The last section contains the DIME Analytics Stata Style Guide. Widely accepted and used style guides are common in most programming languages, but we have never found a sufficiently encompassing style guide for Stata. We created this style guide having in mind practices that, in our experience, greatly improve the quality of research projects coded in Stata. We hope that this guide can help increase the emphasis given to using, improving, sharing, and standardizing code style among the Stata community. We believe these resources can help anyone write more understandable code, and how you, like an architect, can create a blueprint that can be understood and used by everyone in your trade.

Writing good code

“Good” code has two elements: (1) it is correct, in that it doesn’t produce any errors and its outputs are the objects intended, and (2) it is useful and comprehensible to someone who hasn’t seen it before (or even someone who has, weeks, months, or years later). Many researchers have only been trained to code correctly. But we believe that when your code runs on your computer and you get the desired results, you are only half-done writing *good* code. Good code is easy to read and replicate, making it easier to spot mistakes. Good code reduces sampling, randomization, and cleaning errors. Good code can easily be reviewed by others before it’s published and can be re-used afterwards. We always tell people to “code as if a stranger is

reading it”.

You should think of good code in terms of three major elements: **structure**, **syntax**, and **style**. The **structure** is the environment and file organization your code lives in: good structure means that it is easy to find individual pieces of code, within and across files, that correspond to specific tasks and outputs. It also means that functional code blocks are sufficiently independent from each other such that they can be shuffled around, repurposed, and even deleted without affecting the execution of other portions. The **syntax** is the literal language of your code. Good syntax means that your code is readable in terms of how its mechanics implement ideas – it should not require arcane reverse-engineering to figure out what a code chunk is trying to do. It should use common commands in a generally accepted way so others can easily follow and reconstruct your intentions. Finally, **style** is the way that the non-functional elements of your code convey its purpose. Elements like spacing, indentation, and naming conventions (or lack thereof) can make your code much more (or much less) accessible to someone who is reading it for the first time and needs to understand it quickly and accurately.

One key tool for writing good code is using help documentation. Whether or not you are new to the programming language you are using – Stata, R, Python, or one of the many others – you should constantly revisit help files for the most common commands. Often you will learn they can do something you never realized. We cannot emphasize enough how important it is that you get into the habit of reading help files, especially for commands you are very familiar with. Most of us have a help file window open at all times. Similarly, you will always run into commands or uses of commands that you have not seen before or whose functionality you do not remember. Every time that happens, you should look up the help file for that command. We often encounter the conception that help files are only for beginners. We could not disagree with that conception more. The only way to get better at the programming language you use is to constantly read help files.

Using the code examples in this book

Providing some standardization for Stata code style is also a goal of our team. Stata is one of several programming languages used at DIME, but since few high-quality resources based on Stata exist relative to its importance and frequency of use, we decided to use Stata for the examples in this book. This book includes several code blocks where we demonstrate a good code execution of some of the most common tasks in data work. We have ensured that each

code block runs independently, is well-formatted, and uses built-in commands as much as possible. We hope that these snippets will provide a foundation for your code style. We try to comment code examples generously (as you should), and you should reference Stata help files by writing `help [command]` whenever you do not understand the command that is being used. For example, if you are not familiar with `isid`, then write `help isid`, and the help file for the command `isid` will open. You should do this even if you know `isid` but has not read the help file for that command in a while, as there is always something new to learn.

In the book, code examples are presented like the following:

code.do

```

1 * Load the auto dataset
2     sysuse auto.dta , clear
3
4 * Run a simple regression
5     reg price mpg rep78 headroom , coefl
6
7 * Transpose and store the output
8     matrix results = r(table)'
9
10 * Load the results into memory
11    clear
12    svmat results , n(col)

```

You can access the raw code used in examples in this book in several ways. We use GitHub to version control all the content of this book, including the code examples. To see the code examples on GitHub, go to: <https://github.com/worldbank/dime-data-handbook/tree/master/code>. We only use Stata's built-in datasets in our code examples, so you do not need to download any data. If you have Stata installed on your computer, then you will already have the data files used in the code.

A less technical way to access the code is to click the individual file in the URL above, then click the button that says **Raw**. You will then get to a page that looks like the one at: <https://raw.githubusercontent.com/worldbank/dime-data-handbook/master/code/code.do>. There, you can copy the code from your browser window to your do-file editor with the formatting intact. This method is only practical for a single file at a time. If you want to download all code used in this book, you can do that at: <https://github.com/worldbank/dime-data-handbook/archive/master.zip>. That link downloads a .zip file with all the content used in writing this book, including the `LATEX` code used for the book itself. After extracting the .zip-file you will find all the code in a folder called `/code/`.

While we use built-in commands as much as possible in this book, we point to user-written commands when they provide important new functionality. In particular, we point to two suites of Stata commands developed by DIME Analytics, `ietoolkit`¹ and `iefieldkit`,² which DIME Analytics wrote to standardize core data collection, management, and analysis workflows.

When you encounter code that employs user-written commands, you will not be able to run them or read their help files until you have installed the commands. The most common place to distribute user-written commands for Stata is the Boston College Statistical Software Components (SSC) archive.³ The user-written commands in our code example are all from the SSC archive. So, if your installation of Stata does not recognize a command in our code, for example `randtreat`, then type `ssc install randtreat` in Stata.

Some commands on SSC are distributed in packages. This is the case, for example, of `ieboilstart`. That means that you will not be able to install it using `ssc install ieboilstart`. If you do, Stata will suggest that you instead use `findit ieboilstart`, which will search SSC (among other places) and see if there is a package that contains a command called `ieboilstart`. Stata will find `ieboilstart` in the package `ietoolkit`, so to use this command you will type `ssc install ietoolkit` in Stata instead.

We understand that it can be confusing to work with packages for first time, but this is the best way to set up your Stata installation to benefit from other people's work that has been made publicly available. Once you get used to installing commands like this it will not be confusing at all. All code with user-written commands, furthermore, is best written when it installs such commands at the beginning of the master do-file, so that the user does not have to search for packages manually.

The DIME Analytics Stata Style Guide

Programming languages used in computer science always have style guides associated with them. Sometimes they are official guides that are universally agreed upon, such as PEP8 for Python.⁴ More commonly, there are well-recognized but non-official style guides like the JavaScript Standard Style⁵ for JavaScript or Hadley Wickham's style guide for R.⁶ Google, for example, maintains its own style guides for all languages that are used in its projects.⁷

Aesthetics is an important part of style guides, but not the main point. Neither is telling you which commands to use: there are plenty of guides to Stata's extensive functionality.⁸ The important function is to allow programmers who are likely to work together

¹ <https://dimewiki.worldbank.org/ietoolkit>

² <https://dimewiki.worldbank.org/iefieldkit>

³ <https://ideas.repec.org/s/boc/bocode.html>

⁴ <https://www.python.org/dev/peps/pep-0008>

⁵ <https://standardjs.com/#the-rules>

⁶ <https://style.tidyverse.org/syntax.html>

⁷ <https://github.com/google/styleguide>

⁸ See for example McGovern (2012b)

to share conventions and understandings of what the code is doing. Style guides therefore help improve the quality of the code in that language that is produced by all programmers in a community. It is through a shared style that newer programmers can learn from more experienced programmers how certain coding practices are more or less error-prone. Broadly-accepted style conventions make it easier to borrow solutions from each other and from examples online without causing bugs that might only be found too late. Similarly, globally standardized style guides make it easier to solve each others' problems and to collaborate or move from project to project, and from team to team.

There is room for personal preference in style guides, but style guides are first and foremost about quality and standardization – especially when collaborating on code. We believe that a commonly used Stata style guide would improve the quality of all code written in Stata, which is why we have begun the one included here. You do not necessarily need to follow our style guide precisely. We encourage you to write your own style guide if you disagree with us. The best style guide would be the one adopted the most widely. All style rules introduced in this section are the way we suggest to code, but the most important thing is that the way you style your code is *consistent*. This guide allows our team to have a consistent code style.

Commenting code

Comments do not change the output of code, but without them, your code will not be accessible to your colleagues. It will also take you a much longer time to edit code you wrote in the past if you did not comment it well. So, comment a lot: do not only write *what* your code is doing but also *why* you wrote it like the way you did. In general, try to write simpler code that needs less explanation, even if you could use an elegant and complex method in less space, unless the advanced method is a widely accepted one.

There are three types of comments in Stata and they have different purposes:

```
stata-comments.do
```

```

1 TYPE 1:
2
3 /*
4     This is a do-file with examples of comments in Stata. This
5     type of comment is used to document all of the do-file or a large
6     section of it
7 */
8
9 TYPE 2:
10
11 * Standardize settings, explicitly set version, and clear memory
12 * (This comment is used to document a task covering at maximum a few lines of code)
13     ieboilstart, version(13.1)
14     `r(version)'
15
16 TYPE 3:
17
18 * Open the dataset
19     sysuse auto.dta // Built in dataset (This comment is used to document a single line)

```

Abbreviating commands

Stata commands can often be abbreviated in the code. You can tell if a command can be abbreviated if the help file indicates an abbreviation by underlining part of the name in the syntax section at the top. Only built-in commands can be abbreviated; user-written commands cannot. (Many commands additionally allow abbreviations of options: these are always acceptable at the shortest allowed abbreviation.) Although Stata allows some commands to be abbreviated to one or two characters, this can be confusing – two-letter abbreviations can rarely be “pronounced” in an obvious way that connects them to the functionality of the full command. Therefore, command abbreviations in code should not be shorter than three characters, with the exception of `tw` for `twoway` and `di` for `display`, and abbreviations should only be used when widely accepted abbreviation exists. We do not abbreviate `local`, `global`, `save`, `merge`, `append`, or `sort`. The following is a list of accepted abbreviations of common Stata commands:

Abbreviation	Command
tw	two way
di	display
gen	generate
mat	matrix
reg	regress
lab	label
sum	summarize
tab	tabulate
bys	bysort
qui	quietly
noi	noisily
cap	capture
forv	forvalues
prog	program
hist	histogram

Abbreviating variable names

Never abbreviate variable names. Instead, write them out completely. Your code may change if a variable is later introduced that has a name exactly as in the abbreviation. `ieboilstart` executes the command `set varabbrev off` by default, and will therefore break any code using variable abbreviations.

Using wildcards and lists in Stata for variable lists (*, ?, and -) is also discouraged, because the functionality of the code may change if the dataset is changed or even simply reordered. If you intend explicitly to capture all variables of a certain type, prefer `unab` or `lookfor` to build that list in a local macro, which can then be checked to have the right variables in the right order.

Writing loops

In Stata examples and other code languages, it is common for the name of the local generated by `foreach` or `forvalues` to be something as simple as `i` or `j`. In Stata, however, loops generally index a real object, and looping commands should name that index descriptively. One-letter indices are acceptable only for general examples; for looping through **iterations** with `i`; and for looping across matrices with `i`, `j`. Instead, index names should describe what the code is looping over – for example household members, crops, or medicines. Even counters should be explicitly named. This makes code much more readable, particularly in nested loops.

stata-loops.do

```

1  GOOD:
2
3 * Loop over crops
4 foreach crop in potato cassava maize {
5     * do something to `crop'
6 }
7
8 GOOD:
9
10 * Loop over crops
11 local crops potato cassava maize
12 foreach crop of local crops {
13     * Loop over plot number
14     forvalues plot_num = 1/10 {
15         * do something to `crop' in `plot_num'
16     } // End plot loop
17 } // End crop loop
18
19 BAD:
20
21 * Loop over crops
22 foreach i in potato cassava maize {
23     * do something to `i'
24 }
25

```

Using whitespace

In Stata, adding one or many spaces does not make a difference to code execution, and this can be used to make the code much more readable. We are all very well trained in using whitespace in software like PowerPoint and Excel: we would never present a PowerPoint presentation where the text does not align or submit an Excel table with unstructured rows and columns. The same principles apply to coding. In the example below the exact same code is written twice, but in the better example whitespace is used to signal to the reader that the central object of this segment of code is the variable employed. Organizing the code like this makes it much quicker to read, and small typos stand out more, making them easier to spot.

 stata-whitespace-columns.do

1 ACCEPTABLE:

```

2 * Create dummy for being employed
3 gen employed = 1
4 replace employed = 0 if (_merge == 2)
5 lab var employed "Person exists in employment data"
6 lab def yesno 1 "Yes" 0 "No"
7 lab val employed yesno
8
9
```

10 BETTER:

```

11 * Create dummy for being employed
12 gen employed = 1
13 replace employed = 0 if (_merge == 2)
14 lab var employed "Person exists in employment data"
15 lab def yesno 1 "Yes" 0 "No"
16 lab val employed yesno
17
```

Indentation is another type of whitespace that makes code more readable. Any segment of code that is repeated in a loop or conditional on an if-statement should have indentation of 4 spaces relative to both the loop or conditional statement as well as the closing curly brace. Similarly, continuing lines of code should be indented under the initial command. If a segment is in a loop inside a loop, then it should be indented another 4 spaces, making it 8 spaces more indented than the main code. In some code editors this indentation can be achieved by using the tab button. However, the type of tab used in the Stata do-file editor does not always display the same across platforms, such as when publishing the code on GitHub. Therefore we recommend that indentation be 4 manual spaces instead of a tab.

 stata-whitespace-indentation.do

```

1      GOOD:
2
3      * Loop over crops
4      foreach crop in potato cassava maize {
5          * Loop over plot number
6          forvalues plot_num = 1/10 {
7              gen crop_`crop'_`plot_num' = "`crop'"
8          }
9      }
10
11      * or
12      local sampleSize = `c(N)'
13      if (`sampleSize' <= 100) {
14          gen use_sample = 0
15      }
16      else {
17          gen use_sample = 1
18      }
19
20  BAD:
21
22      * Loop over crops
23      foreach crop in potato cassava maize {
24          * Loop over plot number
25          forvalues plot_num = 1/10 {
26              gen crop_`crop'_`plot_num' = "`crop'"
27          }
28      }
29
30      * or
31      local sampleSize = `c(N)'
32      if (`sampleSize' <= 100) {
33          gen use_sample = 0
34      }
35      else {
36          gen use_sample = 1
37      }

```

Writing conditional expressions

All conditional (true/false) expressions should be within at least one set of parentheses. The negation of logical expressions should use bang (!) and not tilde (~). Always use explicit truth checks (`if `value' == 1`) rather than implicits (`if `value'`). Always use the `missing(`var')` function instead of arguments like (`if `var' >= .`). Always consider whether missing values will affect the evaluation of conditional expressions.

 stata-conditional-expressions1.do

```

1 GOOD:
2
3     replace gender_string = "Woman" if (gender == 1)
4     replace gender_string = "Man"    if ((gender != 1) & !missing(gender))
5
6 BAD:
7
8     replace gender_string = "Woman" if gender == 1
9     replace gender_string = "Man"    if (gender ~= 1)

```

Use **if-else** statements when applicable even if you can express the same thing with two separate **if** statements. When using **if-else** statements you are communicating to anyone reading your code that the two cases are mutually exclusive, which makes your code more readable. It is also less error-prone and easier to update if you want to change the condition.

 stata-conditional-expressions2.do

```

1 GOOD:
2
3     if (`sampleSize' <= 100) {
4         * do something
5     }
6     else {
7         * do something else
8     }
9
10 BAD:
11
12    if (`sampleSize' <= 100) {
13        * do something
14    }
15    if (`sampleSize' > 100) {
16        * do something else
17    }

```

Using macros

Stata has several types of **macros** where numbers or text can be stored temporarily, but the two most common macros are **local** and **global**. Never abbreviate the commands for **local** and **global**. Locals should always be the default type and globals should only be used when the information stored is used in a different do-file. Globals are error-prone since they are active as long as Stata is open, which creates a risk that a global from one project is incorrectly used in another, so only use globals where they are necessary. Our recommendation is that globals should only be defined in the **master do-file**. All globals should be referenced using both the the dollar

sign and curly brackets around their name (`${ }`); otherwise, they can cause readability issues when the endpoint of the macro name is unclear.

You should use descriptive names for all macros (up to 32 characters; preferably fewer). There are several naming conventions you can use for macros with long or multi-word names. Which one you use is not as important as whether you and your team are consistent in how you name them. You can use all lower case (`mymacro`), underscores (`my_macro`), or “camel case” (`myMacro`), as long as you are consistent. Simple prefixes are useful and encouraged such as `this_estimate` or `current_var`, or, using `camelCase`, `lastValue`, `allValues`, or `nValues`.

stata-macros.do

```

1 GOOD:
2
3     global myGlobal = "A string global"
4     display "${myGlobal}"
5
6 BAD:
7
8     global my_Global = "A string global" // Do not mix naming styles
9     display "$myGlobal"                // Always use ${} for globals

```

Writing file paths

All file paths should always be enclosed in double quotes, and should always use forward slashes for folder hierarchies (/). File names should be written in lower case with dashes (`my-file.dta`). Mac and Linux computers cannot read file paths with backslashes, and backslashes cannot easily be removed with find-and-replace because the character has other functional uses in code. File paths should always include the file extension (.dta, .do, .csv, etc.). Omitting the extension causes ambiguity if another file with the same name is created (even if there is a default file type).

File paths should also be absolute and dynamic. **Absolute** means that all file paths start at the root folder of the computer, often C:/ on a PC or /Users/ on a Mac. This ensures that you always get the correct file in the correct folder. Do not use cd unless there is a function that requires it. When using cd, it is easy to overwrite a file in another project folder. Many Stata functions use cd and therefore the current directory may change without warning. Relative file paths are common in many other programming languages, but unless they are relative to the location of the file running the code, this is a risky practice. In Stata, relative file paths are relative to the current directory, not to the code file being run.

Dynamic file paths use global macros for the location of the root folder. These globals should be set in a central master do-file. This makes it possible to write file paths that work very similarly to relative paths. It also achieves the functionality that setting cd is often intended to: executing the code on a new system only requires updating file path globals in one location. If global names are unique, there is no risk that files are saved in the incorrect project folder. You can create multiple folder globals as needed and this is encouraged.

 stata-filepaths.do

```

1 GOOD:
2
3   global myDocs    = "C:/Users/username/Documents"
4   global myProject = "${myDocs}/MyProject"
5   use "${myProject}/my-dataset.dta" , clear
6
7 BAD:
8
9   RELATIVE PATHS:
10  cd "C:/Users/username/Documents/MyProject"
11  use MyDataset.dta
12
13  STATIC PATHS:
14  use "C:/Users/username/Documents/MyProject/MyDataset.dta"
```

Line breaks

Long lines of code are difficult to read if you have to scroll left and right to see the full line of code. When your line of code is wider than text on a regular paper, you should introduce a line break. A common line breaking length is around 80 characters. Stata's do-file editor and other code editors provide a visible guide line. Around that length, start a new line using `///`. Using `///` breaks the line in the code editor, while telling Stata that the same line of code continues on the next line. Recent versions of the Stata do-file editor – and many other code editors – automatically wrap code lines that are too long, but you should still use `///` to make sure that the lines breaks are placed where they are most readable. The `///` breaks do not need to be horizontally aligned in code, although you may prefer to if they have comments that read better aligned, since indentations should reflect that the command continues to a new line. Break lines where it makes functional sense. You can write comments after `///` just as with `//`, and that is usually a good thing.

The `#delimit` command should only be used for advanced function programming and is officially discouraged in analytical code.⁹ Never, for any reason, use `/* */` to wrap a line: it is distracting and difficult to follow compared to the use of those characters to write

⁹ Cox (2005)

regular comments. Line breaks and indentations may be used to highlight the placement of the **option comma** or other functional syntax in Stata commands.

stata-linebreak.do

```

1 GOOD:
2     graph hbar invil      /// Proportion in village
3         if (priv == 1)    /// Private facilities only
4             , over(stlename, sort(1) descending)      /// Order states by values
5                 blabel(bar, format(%9.0f))           /// Label the bars
6                 ylab(0 "0%" 25 "25%" 50 "50%" 75 "75%" 100 "100%") ///
7                     ytit("Share of private primary care visits made in own village")
8
9 BAD:
10    #delimit ;
11    graph hbar
12        invil if (priv == 1)
13            , over(stlename, sort(1) descending) blabel(bar, format(%9.0f))
14            ylab(0 "0%" 25 "25%" 50 "50%" 75 "75%" 100 "100%")
15            ytit("Share of private primary care visits made in own village");
16    #delimit cr
17
18 UGLY:
19     graph hbar /*
20     */      invil if (priv == 1)

```

Using boilerplate code

Boilerplate code consists of a few lines of code that always come at the top of the code file, and its purpose is to harmonize settings across users running the same code to the greatest degree possible. There is no way in Stata to guarantee that any two installations will always run code in exactly the same way. In the vast majority of cases it does, but not always, and boilerplate code can mitigate that risk. We have developed the `ieboilstart` command to implement many commonly-used boilerplate settings that are optimized given your installation of Stata. It requires two lines of code to execute the `version` setting, which avoids differences in results due to different versions of Stata. Among other things, it turns the `more` flag off so code never hangs while waiting to display more output; it turns `varabbrev` off so abbreviated variable names are rejected; and it maximizes the allowed memory usage and matrix size so that code is not rejected on other machines for violating system limits. (For example, Stata/SE and Stata/IC, allow for different maximum numbers of variables, and the same happens with Stata 14 and Stata 15, so it may not be able to run code written in one of these version using another.) Finally, it clears all stored information in Stata memory, such as non-installed programs and globals, getting as close as possible to opening Stata fresh.

 stata-boilerplate.do

```

1 GOOD:
2
3 `ieboilstart, version(13.1)
4 `r(version)'
5
6 OK:
7
8 set more off , perm
9 clear all
10 set maxvar 10000
11 version 13.1

```

Saving data

There are good practices that should be followed before saving any dataset. These are to `sort` and `order` the dataset, dropping intermediate variables that are not needed, and compressing the dataset to save disk space and network bandwidth.

If there is a unique ID variable or a set of ID variables, the code should test that they uniquely and fully identify the dataset.¹⁰ ID variables are also perfect variables to sort on, and to `order` first in the dataset.

The command `compress` makes the dataset smaller in terms of memory usage without ever losing any information. It optimizes the storage types for all variables and therefore makes it smaller on your computer and faster to send over a network or the internet.

¹⁰ https://dimewiki.worldbank.org/ID_Variable.Properties

 stata-before-saving.do

```

1 * If the dataset has ID variables, test if they uniquely identifying the observations.
2
3 local idvars household_ID household_member year
4 isid `idvars'
5
6 * Sort and order on the idvars (or any other variables if there are no ID variables)
7
8 sort `idvars'
9 order *, seq // Place all variables in alphanumeric order (optional but useful)
10 order `idvars' , first // Make sure the idvars are the leftmost vars when browsing
11
12 * Drop intermediate variables no longer needed
13
14 * Optimize disk space
15
16 compress
17
18 * Save data
19
20 save    "${myProject}/myDataFile.dta" , replace // The folder global is set in master do-file
21 saveold "${myProject}/myDataFile-13.dta" , replace v(13) // For users with older versions of Stata

```

Miscellaneous notes

Write multiple graphs as `tw (xx)(xx)(xx)`, not `tw xx| |xx| |xx`.

In simple expressions, put spaces around each binary operator except `^`. Therefore write `gen z = x + y` and `x^2`.

When order of operations applies, you may adjust spacing and parentheses: write `hours + (minutes/60) + (seconds/3600)`, not `hours + minutes / 60 + seconds / 3600`. For long expressions, + and - operators should start the new line, but * and / should be used inline. For example:

```
gen newvar = x ///
- (y/2) ///
+ a * (b - c)
```

Make sure your code doesn't print very much to the results window as this is slow. This can be accomplished by using `run file.do` rather than `do file.do`. Interactive commands like `sum` or `tab` should be used sparingly in do-files, unless they are for the purpose of getting `r()`-statistics. In that case, consider using the `qui` prefix to prevent printing output. It is also faster to get outputs from commands like `reg` using the `qui` prefix.

Appendix: DIME Analytics Resource Directory

The resources listed in this appendix are mentioned elsewhere in the chapters of this book, and this appendix includes them all in one place for easy reference. All these resources are made public under generous open-source licenses. This means that you are free to use, reuse, and adapt these resources for any purpose as you see fit, so long as you include an appropriate citation.

Public Resources and Tools

DIME Wiki.¹ One-stop shop for impact evaluation research solutions. The DIME Wiki is a resource focused on practical implementation guidelines rather than theory, open to the public, easily searchable, suitable for users of varying levels of expertise, up-to-date with the latest technological advances in electronic data collection, and curated by a vibrant network of editors with expertise in the field.

¹ <https://dimewiki.worldbank.org>

Stata Visual Library.² A curated, easy-to-browse selection of graphs created in Stata. Clicking on each graph reveals the source code, to allow for easy replication.

² <https://worldbank.github.io/Stata-IE-Visual-Library/>

R Econ Visual Library.³ A curated, easy-to-browse selection of graphs created in R. Clicking on each graph reveals the source code, to allow for easy replication.

³ <https://worldbank.github.io/r-econ-visual-library/>

DIME Analytics Research Standards.⁴ A repository outlining DIME's public commitments to research ethics, transparency, reproducibility, data security and data publication, along with supporting tools and resources.

⁴ <https://github.com/worldbank/dime-standards>

Flagship Courses

Manage Successful Impact Evaluations (MSIE).⁵ DIME Analytics' flagship training is a week-long annual course, held in person in Washington, D.C. MSIE is intended to improve the skills and knowledge of impact evaluation (IE) practitioners, familiarizing them with critical issues in IE implementation, recurring challenges, and cutting-edge technologies. The course consists of lectures and hands-on

⁵ <https://osf.io/h4d8y/>

sessions. Through small group discussions and interactive computer lab sessions, participants work together to apply what they've learned and have a first-hand opportunity to develop skills. Hands-on sessions are offered in parallel tracks, with different options based on software preferences and skill level.

Manage Successful Impact Evaluation Surveys (MSIES).⁶ A fully virtual course, in which participants learn the workflow for primary data collection. The course covers best practices at all stages of the survey workflow, from planning to piloting instruments and monitoring data quality once fieldwork begins. There is a strong focus throughout on research ethics and reproducible workflows. The course uses a combination of virtual lectures, case studies, readings, and hands-on exercises.

⁶ <https://osf.io/resya/>

Research Assistant Onboarding Course.⁷ This course is designed to familiarize Research Assistants and Research Analysts with DIME's standards for data work. By the end of the course's six sessions, participants will have the tools and knowledge to implement best practices for transparent and reproducible research. The course will focus on how to set up a collaborative workflow for code, datasets, and research outputs. Most content is platform-independent and software-agnostic, but participants are expected to be familiar with statistical software.

⁷ <https://osf.io/qtmdp>

Introduction to R for advanced Stata users.⁸ This course is an introduction to the R programming language, building upon knowledge of Stata. The course focuses on common tasks in development research, related to descriptive analysis, data visualization, data processing, and geospatial data work.

⁸ <https://osf.io/nj6bf>

DIME Analytics Trainings.⁹ DIME Analytics' home on the Open Science Framework, with links to materials for all past courses and technical trainings.

⁹ <https://osf.io/wzjtk>

Software tools and trainings

ietoolkit.¹⁰ Suite of Stata commands to routinize common tasks for data management and impact evaluation analysis.

¹⁰ <https://github.com/worldbank/ietoolkit>

iefieldkit.¹¹ Suite of Stata commands to routinize and document common tasks in primary data collection.

¹¹ <https://github.com/worldbank/iefieldkit>

DIME Analytics GitHub Trainings and Resources.¹² A GitHub repository containing all the GitHub training materials and resources

¹² <https://github.com/worldbank/dime-github-trainings>

developed by DIME Analytics. The trainings follow DIME’s model for organizing research teams on GitHub, and are designed for face-to-face delivery, but materials are shared so that they may be used and adapted by others.

DIME Analytics L^AT_EX-training.¹³ A user-friendly guide to getting started with LaTeX. Exercises provide opportunities to practice creating appendices, exporting tables from R or Stata to LaTeX, and formatting tables in LaTeX.

¹³ <https://github.com/worldbank/DIME-LaTeX-Templates>

Appendix: Research design for impact evaluation

Development Research in Practice focuses on tools, workflows, and practical guidance for implementing research projects. While not central to the content of this book, we think it is essential for all research team members – including field staff and research assistants – to understand research design, and specifically how research design choices impact data work. Without going into too much technical detail, as there are many excellent resources on impact evaluation design, this appendix presents a brief overview of the most common causal inference methods, focusing on implications for data structure and analysis. This appendix is intended to be a reference, especially for junior team members, to obtain an understanding of the way in which each causal inference method constructs treatment and control groups, the data structures needed to estimate the corresponding effects, and specific code tools designed for each method.

It is essential for the research team members who will do the data work to understand the study design, for several reasons. If you do not know how to calculate the correct estimator for your study, you will not be able to assess the statistical power of your research design. You will also be unable to make decisions in the field when you inevitably have to allocate scarce resources between tasks like maximizing sample size and ensuring follow-up with specific individuals. You will save time by understanding the way your data needs to be organized to produce meaningful analytics throughout your projects. Just as importantly, familiarity with each of these approaches will allow you to keep your eyes open for research opportunities: many of the most interesting projects occur because people in the field recognize the opportunity to implement one of these methods in response to an unexpected event.

This appendix is split into two sections. The first covers causal inference methods in experimental and quasi-experimental research designs. The second discusses how to measure treatment effects and structure data for specific methods, including cross-sectional randomized control trials, difference-in-difference designs, regression discontinuity, instrumental variables, matching, and synthetic controls.

Causality, inference, and identification

When we are discussing the types of inputs – “treatments” – commonly referred to as “programs” or “interventions”, we are typically attempting to obtain estimates of program-specific **treatment effects**.

These are the changes in outcomes attributable to the treatment.¹

¹ Abadie and Cattaneo (2018)

The primary goal of research design is to establish **causal identification** for an effect. Causal identification means establishing that a change in an input directly altered an outcome. When a study is well-identified, then we can say with confidence that our estimate of the treatment effect would, with an infinite amount of data, give us a

precise estimate of that treatment effect. Under this condition, we can proceed to draw evidence from the limited samples we have access to, using statistical techniques to express the uncertainty of not having infinite data. Without identification, we cannot say that the estimate would be accurate, even with unlimited data, and therefore cannot attribute it to the treatment in the small samples that we typically have access to. More data is not a substitute for a well-identified experimental design. Therefore it is important to understand how exactly your study identifies its estimate of treatment effects, so you can calculate and interpret those estimates appropriately.

All the study designs we discuss here use the potential outcomes framework² to compare a group that received some treatment to another, counterfactual group. Each of these approaches can be used in two types of designs: **experimental** designs, in which the research team is directly responsible for creating the variation in treatment, and **quasi-experimental** designs, in which the team identifies a “natural” source of variation and uses it for identification. Neither type is implicitly better or worse, and both types are capable of achieving causal identification in different contexts.

² Athey and Imbens (2017b)

Estimating treatment effects using control groups

The key assumption behind estimating treatment effects is that every person, facility, or village (or whatever the unit of intervention is) has two possible states: their outcomes if they do not receive some treatment and their outcomes if they do receive that treatment. Each unit’s treatment effect is the individual difference between these two states, and the **average treatment effect (ATE)** is the average of all individual differences across the potentially treated population. This is the parameter that most research designs attempt to estimate, by establishing a **counterfactual**³ for the treatment group against which outcomes can be directly compared. There are several resources that provide more or less mathematically intensive approaches to understanding how various methods do this. *Impact Evaluation in Practice*⁴ is a strong general guide to these methods. *Causal Inference*⁵ and *Causal Inference: The Mixtape*⁶ provides more detailed mathematical approaches to the tools. *Mostly Harmless Econometrics*⁷ and *Mastering Metrics*⁸ are excellent resources on the statistical principles behind all econometric approaches.

Intuitively, the problem is as follows: we can never observe the same unit in both their treated and untreated states simultaneously, so measuring and averaging these effects directly is impossible.⁹ Instead, we typically make inferences from samples. **Causal inference** methods are those in which we are able to estimate the average treat-

³ **Counterfactual:** A statistical description of what would have happened to specific individuals in an alternative scenario, for example, a different treatment assignment outcome.

⁴ Gertler et al. (2016)

⁵ Hernán and Robins (2010)

⁶ Cunningham (2018)

⁷ Angrist and Pischke (2008)

⁸ Angrist and Pischke (2014)

⁹ Rubin (2003)

ment effect without observing individual-level effects, but through some comparison of averages with a **control** group. Every research design is based on a way of comparing another set of observations – the “control” observations – against the treatment group. They all work to establish that the control observations would have been identical *on average* to the treated group in the absence of the treatment. Then, the mathematical properties of averages imply that the calculated difference in averages is equivalent to the average difference: exactly the parameter we are seeking to estimate. Therefore, almost all designs can be accurately described as a series of between-group comparisons.

Most of the methods that you will encounter rely on some variant of this strategy, which is designed to maximize their ability to estimate the effect of an average unit being offered the treatment you want to evaluate. The focus on identification of the treatment effect, however, means there are several essential features of causal identification methods that are not common in other types of statistical and data science work. First, the econometric models and estimating equations used do not attempt to create a predictive or comprehensive model of how the outcome of interest is generated. Typically, causal inference designs are not interested in predictive accuracy, and the estimates and predictions that they produce will not be as good at predicting outcomes or fitting the data as other models. Second, when control variables or other variables are used in estimation, there is no guarantee that the resulting parameters are marginal effects. They can only be interpreted as correlative averages, unless there are additional sources of identification. The models you will construct and estimate are intended to do exactly one thing: to express the intention of your project’s research design, and to accurately estimate the effect of the treatment it is evaluating. In other words, these models tell the story of the research design in a way that clarifies the exact comparison being made between control and treatment.

Experimental and quasi-experimental research designs

Experimental research designs explicitly allow the research team to change the condition of the populations being studied,¹⁰ often in the form of government programs, NGO projects, new regulations, information campaigns, and many more types of interventions.¹¹ The classic experimental causal inference method is the **randomized control trial (RCT)**.¹² In randomized control trials, the treatment group is randomized – that is, from an eligible population, a random group of units are given the treatment. Another way to think about these designs is how they establish the control group: a random subset of

¹⁰ [https://dimewiki.worldbank.org/
Experimental_Methods](https://dimewiki.worldbank.org/Experimental_Methods)

¹¹ Banerjee and Duflo (2009)

¹² [https://dimewiki.worldbank.org/
Randomized_Control_Trials](https://dimewiki.worldbank.org/Randomized_Control_Trials)

units are *not* given access to the treatment, so that they may serve as a counterfactual for those who are. A randomized control group, intuitively, is meant to represent how things would have turned out for the treated group if they had not been treated, and it is particularly effective at doing so as evidenced by its broad credibility in fields ranging from clinical medicine to development. Therefore RCTs are very popular tools for determining the causal impact of specific programs or policy interventions, as evidenced by the awarding of the 2019 Nobel Prize in Economics to Abhijit Banerjee, Esther Duflo and Michael Kremer “for their experimental approach to alleviating global poverty”.¹³ However, there are many other types of interventions that are impractical or unethical to effectively approach using an experimental strategy, and therefore there are limitations to accessing “big questions” through RCT approaches.¹⁴

Randomized designs all share several major statistical concerns. The first is the fact that it is always possible to select a control group, by chance, which is not in fact very similar to the treatment group. This feature is called randomization noise, and all RCTs share the need to assess how randomization noise may impact the estimates that are obtained. (More detail on this later.) Second, take-up and implementation fidelity are extremely important, since programs will by definition have no effect if the population intended to be treated does not accept or does not receive the treatment.¹⁵ Loss of statistical power occurs quickly and is highly nonlinear: 70% take-up or efficacy doubles the required sample, and 50% quadruples it.¹⁶ Such effects are also very hard to correct ex post, since they require strong assumptions about the randomness or non-randomness of take-up. Therefore a large amount of field time and descriptive work must be dedicated to understanding how these effects played out in a given study, and may overshadow the effort put into the econometric design itself.

Quasi-experimental research designs,¹⁷ by contrast, are causal inference methods based on events not controlled by the research team. Instead, they rely on “experiments of nature”, in which natural variation can be argued to approximate the type of exogenous variation in treatment availability that a researcher would attempt to create with an experiment.¹⁸ Unlike carefully planned experimental designs, quasi-experimental designs typically require the extra luck of having access to data collected at the right times and places to exploit events that occurred in the past, or having the ability to collect data in a time and place where an event that produces causal identification occurred or will occur. Therefore, these methods often use either secondary data, or they use primary data in a cross-sectional retrospective method, including administrative data or other new

¹³ NobelPrize.org (2020). The Prize in Economic Sciences 2019. [Online; accessed 10-September-2020]

¹⁴ Deaton (2009)

¹⁵ See Jung and Hasan (2016) for an example.

¹⁶ <https://blogs.worldbank.org/impactevaluations/power-calculations-101-dealing-with-incomplete-take-up>

¹⁷ https://dimewiki.worldbank.org/Quasi-Experimental_Methods

¹⁸ DiNardo (2016)

classes of routinely-collected information.

Quasi-experimental designs therefore can access a much broader range of questions, and with much less effort in terms of executing an intervention. However, they require in-depth understanding of the precise events the researcher wishes to address in order to know what data to use and how to model the underlying natural experiment. Additionally, because the population exposed to such events is limited by the scale of the event, quasi-experimental designs are often power-constrained. Since the research team cannot change the population of the study or the treatment assignment, power is typically maximized by ensuring that sampling for data collection is carefully designed to match the study objectives and that attrition from the sampled groups is minimized.

Obtaining treatment effects from specific research designs

Cross-sectional designs

A cross-sectional research design is any type of study that observes data in only one time period and directly compares treatment and control groups. This type of data is easy to collect and handle because you do not need to track individuals across time. If this point in time is after a treatment has been fully delivered, then the outcome values at that point in time already reflect the effect of the treatment. If the study is experimental, the treatment and control groups are randomly constructed from the population that is eligible to receive each treatment. By construction, each unit's receipt of the treatment is unrelated to any of its other characteristics and the ordinary least squares (OLS) regression of outcome on treatment, without any control variables, is an unbiased estimate of the average treatment effect.

Cross-sectional designs can also exploit variation in non-experimental data to argue that observed correlations do in fact represent causal effects. This can be true unconditionally – which is to say that something random, such as winning the lottery, is a true random process and can tell you about the effect of getting a large amount of money.¹⁹ It can also be true conditionally – which is to say that once the characteristics that would affect both the likelihood of exposure to a treatment and the outcome of interest are controlled for, the process is as good as random: like arguing that once risk preferences are taken into account, exposure to an earthquake is unpredictable and post-event differences are causally related to the event itself.²⁰

¹⁹ Imbens et al. (2001)

For cross-sectional designs, what needs to be carefully maintained in data is the treatment randomization process itself (whether ex-

²⁰ Callen (2015)

perimental or not), as well as detailed information about differences in data quality and attrition across groups.²¹ Only these details are needed to construct the appropriate estimator: clustering of the standard errors is required at the level at which the treatment is assigned to observations, and variables which were used to stratify the treatment must be included as controls (in the form of strata fixed effects).²² **Randomization inference** can be used to estimate the underlying variability in the randomization process (more on this in the next chapter). **Balance checks**²³ are often reported as evidence of an effective randomization, and are particularly important when the design is quasi-experimental (since then the randomization process cannot be simulated explicitly). However, controls for balance variables are usually unnecessary in RCTs, because it is certain that the true data-generating process has no correlation between the treatment and the balance factors.²⁴

Analysis is typically straightforward *once you have a strong understanding of the randomization*. A typical analysis will include a description of the sampling and randomization results, with analyses such as summary statistics for the eligible population, and balance checks for randomization and sample selection. The main results will usually be a primary regression specification (with multiple hypotheses²⁵ appropriately adjusted for), and additional specifications with adjustments for non-response, balance, and other potential contamination. Robustness checks might include randomization-inference analysis or other placebo regression approaches. There are a number of user-written code tools that are also available to help with the complete process of data analysis, including to analyze balance²⁶ and to visualize treatment effects.²⁷ Extensive tools and methods for analyzing selective non-response are available.²⁸

Difference-in-differences

Where cross-sectional designs draw their estimates of treatment effects from differences in outcome levels in a single measurement, **differences-in-differences**²⁹ designs (abbreviated as DD, DiD, diff-in-diff, and other variants) estimate treatment effects from *changes* in outcomes between two or more rounds of measurement. In these designs, three control groups are used – the baseline level of treatment units, the baseline level of non-treatment units, and the endline level of non-treatment units.³⁰ The estimated treatment effect is the excess growth of units that receive the treatment, in the period they receive it: calculating that value is equivalent to taking the difference in means at endline and subtracting the difference in means at baseline (hence the singular “difference-in-differences”).³¹ The regression

²¹ Athey and Imbens (2017a)

²² <https://blogs.worldbank.org/impactevaluations/impactevaluations/how-randomize-using-many-baseline-variables-guest-post-thomas-barrios>

²³ **Balance checks:** Statistical tests of the similarity of treatment and control groups.

²⁴ <https://blogs.worldbank.org/impactevaluations/should-we-require-balance-t-tests-baseline-observables-randomized-experiments>

²⁵ See Armand et al. (2017) for an example.

²⁶ <https://dimewiki.worldbank.org/iebaltab>

²⁷ <https://dimewiki.worldbank.org/iegraph>

²⁸ <https://blogs.worldbank.org/impactevaluations/dealing-attrition-field-experiments>

²⁹ <https://dimewiki.worldbank.org/Difference-in-Differences>

³⁰ Torres-Reyna (2015)

³¹ McKenzie (2012)

model includes a control variable for treatment assignment, and a control variable for time period, but the treatment effect estimate corresponds to an interaction variable for treatment and time: it indicates the group of observations for which the treatment is active. This model depends on the assumption that, in the absense of the treatment, the outcome of the two groups would have changed at the same rate over time, typically referred to as the **parallel trends** assumption.³² Experimental approaches satisfy this requirement in expectation, but a given randomization should still be checked for pre-trends as an extension of balance checking.³³

There are two main types of data structures for differences-in-differences: **repeated cross-sections** and **panel data**. In repeated cross-sections, each successive round of data collection contains a random sample of observations from the treated and untreated groups; as in cross-sectional designs, both the randomization and sampling processes are critically important to maintain alongside the data. In panel data structures, we attempt to observe the exact same units in different points in time, so that we see the same individuals both before and after they have received treatment (or not).³⁴ This allows each unit's baseline outcome (the outcome before the intervention) to be used as an additional control for its endline outcome, which can provide large increases in power and robustness.³⁵ When tracking individuals over time for this purpose, maintaining sampling and tracking records is especially important, because attrition will remove that unit's information from all points in time, not just the one they are unobserved in. Panel-style experiments therefore require a lot more effort in field work for studies that use original data.³⁶ Since baseline and endline may be far apart in time, it is important to create careful records during the first round so that follow-ups can be conducted with the same subjects, and attrition across rounds can be properly taken into account.³⁷

As with cross-sectional designs, difference-in-differences designs are widespread. Therefore there exist a large number of standardized tools for analysis. Our *ietoolkit* Stata package includes the *ieddtab* command which produces standardized tables for reporting results.³⁸ For more complicated versions of the model (and they can get quite complicated quite quickly), you can use an online dashboard to simulate counterfactual results.³⁹ As in cross-sectional designs, these main specifications will always be accompanied by balance checks (using baseline values), as well as randomization, selection, and attrition analysis. In trials of this type, reporting experimental design and execution using the CONSORT style is common in many disciplines and will help you to track your data over time.⁴⁰

³² <https://blogs.worldbank.org/impactevaluations/often-unspoken-assumptions-behind-difference-difference-estimator-practice>

³³ <https://blogs.worldbank.org/impactevaluations/revisiting-difference-differences-parallel-trends-assumption-part-i-pre-trend>

³⁴ <https://blogs.worldbank.org/impactevaluations/what-are-we-estimating-when-we-estimate-difference-differences>

³⁵ <https://blogs.worldbank.org/impactevaluations/another-reason-prefer-ancova-dealing-changes-measurement-between-baseline-and-follow>

³⁶ Torres-Reyna (2007)

³⁷ <https://blogs.worldbank.org/impactevaluations/dealing-attrition-field-experiments>

³⁸ <https://dimewiki.worldbank.org/ieddtab>

³⁹ <https://blogs.worldbank.org/impactevaluations/econometrics-sandbox-event-study-designs-co>

⁴⁰ Schulz et al. (2010)

Regression discontinuity

Regression discontinuity (RD) designs exploit sharp breaks or limits in policy designs to separate a single group of potentially eligible recipients into comparable groups of individuals who do and do not receive a treatment.⁴¹ These designs differ from cross-sectional and diff-in-diff designs in that the group eligible to receive treatment is not defined directly, but instead created during the treatment implementation. In an RD design, there is typically some program or event that has limited availability due to practical considerations or policy choices and is therefore made available only to individuals who meet a certain threshold requirement. The intuition of this design is that there is an underlying **running variable** that serves as the sole determinant of access to the program, and a strict cutoff determines the value of this variable at which eligibility stops.⁴² Common examples are test score thresholds and income thresholds.⁴³ The intuition is that individuals who are just above the threshold will be very nearly indistinguishable from those who are just under it, and their post-treatment outcomes are therefore directly comparable.⁴⁴ The key assumption here is that the running variable cannot be directly manipulated by the potential recipients. If the running variable is time (what is commonly called an “event study”), there are special considerations.⁴⁵ Similarly, spatial discontinuity designs are handled a bit differently due to their multidimensionality.⁴⁶

Regression discontinuity designs are, once implemented, very similar in analysis to cross-sectional or difference-in-differences designs. Depending on the data that is available, the analytical approach will center on the comparison of individuals who are narrowly on the inclusion side of the discontinuity, compared against those who are narrowly on the exclusion side.⁴⁷ The regression model will be identical to the matching research designs, i.e., contingent on whether data has one or more time periods and whether the same units are known to be observed repeatedly. The treatment effect will be identified, however, by the addition of a control for the running variable – meaning that the treatment effect estimate will only be applicable for observations in a small window around the cutoff: in the lingo, the treatment effects estimated will be “local” rather than “average”. In the RD model, the functional form of the running variable control and the size of that window, often referred to as the choice of **bandwidth** for the design, are the critical parameters for the result.⁴⁸ Therefore, RD analysis often includes extensive robustness checking using a variety of both functional forms and bandwidths, as well as placebo testing for non-realized locations of the cutoff.

In the analytical stage, regression discontinuity designs often

⁴¹ https://dimewiki.worldbank.org/Regression_Discontinuity

⁴² Imbens and Lemieux (2008)

⁴³ <https://blogs.worldbank.org/impactevaluations/regression-discontinuity-porn>

⁴⁴ Lee and Lemieux (2010)

⁴⁵ Hausman and Rapson (2018)

⁴⁶ <https://blogs.worldbank.org/impactevaluations/spatial-jumps>

⁴⁷ Cattaneo et al. (2019)

⁴⁸ Calonico et al. (2019)

include a large component of visual evidence presentation. These presentations help to suggest both the functional form of the underlying relationship and the type of change observed at the discontinuity, and help to avoid pitfalls in modeling that are difficult to detect with hypothesis tests.⁴⁹ Because these designs are so flexible compared to others, there is an extensive set of commands that help assess the efficacy and results from these designs under various assumptions.⁵⁰ These packages support the testing and reporting of robust plotting and estimation procedures, tests for manipulation of the running variable, and tests for power, sample size, and randomization inference approaches that will complement the main regression approach used for point estimates.

⁴⁹ Pischke (2018)

⁵⁰ Calonico et al. (2014)

Instrumental variables

Instrumental variables (IV) designs, unlike the previous approaches, begin by assuming that the treatment delivered in the study in question is linked to the outcome in a pattern such that its effect is not directly identifiable. Instead, similar to regression discontinuity designs, IV attempts to focus on a subset of the variation in treatment take-up and assesses that limited window of variation that can be argued to be unrelated to other factors.⁵¹ To do so, the IV approach selects an **instrument** for the treatment status – an otherwise-unrelated predictor of exposure to treatment that affects the take-up status of an individual.⁵² Whereas regression discontinuity designs are “sharp” – treatment status is completely determined by which side of a cutoff an individual is on – IV designs are “fuzzy”, meaning that they do not completely determine the treatment status but instead influence the *probability* of treatment.

⁵¹ Angrist and Krueger (2001)

⁵² https://dimewiki.worldbank.org/instrumental_variables

As in regression discontinuity designs, the fundamental form of the regression is similar to either cross-sectional or difference-in-differences designs. However, instead of controlling for the instrument directly, the IV approach typically uses the **two-stage-least-squares (2SLS)** estimator.⁵³ This estimator forms a prediction of the probability that the unit receives treatment based on a regression against the instrumental variable. That prediction will, by assumption, be the portion of the actual treatment that is due to the instrument and not any other source, and since the instrument is unrelated to all other factors, this portion of the treatment can be used to assess its effects. Unfortunately, these estimators are known to have very high variances relative other methods, particularly when the relationship between the instrument and the treatment is small.⁵⁴ IV designs furthermore rely on strong but untestable assumptions about the relationship between the instrument and the outcome.⁵⁵ Therefore

⁵³ Bond (2020)

⁵⁴ Young (2019)

⁵⁵ Bound et al. (1995)

IV designs face intense scrutiny on the strength and exogeneity of the instrument, and tests for sensitivity to alternative specifications and samples are usually required with an instrumental variables analysis. However, the method has special experimental cases that are significantly easier to assess: for example, a randomized treatment *assignment* can be used as an instrument for the eventual take-up of the treatment itself,⁵⁶ especially in cases where take-up is expected to be low, or in circumstances where the treatment is available to those who are not specifically assigned to it (“encouragement designs”).

In practice, there are a variety of packages that can be used to analyse data and report results from instrumental variables designs. While the built-in Stata command `ivregress` will often be used to create the final results, the built-in packages are not sufficient on their own. The **first stage** of the design should be extensively tested, to demonstrate the strength of the relationship between the instrument and the treatment variable being instrumented.⁵⁷ This can be done using the `weakiv` and `weakivtest` commands.⁵⁸ Additionally, tests should be run that identify and exclude individual observations or clusters that have extreme effects on the estimator, using customized bootstrap or leave-one-out approaches.⁵⁹ Finally, bounds can be constructed allowing for imperfections in the exogeneity of the instrument using loosened assumptions, particularly when the underlying instrument is not directly randomized.⁶⁰

⁵⁶ See Iacovone et al. (2019) for an example.

⁵⁷ Stock and Yogo (2005)

⁵⁸ Pflueger and Wang (2015)

⁵⁹ Young (2019)

⁶⁰ Clarke and Matta (2018)

Matching

Matching methods use observable characteristics of individuals to directly construct treatment and control groups to be as similar as possible to each other, either before a randomization process or after the collection of non-randomized data.⁶¹ Matching observations may be one-to-one or many-to-many; in any case, the result of a matching process is similar in concept to the use of randomization strata in simple randomized control trials. In this way, the method can be conceptualized as averaging across the results of a large number of “micro-experiments” in which the randomized units are verifiably similar aside from the treatment.

When matching is performed before a randomization process, it can be done on any observable characteristics, including outcomes, if they are available. The randomization should then record an indicator for each matching set, as these become equivalent to randomization strata and require controls in analysis. This approach is stratification taken to its most extreme: it reduces the number of potential randomizations dramatically from the possible number that would be available if the matching was not conducted, and therefore

⁶¹ <https://dimewiki.worldbank.org/Matching>

reduces the variance caused by the study design. When matching is done ex post in order to substitute for randomization, it is based on the assertion that within the matched groups, the assignment of treatment is as good as random. However, since most matching models rely on a specific linear model, such as **propensity score matching**,⁶² they are open to the criticism of “specification searching”, meaning that researchers can try different models of matching until one, by chance, leads to the final result that was desired; analytical approaches have shown that the better the fit of the matching model, the more likely it is that it has arisen by chance and is therefore biased.⁶³ Newer methods, such as **coarsened exact matching**,⁶⁴ are designed to remove some of the dependence on linearity. In all ex-post cases, pre-specification of the exact matching model can prevent some of the potential criticisms on this front, but ex-post matching in general is not regarded as a strong identification strategy.

Analysis of data from matching designs is relatively straightforward; the simplest design only requires controls (indicator variables) for each group or, in the case of propensity scoring and similar approaches, weighting the data appropriately in order to balance the analytical samples on the selected variables. The **teffects** suite in Stata provides a wide variety of estimators and analytical tools for various designs.⁶⁵ The coarsened exact matching (**cem**) package applies the nonparametric approach.⁶⁶ DIME’s **iematch** command in the **ietoolkit** package produces matchings based on a single continuous matching variable.⁶⁷ In any of these cases, detailed reporting of the matching model is required, including the resulting effective weights of observations, since in some cases the lack of overlapping supports for treatment and control mean that a large number of observations will be weighted near zero and the estimated effect will be generated based on a subset of the data.

Synthetic controls

Synthetic control is a relatively new method for the case when appropriate counterfactual individuals do not exist in reality and there are very few (often only one) treatment units.⁶⁸ For example, state- or national-level policy changes that can only be analyzed as a single unit are typically very difficult to find valid comparators for, since the set of potential comparators is usually small and diverse and therefore there are no close matches to the treated unit. Intuitively, the synthetic control method works by constructing a counterfactual version of the treated unit using an average of the other units available.⁶⁹ This is a particularly effective approach when the lower-level components of the units would be directly comparable: people,

⁶² **Propensity Score Matching (PSM):** An estimation method that controls for the likelihood that each unit of observation would receive treatment as predicted by observable characteristics.

⁶³ King and Nielsen (2019)

⁶⁴ Iacus et al. (2012)

⁶⁵ Cooperative (2015)

⁶⁶ Blackwell et al. (2009)

⁶⁷ <https://dimewiki.worldbank.org/iematch>

⁶⁸ Abadie et al. (2015)

⁶⁹ Abadie et al. (2010)

households, business, and so on in the case of states and countries; or passengers or cargo shipments in the case of transport corridors, for example.⁷⁰ This is because in those situations the average of the untreated units can be thought of as balancing by matching the composition of the treated unit.

To construct this estimator, the synthetic controls method requires retrospective data on the treatment unit and possible comparators, including historical data on the outcome of interest for all units.⁷¹ The counterfactual blend is chosen by optimizing the prediction of past outcomes based on the potential input characteristics, and typically selects a small set of comparators to weight into the final analysis. These datasets therefore may not have a large number of variables or observations, but the extent of the time series both before and after the implementation of the treatment are key sources of power for the estimate, as are the number of counterfactual units available. Visualizations are often excellent demonstrations of these results. The synth package provides functionality for use in Stata and R, although since there are a large number of possible parameters and implementations of the design it can be complex to operate.⁷²

⁷⁰ Gobillon and Magnac (2016)

⁷¹ See Fernandes et al. (2016) for an example.

⁷² Abadie et al. (2014)

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