# README for "Empirical Bayes When Estimation Precision Predicts Parameters"

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MS-22935 for Econometrica

This replication package is written with the assistance of GitHub Copilot, OpenAI Codex, and Claude Code.

# Quickstart

To regenerate figures and tables without regenerating/re-scoring the Monte Carlo data, simply run after installation:

./generate\_assets.sh

This writes tables and figures to assets/

## **Overview**

The code in this replication package does the following:

- 1. Cleans raw data from Opportunity Atlas (Chetty et al., 2022)
- 2. Generate Monte Carlo samples used in the empirical exercises (**Time consuming**; pre-computed outputs included in simulated\_posterior\_means.zip)
- 3. Scores the Monte Carlo i.e., computes statistics underlying tables and figures in the paper
- 4. Builds figures and tables in the paper from the output of step 3.

For a partial replication, skip step 2 and use the included pre-computed output.

Three main analysis files generate all 9 figures (5 in the main text, 4 in the online appendix) and 1 table (online appendix).

A full replication of the Monte Carlo (step 2) is time-consuming but extremely parallelizable. See <u>a note below on parallelism for details</u>. Full replication is also tricky because of an upstream problem where long-running Monte Carlos appear to fail silently (see <u>NOTE</u>) - though progress is saved and restarting a script resumes the progress. Please reach out to <u>jiafeng@stanford.edu</u> for a copy of the Monte Carlo output.

Specifically, each exercise runs M iterations for V different outcome variables. Right now the code parallelizes over V but not over M. A rough estimate of time taken is in the table below (your mileage may vary on time per iteration):

exercise	time per iteration	# total iterations	V	M	max core	estimated total hours at full parallelization
Calibrated simulation	1 min	15,000	15	1,000	15	16.67
Validation (coupled bootstrap)	~2 min	15,000	15	1,000	15	33.33
Weibull (OA5.3)	0.5 min	600	6	100	6	0.83

exercise	time per iteration	# total iterations	V	M	max core	estimated total hours at full parallelization
Additive model (OA5.4)	0.5 min	600	6	100	6	0.83

In lieu of a full replication of the Monte Carlo generation, each Monte Carlo run can also be checked separately (things are time-consuming overall because there are a lot of runs). I provide code for generating specific runs of the Monte Carlo exercise and checking against the data I provided. See <u>NOTE</u>.

All code is in Python, but NPMLE estimation relies on the package rpy2, R, and Mosek.

## **Data Availability and Provenance Statements**

The data are taken from the published datasets by Chetty et al. (forthcoming) in Chetty et al. (2022) under CC-BY-4.0. They are available at <a href="https://opportunityinsights.org/data/?geographic level=0&topic=0&paper id=1652#resource-listing">https://opportunityinsights.org/data/?geographic level=0&topic=0&paper id=1652#resource-listing</a> (Accessed 2024-09-16).

## **Statement about Rights**

[x] I certify that the author(s) of the manuscript have legitimate access to and permission to use the data used in this manuscript.

[x] I certify that the author(s) of the manuscript have documented permission to redistribute/publish the data contained within this replication package. Appropriate permission are documented in the <u>LICENSE</u> (/LICENSE.txt) file.

#### **License for Data**

The data are licensed under a CC-BY-4.0 license. See LICENSE.txt for details.

## **Summary of Availability**

[x] All data **are** publicly available

#### Details on each dataset and data source

This package builds from the following raw data files, all from Chetty et al. (2022). They are downloaded at <a href="https://opportunityinsights.org/data/?geographic level=0&topic=0&paper id=1652#resource-listing">https://opportunityinsights.org/data/?geographic level=0&topic=0&paper id=1652#resource-listing</a> (Accessed 2024-09-16).

- data/raw/tract\_covariates.dta ("Neighborhood Characteristics by Census Tract"): <a href="https://opportunityinsights.org/wp-content/uploads/2018/10/tract\_outcomes\_dta.zip">https://opportunityinsights.org/wp-content/uploads/2018/10/tract\_outcomes\_dta.zip</a> (Accessed 2024-09-16)
- data/raw/tract\_outcomes\_early.csv ("All Outcomes by Census Tract, Race, Gender and Parental Income Percentile"): <a href="https://opportunityinsights.org/wp-content/uploads/2018/10/tract\_outcomes.zip">https://opportunityinsights.org/wp-content/uploads/2018/10/tract\_outcomes.zip</a> (Accessed 2024-09-16)

# Computational requirements and installation instructions

## **Option 1 (Docker - recommended)**

- 1. Install Docker (see the <u>official guides</u>). *Linux users:* if you hit a permission error, add your user to the docker group (sudo usermod -aG docker \$USER && newgrp docker).
- 2. Place your mosek.lic in the project root.
- 3. Load the pre-built image (docker load < eb-replication.tar.gz)
- 4. Start and enter the container:

```
docker compose up -d eb-replication
```

docker compose exec eb-replication bash

For detailed troubleshooting and examples, see <u>Docker instructions</u> (./docker-instructions.md).

The docker-compose.yml file also provides an eb-replication-test service that mounts only raw data and an empty scratch directory for /app/data/simulated\_posterior\_means, so one can test without touching the already-generated Monte Carlo outputs.

Cleanup: when you're done, stop and remove the containers with

docker compose down

Use docker compose down --volumes if you also want to drop any Docker-managed volumes.

To remove the docker image when one's done: docker rmi replication-eb-replication:latest. Make sure replication-eb-replication:latest matches what's shown in docker images.

## **Option 2 (Source)**

If Docker is not an option, use the host installation walkthrough in <u>source-install.md</u> (./source-install.md). It covers:

- Python 3.10 environment setup with pip requirements
- R 4.4.0 + renv restore and the Rmosek builder steps
- MOSEK 10.2.5 licensing/configuration and GNU Parallel
- Verification commands (python -m rpy2.situation, python test.py, targeted Monte Carlo checks)

Use this path only if you are comfortable managing those dependencies manually.

## **Controlled Randomness**

Random seed is set at:

- Line 173 of covariate\_additive\_model.py
- Line 226 of empirical\_exercise.py

Note: REBayes::GLmix uses REBayes::KWDual to interact with an underlying MOSEK optimizer. This optimizer may introduce additional randomness that cannot be seeded from the Python side. As far as I'm aware, REBayes does not expose a seeding option for this solver.

## Memory, Runtime, Storage Requirements

- <10 minutes (reproducing from scored Monte Carlo outputs—i.e., starting from step 4)
- 10–60 minutes (reproducing from Monte Carlo outputs—i.e., starting from step 3)
- 1–3 days (full reproduction starting from step 1 or 2)

The code was last run on a 2022 Mac Studio, 32 GB RAM, Apple M1 Max.

# Description of programs/code and instruction to replicators

## Overview

The replication process is modularized as follows. Each step saves output to the disk that the next step depends on. These outputs are included in the replication package transmitted to the data editor, and so **partial replications can start from any step**. For instance, directly going to step 4 generates all tables and figures.

(Step 1 - Raw data cleaning) build\_data.py performs basic cleaning on the raw Opportunity Atlas data and saves the processed data in data/processed/oa\_data\_used.feather. The rest of the analysis proceeds only with data/processed/oa\_data\_used.feather.

(Step 2 - Monte Carlo data generation) Most empirical exercises in the paper aggregate over simulation draws that depend on the cleaned data. These simulation draws---which we refer to as the *Monte Carlo data*---are directly saved in data/simulated\_posterior\_means/. The subsequent analysis only depends on the generated Monte Carlo data. This step is time consuming and error-prone: For a partial replication, one could directly proceed from the next step or step 4, without re-generating the Monte Carlo data (pre-computed Monte Carlo data from a previous run should be included in this replication package - contact <u>jiafeng@stanford.edu</u> if not).

One could also selectively verify subsets of the output generated by this step, as opposed to regenerating every simulation run: For instance,

See <u>NOTE</u> below for detailed instructions.

The Monte Carlo data are generated by the following bash scripts:

- ./monte\_carlo.sh runs the calibrated simulation exercise
- ./coupled\_bootstrap.sh runs the coupled bootstrap exercise
- ./weibull model.sh runs the Weibull distribution exercise in online appendix
- ./additive\_model.sh runs an additional model exercise in online appendix
- (Step 3 Computations on the Monte Carlo data whose output underlies the figures) Various statistics underlying tables and figures are computed on the Monte Carlo data (We call this *scoring* the Monte Carlo data).
- ./generate\_scores.py takes Monte Carlo results in data/simulated\_posterior\_means/ and scores them. It saves results in results/[SimulatorName]/\*.csv, where SimulatorName is one of {coupled\_bootstrap-0.9, covariate\_additive\_model, npmle\_by\_bins, weibull}
- (**Step 4 Tables and figures**) Finally, the tables and figures are generated directly from the output of the last step. How each table/figure links to each python script is detailed below. generate\_assets.sh simply runs all of them.

**Note on figure output:** Figures with many scatter points (Figures 1-3) are saved in both PDF and PNG formats. PDF files use rasterized scatter points for smaller file sizes but may display inconsistently across PDF viewers (e.g., missing points in Chrome, artifacts in Preview). PNG files provide reliable viewing across all platforms.

Content	Script
Figures 1–3	./assets_introduction.py
Footnote 6 voice over	./assets_introduction.py
Figures 4–5	./assets_empirical.py
Table OA5.1	./assets_appendix.py
Figures OA5.1–OA5.4	./assets_appendix.py

## Lower-level files and additional dependencies

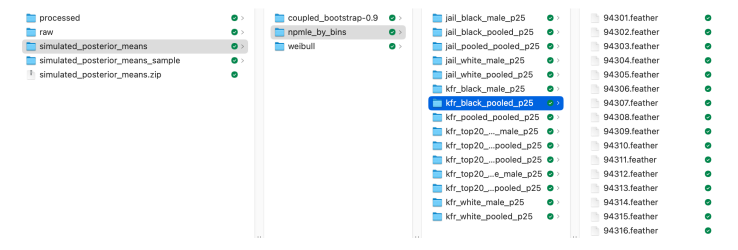
These steps depend on the following lower-level files:

- 1. empirical\_exercise.py samples either calibrated Monte Carlo simulation or coupled bootstrap simulation. It then computes various posterior mean estimates using various empirical Bayes or non-empirical Bayes methods. The .sh files in Step 2 are essentially wrappers that call empirical\_exercise.py.
- 2. covariate\_additive\_model.py runs CLOSE-NPMLE with a flexible additive model for covariates. This is only relevant for ./additive\_model.sh
- 3. Helpers:
- residualize.py implements linear residualization by covariates
- conditional\_means/ contains methods for estimating conditional means
- empirical bayes/ contains methods for implementing empirical Bayes methods
- postprocessing/ contains methods for computing various metrics and visualization
- simulator/ contains code for implementing various methods for simulation synthetic data from raw data

## **Instructions to replicators**

Below are code snippets for reproducing each step described in the list above. I recommend starting with Steps 3 or 4 to check if everything replicates given the Monte Carlo output. Run all code from the top level of the directory.

The Monte Carlo output should be included as simulated\_posterior\_means.zip (please contact <a href="mailto:jiafeng@stanford.edu">jiafeng@stanford.edu</a> if not). This file can be unzipped into data/, where data/simulated\_posterior\_means/ should include three directories coupled\_bootstrap-0.9, npmle\_by\_bins, and weibull. Each of these three directories should then contain subdirectories with variable names like kfr\_black\_male\_p25. There should be 1000 or 100 .feather files in these directories, with number names:



A full replication of Step 2 is time-consuming, but selective subsets of the output can be checked easily.

#### **Step 0: Change permissions**

Run chmod +x \*.sh to allow the .sh files to be run as executables. Alternatively one may run bash script.sh to execute them.

#### **Step 1: Raw data cleaning**

```
# Builds the raw analysis dataset from raw data
python build_data.py
```

Note: if run inside the eb-replication service, then this overwrites the current file in data/processed. If one works with the container eb-replication-test instead of eb-replication, then app/data/processed in the container is mounted to ./data/processed\_empty.

## **Step 2: Monte Carlo data generation**

The following bash commands runs each bash script and generates the Monte Carlo data.

NB: This is the most time-consuming and error-prone step (see <u>NOTE</u> below): The output of this step is included in the replication package directly. Moreover, optionally, instead of fully replicating this step, one could verify a small subset of the Monte Carlo data. The <u>NOTE</u> below includes instructions for doing so.

If one decides to run a full replication, before starting to replicate, **ensure that data/simulated\_posterior\_means** is **empty**.

To enforce data/simulated\_posterior\_means is empty, start the test service with docker compose up -d eb-replication-test and work within the container eb-replication-test instead of eb-replication; eb-replication-test mounts an empty scratch directory at /app/data/simulated\_posterior\_means (in container) linked to data/simulated\_posterior\_means\_empty/ (local).

```
# Run the Monte Carlo
# The following generates results/[simulator-name]
# where [simulator-name] is one of "coupled_bootstrap-0.9",
# "covariate_additive_model", "npmle_by_bins", "weibull"
rm −f logs/*
              # Clear logs
# Time consuming, the progress bar is written to std.err, which is written to logs/.
# Can check corresponding files in logs/ to monitor
# (monte_carlo.sh coupled_bootstrap.sh weibull_model.sh) do not need to be run sequentially.
# They can be run concurrently
# With &, scripts run in the background of the terminal session and print a PID for the
        wrapper process.
# That PID does not own the python workers; use `kill -- -[pid]` (note leading minus) or
        `pkill -P [pid]` to stop everything.
# Calibrated simulation exercise
# Time estimate: (1 minute per iteration x 15000 iterations) / min(#cores, 15)
# The NUM_CORES options below are by default the maximum the code would benefit from.
# Higher core counts won't break anything - the code takes a minimum.
```

NUM\_CORES=15 ./monte\_carlo.sh & # To monitor: tail -F logs/mc\_error\*

## Step 3: Computing table/figure-relevant statistics

```
### Assumes that data/simulated_posterior_means/ is populated,
### either because Step 2 is run or because the included Monte Carlo
### output simulated_posterior_means.zip is unzipped.

# Clean up the generated raw Monte Carlo results

python generate_scores.py --simulator-name coupled_bootstrap-0.9 --nsim 1000 # ~20 minutes

python generate_scores.py --simulator-name npmle_by_bins --nsim 1000 # ~1.5 minute

python generate_scores.py --simulator-name weibull --nsim 100 # ~5 seconds

# The additive model results are scored directly in additive_model.sh
```

#### **Step 4: Creating tables and figures**

```
# Generate figures and tables in assets/
# Assumes results/ is correctly populated with scored outputs
./generate_assets.sh
```

## **Parallelism**

The .sh files in step 2 runs the following script in parallel **over est\_var** 

Across the various empirical exercises, est\_var ranges over 6 to 15 choices. As a result, we would only benefit from at most 15 cores.

Alternatively, one could further parallelize by running, e.g., the following in parallel. This would parallelize within a single est\_var.

```
# Runs seeds 94301-94800
python empirical_exercise.py --simulator-name [simulator-name] --methods [methods-for-simulator] --nsim 500 --starting_seed 94301 --est_var [est_var]

# Runs seeds 94801-95300
python empirical_exercise.py --simulator-name [simulator-name] --methods [methods-for-simulator] --nsim 500 --starting_seed 94801 --est_var [est_var]
```

Doing so is a little memory inefficient because runs using the same est\_var share underlying data.

Simulator name	Outcome variable names	Seed range	Methods
coupled_bootstrap- 0.9	One of ("kfr_pooled_pooled_p25"  "kfr_white_male_p25"  "kfr_black_male_p25"  "kfr_black_pooled_p25"  "kfr_white_pooled_p25"  "jail_black_male_p25"  "jail_white_male_p25"  "jail_white_pooled_p25"  "jail_whooled_pooled_p25"  "kfr_top20_black_male_p25"  "kfr_top20_white_male_p25"  "kfr_top20_white_pooled_p25"  "kfr_top20_white_pooled_p25"  "kfr_top20_white_pooled_p25"  "kfr_top20_white_pooled_p25"  "kfr_top20_white_pooled_p25"	94301- 95300	all
npmle_by_bins	One of ("kfr_pooled_pooled_p25" "kfr_white_male_p25" "kfr_black_male_p25" "kfr_black_pooled_p25" "kfr_white_pooled_p25" "jail_black_male_p25" "jail_white_male_p25" "jail_white_pooled_p25" "jail_white_pooled_p25" "jail_pooled_pooled_p25" "kfr_top20_black_male_p25" "kfr_top20_white_male_p25" "kfr_top20_white_pooled_p25" "kfr_top20_white_pooled_p25" "kfr_top20_white_pooled_p25" "kfr_top20_white_pooled_p25" "kfr_top20_pooled_pooled_p25")	94301- 95300	all
weibull	One of ("kfr_pooled_pooled_p25" "kfr_black_pooled_p25" "jail_black_pooled_p25" "jail_pooled_pooled_p25" "kfr_top20_black_pooled_p25" "kfr_top20_pooled_pooled_p25")	94301 - 94400	indep_gauss,close_npmle,close_gau

## **NOTE:** on replicating Monte Carlo data

For some upstream reason having to do with MOSEK or REBayes, running monte\_carlo.sh for many iterations might silently fail, due to a memory leak. When it fails, the code would appear to run but resource consumption is low and no new output is generated. Interrupting the code prints Segmentation fault. I find it quite difficult to reproduce the issue, as there's no fixed data seed causing a problem. When this happens, interrupting and restarting resolves the issue. This has only happened when I repeatedly apply NPMLE to sample new data and to estimate various methods.

Each Monte Carlo draw results in a file of the form

data/simulated\_posterior\_means/[SimulatorName]/[VariableName]/[Seed].feather.

It is not time-consuming to regenerate and verify a given file - the *number* of these files makes it time-consuming overall. I have included a script that generates a new Monte Carlo draw for a particular seed seed number and a particular outcome variable est var

```
# See the table below for valid options for --est var and --simulator name
# --est var: Outcome variable being P(top 20 | Black, pooled, parents at 25th percentile)
# --seed_number: The seed used is 94301 + (seed_number mod seed_range). In this case we
        check seed 94682
# --simulator name: Simulator name = npmle by bins, coupled bootstrap-0.9, weibull
python check_monte_carlo.py --est_var kfr_top20_black_pooled_p25 --seed_number 381 --
        simulator_name npmle_by_bins
```

check monte carlo py works by generating a specific draw of the Monte Carlo data, saves it in data/simulated\_posterior\_means\_sample, and compares it against its counterpart in data/simulated posterior means. I have found that different hardware/version would only agree up to something like 1e-6, and so I check agreement between two files by regressing one on the other. A typical output is as follows across two machines - the regression fit is essentially perfect.

```
- Project '~/Library/CloudStorage/Dropbox/research/empirical-bayes/replication' loaded.
[renv 1.1.4]
- The project is out-of-sync -- use `renv::status()` for details.
Checking Monte Carlo outputs...
```

kfr top20 black pooled p25: 100%|

1/1 [00:00<00:00, 14315.03it/s]

Seed: 94682

Outcome variable: kfr\_top20\_black\_pooled\_p25

Simulator name: npmle\_by\_bins

Correlation between original and new Monte Carlo samples (some differences may exist due to hardware or version):

	Correlation	Intercept	Regression Coef
naive	1.000000	-1.370503e-07	1.000000
indep_npmle	0.999999	5.062243e-07	0.999980
indep_gauss	1.000000	1.034661e-06	0.999973
close_npmle	1.000000	-8.168789e-08	1.000002
close_gauss	1.000000	1.060407e-07	0.999997
close_gauss_parametric	1.000000	3.004981e-07	0.999993
oracle	1.000000	-2.953876e-07	1.000003
truth	1.000000	-2.181302e-07	1.000002
<pre>indep_npmle_nocov</pre>	1.000000	-4.812911e-07	1.000013
indep_gauss_nocov	0.999999	-5.065023e-07	1.000019
close_npmle_nocov	1.000000	-1.943102e-07	1.000002

close_gauss_nocov	1.000000 -1.923112e-07	1.000001
close_gauss_parametric_nocov	1.000000 -1.189520e-07	1.000000
true_covariate_fn	1.000000 -2.741553e-14	1.000000
truth_residualized	1.000000 -2.106331e-07	1.000003

\_\_\_\_\_

# References

Chetty, Raj, John Friedman, Nathaniel Hendren, Maggie R. Jones, and Sonya R. Porter, "Replication Data for: The Opportunity Atlas: Mapping the Childhood Roots of Social Mobility," 2022.

Chetty, Raj, John Friedman, Nathaniel Hendren, Maggie R. Jones, and Sonya R. Porter, "The opportunity atlas: Mapping the childhood roots of social mobility," American Economic Review, forthcoming.

O. Tange (2018): GNU Parallel 2018, March 2018, https://doi.org/10.5281/zenodo.1146014.