## Matching for Causal Inference Without Balance Checking

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(talk at the University of Rochester, 1/19/09)

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# Preview Slide: Coarsened Exact Matching (CEM)

A simple (and ancient) method of causal inference, with surprisingly powerful properties

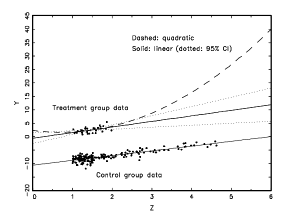
- **1** Preprocess (X, T) with CEM:
  - (A) Temporarily coarsen X as much as you're willing
    - e.g., Education (grade school, high school, college, graduate)
    - Easy to understand, or can be automated as for a histogram
  - (B) Perform exact matching on the coarsened X, C(X)
    - Sort observations into strata, each with unique values of C(X)
    - Prune any stratum with 0 treated or 0 control units
  - (C) Pass on original (uncoarsened) units except those pruned
- Analyze as without matching (adding weights for stratum-size) (Or apply other matching methods within CEM strata & they inherert CEM's properties)
- → A version of CEM: Last studied 40 years ago by Cochran
- → First used many decades before that
- → We prove: many new properties, uses, & extensions, and show how it resolves many problems in the literature

#### Characteristics of Observational Data

- Lots of data
- Data is of uncertain origin. Treatment assignment: not random, not controlled by investigator, not known
- Bias-Variance Tradeoff Bias-Variance Tradeoff
- The idea of matching: sacrifice some data to avoid bias
- Removing heterogeneous data will often reduce variance too
- (Medical experiments are the reverse: small-n with random treatment assignment; don't match unless something goes wrong)

#### Model Dependence

(King and Zeng, 2006: fig.4 Political Analysis)

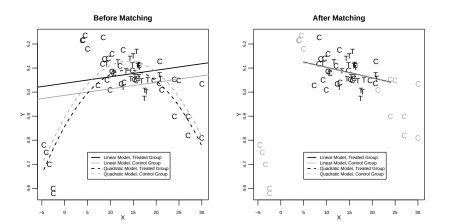


#### What to do?

- Preprocess I: Eliminate extrapolation region (a separate step)
- Preprocess II: Match (prune bad matches) within interpolation region
- Model remaining imbalance

#### Matching within the Interpolation Region

(Ho, Imai, King, Stuart, 2007: fig.1, Political Analysis)



Matching reduces model dependence, bias, and variance

## The Goals, with some more precision

Notation:

 $Y_i$  Dependent variable

 $T_i$  Treatment variable (0/1)

 $X_i$  pre-treatment covariates

• Treatment Effect for treated  $(T_i = 1)$  observation i:

$$TE_i = Y_i(T_i = 1) - Y_i(T_i = 0)$$
  
= observed -unobserved

- Estimate  $Y_i(0)$  with  $Y_j$  from matched  $(X_i \approx X_j)$  controls  $\hat{Y}_i(0) = Y_j(0)$  or a model  $\hat{Y}_i(0) = \hat{g}_0(X_j)$
- Prune unmatched units to improve balance (so X is unimportant)
- Sample Average Treatment effect on the Treated:

$$\mathsf{SATT} = \frac{1}{n_T} \sum_{i \in \{T_i = 1\}} \mathsf{TE}_i$$

## Problems With Existing Matching Methods

- Don't eliminate extrapolation region
- Don't work with multiply imputed data
- Most violate the congruence principle
- Largest class of matching methods (EPBR, e.g., propensity scores, Mahalanobis distance): requires normal data (or DMPES); all X's must have same effect on Y; Y must be a linear function of X; aims only for expected (not in-sample) imbalance; → in practice, we're lucky if mean imbalance is reduced
- Not well designed for observational data:
  - Least important (variance): matched *n* chosen ex ante
  - Most important (bias): imbalance reduction checked ex post
- Hard to use: Improving balance on 1 variable can reduce it on others
  - Best practice: choose *n*-match-check, tweak-match-check, tweak-match-check, tweak-match-check, · · ·
  - Actual practice: choose n, match, publish, STOP.
     (Is balance even improved?)

### A New Class of Methods: Monotonic Imbalance Bounding

- No restrictions on data types
- Designed for observational data (reversing EPBR):
  - Most important (bias): degree of balance chosen ex ante
  - Least important (variance): matched *n* checked ex post
- Balance is measured in sample (like blocked designs), not merely in expectation (like complete randomization)
- Covers all forms of imbalance: means, interactions, nonlinearities, moments, multivariate histograms, etc.
- One adjustable tuning parameter per variable
- Convenient monotonicity property: Reducing maximum imbalance on one *X*: no effect on others

MIB Formally (simplifying for this talk):

$$\begin{array}{ll} D(\mathbf{X}_T^\epsilon, \mathbf{X}_C^\epsilon) \leq \boldsymbol{\gamma}(\epsilon) & \text{vars to adjust} \\ D(X_T^\epsilon, X_C^\epsilon) \leq \boldsymbol{\gamma}(\epsilon) & \text{remaining vars} \end{array}$$

Treated and control X variables to adjust Remaining treated and control X variables "Imbalance" given chosen distance metric Bounds

## What's Coarsening?

- Coarsening is intrinsic to measurement
  - We think of measurement as clarity between categories
  - But measurement also involves homogeneity within categories
  - Examples: male/female, rich/middle/poor, black/white, war/nonwar.
  - Better measurement devices (e.g., telescopes) produce more detail
- Data analysts routinely coarsen, thinking grouping error is less risky than measurement error. E.g.:
  - 7 point Party ID → Democrat/Independent/Republican
  - Likert Issue questions → agree/{neutral,no opinion}/disagree
  - multiparty voting → winner/losers
  - Religion, Occupation, SEC industries, ICD codes, etc.
- Temporary Coarsening for CEM; e.g.:
  - Education: grade school, middle school, high school, college, graduate
  - Income: poverty level threshold, or larger bins for higher income
  - Age: infant, child, adolescent, young adult, middle age, elderly

#### CEM as an MIB Method

- ullet Define:  $\epsilon$  as largest (coarsened) bin size ( $\epsilon=0$  is exact matching)
- We Prove: setting  $\epsilon$  bounds the treated-control group difference, within strata and globally, for: means, variances, skewness, covariances, comoments, coskewness, co-kurtosis, quantiles, and full multivariate histogram.
  - $\implies$  Setting  $\epsilon$  controls all multivariate treatment-control differences, interactions, and nonlinearities, up to the chosen level (matched n is determined ex post)
- By default, both treated and control units are pruned: CEM estimates a quantity that can be estimated without model dependence
- What if  $\epsilon$  is set . . .
  - too large? → You're left modeling remaining imbalances
  - too small?  $\rightsquigarrow$  *n* may be too small
  - as large as you're comfortable with, but n is still too small?
    - → No magic method of matching can save you;
    - → You're stuck modeling or collecting better data

## Other CEM properties we prove

- Automatically eliminates extrapolation region (no separate step)
- Bounds model dependence
- Bounds causal effect estimation error
- Meets the congruence principle
  - The principle: data space = analysis space
  - Estimators that violate it are nonrobust and counterintuitive
  - CEM:  $\epsilon_i$  is set using each variable's units
  - E.g., calipers (strata centered on each unit): would bin college drop out with 1st year grad student; and not bin Bill Gates & Warren Buffett
- Approximate invariance to measurement error:

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CEM pscore Mahalanobis Genetic % Common Units 96.5 70.2 80.9 80.0
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- Fast and memory-efficient even for large n; can be fully automated
- Simple to teach: coarsen, then exact match

#### Imbalance Measures

Variable-by-Variable Difference in Global Means

$$I_1^{(j)} = \left| \bar{X}_{m_T}^{(j)} - \bar{X}_{m_C}^{(j)} \right|, \quad j = 1, \dots, k$$

Multivariate Imbalance: difference in histograms (bins fixed ex ante)

$$\mathcal{L}_1(f,g) = \sum_{\ell_1 \cdots \ell_k} |f_{\ell_1 \cdots \ell_k} - g_{\ell_1 \cdots \ell_k}|$$

Local Imbalance by Variable (given strata fixed by matching method)

$$I_2^{(j)} = \frac{1}{S} \sum_{s=1}^{S} \left| \bar{X}_{m_T^s}^{(j)} - \bar{X}_{m_C^s}^{(j)} \right|, \quad j = 1, \dots, k$$

### CEM in Practice: EPBR-Compliant Data

Monte Carlo:  $\mathbf{X}_T \sim N_5(\mathbf{0}, \Sigma)$  and  $\mathbf{X}_C \sim N_5(\mathbf{1}, \Sigma)$ . n = 2,000, reps=5,000 Allow MAH & PSC to match with replacement; use automated CEM Difference in means ( $I_1$ ):

	$X_1$	$X_2$	$X_3$	$X_4$	$X_5$	Seconds
initial	1.00	1.00	1.00	1.00	1.00	
MAH	.20	.20	.20	.20	.20	.28
PSC	.11	.06	.03	.06	.03	.16
CEM	.04	.02	.06	.06	.04	.08

Local  $(I_2)$  and multivariate  $\mathcal{L}_1$  imbalance:

	$X_1$	$X_2$	$X_3$	$X_4$	$X_5$	$\mathcal{L}_1$
initial						1.24
PSC	2.38	1.25	.74	1.25	.74	1.18
MAH	.56	.36	.29	.36	.29	1.13
CEM	.42	.26	.17	.22	.19	.78

→ CEM dominates EPBR-methods in EPBR Data

#### CEM in Practice: Non-EPBR Data

Monte Carlo: Exact replication of Diamond and Sekhon (2005), using data from Dehejia and Wahba (1999). CEM coarsening automated.

	BIAS	SD	RMSE	Seconds	$\mathcal{L}_1$
initial	-423.7	1566.5	1622.6	.00	1.28
MAH	784.8	737.9	1077.2	.03	1.08
PSC	260.5	1025.8	1058.4	.02	1.23
GEN	78.3	499.5	505.6	27.38	1.12
CEM	.8	111.4	111.4	.03	.76

#### CEM Extensions I

- CEM and Multiple Imputation for Missing Data
  - put missing observation in stratum where plurality of imputations fall
  - 2 pass on uncoarsened imputations to analysis stage
  - Use the usual MI combining rules to analyze
- ullet Multicategory treatments: No modification necessary; keep all strata with  $\geq 1$  unit having each value of T
- Blocking in Randomized Experiments: no modification needed; randomly assign T within CEM strata
- Automating user choices Histogram bin size calculations, Estimated SATT error bound, Progressive Coarsening
- Detecting Extreme Counterfactuals

## CEM Extensions II: Improving Existing Matching Methods

- Most commonly used methods:
  - cannot be used to eliminate extrapolation region
  - don't possess most other CEM properties
  - but inherent CEM properties if applied within CEM strata
- Propensity Score matching:
  - requires correct specification or balance can drop (the usual specification tests are irrelevant; must check balance)
  - CEM strata can bound bias in pscore matching
  - may be good for applications with many covariates we know little about (so we're willing to take balance on any subset)
- Mahalanobis distance: can apply within CEM strata
- Genetic Matching: can constrain results to CEM strata
- Synthetic Matching, or Robins' weights: CEM can identify region to apply weights, increasing efficiency/robustness
- Nonparametric Adjustments: can apply within CEM strata

For papers, software (for R and Stata), tutorials, etc.

http://GKing.Harvard.edu/cem