

Experiments & Observational Studies:

Causal Inference in Statistics

Paul R. Rosenbaum

Department of Statistics

University of Pennsylvania

Philadelphia, PA 19104-6340

1 My Concept of a 'Tutorial'

- In the computer era, we often receive compressed files, .zip. Minimize redundancy, minimize storage, at the expense of intelligibility.
- Sometimes scientific journals seemed to have been compressed.
- Tutorial goal is: 'uncompress'. Make it possible to read a current article or use current software without going back to dozens of earlier articles.

2 A Causal Question

- At age 45, Ms. Smith is diagnosed with stage II breast cancer.
- Her oncologist discusses with her two possible treatments: (i) lumpectomy alone, or (ii) lumpectomy plus irradiation. They decide on (ii).
- Ten years later, Ms. Smith is alive and the tumor has not recurred.
- Her surgeon, Steve, and her radiologist, Rachael debate.
- Rachael says: “The irradiation prevented the recurrence — without it, the tumor would have recurred.”
- Steve says: “You can’t know that. It’s a fantasy — you’re making it up. We’ll never know.”

3 Many Causal Questions

- Steve and Rachael have this debate all the time. About Ms. Jones, who had lumpectomy alone. About Ms. Davis, whose tumor recurred after a year.
- Whenever a patient treated with irradiation remains disease free, Rachael says: “It was the irradiation.” Steve says: “You can’t know that. It’s a fantasy. We’ll never know.”
- Rachael says: “Let’s keep score, add ’em up.” Steve says: “You don’t know what would have happened to Ms. Smith, or Ms. Jones, or Ms Davis — you just made it all up, it’s all fantasy. Common sense says: ‘A sum of fantasies is total fantasy.’ Common sense says: ‘You can’t add fantasies and get facts.’ Common sense says: ‘You can’t prove causality with statistics.’”

4 Fred Mosteller's Comment

- Mosteller like to say: “You can *only* prove causality with statistics.”
- He was thinking about a particular statistical method and a particular statistician.
- Not Gauss and least squares, or Yule and Yule's Q (a function of the odds ratio), or Wright and path analysis, or Student and the t-test.
- Rather, Sir Ronald Fisher and randomized experiments.

5 Fisher & Randomized Experiments

- Fisher's biographer (and daughter) Joan Fisher Box (1978, p. 147) suggests that Fisher invented randomized experiments around 1920, noting that his paper about ANOVA in experiments of 1918 made no reference to randomization, referring to Normal theory instead, but by 1923, his papers used randomization, not Normal theory, as the justification for ANOVA.
- Fisher's clearest and most forceful discussion of randomization as 'the reasoned basis for inference' in experiments came in his book of 1935, *Design of Experiments*.

6 15 Pages

- In particular, the 15 pages of Chapter 2 discuss what came to be known as Fisher's exact test for a 2×2 table. The hypergeometric distribution is dispatched in half a paragraph, and Fisher hammers away in English for $14\frac{1}{2}$ pages about something else.
- Of Fisher's method of randomization and randomization, Yule would write: "I simply cannot make head or tail of what the man is doing." (Box 1978, p. 150). But Neyman (1942, p. 311) would describe it as "a very brilliant method."

7 Lumpectomy and Irradiation

- Actually, Rachael was right, Steve was wrong. Perhaps not in every case, but in many cases. The addition of irradiation to lumpectomy causes there to be fewer recurrences of breast cancer.
- On 17 October 2002, the *New England Journal of Medicine* published a paper by Bernard Fisher, et al. describing 20 year follow-up of a randomized trial comparing lumpectomy alone and lumpectomy plus irradiation.
- There were 634 women randomly assigned to lumpectomy, 628 to lumpectomy plus irradiation.
- Over 20 years of follow-up, 39% of those who had lumpectomy alone had a recurrence of cancer, as opposed to 14% of those who had lumpectomy plus irradiation ($P < 0.001$).

8 Outline: Causal Inference

... in randomized experiments.

- Causal effects. ■ Randomization tests of no effect.
- Inference about magnitudes of effect.

... in observational studies.

- What happens when randomized experiments are not possible? ■ Adjustments for overt biases: How to do it. When does it work or fail. ■ Sensitivity to hidden bias. ■ Reducing sensitivity to hidden bias.

9 Finite Population

- In Fisher's formulation, randomization inference concerns a finite population of n subjects, the n subjects actually included in the experiment, $i = 1, \dots, n$.
- Say $n = 1,262$, in the randomized experiment comparing lumpectomy (634) vs lumpectomy plus irradiation (628).
- The inference is *not* to some other population. The inference is to how these n people would have responded under treatments they did not receive.
- We are not sampling people. We are sampling possible futures for n fixed people.
- Donald Campbell would emphasize the distinction between *internal* and *external* validity.

10 Causal Effects: Potential Outcomes

- Key references: Neyman (1923), Rubin (1974).
- Each person i has two potential responses, a response that would be observed under the 'treatment' condition T and a response that would be observed under the 'control' condition C .

$$r_{Ti} = \begin{cases} 1 & \text{if woman } i \text{ would have cancer} \\ & \text{recurrence with lumpectomy alone} \\ 0 & \text{if woman } i \text{ would not have cancer} \\ & \text{recurrence with lumpectomy alone} \end{cases}$$

$$r_{Ci} = \begin{cases} 1 & \text{if woman } i \text{ would have cancer} \\ & \text{recurrence with lumpectomy+irradiation} \\ 0 & \text{if woman } i \text{ would not have cancer} \\ & \text{recurrence with lumpectomy+irradiation} \end{cases}$$

- We see r_{Ti} or r_{Ci} , but never both. For Ms. Smith, we saw r_{Ci} .

11 Comparing Potential Outcomes

- r_{Ti} is the response observed from i under lumpectomy alone, and r_{Ci} is observed from i under lumpectomy plus irradiation.
- The effect of the treatment is a comparisons of r_{Ti} and r_{Ci} , such as $\delta_i = r_{Ti} - r_{Ci}$. Possibilities:

r_{Ti}	r_{Ci}	δ_i	
1	1	0	cancer recurrence either way
1	0	1	irradiation prevents recurrence
0	1	-1	irradiation causes recurrence
0	0	0	no recurrence either way

- If someone gave us (r_{Ti}, r_{Ci}) , $i = 1, \dots, n$, causal inference would be arithmetic, not inference. But we never see δ_i for any i . We don't know δ_i for $i = Ms. Smith$.

12 Recap

- A finite population of $n = 1,262$ women.
- Each woman has two potential responses, (r_{Ti}, r_{Ci}) , but we see only one of them. Never see $\delta_i = r_{Ti} - r_{Ci}$, $i = 1, \dots, n$.
- Is it plausible that irradiation does nothing? *Null hypothesis of no effect.* $H_0 : \delta_i = 0, i = 1, \dots, n$.
- Estimate the *average treatment effect*: $\frac{1}{n} \sum_{i=1}^n \delta_i$.
- How many more women did not have a recurrence of cancer because they received irradiation? (*Attributable effect*)
- The (r_{Ti}, r_{Ci}) are $2n$ fixed numbers describing the finite population. Nothing is random.

13 Fisher's Idea: Randomization

- Randomization converts impossible arithmetic into feasible statistical inference.
- Pick m of the n people at random and give them treatment condition T . In the experiment, $m = 634$, $n = 1,262$. That is, assign treatments

“in a random order, that is in an order not determined arbitrarily by human choice, but by the actual manipulation of the physical apparatus used in games of chance, cards, dice, roulettes, etc., or, more expeditiously, from a published collections of random sampling numbers. . .” (Fisher, 1935, Chapter 2)

- This means that each of the $\binom{n}{m} = \binom{1,262}{634}$ treatment assignments has the same probability, $\binom{1,262}{634}^{-1}$. The only probabilities that enter Fisher's randomization inference are created by randomization.

14 Observable Quantities

- Write $Z_i = 1$ if i is assigned to T and $Z_i = 0$ if i is assigned to C . Then $m = \sum_{i=1}^n Z_i$.

- Write R_i for the observed response from i . Then:

$$R_i = \begin{cases} r_{Ti} & \text{if } Z_i = 1 \text{ (randomly assigned to} \\ & \text{lumpectomy)} \\ r_{Ci} & \text{if } Z_i = 0 \text{ (randomly assigned to} \\ & \text{lumpectomy+irradiation)} \end{cases}$$

or formally

$$R_i = Z_i r_{Ti} + (1 - Z_i) r_{Ci} = r_{Ci} + Z_i \delta_i.$$

- Unlike the causal effect, δ_i , which are fixed but unobservable features of the finite population, the Z_i and R_i are observable random variables.

15 The Observable 2×2 Table

	<i>Recurrence</i> $R_i = 1$	<i>No recurrence</i> $R_i = 0$	Total
No rads $Z_i = 1$	$\sum Z_i R_i$	$\sum Z_i (1 - R_i)$	m
Rads $Z_i = 0$	$\sum (1 - Z_i) R_i$	$\sum (1 - Z_i) (1 - R_i)$	$n-m$

	<i>Recurrence</i> $R_i = 1$	<i>No recurrence</i> $R_i = 0$	Total
No rads $Z_i = 1$	220	414	634
Rads $Z_i = 0$	78	550	628
Total	298	964	1,262

16 Observable Table in Terms of Potential Responses

	<i>Recurrence</i> $R_i = 1$	<i>No recurrence</i> $R_i = 0$	Total
No rads $Z_i = 1$	$\sum Z_i R_i$	$\sum Z_i (1 - R_i)$	m
Rads $Z_i = 0$	$\sum (1 - Z_i) R_i$	$\sum (1 - Z_i) (1 - R_i)$	$n-m$

is the same as the following table because $R_i = r_{Ti}$ when $Z_i = 1$ and $R_i = r_{Ci}$ when $Z_i = 0$:

	<i>Recurrence</i> $R_i = 1$	<i>No recurrence</i> $R_i = 0$	Total
No rads $Z_i = 1$	$\sum Z_i r_{Ti}$	$\sum Z_i (1 - r_{Ti})$	m
Rads $Z_i = 0$	$\sum (1 - Z_i) r_{Ci}$	$\sum (1 - Z_i) (1 - r_{Ci})$	$n-m$

17 Testing No Effect

- If the treatment has no effect, $H_0 : \delta_i = 0$ for $i = 1, \dots, n$, then

$$0 = \delta_i = r_{Ti} - r_{Ci}$$

or $r_{Ti} = r_{Ci}, i = 1, \dots, n.$

- The observed response is then

$$R_i = r_{Ci} + Z_i \delta_i = r_{Ci}$$

is just r_{Ci} , which is fixed, not varying with the treatment assignment Z_i .

- If the null hypothesis were true, then irradiation doesn't affect whether cancer recurs — we observe $R_i = r_{Ci}$ with or without irradiation.

If the null hypothesis were true, the responses in the lumpectomy-alone group are just a simple random sample (without replacement) of size m from a finite populations of size n consisting of the n binary r_{Ci} 's.

18 2×2 Table Under No effect: Fisher's Exact Test

- If the treatment has no effect, $H_0 : \delta_i = 0$ for $i = 1, \dots, n$, then $R_i = r_{Ci} + Z_i \delta_i = r_{Ci}$, and the observable table of Z_i by R_i equals the table of Z_i by r_{Ci} :

	<i>Recurrence</i> $r_{Ci} = 1$	<i>No recurrence</i> $r_{Ci} = 0$
No rads $Z_i = 1$	$\sum Z_i r_{Ci}$	$\sum Z_i (1 - r_{Ci})$
Rads $Z_i = 0$	$\sum (1 - Z_i) r_{Ci}$	$\sum (1 - Z_i) (1 - r_{Ci})$

which has the hypergeometric distribution from the randomization.

- That is, under the null hypothesis, $\sum_{i=1}^n Z_i r_{Ci}$ is the total in a simple random sample without replacement of size m from a population of size n containing $\sum_{i=1}^n r_{Ci}$ 1's and $\sum_{i=1}^n (1 - r_{Ci})$ 0's.

19 Fisher's Exact Test

	<i>Recurrence</i> $R_i = 1$	<i>No recurrence</i> $R_i = 0$	Total
No rads $Z_i = 1$	220	414	634
Rads $Z_i = 0$	78	550	628
Total	298	964	1,262

- If the null hypothesis were true, so the corner cell had the hypergeometric distribution, then $\Pr(T \geq 220) = 2.7 \times 10^{-21}$.
- That is, if irradiation changed nothing, then the experiment randomly split 1,262 people into 634 and 628.
- A random split would produce the 220/78 split (or larger) of recurrences by chance with probability 2.7×10^{-21} .

20 How far have we come?

- We never see any causal effects, δ_i .
- Yet we are $100 \left(1 - 2.7 \times 10^{-21}\right)$ % confident that some $\delta_i > 0$.
- Causal inference is impossible at the level of an individual, i , but it is straightforward for a population of n individuals if treatments are randomly assigned.
- Mosteller's comment: "You can only prove causality with statistics."

21 Testing other hypotheses

- Recall that $\delta_i = r_{Ti} - r_{Ci}$, and Fisher's exact test rejected $H_0 : \delta_i = 0, i = 1, \dots, n = 1262$.
- Consider testing instead $H_0 : \delta_i = \delta_{0i}, i = 1, \dots, n = 1262$ with the δ_{0i} as *possible* specified values of δ_i .
- Since $R_i = r_{Ci} + Z_i \delta_i$, if the hypothesis H_0 were true, then $R_i - Z_i \delta_{0i}$ would equal r_{Ci} .
- But R_i and Z_i are observed and δ_{0i} is specified by the hypothesis, so if the hypothesis were true, we could calculate the r_{Ci} .
- Under the null hypothesis, the 2×2 table recording r_{Ci} by Z_i has the hypergeometric distribution, yielding a test.

22 Procedure

- If $H_0 : \delta_i = \delta_{0i}, i = 1, \dots, n = 1262$ were true, then $r_{Ci} = R_i - Z_i \delta_{0i}$, so the the 2×2 table recording r_{Ci} by Z_i would be:

	<i>Recurrence</i> $R_i = 1$	<i>No recurrence</i> $R_i = 0$
No Rads $Z_i = 1$	$\sum Z_i (R_i - Z_i \delta_{0i})$	$\sum Z_i (1 - R_i + Z_i \delta_{0i})$
Rads $Z_i = 0$	$\sum (1 - Z_i) R_i$	$\sum (1 - Z_i) (1 - R_i)$

	<i>Recurrence</i> $r_{Ci} = 1$	<i>No recurrence</i> $r_{Ci} = 0$
No Rads $Z_i = 1$	$\sum Z_i r_{Ci}$	$\sum Z_i (1 - r_{Ci})$
Rads $Z_i = 0$	$\sum (1 - Z_i) r_{Ci}$	$\sum (1 - Z_i) (1 - r_{Ci})$

which would have the hypergeometric distribution.

23 Attributable effect

- The procedure shifts a count of $A_0 = \sum Z_i \delta_{0i}$, which, if the null hypothesis is true, equals

$$A = \sum Z_i \delta_i = \sum Z_i (r_{Ti} - r_{Ci}),$$

that is the net number of additional women caused to have a recurrence by the use of lumpectomy alone rather than lumpectomy plus irradiation.

- Although I can calculate $A_0 = \sum Z_i \delta_{0i}$ from the hypothesis and the data, the true $A = \sum Z_i \delta_i$ is an unobservable random variable.

24 Example

- If a possible hypothesis $H_0 : \delta_i = \delta_{0i}, i = 1, \dots, n = 1262$ yields $A_0 = \sum Z_i \delta_{0i} = 119$, compute:

	<i>Recurrence</i> $R_i = 1$	<i>No recurrence</i> $R_i = 0$	Total
No rads $Z_i = 1$	220 – 119	414 + 119	634
Rads $Z_i = 0$	78	550	628
Total	179	1,083	1,262

and the hypergeometric tail probability $\Pr(T \geq 220 - 119) = \Pr(T \geq 101) = 0.0438$, so H_0 is not quite plausible. If we do the same for a possible hypothesis $H_0 : \delta_i = \delta_{0i}, i = 1, \dots, n = 1262$ yielded $A_0 = \sum Z_i \delta_{0i} = 120$, then the tail probability is 0.0514, and so barely plausible.

- That is, we are 95% confident that, net, at least 120 more of the 634 women treated with lumpectomy alone had recurrence of cancer caused by the failure to combine lumpectomy with irradiation.

25 Randomization Inference in General

- Randomization inference was illustrated in a simple situation: Fisher's exact test for a 2×2 table.
- The concept is very general, however. Among the tests commonly derived as randomization tests are: Wilcoxon's signed rank and rank sum tests, the Mantel-Haenszel-Birch test for $2 \times 2 \times S$ tables, the Mantel-extension test for integer scores, the Gehan and log-rank tests for censored survival data. (eg Lehmann 1999)
- In a randomized experiment, ANOVA-based tests can be derived as approximations to randomization tests. (eg. Welch 1937).
- Can be combined with models (Gail, et al 1988), smoothers (Raz 1990), and nonparametric covariance adjustments (Rosenbaum 2002).

26 Wilcoxon's Signed Rank Statistics

- Partly to illustrate, partly as a transition to observational studies, will illustrate randomization inference with Wilcoxon's signed rank statistic.
- Do with data from an observational study, a nonrandomized study of treatment effects, at first acting as if it were a randomized experiment, then considering the absence of randomization.
- Matched pairs: treated, control. Rank the absolute differences in responses within pairs. Sum ranks of positive differences.

27 Example: A Matched Observational Study

- From Morton, et al. (1982) Lead absorption in children of employees in a lead-related industry. *American Journal of Epidemiology*, 115, 549-
- Study of one child of each of 33 workers in a battery factory in Oklahoma in 1978. Concern was that they might bring lead home, exposing their children.
- 33 control children were individually selected and matched to the exposed children. They were matched for neighborhood and age (± 1 year). Neighborhood: (i) if an apartment, then another apartment from same complex, (ii) if facing a main road, then a nearby house facing the same road, etc.
- Outcome: child's blood lead level, μg of lead per *dl* blood.

28 Notation for a Paired Experiment

Pair s , Subject i : $S = 33$ pairs, $s = 1, \dots, S = 33$, with 2 subjects in each pair, $i = 1, 2$.

One treated, one control in each pair: Write $Z_{si} = 1$ if the i^{th} subject in pair s is treated, $Z_{si} = 0$ if control, so $Z_{s1} + Z_{s2} = 1$ for every s , or $Z_{s2} = 1 - Z_{s1}$. For all $2S$ subjects,

$$\mathbf{Z} = (Z_{11}, Z_{12}, \dots, Z_{S1}, Z_{S2})^T.$$

Random assignment of treatments within pairs: Ω is the set of the $K = 2^S$ possible values \mathbf{z} of \mathbf{Z} , and randomization picks one of these at random,

$$\Pr(\mathbf{Z} = \mathbf{z}) = \frac{1}{K} \text{ for each } \mathbf{z} \in \Omega.$$

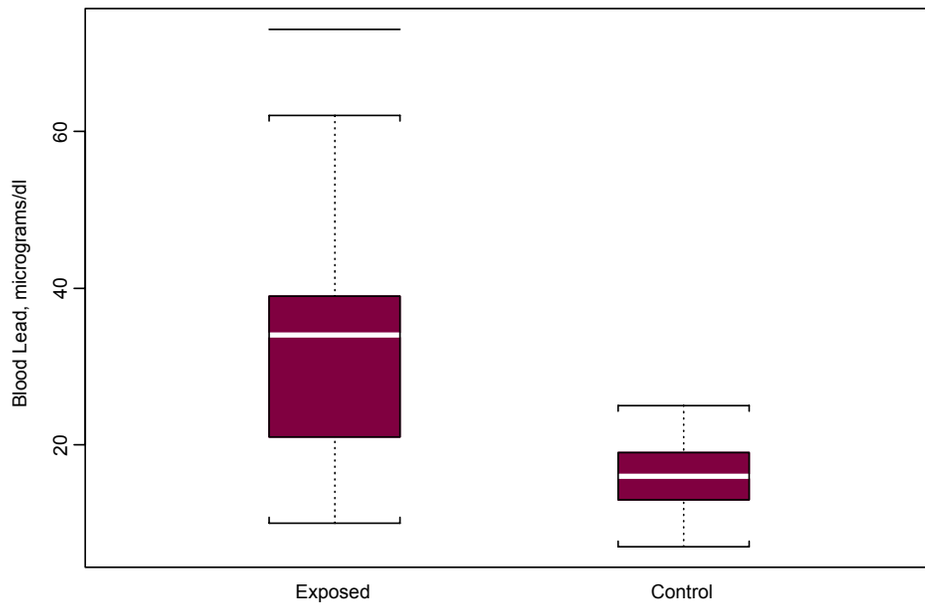


Figure 1: Lead levels in children whose fathers were exposed to lead at work and matched, unexposed control children.

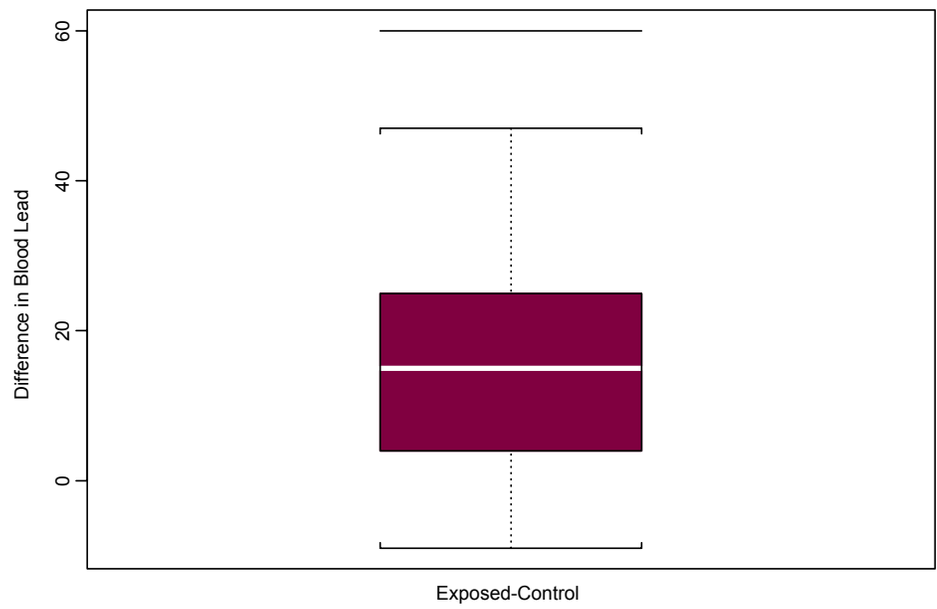


Figure 2: Matched pair differences in lead levels.

29 Responses, Causal Effects

Potential responses, causal effects, as before. Each of the $2S$ subjects (s, i) has two potential responses, a response r_{Tsi} that would be seen under treatment and a response r_{Csi} that would be seen under control. (Neyman 1923, Rubin 1974). Treatment effect is $\delta_{si} = r_{Tsi} - r_{Csi}$. *Additive effect*, $r_{Tsi} - r_{Csi} = \tau$ or $\delta_{si} = \tau$ for all s, i .

Finite population, as before. The (r_{Tsi}, r_{Csi}) , $s = 1, \dots, S$, $i = 1, 2$, are again fixed features of the finite population of $2S$ subjects.

Observed responses, as before. Observed response is $R_{si} = r_{Tsi}$ if $Z_{si} = 1$ or $R_{si} = r_{Csi}$ if $Z_{si} = 0$, that is, $R_{si} = Z_{si} r_{Tsi} + (1 - Z_{si}) r_{Csi} = r_{Csi} + Z_{si} \delta_{si}$. If effect is additive, $R_{si} = r_{Csi} + Z_{si} \tau$.

Vectors. $2S$ -dimensional vectors \mathbf{r}_T , \mathbf{r}_C , $\boldsymbol{\delta}$, \mathbf{R} ; e.g., $\mathbf{R} = (R_{11}, \dots, R_{S2})^T$.

30 Treated-Minus-Control Differences

Who is treated in pair s ? If $Z_{s1} = 1$, then $(s, 1)$ is treated and $(s, 2)$ is control, but if $Z_{s2} = 1$ then $(s, 2)$ is treated and $(s, 1)$ is control.

Treated-minus-control differences with additive effects:

If $r_{Tsi} - r_{Csi} = \tau$, then a little algebra shows the *treated-minus-control difference* in observed responses in pair s is:

$$D_s = (Z_{s1} - Z_{s2})(r_{Cs1} - r_{Cs2}) + \tau.$$

In general, without additive effects: In general, with

$$\delta_{si} = r_{Tsi} - r_{Csi}:$$

$$D_s = (Z_{s1} - Z_{s2})(r_{Cs1} - r_{Cs2}) + Z_{s1} \delta_{s1} + Z_{s2} \delta_{s2}.$$

31 Signed Rank Statistic

Usual form. Wilcoxon's *signed rank statistic* W ranks the $|D_s|$ from 1 to S , and sums the ranks of the positive D_s . (Ties ignored today.)

Equivalent alternative form. Another expression for W uses the $S(S+1)/2$ Walsh averages,

$$\frac{D_s + D_{s'}}{2}, \text{ with } 1 \leq s \leq s' \leq S;$$

then W is the number of positive Walsh averages. (Lehmann 1998)

Offsets. A Walsh average $(D_s + D_{s'})/2$ is positive if the more affected pair of s and s' was sufficiently affected to offset whatever happened to the less affected pair, that is, if $\max(D_s, D_{s'})$ is sufficiently large that it averages with $\min(D_s, D_{s'})$ to be positive. Then W is the number of offsets.

32 No Effect in an Experiment

Null hypothesis. $H_0 : \delta_{si} = 0$, for $s = 1, \dots, S$,
 $i = 1, 2$ where $\delta_{si} = r_{Tsi} - r_{Csi}$.

Differences. If H_0 is true, then the treated-minus-control difference is:

$$D_s = (Z_{s1} - Z_{s2})(r_{Cs1} - r_{Cs2})$$

where $Z_{s1} - Z_{s2}$ is ± 1 where randomization ensures $\Pr(Z_{s1} - Z_{s2} = 1) = \frac{1}{2}$, independently in different pairs, and $r_{Cs1} - r_{Cs2}$ is fixed in Fisher's finite population.

Signed rank statistic. If H_0 is true, D_s is $\pm (r_{Cs1} - r_{Cs2})$ with probability $\frac{1}{2}$, so $|D_s| = |r_{Cs1} - r_{Cs2}|$ is fixed, as is its rank, so ranks independently add to W with probability $\frac{1}{2}$, generating W 's distribution.

Randomization. Uses just fact of randomization and null hypothesis, so forms the "reasoned basis for inference," in Fisher's phrase.

33 Randomization Test for an Additive Effect

Additive effect. $H_0 : \delta_{si} = \tau_0$, for $s = 1, \dots, S$, $i = 1, 2$ where $\delta_{si} = r_{Tsi} - r_{Csi}$.

Matched pair differences. If H_0 were true, then

$$D_s = (Z_{s1} - Z_{s2})(r_{Cs1} - r_{Cs2}) + \tau_0$$

so the adjusted differences

$$D_s - \tau_0 = (Z_{s1} - Z_{s2})(r_{Cs1} - r_{Cs2})$$

satisfy the hypothesis of no effect, and W computed from $D_s - \tau_0$ has the usual null distribution of the signed rank statistic.

Randomization. Again, the inference uses only the fact of randomization and the null hypothesis being tested.

34 Confidence Interval for Additive Effect

Additive effects. $\delta_{si} = \tau$, for all s, i where $\delta_{si} = r_{Tsi} - r_{Csi}$

Inverting tests. The 95% interval for τ is the set of all τ_0 not rejected in a 0.05 level test.

Confidence intervals. Test every τ_0 by computing W from the adjusted differences, $D_s - \tau_0$, retaining values τ_0 not rejected at the 0.05 level.

Hodges-Lehmann estimates. Find $\hat{\tau}$ so that W computed from $D_s - \hat{\tau}$ equals its null expectation.

35 Example: Lead Exposure

Morton, et al. 33 matched pairs of children, exposed-control, D_s is the difference in blood lead levels.

Not randomized. First, will perform analysis appropriate for a randomized experiment, then return to the example several times to think about consequences of nonrandom assignment to treatment.

Test of no effect. Signed rank statistic is $W = 527$, with randomization based $P - value = 10^{-5}$.

Confidence interval. 95% for an additive effect is $[9.5, 20.5]$ $\mu g/dl$. The two-sided $P - value$ is ≥ 0.05 if W is computed from $D_s - \tau_0$ for $\tau_0 \in (9.5, 20.5)$ and is less than 0.05 for $\tau_0 \notin [9.5, 20.5]$.

HL estimate. $\hat{\tau} = 15 \mu g/dl$ as $D_s - 15$ (effectively) equates W to its null expectation.

36 Nonadditive effects

Effects that vary. If $\delta_{si} = r_{Tsi} - r_{Csi}$ vary from person to person, cannot estimate a single effect τ .

Walsh averages. W is the number of positive offsets or Walsh averages:

$$\frac{D_s + D_{s'}}{2}, \text{ with } 1 \leq s \leq s' \leq S;$$

the number of times a large D_s offset whatever happened to $D_{s'}$ to yield a positive average.

It could be luck. Some are positive because $\delta_{si} > 0$ or $\delta_{s'i} > 0$, and others by the luck. Under the null hypothesis of no effect, expect half (or $S(S+1)/4$) of the Walsh averages to be positive by chance in a randomized experiment.

Attributable effect. How many are positive because of treatment effects, say $\delta_{si} > 0$ or $\delta_{s'i} > 0$, and not by luck.

37 Attributable Effect

Walsh averages without treatment. If lead exposure had been prevented for all children, then the S differences would not have been

$$D_s = (Z_{s1} - Z_{s2})(r_{Cs1} - r_{Cs2}) + Z_{s1} \delta_{s1} + Z_{s2} \delta_{s2}.$$

but rather

$$\tilde{D}_s = (Z_{s1} - Z_{s2})(r_{Cs1} - r_{Cs2}),$$

and we would have computed their Walsh averages

$$\frac{\tilde{D}_s + \tilde{D}_{s'}}{2}, \text{ with } 1 \leq s \leq s' \leq S.$$

Attributable effect. The number of times A that

$$\frac{D_s + D_{s'}}{2} > 0, \text{ but } \frac{\tilde{D}_s + \tilde{D}_{s'}}{2} \leq 0, \text{ with } 1 \leq s \leq s' \leq S.$$

38 Attributable Effect: Inference

Null distribution. Under H_0 of no effect, $\Pr(W \geq 374) = 0.0485 \leq 0.05$, from standard tables with $S = 33$ pairs. We observed $W = 527$.

Confidence statement. In a randomized experiment, 95% confidence that:

$$A \geq 527 - 374 + 1 = 154$$

because generally

$$\Pr(A \geq W - k_\alpha + 1) \geq 1 - \alpha$$

where $\Pr(W \geq k_\alpha) = \alpha$ from tables of the signed rank statistic.

Yardstick. In a randomized experiment, we expected half, or $S(S + 1)/4 = 280.5$ to be positive by luck, but observed $W = 527$ positive, or $(527 - 280.5)/280.5 = 88\%$ more than expected. 95% confident that $154/280.5 = 55\%$ are due to effects of the treatment.

39 But the study was not randomized . . .

Not randomized. The analysis would have been justified by randomization in a randomized experiment.

Unknown assignment probabilities. An observational study is a study of treatment effects in which each person has an unknown probability of treatment, typically different probabilities for different people.

Simple model. In some finite population of people, $j = 1, \dots, J$, person j has probability $\pi_j = \Pr(Z_j = 1)$ of exposure to treatment, where π_j is not known. Probabilities are always *conditional on things we regard as fixed*, usually measured and unmeasured covariates, potential outcomes, (r_{Tj}, r_{Cj}) , etc.

40 Simple model continued . . .

Covariates. The people, $j = 1, \dots, J$, in the finite population have observed covariates \mathbf{x}_j and unobserved covariate u_j . In the example, \mathbf{x}_j describes child's age and neighborhood.

Absolutely simplest case: Select S pairs, $i = 1, 2$, one treated, one control, from the J people in the population. Match exactly for \mathbf{x} , so that $\mathbf{x}_{s1} = \mathbf{x}_{s2}$ for each s , $s = 1, \dots, S$.

Matching algorithm: In this simplest case, the matching algorithm is permitted to use only \mathbf{x} and $\mathbf{1} = Z_{s1} + Z_{s2}$.

41 Minor Issue: Dependence Among Assignments

Experiments. In most experiments, treatment assignments are dependent in simple ways; eg, half the people get treatment. Or one person in each pair gets treatment. Shapes randomization distributions.

Goal: Keep the model for observational studies parallel to experiments, but keep it simple.

Solution: Dependence introduced by conditioning.

Nice way to handle this is to make the Z_j independent and then introduce dependence by conditioning, eg on $1 = Z_{s1} + Z_{s2}$.

42 Is the randomization analysis ever appropriate?

Ignorability, etc. Cluster of concepts — ignorability, strong ignorability, free of hidden bias, no unmeasured confounders, etc. — .with similar intuition.

Intuition: Answers the *theoretical* question: When is it enough to adjust for the observed covariates you have, \mathbf{x}_j ?

A theoretical question. Frames the discussion, but doesn't tell you what to make of data. In observational studies you always: (i) adjust for the covariates you have \mathbf{x}_j , (ii) worry that you omitted some important (unobserved covariate) u_j , and try to find ways to address this possibility.

43 Free of hidden bias

Definition. Treatment assignment is free of hidden bias if π_j is a (typically unknown) function of \mathbf{x}_j — two people with the same \mathbf{x}_j have the same π_j .

Intuition. A kid j who lives 30 miles from the battery factory is less likely to have a dad working in factory than a kid k who lives two miles from the factory, $\pi_j < \pi_k$, but two kids of the same age who next door are equally likely to have a dad in the factory.

But they didn't match on kid's gender. If gender were not recorded, it would violate 'free of hidden bias' if (roughly) boys were more likely (or less likely) than girls to have a dad working in the battery factor.

44 If free of hidden bias . . .

Problem: Unlike an experiment, π_j are unknown.

If free of hidden bias: Two people with the same \mathbf{x}_j have the same π_j , which is typically unknown.

Eliminate unknowns by conditioning: If we match exactly for \mathbf{x} , so $\mathbf{x}_{s1} = \mathbf{x}_{s2}$, then

$$\begin{aligned} \Pr(Z_{s1} = 1 \mid Z_{s1} + Z_{s2}) \\ = \frac{\pi_{s1}(1 - \pi_{s2})}{\pi_{s1}(1 - \pi_{s2}) + \pi_{s2}(1 - \pi_{s1})} = \frac{1}{2} \end{aligned}$$

because $\pi_{s1} = \pi_{s2}$. A little more work shows that we get the randomization distribution by conditioning.

More generally, This argument is quite general, working for matched sets, strata, and more complex problems.

45 Interpretation

If free of hidden bias: Two people with the same \mathbf{x}_j have the same π_j , which is typically unknown.

When do adjustments work? If a study is free of hidden bias, if the only bias is due to observed covariates \mathbf{x}_j , even if the bias is unknown, the bias can be removed in various ways, such as matching on \mathbf{x}_j , and conventional randomization inferences yield appropriate inferences about treatment effect.

Key, if problematic, assumption. Identifies the key assumption, but of course, doesn't make it true. Focuses attention, frames discussion. In contrast, in an experiment, randomization makes it true.

Divides methods. Methods of adjustment for \mathbf{x} should work when study is free of hidden bias. Need other methods to address concerns about whether the study is free of hidden bias.

46 Propensity Scores

Many observed covariates. If \mathbf{x} is of high dimension, it's hard to match. With just 20 binary covariates, there are 2^{20} or about a million covariate patterns.

If free of hidden bias: Two people with the same \mathbf{x}_j have the same π_j , so π_j is a function of \mathbf{x}_j , say $\pi_j = e(\mathbf{x}_j)$, which is then called the propensity score. .

Old argument again: Match exactly for \mathbf{x} , so $\mathbf{x}_{s1} = \mathbf{x}_{s2}$, then

$$\begin{aligned} \Pr(Z_{s1} = 1 \mid Z_{s1} + Z_{s2}) \\ = \frac{\pi_{s1}(1 - \pi_{s2})}{\pi_{s1}(1 - \pi_{s2}) + \pi_{s2}(1 - \pi_{s1})} = \frac{1}{2} \end{aligned}$$

because $\pi_{s1} = \pi_{s2}$ or $e(\mathbf{x}_{s1}) = e(\mathbf{x}_{s2})$

Key point: Don't need to match on high dimension \mathbf{x} , just need to match on the scalar $e(\mathbf{x})$.

47 Balancing with Propensity Scores

Whether or not the study is free of hidden bias, matching on propensity scores $e = e(\mathbf{x})$ tends to balance the observed covariates \mathbf{x} used in the score. Define $e = e(\mathbf{x}) = \Pr(Z = 1 | \mathbf{x})$, so the study is free of hidden bias if $\pi_j = e(\mathbf{x}_j)$ for all j , but $e(\mathbf{x})$ is defined even if π_j depends on things besides \mathbf{x} .

That is:

$$\Pr(\mathbf{x} | Z = 1, e) = \Pr(\mathbf{x} | Z = 0, e)$$

$$\text{or } \mathbf{x} \perp\!\!\!\perp Z | e(\mathbf{x})$$

Proof: Suffices to show $\Pr\{Z = 1 | \mathbf{x}, e(\mathbf{x})\}$ equals $\Pr\{Z = 1 | e(\mathbf{x})\}$. But $\Pr\{Z = 1 | \mathbf{x}, e(\mathbf{x})\} = \Pr(Z = 1 | \mathbf{x})$ which is just $e(\mathbf{x})$. Also, $\Pr\{Z = 1 | e(\mathbf{x})\}$ equals $E[\Pr\{Z = 1 | \mathbf{x}, e(\mathbf{x})\} | e(\mathbf{x})] = E[\Pr\{Z = 1 | \mathbf{x}\} | e(\mathbf{x})] = E[e(\mathbf{x}) | e(\mathbf{x})] = e(\mathbf{x})$.

48 Propensity Scores: Example

Source: From Rosenbaum and Rubin (1984) *JASA*.

Data: Database describing 1,515 patients with coronary artery disease, treated either with CABG or drugs. Interest in effects of CABG vs drugs on survival, pain, etc.

Many covariates: CABG and drug patients differed significantly on 74 covariates. Drug patients were either too sick or too healthy for surgery.

Covariate	t-statistic	F-statistic
Ejection fraction	4.4	19.4
Poor left ventricle function	7.2	51.8
Left main artery occluded	4.7	22.1
Progressing Chest Pain	6.6	43.6

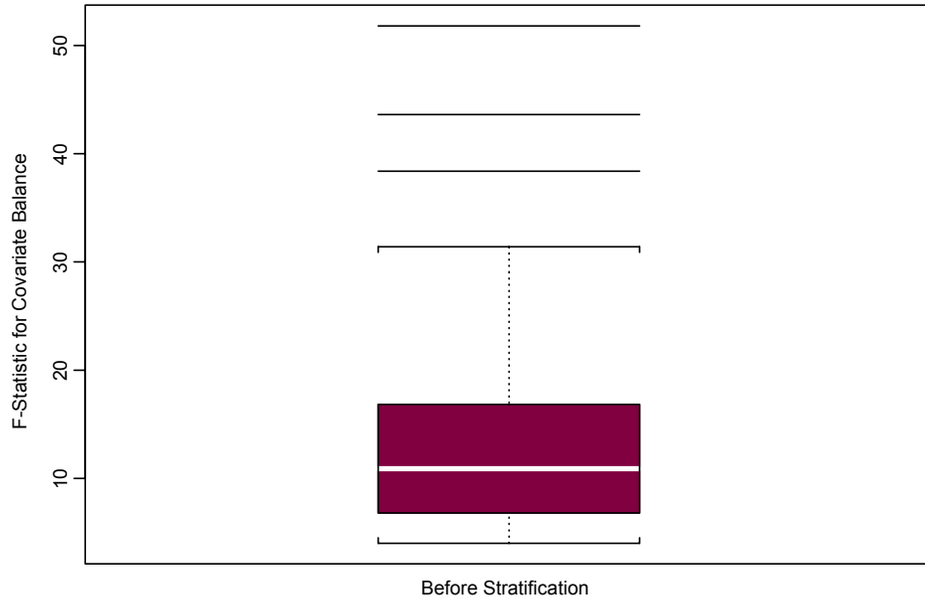


Figure 3:

49 Boxplot Before Stratification

Covariate Imbalance. Covariate imbalance for 74 covariates before stratification on the propensity score. Display is $F = t^2$ for 74 covariates.

50 Procedure

Propensity score: Estimated using logit regression of treatment (CABG or drugs) on covariates, some quadratics, some interactions.

Five strata: Five groups formed at quintiles of the estimated propensity score.

Counts of Patients in Strata

Propensity Score Stratum	Medical	Surgical
1 = <i>lowest = most medical</i>	277	26
2	235	68
3	205	98
4	139	164
5 = <i>highest = most surgical</i>	69	234

51 Checking balance

2-Way 5×2 Anova for Each Covariate

Propensity Score Stratum	Medical	Surgical
1 = <i>lowest</i> = <i>most medical</i>		
2		
3		
4		
5 = <i>highest</i> = <i>most surgical</i>		

Balance check. Main effect and interaction F –statistics.

52 F-statistics Before and After Stratification

Covariate	Before	After Main Effect	After Interaction
Ejection fraction	19.4	0.0	0.3
Poor LV function	51.8	0.4	0.9
Left main occluded	22.1	0.3	0.2
Progressing Pain	43.6	0.1	1.4

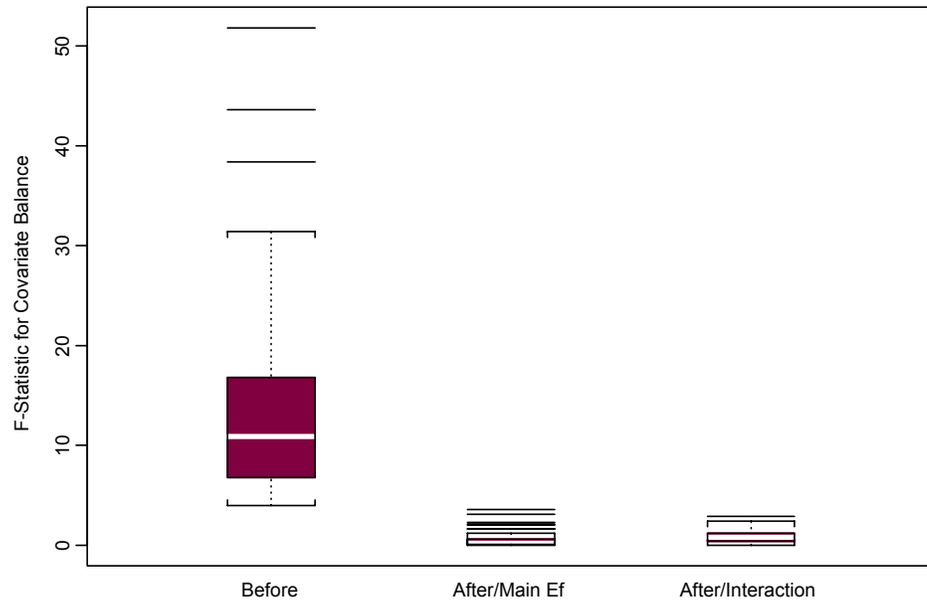


Figure 4:

53 Is there covariate balance within strata?

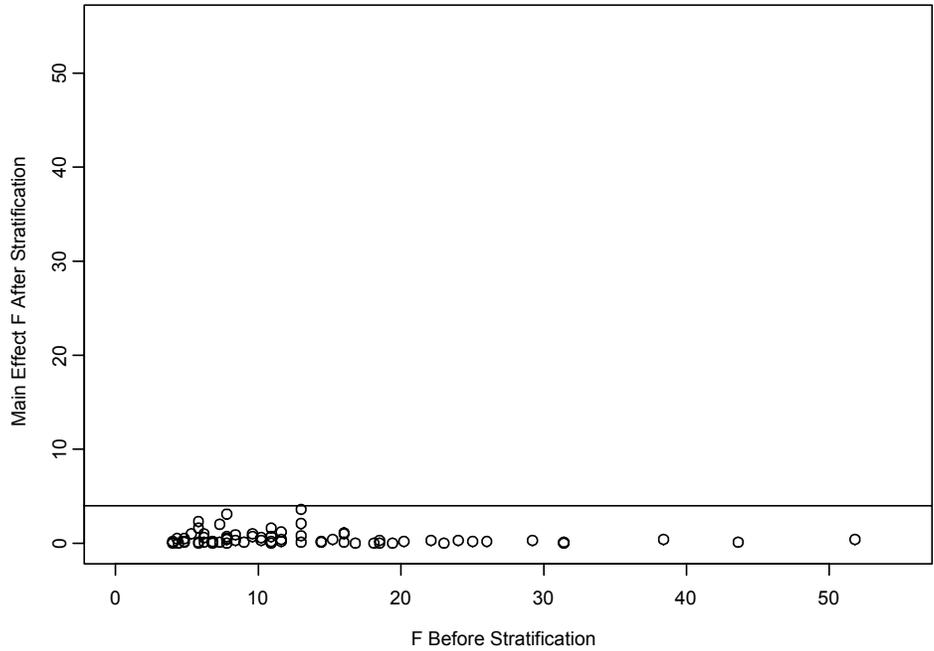


Figure 5:

54 Covariate balance: Alternative view

55 Last words about propensity scores

Balancing. Stratifying or matching on a scalar propensity score tends to balance many observed covariates.

Effects of estimating the score. Examples, simulations, limited theory suggest estimated scores provide slightly *more* than true propensity scores.

Other methods. Various methods permit explicit acknowledgement of use of estimated scores.

Key limitation. Propensity scores balance only observed covariates, whereas randomization also balances unobserved covariates.

56 Addressing hidden bias

If free of hidden bias: Two people with the same observed \mathbf{x}_j have the same π_j , which is typically unknown. Can remove the overt biases due to \mathbf{x}_j .

Common objection: Critic says: “Adjusting for \mathbf{x}_j is not sufficient, because there is an unobserved u_j , and adjustments for (\mathbf{x}_j, u_j) were needed.”

That is, the objection asserts that, or raises the possibility that, the observed association between treatment Z_j and response R_j is not an effect caused by the treatment, but rather due to hidden bias from their shared relationship with u_j .

Formally, treatment assignment Z_j and response $R_j = r_{Cj} + Z_j (r_{Tj} - r_{Cj})$ may be associated because $r_{Tj} - r_{Cj} \neq 0$ (a treatment effect) or because $r_{Tj} - r_{Cj} = 0$ but π_j and r_{Cj} both vary with u_j (a hidden bias due to u_j).

57 Sensitivity analysis

Question answered by a sensitivity analysis: If the objection were true, if the association between treatment Z_j and response R_j were due to hidden bias from u_j , then what would u_j have to be like?

What does the counter-claim actually claim? A sensitivity analysis looks at the observed data and uses it to clarify what the critic's counter claim is actually claiming.

Sensitivity varies. Studies vary markedly in how sensitive they are to hidden bias.

58 First Sensitivity Analysis

Cornfield, et al. (1959): they write:

“If an agent, A , with no causal effect upon the risk of a disease, nevertheless, because of a positive correlation with some other causal agent, B , shows an apparent risk, r , for those exposed to A , relative to those not so exposed, then the prevalence of B , among those exposed to A , relative to the prevalence among those not so exposed, must be greater than r .

Thus, if cigarette smokers have 9 times the risk of non-smokers for developing lung cancer, and this is not because cigarette smoke is a causal agent, but only because cigarette smokers produce hormone X , then the proportion of hormone X -producers among cigarette smokers must be at least 9 times greater than that of nonsmokers. If the relative prevalence of hormone X -producers is considerably less than ninefold, then hormone X cannot account for the magnitude of the apparent effect.”

59 The Cornfield, et al Inequality

The Cornfield, et al sensitivity analysis is an important conceptual advance:

“Association does not imply causation
— hidden bias can produce associations,”

is replaced by

“To explain away the association actually seen,
hidden biases would have to be of such and
such a magnitude.”

Provides a quantitative measure of uncertainty in light
of data.

As a confidence interval measures sampling uncertainty
without making it go away, a sensitivity analysis mea-
sure uncertainty due to hidden bias without making
the uncertainty go away.

60 Alternative sensitivity analysis

Limitations. Cornfield's inequality concerns binary responses only and ignores sampling variability. Not explicit about observed covariates.

Alternative formulation. Two subjects, j and k , with the same observed covariates, $\mathbf{x}_j = \mathbf{x}_k$, may differ in terms of u_j and u_k so that their odds of exposure to treatment differ by a factor of $\Gamma \geq 1$,

$$\frac{1}{\Gamma} \leq \frac{\pi_j (1 - \pi_k)}{\pi_k (1 - \pi_j)} \leq \Gamma.$$

Free of hidden bias is then $\Gamma = 1$.

When bias is present, when $\Gamma > 1$, the unknown π_j cannot be eliminated, as before, by matching on \mathbf{x}_j , so the randomization distribution is no longer justified.

61 Alternative sensitivity analysis, continued

Model. Two subjects, j and k , with $\mathbf{x}_j = \mathbf{x}_k$, may differ their odds of exposure to treatment differ by a factor of $\Gamma \geq 1$,

$$\frac{1}{\Gamma} \leq \frac{\pi_j (1 - \pi_k)}{\pi_k (1 - \pi_j)} \leq \Gamma \quad (1)$$

so Γ provides measured departure from “no hidden bias.”

Intuition: If $\Gamma = 1.001$, the π_j are unknown, but almost the same. If $\Gamma = 5$, π_j are unknown and could be very different.

Plan. For each $\Gamma \geq 1$, find upper and lower bounds on inference quantities, like P-values (or endpoints of confidence intervals), for π_j 's satisfying (1). Report these for several Γ . When do conclusions begin to change?

62 Signed Rank Statistic

Model. If $\mathbf{x}_j = \mathbf{x}_k$, then

$$\frac{1}{\Gamma} \leq \frac{\pi_j (1 - \pi_k)}{\pi_k (1 - \pi_j)} \leq \Gamma. \quad (2)$$

Structure: As before, match on observed covariates \mathbf{x} , to form S pairs, $s = 1, \dots, S$, $i = 1, 2$, with $\mathbf{x}_{s1} = \mathbf{x}_{s2}$, one treated, one control, $Z_{s1} + Z_{s2} = 1$.

Free of hidden bias: If $\Gamma = 1$, obtained the randomization distribution of Wilcoxon's signed rank statistic W , as $\Pr(Z_{s1} = 1 \mid Z_{s1} + Z_{s2}) = \frac{1}{2}$.

Fact: Then (2) implies:

$$\frac{1}{1 + \Gamma} \leq \Pr(Z_{s1} = 1 \mid Z_{s1} + Z_{s2}) \leq \frac{\Gamma}{1 + \Gamma}$$

which places sharp upper and lower bounds on the distribution of W and resulting inferences.

63 Lead Exposure: Significance Levels

Data: $S = 33$ pairs of children matched for age and neighborhood, one having a parent exposed to lead, the other a control. Measured lead levels in the children's blood. Used Wilcoxon's signed rank test, W .

Sensitivity analysis. One sided significance levels for testing no effect.

Γ	min	max
1	<0.0001	<0.0001
2	<0.0001	0.0018
3	<0.0001	0.0136
4	<0.0001	0.0388
4.25	<0.0001	0.0468
5	<0.0001	0.0740

64 One Sided Confidence Intervals

95% CI. For an additive effect, $r_{Tsi} = r_{Csi} + \tau$, the signed rank test may be inverted to yield a one-sided 95% confidence interval.

Range of values: For $\Gamma > 1$, the endpoint $\hat{\tau}_{low}$ of the one-sided 95% interval $[\hat{\tau}_{low}, \infty)$ for τ has a range of values. Table gives the smallest value in the range — the smallest plausible effect for the given quantity of hidden bias.

Sensitivity analysis.

Γ	$\min \hat{\tau}_{low}$
1	10.5
2	5.5
3	2.5
4	0.5
4.25	0.0
5	-1.0

65 Comparing Different Studies

Studies vary markedly in their sensitivity to hidden bias.

Treatment	$\Gamma = 1$	$(\Gamma, \max P - \text{value})$
Smoking/Lung Cancer Hammond 1964	< 0.0001	(5, 0.03)
DES/vaginal cancer Herbst, et al. 1976	< 0.0001	(7, 0.054)
Lead/Blood lead Morton, et al. 1982	< 0.0001	(4.25, 0.047)
Coffee/MI Jick, et al. 1973	0.0038	(1.3, 0.056)

Small biases could explain Coffee/MI association. Very large biases would be needed to explain DES/vaginal cancer association.

66 Sensitivity Analysis: Interpretation

Uses data, says something tangible. Replaces qualitative “association does not imply causation,” by a quantitative statement based on observed data, “to explain away observed associations as noncausal, hidden biases would have to be of such and such a magnitude.”

Measures uncertainty. Measures uncertainty due to hidden bias, but does not dispel it. (As a confidence interval measures sampling uncertainty but does not dispel it.)

Fact of the matter. Your opinion about how much hidden bias is present is your opinion. But the degree of sensitivity to hidden bias is a fact of the matter, something visible in observed data.

67 Hill on Causality

Who was Hill? With Sir Richard Doll, Sir Austin Bradford Hill did some of the most careful and influential studies of smoking and health.

Hill's Aspects: In 1965, Hill wrote a paper "The environment and disease: association or causation." He asked:

"Our observations reveal an association between two variables, perfectly clear-cut and beyond what we would care to attribute to the play of chance. What aspects of that association should we especially consider before deciding that the mostly likely interpretation of it is causation?"

i.e., after adjusting for biases we can see, what aspects of the observed association provide information about cause and effect vs hidden bias?

68 Hill on Causality, continued.

Nine aspects: Strength, consistency, specificity, temporality, biological gradient, plausibility, coherence, experiment, analogy.

Consistency: “Has [the association] been repeatedly observed by different persons, in different places, circumstances and times?”

Biological gradient: “[I]f the association is one which can reveal a biological gradient, or dose-response curve, then we should look most carefully for such evidence.”

Coherence: “[T]he cause-and-effect interpretation . . . should not seriously conflict with generally known facts of the natural history and biology of the disease.”

69 Reactions to Hill's Aspects

Influence: Discussed in many textbooks in epidemiology, often mentioned in empirical papers.

Critics: Rothman, Sartwell, others have been critical of Hill's aspects, perhaps not so much in the form Hill described them, but rather in the more rigid way they are sometimes described in textbooks.

Was Hill correct? As Hill often adjusted for observed covariates, it is clear that his aspects are intended to provide information about hidden bias. Do they?

70 Pattern matching

Social sciences: Similar considerations are referred to as “pattern matching.”

Cook and Shadish 1994, p. 565: ‘Successful prediction of a complex pattern of multivariate results often leaves few plausible alternative explanations.’

Trochim 1985, p. 580: ‘... with more pattern specificity it is generally less likely that plausible alternative explanations for the observed effect pattern will be forthcoming.’

Campbell 1988, p. 33: ‘... great inferential strength is added when each theoretical parameter is exemplified in two or more ways, each mode being as independent as possible of the other, as far as the theoretically irrelevant components are concerned’

71 More on pattern matching

Cook, Campbell, Peracchio (1990): "... the warrant for causal inferences from quasi-experiments rests [on] structural elements of design other than random assignment – pretests, comparison groups, the way treatments are scheduled across groups ... — [which] provide the best way of ruling out threats to internal validity ... [C]onclusions are more plausible if they are based on evidence that corroborates numerous, complex, or numerically precise predictions drawn from a descriptive causal hypothesis."

72 Fisher's Comment: Elaborate Theories

Cochran (1965, §5): “About 20 years ago, when asked in a meeting what can be done in observational studies to clarify the step from association to causation, Sir Ronald Fisher replied: ‘Make your theories elaborate.’ The reply puzzled me at first, since by Occam’s razor, the advice usually given is to make theories as simple as is consistent with known data. What Sir Ronald meant, as subsequent discussion showed, was that when constructing a causal hypothesis one should envisage as many different consequences of its truth as possible, and plan observational studies to discover whether each of these consequences is found to hold.

... this multi-phasic attack is one of the most potent weapons in observational studies.”

73 Lead Example

Two doses. Morton, et al. measured two 'doses' of lead exposure. Father's lead exposure at work: high, medium or low. Father's hygiene upon leaving the factory: poor, moderate or good.

Three groups. Roughly equal sizes.

Exposure, Hygiene	Count
High and Poor	13
Intermediate	8
Low or Good	12
Controls	33

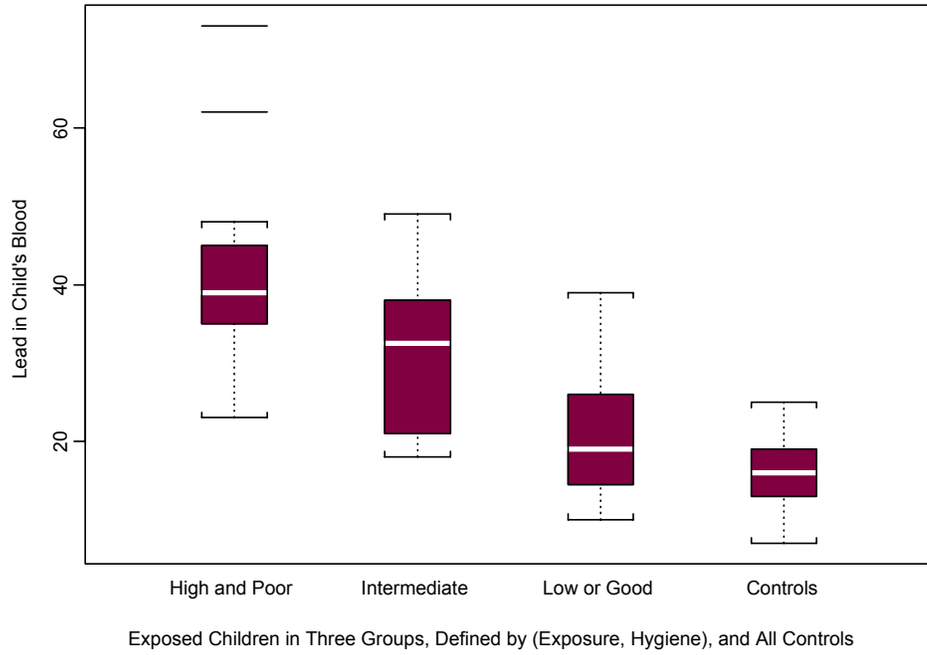


Figure 6:

74 Plot By Dose

75 Coherent signed rank statistic

Pattern weighted. Matched pairs are weighted to reflect anticipated pattern. May use doses (as here) or multivariate patterns, or both.

Here: (low exposure or good hygiene) pairs get weight 1, (high exposure, poor hygiene) pairs get weight 2, intermediate pairs get 1.5.

Statistic: Sum of $weight \times rank(|D_s|)$ if $D_s > 0$.

Properties: If anticipated pattern is correct, greater power than W . If pattern is contradicted, typically lower power.

76 Pattern Specificity and Sensitivity to Hidden Bias

Question: Does pattern specificity (e.g., a dose-response relationship) reduce sensitivity to hidden bias?

Sensitivity analysis. Upper bounds on one sided significance levels for testing no effect.

Γ	Wilcoxon SR	Coherent SR
1	<0.0001	<0.0001
3	0.0136	0.0119
4.35	0.0502	0.0398
4.75	0.0645	0.0503

Reduced sensitivity. Measured reduction in sensitivity to hidden bias, from $\Gamma = 4.35$ to $\Gamma = 4.75$.

77 Reduced Sensitivity by Design

Can measure gain in term of Γ . Gains can be larger or small than in example, or no gain.

Design strategies. Moreover, can study theoretical situations. Try to gain understand whether and when strategies reduce sensitivity to hidden bias.

Could look at 'power'. Power of a sensitivity analysis for specified Γ is the probability of an upper bound on the significance level of 0.05 or less.

Feasible but untidy. Power calculations of this sort depend on many things. Nicer to have something analogous to Pitman efficiency.

78 Design Sensitivity

Relative performance in large samples. Limiting value of Γ as sample size increases for competing strategies.

3 doses vs no doses. We used information on 3 dose levels vs ignoring doses in one data set.

With theory, We could make definitive comparisons of various strategies under various conditions.

79 Some Strategies

Dose response. Hill's suggestion: look for a dose-response relationship.

Coherence among multiple outcomes. Campbell's suggestion of multiple operationalism.

Clinical trials idea: Just two treatments that are as different as possible. (Peto, et al. 1976)

80 Simple setting

Structure. S matched sets, match one treated to k controls, $k + 1$ in each set.

Distribution. p -dimensional multivariate Normal responses, with errors that have constant intercorrelation ρ . (Campbell's multiple operationalism)

Treatment effects. Proportional to dose, multiplier β . (Hill's dose-response)

Statistics. Stratified Wilcoxon rank sum statistic applied to weighted combination of responses (Dawson and Lagaokos 1993). Same weighted by doses.

81 Doses

Three patterns.

$\left(\frac{1}{2}, 1, \frac{3}{2}\right)$ produces a dose response

$(1, 1, 1)$ same average dose, no dose response

$\left(\frac{3}{2}, \frac{3}{2}, \frac{3}{2}\right)$ different as possible, no dose response

Effect size. In computations shown here, effect at dose 1 is half a standard deviation.

82 Design Sensitivity in a Simple Setting

Setting. Table describes three dose patterns, $p = 1$ or $p = 3$ coherent outcomes, with intercorrelation $\rho = 0$ or $\rho = \frac{1}{2}$, with $k = 5$ controls matched to each treated subject. Dose 1 is half standard deviation effect, and effect proportional to dose.

Design sensitivity. Table gives limiting value of Γ as $S \rightarrow \infty$. Bigger is better.

Doses	p	$\rho = 0$	$\rho = \frac{1}{2}$
$(\frac{1}{2}, 1, \frac{3}{2})$	1	2.97	2.97
	3	6.40	3.75
$(1, 1, 1)$	1	2.58	2.58
	3	5.05	3.16
$(\frac{3}{2}, \frac{3}{2}, \frac{3}{2})$	1	4.06	4.06
	3	11.74	5.56

83 Observations

Summary: Strategies can strongly affect sensitivity to hidden bias.

Specifics: Larger doses does most, then coherence among outcomes when error correlations are low, then dose response.

Traditional qualitative advice: Seems correct so far as it goes, but the relative (i.e. quantitative) importance and effectiveness of different strategies is only beginning to be understood.

84 Summary

Causal effects. Comparison of potential outcomes under competing treatments — not jointly observable (Neyman 1923, Rubin 1974).

Randomized experiments. Permit inference about the effects caused by treatments (Fisher 1935).

Observational studies: Adjustments. Without randomization, adjustments are required. Straightforward for observed covariates, but there might be important covariates that you did not observe.

Observational studies: Sensitivity analysis. What would unobserved covariates have to be like to alter conclusions? (Cornfield, et al.)

Observational studies: Pattern specificity. Reducing sensitivity to hidden bias. (Campbell 1988, Hill 1965)