

# Grant Writing Boot Camp Cross-Sectional and Cohort Studies

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#### Study Design I – Cross Sectional Studies

### **Outline**

- Overview of Epidemiological Study Designs
- Descriptive Studies
- Cross-Sectional
  - Design; Analytical approach; Strengths; Weakness
  - Random error, Systematic error, and Confounding
- Observational Studies
  - Cohort Study
    - Design; Analytical
  - Case-Control Study (Dr. Zhao)











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# Cross-Sectional Studies – When to use

**Cross Sectional** 

- Goal is to describe variables and their distribution pattern
  - Example: National Health and Nutrition Examination Survey (NHANES study)
    - Sample designed to represent the US population -- interviewed and examined
    - Each cross-sectional study -- major source of information on health and habits of the US population (e.g., prevalence of smoking in various demographic groups)
- Can be used to examine associations
  - Which variables to label as predictors and outcome depends on the investigator hypothesis
  - Temporal relationship usually cannot be established

Cross Sectional				<b>L'HSC</b>
Cross-Sectional Studies Analytical Approach		Outco	ome	Total
	Exposure	Present	Absent	
	Yes	а	b	a + b
	Νο	С	d	c + d
	Total	a + c	b + d	a + b + c + d
Pre	valence <sub>total</sub> = ((a+c	c) / (a+b+c+d)) x ı / (a+b)) x 10 <sup>n</sup>	10 <sup>n</sup>	
Prev	alence non-exposed	= (c / (c+d)) x 10	ŋn	
Mea asso	sure of ociation	revalence Rat	io = P <sub>exposed</sub>	/P non-exposed

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#### Example of a cross-sectional study

### Cross-Sectional Studies: Example 7.1 Analytical Approach

Exposure to Smoking Depictions in Movies: Its Association With Established Adolescent Smoking

Sargent et al. (2007) sought to determine whether exposure to movies in which the actors smoke is associated with smoking initiation. The steps in performing the study were to:

I. Define selection criteria and recruit the population sample. The investigators did a **random**-digit-dial survey of 6,522 U.S. children aged **10 to 14 years**.

2. Measure the predictor and outcome variables. They quantified smoking in **532 popular movies** and for each subject asked which of a <u>randomly selected subset of 50 movies</u> they had seen (predictor variable). Subjects were also asked about a variety of covariates such as age, race, gender, parent smoking and education, sensation-seeking (e.g., "I like to do dangerous things"), and self-esteem (e.g., "I wish I were someone else"). The outcome variable was whether the child had ever tried smoking a cigarette.

3. Results and conclusion: 1) Overall, 10% of the population had tried smoking. Quartile (Q) of movie smoking exposure was significantly associated with the prevalence of smoking initiation; 2) This association did not differ significantly by race/ethnicity or census region. 3) After controlling for sociodemographics, friend/sibling/parent smoking, school performance, personality characteristics, and parenting style, the adjusted odds ratio for having tried smoking were 1.7 (95% confidence interval [CI]: 1.1, 2.7) for Q2, 1.8 (95% CI: 1.2, 2.9) for Q3, and 2.6 (95% CI: 1.7, 4.1) for Q4 compared with adolescents in Q1. 4) The covariate-adjusted attributable fraction was 0.38 (95% CI: 0.20, 0.56), suggesting that exposure to movie smoking is the primary independent risk factor for smoking initiation in US adolescents in this age group.



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## **Cross-Sectional Studies – Random and Systematic Error**

Random error – by chance – may affect precision in both outcome and exposure measures (frequencies or relationship) – solution: increase the sample size

Systematic error (bias) -- can happen in design, conduct, analysis or reporting of a study

Selection bias:

**Cross Sectional** 

Sampling Bias – Not using representative sample of the source population Incidence-Prevalence Bias – Inclusion of prevalent cases in a study (overrepresentation of those who have lived the longest)

#### Information bias:

Recall bias – use of self-reporting – differences in accuracy or completeness of recall of past events/experiences

More error details refer to :https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7318122/



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#### **Cross Sectional**

### **Cross-Sectional Studies**

### Strengths

- No waiting for the outcome to occur
  - · Fast; Inexpensive; No loss of follow-up
- Can be a first step in a cohort or a clinical trial

#### Weakness

- Impractical for studies of rare diseases (if collecting data from the general population)
- Not suited for diseases of short-duration
- Difficult to establish causal relationship



























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### **Case-Control Studies**

### **Selection of Controls**

- One of the major challenges in a case-control studies
- Controls should be similar to the cases in all respects other than having the disease (event) in question
- Controls should be representative of all persons without the disease in the population from which the cases are selected





Case-Control Studies – Strengths
Efficient for rare outcomes
Require fewer participants than cohort studies, which means that more expensive and rigorous tests can be used
There is no problem with losses to follow-up

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### **Case-Control Studies – Weakness**

- · Cannot estimate the incidence or prevalence of the diseases
- Information on the exposure or risk factor is obtained <u>after</u> the occurrence of disease, so there is not a clear way to estimate the time between exposure and start of disease
- Only one outcome can be studied
- Susceptibility to bias



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### **Case-Control Studies**

### Confounding

- Matching
  - To increase the comparability of cases and controls by controlling a confounding variable in the study design: controls are matched to cases based on having the same value of the confounder (e.g. age)
  - $\circ~$  More than one control may be matched to each case





Nes	ed Case-Control Studies
• St	engths
•	Useful for costly measurements on specimens that have been archived at the beginning of the study
•	Avoids the potential biases of conventional case–control studies that cannot make measurements on fatal cases and that draw cases and controls from different populations
•	Retains the advantages of cohort studies collect predictor variables before the outcomes have happened
• We	akness
•	Same as other observational studies including potential for confounding

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## **Considerations in Grant Application**

#### Bias

- 1) Study design: e.g., nested case-control study; case or control selection; multiple control groups
- 2) Data collection: e.g., staff training, blinded to case and control status; additional data collection for evaluating potential bias
- 3) Data analysis plan: e.g., analyze additional data

#### Confounding

- 1) Study design (study population): e.g., matched study design
- 2) Data collection: e.g., collect potential confounding factors
- 3) Data analysis plan: e.g., stratification analysis; multivariable modeling

