

Showing Cause, Introduction to Study Design

Principles of Epidemiology



Theories of Disease Causation

- **Supernatural Theories**
- **Hippocratic Theory**
- **Miasma**
- **Theory of Contagion**
- **Germ Theory (cause shown via Henle-Koch postulates)**
- **Classic Epidemiologic Theory**
- **Multicausality and Webs of Causation (cause shown via Hill's postulates)**



Henle-Koch Postulates

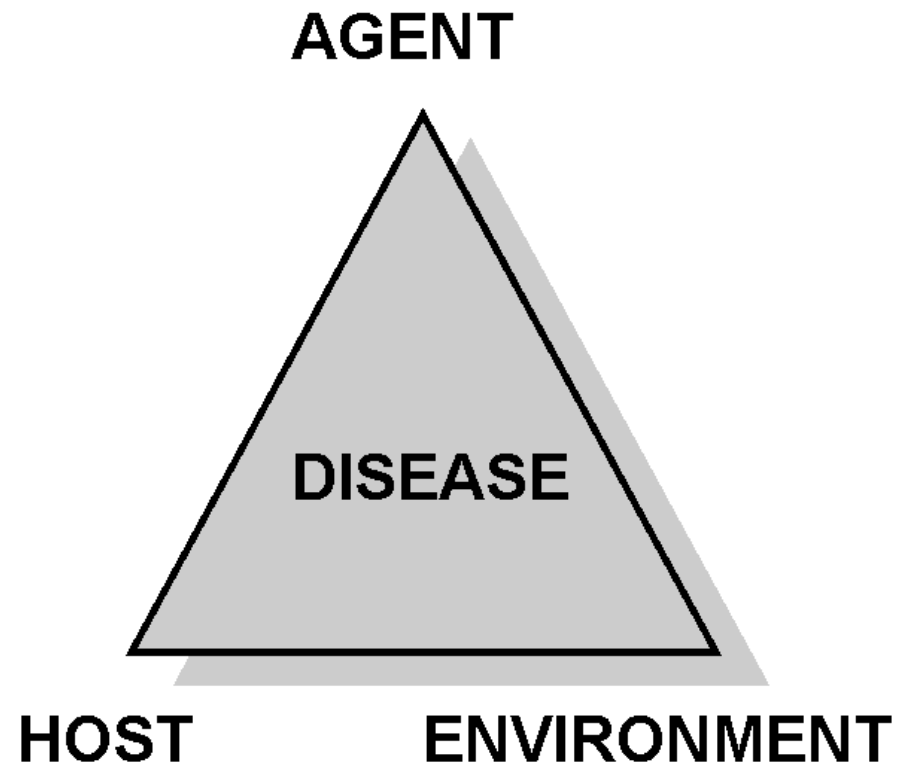
■ Sometimes called “pure determinism”

1. The agent is present in every case of the disease
2. It does not occur in any other disease as a chance or nonpathogenic parasite (one agent one disease)
3. It can be isolated and if exposed to healthy subjects will cause the related disease

Epidemiologic Triad

Disease is the result of forces within a dynamic system consisting of:

- ◆ **agent of infection**
- ◆ **host**
- ◆ **environment**



Classic Epidemiologic Theory

- **Agents**
 - Living organisms
 - Exogenous chemicals
 - Genetic traits
 - Psychological factors and stress
 - Nutritive elements
 - Endogenous chemicals
 - Physical forces
- **Agents have characteristics such as infectivity, pathogenicity and virulence (ability to cause serious disease)**
 - They may be transmitted to hosts via vectors

Classic Epidemiologic Theory (cont.)

- **Host factors:**

- Immunity and immunologic response
- Host behavior

- **Environmental factors:**

- Physical environment (heat, cold, moisture)
- Biologic environment (flora, fauna)
- Social environment (economic, political, culture)

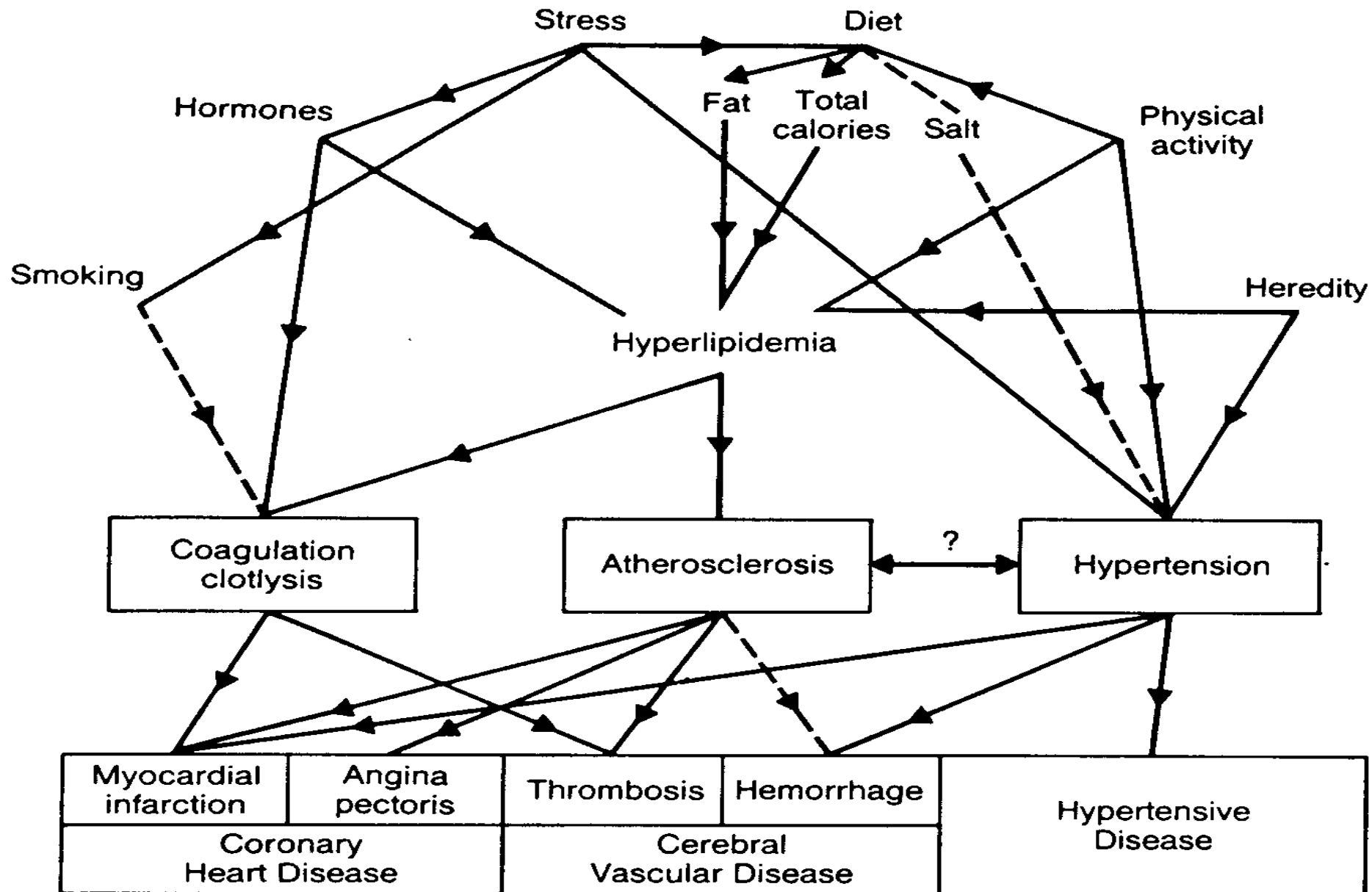
Hill's Postulates

1. **Strength of Association** – the stronger the association, the less likely the relationship is due to chance or a confounding variable
2. **Consistency of the Observed Association** – has the association been observed by different persons, in different places, circumstances, and times? (similar to the replication of laboratory experiments)
3. **Specificity** – if an association is limited to specific persons, sites and types of disease, and if there is no association between the exposure and other modes of dying, then the relationship supports causation
4. **Temporality** – the exposure of interest must precede the outcome by a period of time consistent with any proposed biologic mechanism
5. **Biologic Gradient** – there is a gradient of risk associated with the degree of exposure (dose-response relationship)

Hill's Postulates (cont)

6. **Biologic Plausibility** – there is a known or postulated mechanism by which the exposure might reasonably alter the risk of developing the disease
7. **Coherence** – the observed data should not conflict with known facts about the natural history and biology of the disease
8. **Experiment** – the strongest support for causation may be obtained through controlled experiments (clinical trials, intervention studies, animal experiments)
9. **Analogy** – in some cases, it is fair to judge cause-effect relationships by analogy – “With the effects of thalidomide and rubella before us, it is fair to accept slighter but similar evidence with another drug or another viral disease in pregnancy”

Web of Causation for the Major Cardiovascular Diseases



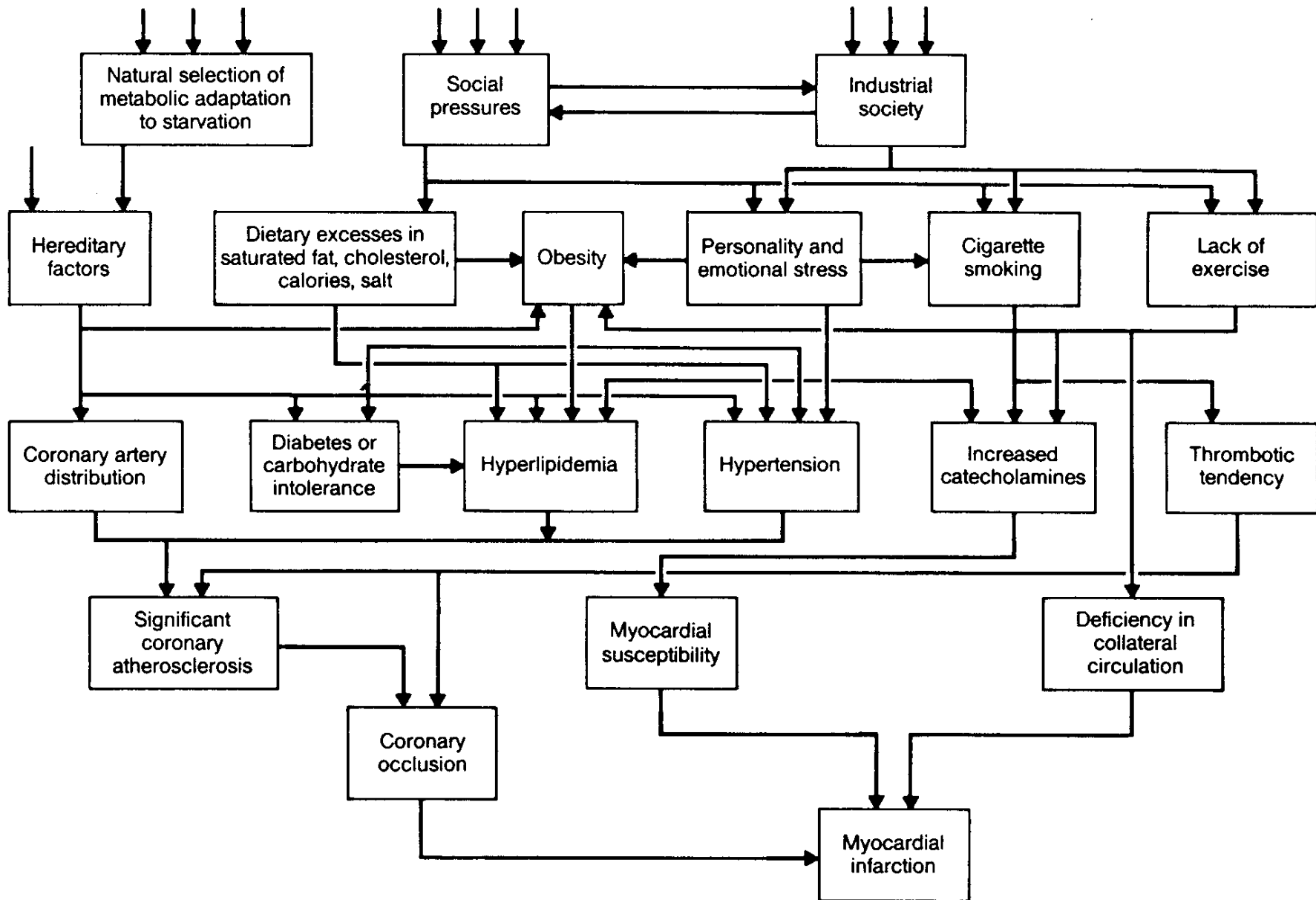


Figure 3.4 A Web of Causation for Myocardial Infarction. (Source: Friedman, G. D. (1980) *Primer of epidemiology*. New York: McGraw-Hill p. 4. Reprinted with permission.)



Causal Relationships

- A causal pathway may be direct or indirect
- In **direct causation**, A causes B without intermediate effects
- In **indirect causation**, A causes B, but with intermediate effects
- In human biology, intermediate steps are virtually always present in any causal process

Types of Causal Relationships

- **Necessary and sufficient** – without the factor, disease never develops
 - With the factor, disease always develops (this situation rarely occurs)
- **Necessary but not sufficient** – the factor in and of itself is not enough to cause disease
 - Multiple factors are required, usually in a specific temporal sequence (such as carcinogenesis)
- **Sufficient but not necessary** – the factor alone can cause disease, but so can other factors in its absence
 - Benzene or radiation can cause leukemia without the presence of the other
- **Neither sufficient nor necessary** – the factor cannot cause disease on its own, nor is it the only factor that can cause that disease
 - This is the probable model for chronic disease relationships



Factors in Causation

- All may be necessary but rarely sufficient to cause a particular disease or state
 - Predisposing – age, sex or previous illness may create a state of susceptibility to a disease agent
 - Enabling – low income, poor nutrition, bad housing or inadequate medical care may favor the development of disease
 - Conversely, circumstances that assist in recovery or in health maintenance may be enabling
 - Precipitating – exposure to a disease or noxious agent
 - Reinforcing – repeated exposure or undue work or stress may aggravate an established disease or state

Comparing Rules of Evidence

Criminal Law	Causation
Criminal present at scene of crime	Agent present in the disease
Premeditation	Causal events precede onset of disease
Accessories involved in the crime	Cofactors and/or multiple causality involved
Severity of crime related to state of victim	Susceptibility and host response determine severity
Motivation – there must be gain to the criminal	The role of the agent in the disease must make biologic and common sense
No other suspect could have committed the crime	No other agent could have caused the disease under the circumstances given
Proof of guilt must be established beyond a reasonable doubt	Proof of causation must be established beyond reasonable doubt or role of chance



Study Designs

Means to assess possible causes by gathering and analyzing evidence



Types of Study Designs

- Descriptive studies (to generate hypotheses)
 - Case-reports
 - Cross-sectional studies (prevalence studies) measure exposure and disease at the same time
 - Ecological studies (correlational studies) use group data rather than data on individuals
 - These data cannot be used to assess individual risk
 - To do this is to commit *ecological fallacy*



Types of Study Designs (cont.)

- **Analytic studies (to test hypotheses)**
 - **Experimental studies**
 - **Clinical trials**
 - **Field trials**
 - **Intervention studies**
 - **Observational studies**
 - **Case-control studies**
 - **Cohort studies**



The Key to Study Design

- The key to any epidemiologic study is in the definition of what constitutes a case and what constitutes exposure
- Definitions must be exclusive, categorical
- Failure to effectively define a case may lead to misclassification bias



Sources of Error in Epidemiologic Studies

- **Misclassification – wrongful classification of status for either disease or exposure**
- **Random variation - chance**



Sources or Error in Epidemiologic Studies

- **Bias** – systematic preferences built into the study design
- **Confounding** – occurs when a variable is included in the study design that is related to both the outcome of interest and the exposure, leading to false conclusions
 - Example: gambling and lung cancer
- **Effect modification** – occurs when the magnitude of the association between the outcome of interest and the exposure differ according to the level of a third variable
 - The effect may be to nullify or heighten the association
 - Example: gender and hip fracture modified by age



Contingency Tables

The findings for most epidemiologic studies can be presented in the 2x2 table

	Disease		
	Yes	No	Total
Exposure			
Yes	a	b	a+b
No	c	d	c+d
Total	a+c	b+d	a+b+c+d

Measures of Association from the 2x2 Table

Cohort Study: the outcome measure is the *relative risk* (or risk ratio or rate ratio)

- In cohort studies you begin with the exposure of interest and then determine the rate of developing disease
- RR measures the likelihood of getting the disease if you are exposed relative to those who are unexposed
 - RR = incidence in the exposed/incidence in the unexposed

$$RR = \frac{a/(a+b)}{c/(c+d)}$$

Measures of Association from the 2X2 Table

Case-control study: the outcome measure is an estimate of the relative risk or the ***odds ratio*** (relative odds)

- In a case-control study, you begin with disease status and then estimate exposure
 - RR is estimated because patients are selected on disease status and we cannot calculate incidence based on exposure
 - The estimate is the odds ratio (OR) or the likelihood of having the exposure if you have the disease relative to those who do not have the disease

$$\sim \text{RR} = \text{OR} = \frac{a/c}{b/d} = \frac{ad}{bc}$$

Attributable Risk or Risk Difference

- In a *cohort study*, we may want to know the risk of disease attributable to the exposure in the exposed group, that is, the difference between the incidence of disease in the exposed and unexposed groups (**excess risk**)

$$AR = a/(a+b) - c/(c+d)$$

AR = 0: No association between exposure and disease

AR > 0: Excess risk attributable to the exposure

AR < 0: The exposure carries a protective effect

Attributable Risk Percent

- In a *cohort study*, we may want to know the proportion of the disease that could be prevented by eliminating the exposure in the exposed group (**attributable fraction or etiologic fraction**)

$$AR\% = AR/[a/(a+b)] \times 100$$

If the exposure is preventive,
calculate the **preventive fraction**

Population Attributable Risk

- In a *cohort study*, we may want to know the risk of disease attributable to exposure in the total study population or the difference between the incidence of disease in the total study population and that of the unexposed group

$$\text{PAR} = (a+c)/(a+b+c+d) - c/(c+d)$$

To estimate the PAR for a population beyond the study group you must know the prevalence of disease in the total population

Population Attributable Risk Percent

- In a *cohort study*, we may want to know the proportion of the disease that could be prevented by eliminating the exposure in the entire study population

$$\text{PAR}\% = \text{PAR} / [(a+c)/(a+b+c+d)] \times 100$$

Summary of Attributable Risk Calculations

	In exposed group	In total population
Incidence attributable to exposure	$I_e - I_n$ AR	$I_p - I_n$ PAR
Proportion of incidence attributable to exposure	$\frac{I_e - I_n}{I_e} \times 100$ AR%	$\frac{I_p - I_n}{I_p} \times 100$ PAR%

Comparing Relative Risks

Age-Adjusted Death Rates per 100,000 for Male British Physicians

	Smokers	Non-smokers
Lung cancer	140	10
CHD	669	413

Source: Doll and Peto. Mortality in relation to smoking: Twenty years' observations on male British doctors. BMJ 1976;2:1525-36

Relative risk (relative risk, risk ratio) I_e/I_n : LC = 14.0; CHD = 1.6

Smokers are **14** times as likely as non-smokers to develop LC

Smokers are **1.6** times as likely as non-smokers to develop CHD

Smoking is a stronger risk factor for lung cancer than for CHD

Comparing Attributable Risks

Age-Adjusted Death Rates per 100,000 for Male British Physicians

	Smokers	Non-smokers
Lung cancer	140	10
CHD	669	413

Source: Doll and Peto. Mortality in relation to smoking: Twenty years' observations on male British doctors. BMJ 1976;2:1525-36

Attributable risk (risk difference, etiologic fraction) $I_e - I_n$:

LC = 130; CHD = 256

The excess of lung cancer attributable to smoking is **130**
per 100,000

The excess of
CHD attributable to smoking is **256** per 100,000

If smoking is causal, eliminating cigarettes would save more smokers from CHD than from LC

Comparing Attributable Risk Percents

Age-Adjusted Death Rates per 100,000 for Male British Physicians

	Smokers	Non-smokers
Lung cancer	140	10
CHD	669	413

Source: Doll and Peto. Mortality in relation to smoking: Twenty years' observations on male British doctors. BMJ 1976;2:1525-36

Attributable Risk % = $[(I_e - I_n) / I_e] \times 100$: **LC = 92%; CHD = 38%**

About **92%** of LC could be eliminated if the smokers in this study did not smoke

About **38%** of CHD could be eliminated if the smokers in this study did not smoke

If smoking is causal, eliminating cigarettes would save double the proportion of smokers from LC than CHD