

Evidence Based Dentistry

# Clinical problems and choice of study designs

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**Manipulation  
with intervention**

**Experimental  
study**

**Non-experimental  
study / observational**

**Random  
allocation**

**Sampling according  
to exposition  
characteristics**

**Sampling according  
to (case) effect  
characteristics**

**Experimental  
study (RCT)**

**Quasi-  
experimental  
study (CCT)**

**Case series /  
cohort study**

**Case-control study**

# Clinical trial terminology - tower of Bable?

analytical study

case control study (89)

case serie

case study, case report

cause-effect study

clinical trial (79)

cohort study (89)

cohort study with historical controls

controlled clinical trial (95)

cross-sectional study (89)

descriptive study

diagnostic meta-analysis

diagnostic study

double blind randomized therapeutical trial with cross-over design

ecological study

etiological study

experimental study

explorative study

feasibility study (79)

follow-up study (67)

historical cohort study

incidence study

intervention study

longitudinal study (79)

N=1 trial

non-randomized trial with

contemporaneous controls

non-randomized trial with

historical controls

observational study

prospective cohort study

prospective follow-up study, observational or experimental

prospective study (67)

quasi-experimental study

randomized clinical trial, RTC

randomized controlled trial, RCT (89)

retrospective cohort study

retrospective follow-up study

retrospective study (67)

surveillance study

survey, descriptive survey

therapeutic meta-analysis

trohoc study

Descriptions of clinical studies can be reduced to three questions

1. Study objective?

Descriptive, no comparison conducted  
Comparison as process research  
Comparison as cause-effect research

2. Procedure, intervention?

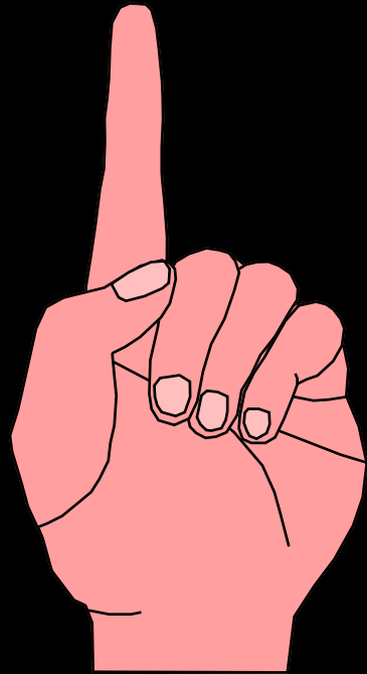
Experimental allocation of procedure  
Survey

3. Data collection?

Retrospective  
Cross-sectional  
Prospective / Cohort / Longitudinal

## Clinical study designs (MESH terms):

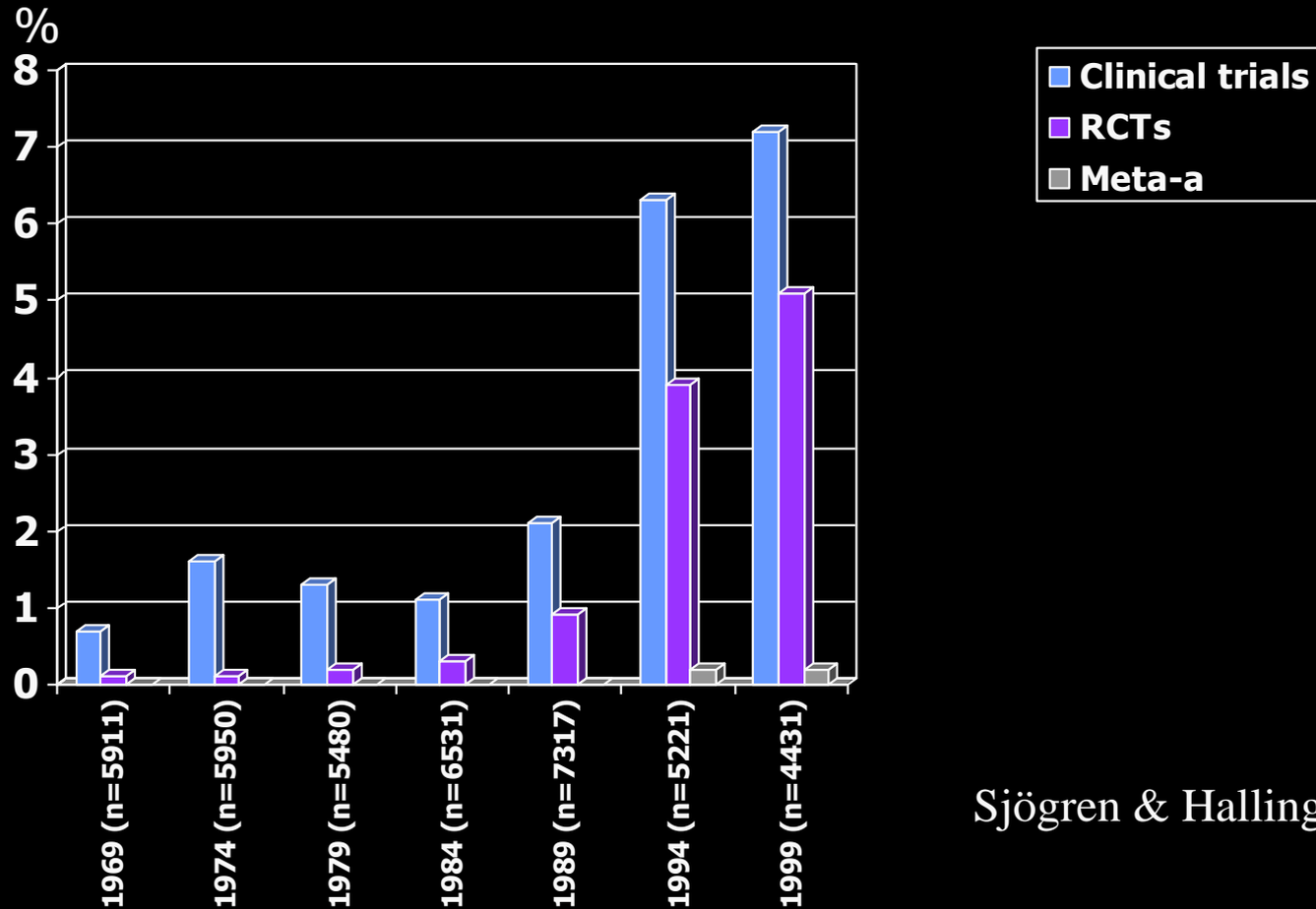
- (Case study/series)
- Case-Control Study
- Cohort Study
- Cross-Sectional Survey
- Randomised Controlled Trial



Most publications  
in the dental  
literature are not  
RCTs

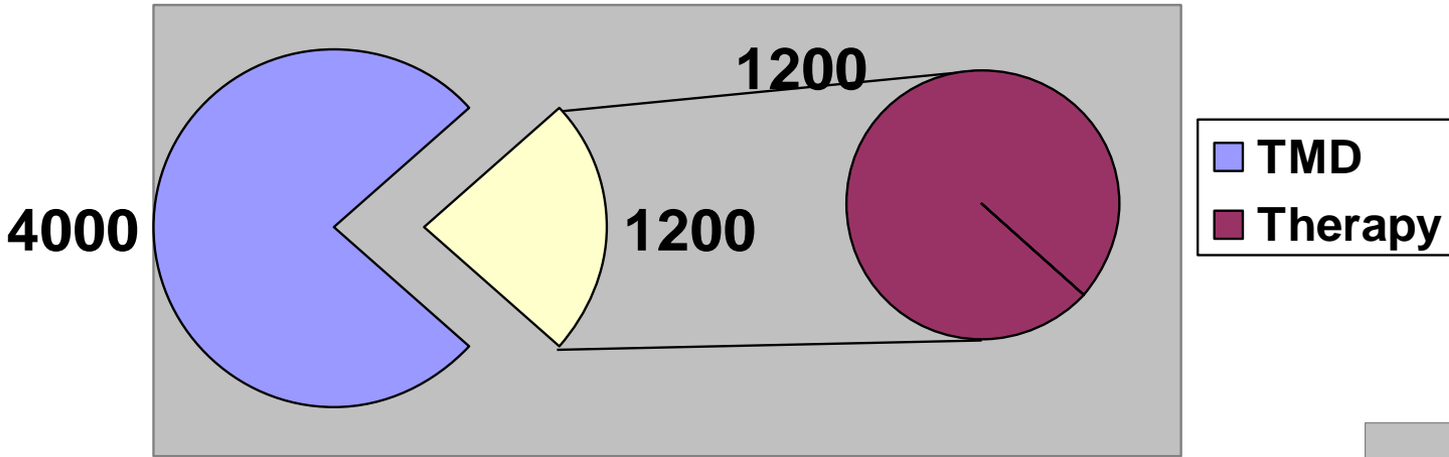
# Dental research - medline

- Medline search 1969-99
  - 7% clinical research, 5% RCT

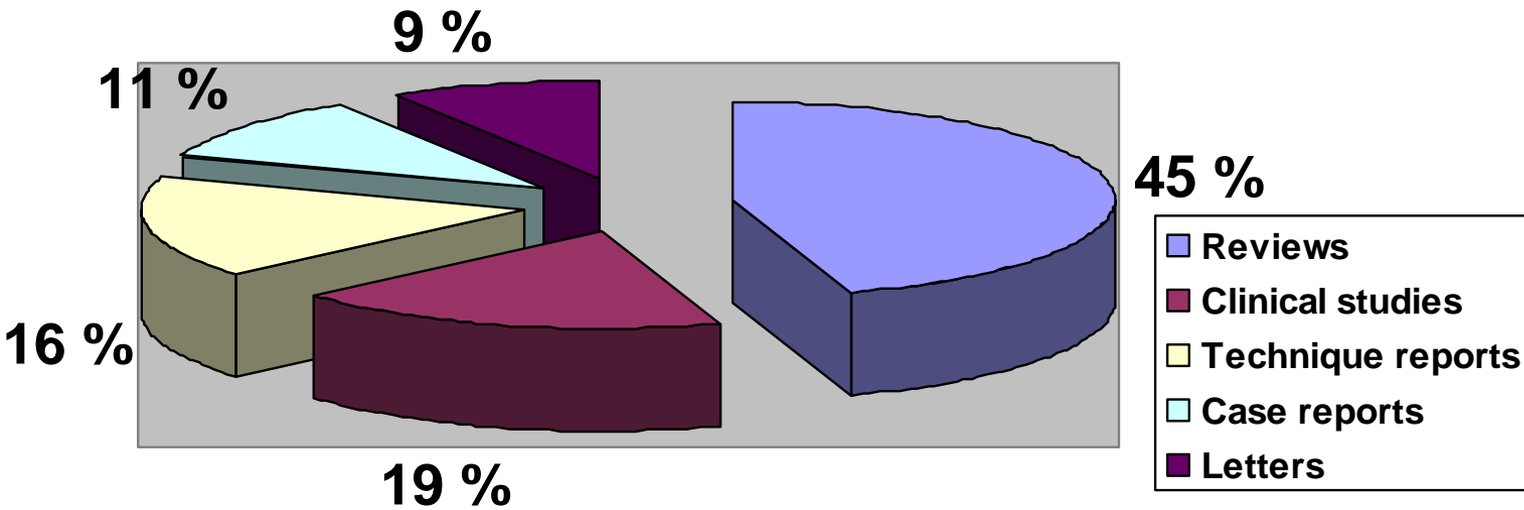
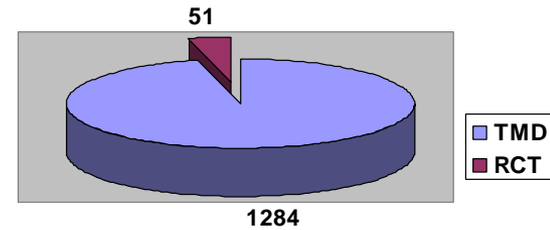


Sjögren & Halling, 2000

# TMD studies 1980-92

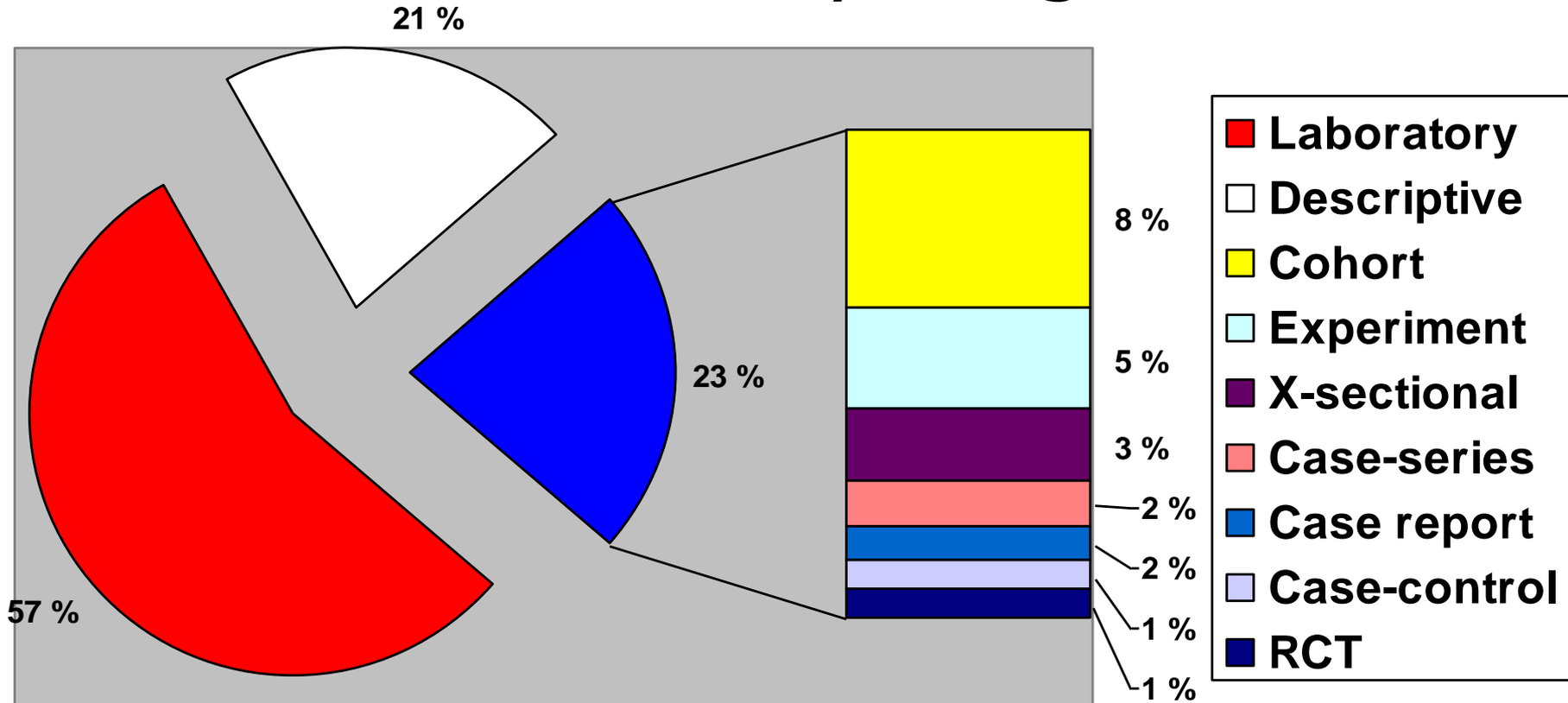


## RCT studies

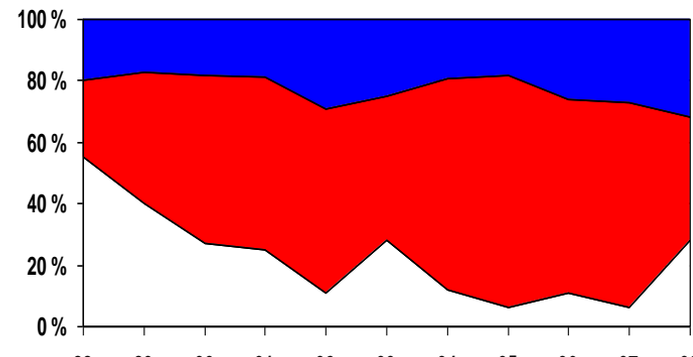


Antcak-Bouckoms, 1995

# Study design



Only 23% of the 726 papers that have been published in International Journal Prosthodontics describe in vivo study findings. (Jokstad, ICP, 1999)



# The central tasks of clinical work

## 1. Clinical findings:

How to properly gather the most relevant findings from the history and physical examination, and interpret these correctly?



## 2. Etiology:

How to identify causes for disease (including its iatrogenic forms) ?



# The central tasks of clinical work

## 3. Differential diagnosis:

When considering the possible causes of a patient's clinical problem, how to rank them by likelihood, seriousness and treatability ?

## 4. Diagnostic tests

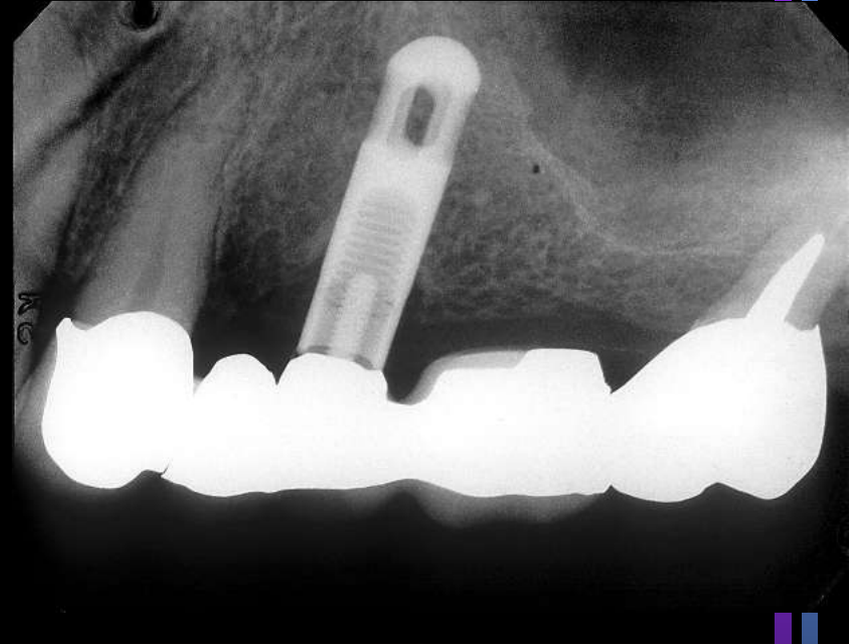
How to select and interpret diagnostic tests, in order to confirm or exclude a diagnosis, based on considering precision, accuracy, acceptability, expense, safety,



# The central tasks of clinical work

## 5. Prognosis:

How to estimate the patient's likely clinical course over time and anticipate likely complications?



## 6. Therapy:

How to select treatments to offer patients that do more good than harm and that are worth the efforts and costs of using them?



# The central tasks of clinical work

## 7. Prevention:

How to reduce the chance of disease by identifying and modifying risk factors and how do we diagnose disease early by screening?

## 8. Self-improvement:

How to keep up to date, improve our clinical skills and run a better, more efficient clinical practice?



**OXYFRESH vs OTHER LEADING MOUTHRINSES**

	DETERGENTS	CPG	DYES	ALCOHOL	SACCHARIN	SALICYLATE	FLUORIDE ENAMEL COLORS	FLUORIDE OILS	FLUORIDE GELS
<i>Oxyfresh</i>	NO	NO	NO	NO	NO	NO	YES	YES	YES
Listerine	YES		YES	YES	YES	YES	NO		NO
Scope	YES	YES	YES	YES	YES		NO	NO	NO
Act	YES	YES	YES		YES		NO	NO	NO
Clear Choice	YES	YES			YES		NO	NO	NO
Plax	YES		YES	YES	YES		NO	NO	NO
Oral B	YES	YES	YES		YES		NO	NO	NO
Viadent	YES			YES	YES		NO	NO	NO
Fluoriguard	YES		YES	YES	YES		NO		NO
Lavoris	YES		YES	YES	YES		NO	NO	NO
Cepacol	YES	YES	YES	YES	YES		NO	NO	NO
Peridex	YES		YES	YES	YES		NO	NO	NO

**ADDITIONS**      **BENEFITS**

# Appropriate Study Designs

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				★	★ ★
Therapy				★	★ ★
Prognosis				★ ★ ★	
Screening			★	★	★ ★
Views/beliefs perceptions	★ ★ ★				
Prevalence/hypothesis generation	★ ★ ★	★ ★ ★			

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆	☆☆
Therapy				☆	☆☆
Prognosis				☆☆☆	
Screening			☆	☆	☆☆
Views/beliefs perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

# Qualitative research

- Aim to make sense of, or interpret, phenomena in terms of the meanings people bring to them
- May define preliminary questions which can then be addressed in quantitative studies
- Address a clinical problem through a clearly formulated question and using more than one research method (triangulation)
- Analysis of qualitative data can and should be done using explicit, systematic, and reproducible methods

# Qualitative research methods -examples

- Documents - Study of documentary accounts of events
- Passive observation - Systematic watching of behaviour and talk in natural occurring settings
- Participant observation - Observation in which the researcher also occupies a role or part in the setting, in addition to observing
- In depth interviews - Face to face conversation with the purpose of exploring issues or topics in detail. Does not use preset questions, but is shaped by a defined set of topics
- Focus groups - Method of group interview which explicitly includes and uses the group interaction to generate data

# Cross-Sectional Survey

## Advantages

1. Cheap and simple
2. Ethically safe

## Disadvantages

1. Establishes association at most, not causality
2. Recall bias susceptibility
3. Confounders may be unequally distributed
4. Group sizes may be unequal

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆	☆☆
Therapy				☆	☆☆
Prognosis				☆☆☆	
Screening			☆	☆	☆☆
Views/beliefs perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

# Case-Control Studies

## Advantages:

1. Quick and cheap
2. Only feasible method for very rare disorders or those with long lag between exposure and outcome
3. Fewer individuals needed than cross-sectional studies

## Disadvantages:

1. Reliance on recall or records to determine exposure status
2. Confounders
3. selection of control groups is difficult

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆	☆☆
Therapy				☆	☆☆
Prognosis				☆☆☆	
Screening			☆	☆	☆☆
Views/beliefs perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

# Characteristics of a poor case-control study:

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆	☆☆
Therapy				☆	☆☆
Prognosis				☆☆☆	
Screening			☆	☆	☆☆
Views/beliefs perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

## Fail to:

- clearly define comparison groups
- and/or fail to measure exposures and outcomes in the same (preferably blinded), objective way in both cases and controls
- and/or fail to identify or appropriately control known confounders.

# Cohort Study

## Advantages:

1. Ethically safe
2. individuals can be matched
3. Can establish timing and directionality of events
4. Eligibility criteria and outcome assessments can be standardised
5. Administratively easier and cheaper than RCT

## Disadvantages:

1. Controls may be difficult to identify
2. Exposure may be linked to a hidden confounder
3. Blinding is difficult
4. Randomisation not present
5. For rare disease, large sample sizes or long follow-up necessary

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆☆	☆☆☆
Therapy				☆☆	☆☆☆
Prognosis				☆☆☆	
Screening			☆☆	☆☆	☆☆
Views/beliefs/perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

# Characteristics of a poor cohort study:

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆	☆☆
Therapy				☆	☆☆
Prognosis				☆☆☆	
Screening			☆	☆	☆☆
Views/beliefs/perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

Fail to :

- clearly define comparison groups and/or
- measure exposures and outcomes in the same (preferably blinded), objective way in both exposed and non-exposed individuals and/or
- identify or appropriately control known confounders and/or
- carry out a sufficiently long and complete follow-up of patients.

# Randomised Controlled Trial - RCT

## Advantages

1. Unbiased distribution of confounders
2. Blinding more likely
3. Randomisation facilitates statistical analysis

## Disadvantages

1. Size, time and money - Expensive!
2. Volunteer bias
3. Ethically problematic at times

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆	☆☆
Therapy				☆	☆☆
Prognosis				☆☆☆	
Screening			☆	☆	☆☆
Views/beliefs/perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

# Cohort & RCT Crossover Design

## Advantages

1. All individuals serve as own controls -> error variance is reduced -> reduced need of large sample size
2. All individuals receive treatment (at least some of the time)
3. Statistical tests assuming randomisation can be used
4. Blinding can be maintained

## Disadvantages

1. All individuals receive placebo or alternative treatment at some point
2. Washout period lengthy or unknown
3. Cannot be used for treatments with permanent effects

Scientific studies can be graded  
according to the  
theoretical possibility  
of an  
incorrect conclusion.

This is reflected by the  
design of the study.

...we will never know exact answers in science....

# Internal and external validity

**Internal validity:** extent to which systematic error (bias) is minimised in clinical trials

**External validity:** extent to which results of trials provide a correct basis for generalisation to other circumstances

# Internal validity - systematic bias

- Selection bias: biased allocation to comparison groups
- Performance bias: unequal provision of care apart from treatment under evaluation
- Detection bias: biased assessment of outcome
- Attrition bias: biased occurrence and handling of deviations from protocol and loss to follow up

# External validity

- Patients: age, sex, severity of disease and risk factors, co-morbidity
- Treatment regimens: dosage, timing and route of administration, type of treatment within a class of treatments, concomitant treatments
- Settings: level of care (primary to tertiary) and experience and specialisation of care provider
- Modalities of outcomes: type or definition of outcomes and duration of follow up

# Diagnostic tests, Differential diagnosis

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆	☆☆
Therapy				☆	☆☆
Prognosis				☆☆☆	
Screening			☆	☆	☆☆
Views/beliefs perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

- Clearly identified comparison groups, at least one of which is free of the target disorder
- Either an objective diagnostic standard/contemporary clinical diagnostic standard with reproducible criteria for any objectively interpreted component
- Interpretation of the test without knowledge of the diagnostic standard result
- Interpretation of the diagnostic standard without knowledge of the test result
- A statistical analysis consistent with study design

# Therapy / Prevention / Education

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆	☆☆
Therapy				☆	☆☆
Prognosis				☆☆☆	
Screening			☆	☆	☆☆
Views/beliefs perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

- Random allocation of the participants to the different interventions
- Outcome measures of known or probably clinical importance for at least 80 per cent of participants who entered the investigation
- A statistical analysis consistent with the study design.

# Prognosis

	Qualitative	Cross-Sectional	Case Control	Cohort	RCT
Diagnosis				☆☆	☆☆☆
Therapy				☆☆	☆☆☆
Prognosis				☆☆☆	
Screening			☆	☆☆	☆☆☆
Views/beliefs perceptions	☆☆☆				
Prevalence/hypothesis generation	☆☆☆	☆☆☆			

- An inception cohort of persons, all initially free of the outcome of interest
- Follow-up of at least 80 per cent of patients until the occurrence of either a major study criteria or the end of the study
- A statistical analysis consistent with the study design.

# Etiology - Harm - Causation

- Clearly identified comparison group for those at risk for, or having, the outcome of interest
- Masking of observers of outcomes to exposures
- Observers of exposures masked to outcomes for case-control studies and individuals masked to exposure for all other study designs
- A statistical analysis consistent with the study design.

# Critical Appraisal Criteria

Exists for studies focused on e.g. :

- therapy
- diagnosis
- screening
- harm
- prognosis
- causation of disease (etiology)
- quality of care
- economic analyses

# Three general questions

1. Is the study valid?
2. What are the results ?
3. Are the results relevant to my question / problem?

# 1. Is the Study Valid ?

- Is there a clear question?
- Most appropriate study design to answer the question?
- Was study conducted reliably?
- Can you follow what the authors did?

## 2. What are the results?

- Are the results presented in a clear and simple manner ?
- Is there a clear bottom line ?
- Are they clinically important ?

### 3. Are the results relevant to my question / problem ?

- Are the participants similar to my patients ?
- Is it realistic for me to apply the study methodology / - result to my patients ?