Transportability analyses in Sweden



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Overview

Negistry based randomized trials

➤ Why do a transportability/generalizability analysis in Swedish data?



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Registry based randomized trials (R-RCTs)



Sweden

- Sweden has a tax payer funded universal healthcare system
- Everyone that lives in Sweden has a personal number, which is used to capture data every time they use the healthcare system
- ➤ There are several mandatory national registers, such as population register, patient register, cause of death register, prescribed drug register etc.
- Also a selection (>100) healthcare quality registries that contain more detailed information on specific diseases. E.g., SWEDEHEART
- Can all be linked using the personal number



Pragmatic randomized trial that uses a clinical registry for one or several major functions for trial conduct and outcomes reporting



➤ When a clinician is collecting data for a healthcare quality registry, they are notified if the patient is eligible for an ongoing R-RCT



The registry helps to:

- **N** Identify eligible patients
- **N** Randomize
- Collect baseline and procedure characteristics
- ▲ Assist with collecting consent forms
- ► Identify clinical endpoints (there is no "active" follow up)



WHAT R-RCT'S CAN DO

Evaluate therapeutic options that are available in routine clinical care. We can find out what works best.

WHAT R-RCT'S CANNOT DO

Cannot design RCTs to experiment new pharmaceutical agents or medical devices.



Example

Thrombus Aspiration in ST- Elevation myocardial infarction in Scandinavia (TASTE trial):



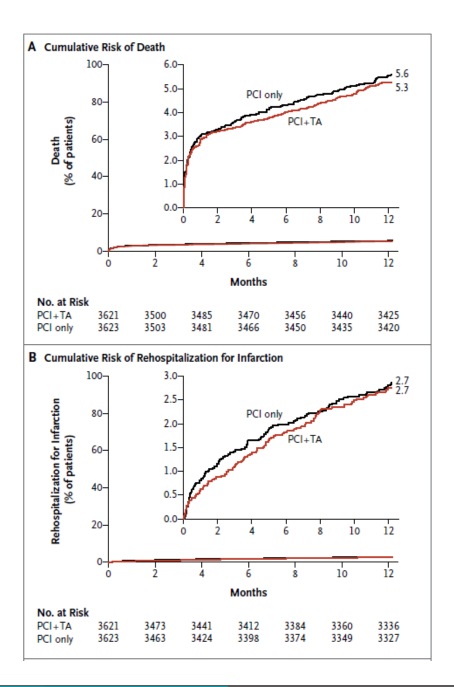
The NEW ENGLAND JOURNAL of MEDICINE

ORIGINAL ARTICLE

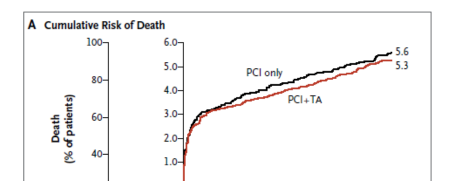
Outcomes 1 Year after Thrombus Aspiration for Myocardial Infarction

Bo Lagerqvist, M.D., Ph.D., Ole Fröbert, M.D., Ph.D., Göran K. Olivecrona, M.D., Ph.D., Thórarinn Gudnason, M.D., Ph.D., Michael Maeng, M.D., Ph.D., Patrik Alström, M.D., Jonas Andersson, M.D., Ph.D., Fredrik Calais, M.D., Jörg Carlsson, M.D., Ph.D., Olov Collste, M.D., Matthias Götberg, M.D., Ph.D., Peter Hårdhammar, M.D., Dan Ioanes, M.D., Anders Kallryd, M.D., Rickard Linder, M.D., Ph.D., Anders Lundin, M.D., Jacob Odenstedt, M.D., Elmir Omerovic, M.D., Ph.D., Verner Puskar, M.D., Tim Tödt, M.D., Ph.D., Eva Zelleroth, M.D., Ollie Östlund, Ph.D., and Stefan K. James, M.D., Ph.D.

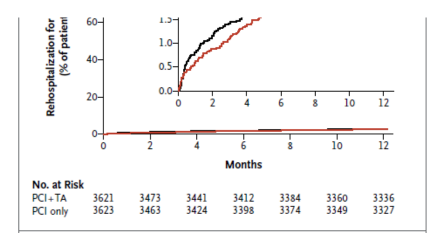








NO DIFFERENCE IN RISK OF DEATH OR MOCARDIAL INFARCTION BY 1 YEAR BETWEEN THOSE RANDOMIZED TO THROMBUS ASPIRATION OR NO THROMBUS ASPIRATION





TASTE used the SWEDHEART registry to:

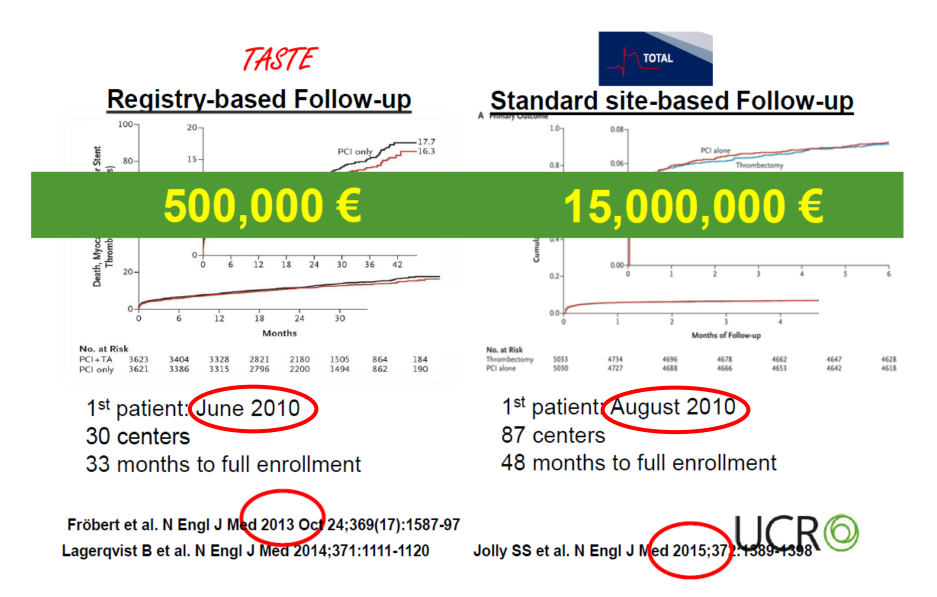
- Identify eligible patients that had an MI and underwent PCI
- Nandomize individuals to thrombus aspiration or no thrombus aspiration
- Collect baseline characteristics
- ▲ Assist with collecting consent forms
- Identify clinical endpoints of death and myocardial infarction



Embedding the trial within a registry meant TASTE was

CHEAP & QUICK





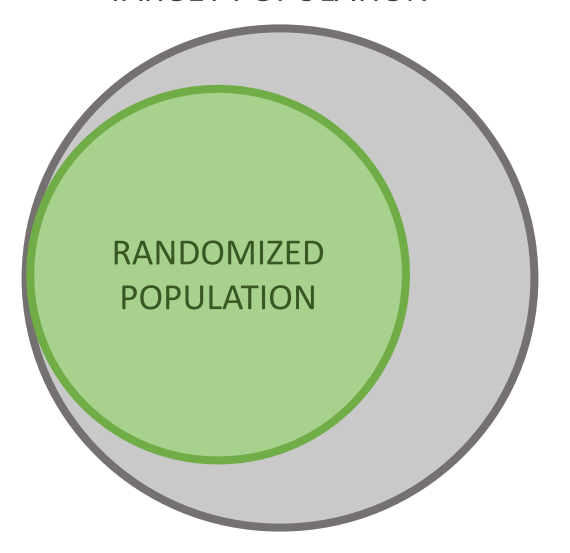
REF: Stefan James

Why undertake a generalizability/transportability analysis in Sweden?

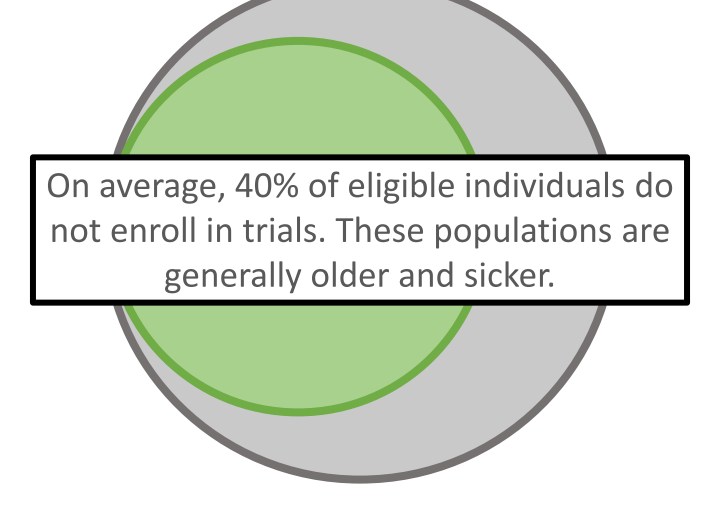


There remain questions around external validity when we undertake R-RCTs











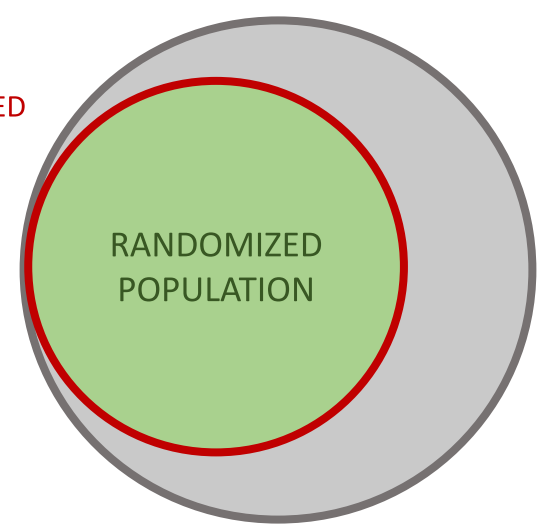
If the characteristic differences of those that do and do not enroll in the trial are also modifiers of the treatment effect

Then, we may not have external validity and our treatment effect may differ between the trial and target populations



TRADITIONAL RCT

BASELINE DATA COLLECTED IN RANDOMIZED POPULATION





TRADITIONAL RCT

BASELINE DATA COLLECTED IN RANDOMIZED POPULATION



R-RCT
BASELINE DATA COLLECTED
IN TARGET POULATION

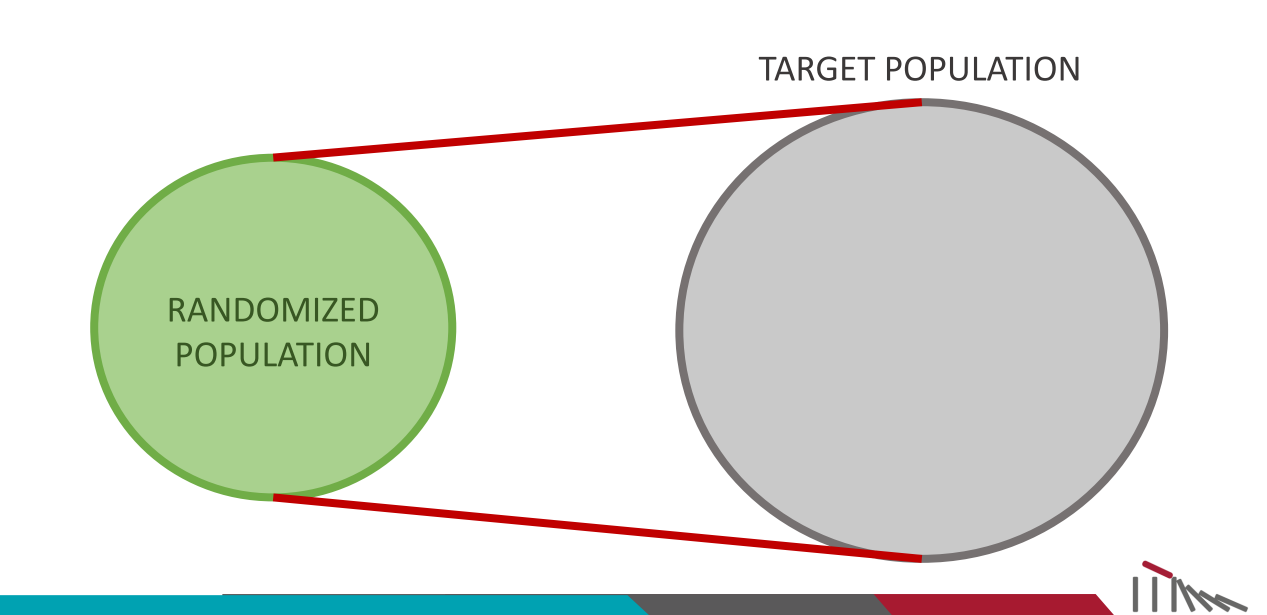


When a trial is embedded within a registry:

- Note that is a second control of the standard data collection for the registry
- The data generating mechanism is the same for everyone, regardless of if they take part in the trial
- Ne can get **the same** baseline data for individuals that were and were not randomized



TRANPORTABILITY ANALYIS - TRADITIONAL RCT



TRANPORTABILITY ANALYIS - TRADITIONAL RCT

TARGET POPULATION

RANDOMIZED POPULATION

Covariate data collected using different mechanisms



TRANPORTABILITY ANALYIS - TRADITIONAL RCT

TARGET POPLII ATION

We make an assumption that the covariates from the trial and the target population are measuring the same thing.

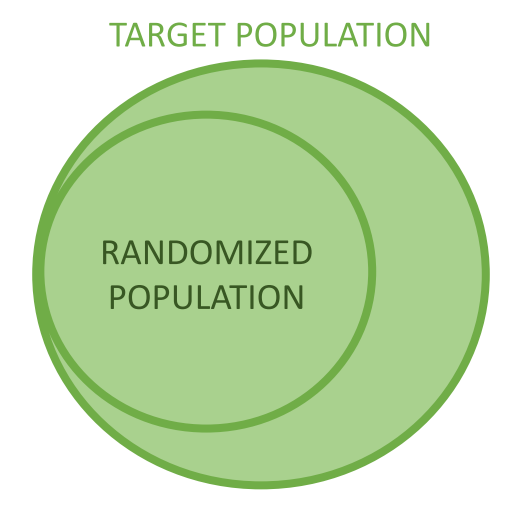
But, if you are using data from different countries/healthcare systems they may not be

e.g., there may be different diagnosis or prescribing thresholds



TRANSPORTABILITY ANALYSIS - R-RCT

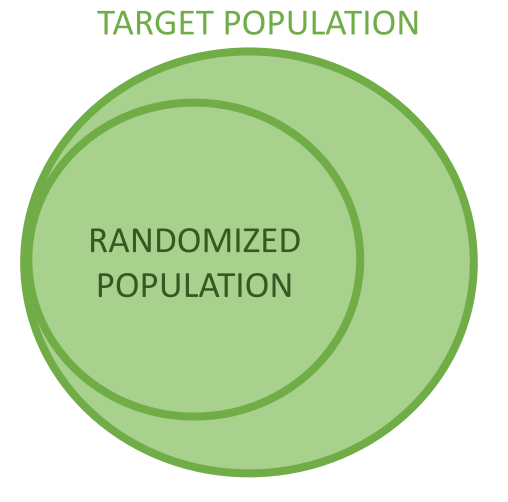
As always, we use baseline covariate data on the potential effect modifiers to transport effect estimates from the randomized to the target population





TRANSPORTABILITY ANALYSIS - R-RCT

As always, we use baseline covariate data on the potential effect modifiers to transport effect estimates from the randomized to the target population



As data from both the trial and target population are collected from the underling healthcare registry, data are collected in the same way.

We don't have to make assumptions that the trial data and observational data are collecting the same info, because THEY ARE



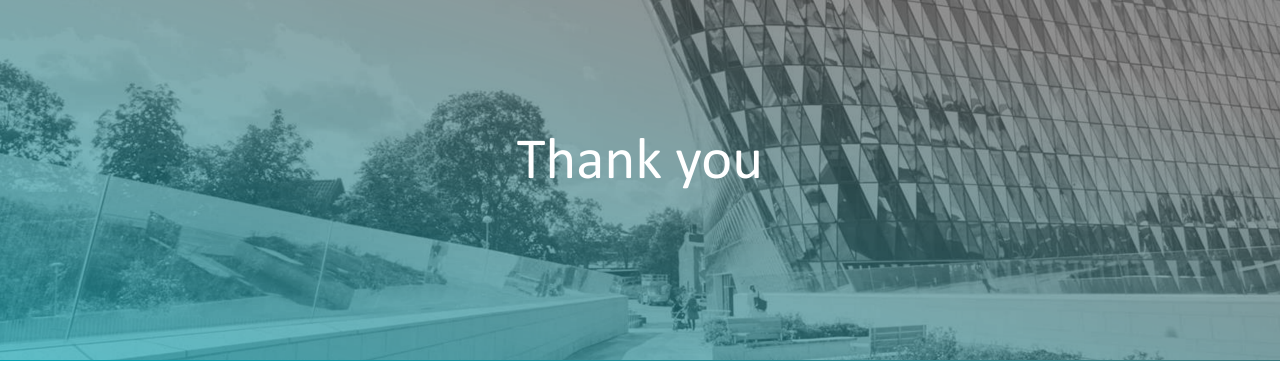
Why undertake a transportability/generalizability analysis in Sweden?

- ➤ There are currently a lot of R-RCTs being undertaken in Sweden (cardiology, cancer, respiratory diseases etc.)
- But these trials still cannot experiment on everyone eligible for the treatments under study – our target population
- ➤ We can use data from the SAME registries and perform transportability analyses to estimate the effects in those not included in the R-RCTs
- Note that are collected from the same registry, we do not have to make assumptions about the data generating mechanisms











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