APAC Community Call

Incidence and Prevalence

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Agenda

• OHDSI News
• Incidence and Prevalence by Patrick Ryan
Patrick Ryan, PhD is Vice President, Observational Health Data Analytics at Janssen Research and Development, where he is leading efforts to develop and apply analysis methods to better understand the real-world effects of medical products. He is an original collaborator in Observational Health Data Sciences and Informatics (OHDSI), a multi-stakeholder, interdisciplinary collaborative to create open-source solutions that bring out the value of observational health data through large-scale analytics. He served as a principal investigator of the Observational Medical Outcomes Partnership (OMOP), a public-private partnership chaired by the Food and Drug Administration, where he led methodological research to assess the appropriate use of observational health care data to identify and evaluate drug safety issues.
Incidence and prevalence: Concepts and implementation in OHDSI

Patrick Ryan
Incidence Rate = \frac{\text{# of new outcomes within a defined time-at-risk}}{\text{Person-time within the population at-risk}}
Incidence proportion != incidence rate

Incidence Rate = \frac{\text{# of new outcomes within a defined time-at-risk}}{\text{Person-time within the population at-risk}}

Incidence Proportion = \frac{\text{# of persons with a new outcome within a defined time-at-risk}}{\text{# of persons with at least 1 day at-risk within the population at-risk}}
Constructing incidence from patient experience

- Variable start and end dates
- Multiple entry entries
- Outcome caused ‘clean window’
- Outcome recurrence
- Prior event impacts risk start
- Outcomes post-TAR
- Different people with varied baseline characteristics:
  - Demographics (age, sex, race)
  - Prior conditions (risk factors)
  - Health behaviors

Incidence rate = 1 new outcome during 3-year time-at-risk / (3 + 3 + 2) person-years of population-at-risk

Incidence rate = 4 new outcomes (from 3 persons) during 3-year time-at-risk / 22 person-years (from 9 persons) of population-at-risk
One desired use case for incidence rates: Comparing observed vs. expected

\[ \text{Incidence Rate Ratio} = \frac{\text{Observed (O) Incidence Rate}}{\text{Expected (E) Incidence Rate}} \]

Potential sources of bias:
- Measurement error – do the target and outcome cohorts have false positives, false negatives, or miscalibration of index date?
- Selection bias – is sampling of population(s) representative?
- Confounding – is the population used to infer ‘expected’ sufficiently comparable to the ‘observed’, as a proxy for the counterfactual?

Epidemiologic study designs:
- Comparative cohort: ‘expected’ = comparator cohort satisfying some criteria (e.g. ‘unexposed’ or active comparator)
- Self-controlled: ‘expected’ = unexposed time within exposed patients
Incidence proportion $\neq$ prevalence

Incidence Proportion
\[
\text{# of persons with a new outcome within a defined time-at-risk} \div \text{# of persons with at least 1 day at-risk within the population at-risk}
\]

Prevalence (proportion)
\[
\text{# of persons who have a condition within a defined time-at-risk} \div \text{# of persons with at least 1 day at-risk within the population at-risk}
\]

persons who have a condition = persons with pre-existing disease + persons with a new outcome

population at-risk excludes persons not eligible for numerator; this can mean excluding persons with pre-existing disease (if disease is chronic or cannot recur as ‘new outcome’) when characterizing incidence but not excluding those persons when characterizing prevalence
• Incidence rates and proportions
  – If outcome is chronic/non-recurrent condition (ex: Multiple sclerosis):
    • Web app: ATLAS/Incidence Rates
    • R package: CohortIncidence
  – If outcome is acute/recurrent condition (ex: influenza):
    • R package: CohortIncidence
      – Handles target and outcome cohorts with multiple entries per person
      – Provides stratification by age/sex/year

• Prevalence
  – Web app: ATLAS / Cohort Definitions
ATLAS / Cohort Definitions:
Condition = ‘pre-existing disease + new outcomes’
ATLAS / Cohort Definitions: Population-at-risk for prevalence
ATLAS / Cohort Definitions:
Population-at-risk for incidence
ATLAS / Cohort Definitions:
Incidence proportion
(not advised as best practice, but shown to illustrate concept)
ATLAS/Incidence Rates design
ATLAS/Incidence Rates results
ATLAS/Cohort definition: Prevalence design
ATLAS/Cohort definition: Prevalence results