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CLINICAL EPIDEMIOLOGY NOTES

What is heterogeneity and is it important?

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Three simple examples from recent BMJ papers illustrate the importance of heterogeneity in a systematic review and how readers can assess it.

Three systematic reviews published in the BMJ, including one in this issue, have referred to heterogeneity and dealt with it in three different ways.¹ ² ³ So what is heterogeneity, and how do we assess its importance in a systematic review?

Clinical heterogeneity

Sometimes trials are just looking at different concepts. Reviewers might set out to summarise interventions for improving patients’ ability to make treatment choices; the trials, however, might have covered diverse interventions, such as information leaflets, CD Rom sessions, and training in consultation techniques for doctors. Although the interventions try to achieve the same end result (to improve patients’ ability to make choices), they are different in nature.

In theory, we could add all the trials in this review together and come up with a number, but would this be useful? Would the averaged number apply to all these diverse interventions? The interventions are so different that combining them does not make clinical sense. This is an example of clinical heterogeneity.

Other circumstances that may give rise to clinical heterogeneity include differences in selection of patients, severity of disease, and management. Judgments about clinical heterogeneity are qualitative, do not involve any calculations, and can be made by putting forward a convincing argument about similarities (or differences) between the trials.

Statistical heterogeneity

Individual trials in a systematic review may seem to measure the same outcome, but may have results that are not consistent with each other. Some trials show a benefit while others show harm, or the trials are inconsistent in the size of benefit or harm. This is the case in the systematic review of medications to prevent allergic reactions caused by contrast media.¹ The trials that measured effects on cutaneous symptoms of allergy showed a range of odds ratios from 0.12 favouring the medication to 1.02 favouring the control (fig 1). This is an example of statistical heterogeneity.

How can you detect it and does it matter?

Statistical heterogeneity is apparent only after the analysis of the results. Heterogeneity may be judged graphically (by looking at the forest plot) and be measured statistically. In a forest plot from the systematic review of calcium supplementation,⁴ the error bars for each trial include the summary result, which suggests that statistical heterogeneity is not a problem and that the message is a consistent one (fig 2).

To determine whether significant heterogeneity exists, look for the P value for the χ² test of heterogeneity. A high P value is good news because it suggests that the heterogeneity is insignificant and that one can go ahead and summarise the results. Because statistical tests for heterogeneity are not very powerful it is sensible to use a higher P value than usual (say, P>0.1) as the cut-off for a decision and to think about clinical heterogeneity anyway.

The systematic review of calcium supplementation passes the test, and the authors have rightly summarised the effects on bone density using a simple fixed effects model. This model assumes that all trials...
Fig 1 | Forest plot adapted from Tramèr et al1 showing statistical heterogeneity in the odds ratios for medications to prevent cutaneous allergic reactions (P for 2 test for heterogeneity for anti-H1 combined was 0.03)

Fig 2 | Forest plot adapted from Winzenberg et al2 showing absence of statistical heterogeneity in the odds ratios for the effect of calcium supplementation on bone mineral density. SMD=standardised mean difference
are trying to measure the same thing and that more influence should be given to larger trials when computing an average effect. But what if the P value for the χ² test of heterogeneity is low, suggesting significant heterogeneity? What can be done? Two approaches are possible. We can either avoid summarising the result and look for reasons for the heterogeneity, or we can summarise the effects using another method—the random effects model. Reasons for heterogeneity, other than clinical differences, could include methodological issues such as problems with randomisation, early termination of trials, use of absolute rather than relative measures of risk, and publication bias.

The authors of the systematic review of medications used to prevent allergic reactions caused by contrast media took the first approach. The forest plots suggest that the two classes of drugs have different effects, particularly for skin reactions, and the P value for the statistical test for heterogeneity was significant at 0.03. They decided not to summarise an average effect and felt that the difference between treatments was part of the message of the review.

The authors of the review of interventions to prevent falls and fractures took the second approach. The forest plot for falls in hospital shows a wide spread of results (fig 3). Some trials suggest benefit and others suggest harm from the multifaceted interventions. The authors present the I² statistic, which measures the percentage of variation that is not due to chance. A high percentage, such as the 80% seen here, suggests important heterogeneity. (An I² value of <25% is considered low.)

Nevertheless, the authors felt that all the trials were trying to measure essentially the same thing and that it was worth summarising the results. They used the random effects model, which uses a different formula to calculate more conservative 95% confidence intervals. The effects of treatment are assumed to vary around some overall average treatment effect, as opposed to a fixed effects model, in which it is assumed that each study has the same fixed common treatment effect.

Systematic reviews with a meta-analysis try to provide better numerical answers to the questions, “what is the effect of this intervention and how sure are we about that?” But before believing the results of this method, it might be useful to consider four questions (see box).

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CORRECTIONS AND CLARIFICATIONS

Etoricoxib and diclofenac are associated with similar cardiovascular risks
A momentary lapse in concentration led to a transposition of data when reporting the findings of a recent study in this Short Cuts item by Alison Tonks (BMJ 2006;333:1113-11). The event rates (of thrombotic cardiovascular events such as heart attack) according to the pooled analysis from three large clinical trials in patients with arthritis should have been given as 1.30 [not 1.24] per 100 patient years for diclofenac and 1.24 [not 1.30] for etoricoxib. The hazard ratio was correct: 0.95 (95% confidence interval 0.81 to 1.11) for etoricoxib vs diclofenac, with 0.81 being etoricoxib.

GMC strikes off expert in drug addiction
In this news article by Owen Dyer we wrongly stated that Colin Brewer “took over the Stapleford Centre […] in 1987 after the clinic’s previous director was found guilty of overprescribing” (BMJ 2006;333:1035, 18 Nov, doi: 10.1136/bmj.39034.335278.DB). In fact, it was Brewer who set up the Stapleford Centre, so the centre had no previous director.

Preventing and treating hepatitis B infection
Two years on, we have been alerted to an error in a box in this clinical review by Rakesh Aggarwal and Piyush Ranjan (BMJ 2004;329:1080-6, doi: 10.1136/bmj.329.7474.1080). The error occurs only in the “Full Text” (html) version of the article (not in the pdf or the printed journal). In box 5, a vital superscript value was missing: the second bullet point should read “Virological response—Decline in hepatitis B virus DNA to <10⁷ copies/mL.”

We have already published these corrections on bmj.com

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