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Does animal experimentation inform human healthcare? Observations from a systematic review of international animal experiments on fluid resuscitation

Ian Roberts, Irene Kwan, Phillip Evans, Steven Haig

Animal models are often used to test the effectiveness of a drug or procedure before proceeding to clinical trials. One reason for use of animal models is that they allow researchers to focus on particular pathological processes without the confounding effects of other injuries and treatments. However, it is essential that their results are valid and precise. Biased or imprecise results from animal experiments may result in clinical trials of biologically inert or even harmful substances, thus exposing patients to unnecessary risk and wasting scarce research resources. Moreover, if animal experiments fail to inform medical research then the animals suffer unnecessarily.

The Italian pathologist Pietro Croce criticised vivisection on scientific grounds. He argued that results from animal experiments cannot be applied to humans because of the biological differences between animals and humans and because the results of animal experiments are too dependent on the type of animal model used. Croce's arguments were based on insights into zoology and pathophysiology. In this paper, we make some methodological observations on animal experiments. Our observations were made in the context of a systematic review of all available randomised controlled trials of fluid resuscitation in animal models of uncontrolled bleeding. We conducted this review because we wanted to assess the scientific basis for fluid resuscitation. A previous systematic review of randomised trials of fluid resuscitation in bleeding trauma patients had provided no evidence that fluid resuscitation improved outcome.

Summary points

- New drugs and procedures are usually tested in animals before conducting clinical trials
- Validity of animal experiments is essential for human health care and fundamental to animal welfare
- A systematic review of animal experiments on fluid resuscitation found that most studies were underpowered and provided little information on possible bias
- Systematic reviews of animal experiments allow a more objective appraisal of the evidence and reduce the chance of false negatives results
- Systematic reviews across species would help determine whether the results could be generalised to humans

Systematic review of fluid resuscitation in uncontrolled haemorrhage

We did a systematic review of randomised controlled trials of the timing or volume of fluid administration in animal models of uncontrolled haemorrhage. Details of the review methods, search strategy, and included trials are available on bmj.com. The combined electronic search strategies identified 3193 potentially eligible reports. Two reviewers examined each of these records and 104 reports were retrieved in full. From these, we identified 44 randomised controlled trials meeting the inclusion criteria. The 44 trials included a total of 2039 experimental animals (1772 rats, 251 pigs, and 16 sheep). Mortality data were reported in 42 trials, of which 31 were in rats, 10 in pigs, and one in sheep. In most of the rat experiments uncontrolled bleeding was induced by resecting the tail. Three trials in large animals (pigs and sheep) could not be included in the meta-analysis because they did not include a no fluid resuscitation group; one compared early and late resuscitation and two compared different blood pressure resuscitation targets. Three trials in rats could not be included in the meta-analysis: one compared early and late fluid resuscitation, one compared different blood pressure resuscitation targets, and one presented time to death data only.

The pooled odds ratio (fixed effect) for death in large animals (pigs and sheep) with fluid resuscitation was 0.63 (95% confidence interval 0.15 to 2.61) but there was statistical heterogeneity ($\chi^2 = 16.84$, df = 7, $P = 0.018$). The pooled odds ratio (fixed effect) for death in small animals with fluid resuscitation was 1.14 (0.65 to 2.02). Again, there was substantial heterogeneity ($\chi^2 = 93.40$, df = 27, $P < 0.0001$). When the meta-analysis was stratified according to how uncontrolled bleeding was induced, a large amount of the heterogeneity was accounted for. Figure 1 shows the...
results of meta-analysis of the 16 randomised controlled trials of fluid resuscitation in rats in which bleeding was induced by resecting the tail. The meta-analysis is stratified according to where the tail was cut. Fluid resuscitation seems to be harmful (odds ratio = 2.88, 95% confidence interval 1.72 to 4.80) with less than 50% tail resection ($\chi^2 = 5.57, df = 7, P = 0.59$) but beneficial (odds ratio = 0.25, 0.15 to 0.42) with greater than 50% tail resection ($\chi^2 = 6.14, df = 7, P = 0.52$).

### Are the individual experiments valid?

In clinical trials, systematic error can arise from problems with the study design, especially if allocation of treatment is inadequately concealed. Bias is avoided by ensuring strict randomisation with well concealed treatment allocation. The extent to which inadequate concealment of allocation might introduce bias in animal experiments is uncertain. However, it is easy to imagine how bias could arise. For example, weaker animals may be easier to catch than healthy animals, and this could result in systematic differences between the intervention and control groups on baseline prognostic factors. Of the 44 randomised controlled trials meeting the inclusion criteria, only two described how the animals were divided into treatment groups; both of these trials used alternation.

Random error in clinical trials is minimised by increasing the number of randomised participants. However, animal researchers are encouraged to reduce the number of experimental animals to a minimum. Indeed, the need to use the minimum number of animals to obtain valid results is embodied in the Animals (Scientific Procedures) Act 1986 and European legislation. As a result, some animal experiments are underpowered and provide little reliable information. All of the animal experiments in our systematic review were small (fig 2). The average number of animals per trial was 46 (2039/44), and the largest trial included only 207 animals (rats). None of the trials would have been large enough to detect reliably a 10% absolute difference (halving) in the risk of death between the intervention and comparison groups. Moreover, many of the trials included several different fluid resuscitation groups, which we combined for our analyses. The average number of experimental animals per treatment group was only 13 (160 groups). If, as was the case in most trials, the aim was to compare the effects of different fluid resuscitation regimens, the studies had little power.

### Has all the evidence been assessed?

Although each individual animal experiment provides little reliable information on the effectiveness of fluid resuscitation, each contributes to the total body of evidence. Any inferences should be based on all the evidence. A 1996 narrative review of fluid resuscitation in animal experiments included only nine of the 24 trials (38%) that were available at that time.

Systematic reviews and meta-analyses of animal experiments are uncommon. About 1 in 1000 Medline records pertaining to human research is tagged as a meta-analysis compared with 1 in 10 000 records pertaining to animal research. In his book *The Principles of Humane Experimental Technique*, William Russell posed the principle of reduction—that is, the use of methods to “reduce the number of animals needed to obtain information of a given amount and precision.” Meta-analyses of the results of previous animal experiments would increase the precision of estimates of treatment effects and therefore reduce the number of animals needed in future experiments.

Publication bias may be as potent a threat to validity in systematic reviews of animal experiments as it is in systematic reviews of clinical trials. We contacted the authors of included trials to ask about unpublished studies but none were identified. However, it would be surprising if there were no unpublished trials meeting our inclusion criteria. Prospective registration of

### Table 1

<table>
<thead>
<tr>
<th>Study</th>
<th>No of deaths</th>
<th>Odds ratio (95% CI fixed)</th>
<th>Weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fluids</td>
<td>No fluids</td>
<td></td>
</tr>
<tr>
<td>&lt;= 50% tail resection:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bilynsky 1992</td>
<td>13/60</td>
<td>9/60</td>
<td>2.88 (1.72 to 4.80)</td>
</tr>
<tr>
<td>Gross 1989</td>
<td>17/68</td>
<td>3/15</td>
<td>1.72 (0.63 to 4.80)</td>
</tr>
<tr>
<td>Krausz 1991</td>
<td>7/20</td>
<td>0/17</td>
<td>0.63 (0.25 to 1.55)</td>
</tr>
<tr>
<td>Krausz 1992</td>
<td>6/24</td>
<td>2/9</td>
<td>0.25 (0.15 to 0.42)</td>
</tr>
<tr>
<td>Krausz 1992</td>
<td>21/60</td>
<td>2/15</td>
<td>0.15 (0.05 to 0.42)</td>
</tr>
<tr>
<td>Krausz 1992</td>
<td>8/12</td>
<td>4/13</td>
<td>0.13 (0.05 to 0.34)</td>
</tr>
<tr>
<td>Rabinovic 1989</td>
<td>19/40</td>
<td>1/10</td>
<td>0.05 (0.01 to 0.27)</td>
</tr>
<tr>
<td>Talmor 1999</td>
<td>25/68</td>
<td>3/19</td>
<td>0.19 (0.07 to 0.49)</td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td>111/320</td>
<td>24/158</td>
<td>2.88 (1.72 to 4.80)</td>
</tr>
<tr>
<td>&gt; 50% tail resection:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Capone 1995</td>
<td>14/20</td>
<td>9/10</td>
<td>0.15 (0.05 to 0.42)</td>
</tr>
<tr>
<td>Capone 1995</td>
<td>3/20</td>
<td>1/10</td>
<td>0.10 (0.03 to 0.34)</td>
</tr>
<tr>
<td>Capone 1995</td>
<td>9/10</td>
<td>10/10</td>
<td>1.00 (0.50 to 2.00)</td>
</tr>
<tr>
<td>Greene 1998</td>
<td>20/42</td>
<td>16/21</td>
<td>0.86 (0.63 to 1.18)</td>
</tr>
<tr>
<td>Kim 1997</td>
<td>12/20</td>
<td>17/20</td>
<td>1.70 (1.20 to 2.40)</td>
</tr>
<tr>
<td>Sindlinger 1993</td>
<td>14/30</td>
<td>11/15</td>
<td>0.83 (0.50 to 1.40)</td>
</tr>
<tr>
<td>Soucy 1995</td>
<td>45/90</td>
<td>40/45</td>
<td>0.90 (0.63 to 1.28)</td>
</tr>
<tr>
<td>Soucy 1995</td>
<td>25/31</td>
<td>10/12</td>
<td>0.40 (0.20 to 0.80)</td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td>137/263</td>
<td>114/143</td>
<td>0.25 (0.15 to 0.42)</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>248/583</td>
<td>138/301</td>
<td>0.86 (0.63 to 1.18)</td>
</tr>
</tbody>
</table>

**Fig 1** Meta-analysis of 16 randomised controlled trials of fluid resuscitation in rats with uncontrolled haemorrhage by tail resection. "Capone" reported two trials
animal experiments at inception may help to avoid the problem of publication bias. In the United Kingdom, the Animals (Scientific Procedures) Act 1986 regulates “any experimental or other scientific procedure applied to a protected animal which may have the effect of causing that animal pain, suffering, distress, or lasting harm.” Researchers must have a project licence from the Home Office before conducting any animal research, and the licence application describes the experimental protocol. These data could be used for prospective registration of all animal experiments.

Systematic reviews of animal models could, like ours, include a range of animal species and models. If the results were consistent across species and models this would indicate that they might also apply in humans. Since the primary aim of animal experimentation is to inform human experimentation, this would be valuable information.

We found substantial statistical heterogeneity in our meta-analysis, making it impossible to interpret the odds ratios. Investigation of heterogeneity is essential and can increase the scientific and clinical relevance of their results. In our meta-analysis, stratification according to uncontrolled bleeding was induced accounted for a large amount of the heterogeneity, but these results need to be interpreted with caution. Meta-analytic subgroup analyses are akin to subgroup analyses within trials and are prone to bias. Although we specified in our protocol that the analyses would be stratified according to the animal model used, we did not specify that we would stratify according to where the tail was cut. Nevertheless, the meta-analysis provides an insight into model dependency that could be taken into account in future animal experiments and when considering whether the results can be generalised to humans.

Implications for human health
Animal experiments can inform human health care only if their results are valid and can be generalised. However, little information is available on the methodological determinants of bias in animal experiments, and in our example the sample sizes were too small to obtain precise estimates of the effects of the interventions. Systematic reviews of animal experiments would help to ensure that animal experiments do not set out to answer questions that have already been answered, reduce bias and increase precision, and provide reassurance about whether the results can be generalised.

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Contributors: IR and PE proposed the study. IR drafted the protocol that was revised following comments from all authors. IR and SH examined the electronic search results for reports of possibly relevant randomised controlled trials. IR, PE, and SH applied the selection criteria independently to the trial reports. IR and IK extracted information from the included trials. IK contacted authors for further information and IK and IR conducted the analyses. IR drafted the paper that was revised on the basis of comments from IK, PE, SH, IR will act as guarantor. Funding: None. Competing interests: None declared.

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