LETTERS TO THE EDITOR

Need to avoid bias in controlled trials

EDITOR—It is accepted practice that in all quantitative research statistical tests should be used to estimate the likelihood that findings arose by chance and that authors should avoid overinterpreting data where these tests apply, do not show significance. It is also accepted that to avoid bias in the analysis of randomised controlled trials clinically important endpoints must be specified.

A recent report of a clinical trial of two different surfactant preparations illustrates the dangers of neglecting these guidelines. Although infants in the group treated with one of the preparations appeared to fair better on the pre-determined clinical endpoints, none of the differences approached conventional levels of significance. The paper also reports data on a number of intermediate measures of oxygenation and ventilatory requirements, outcomes which the authors specifically state were not established at the design stage, where on some occasions, although statistically, differences do reach conventional levels of significance.

In the discussion the authors begin by noting the need for caution in the interpretation of the results from this small study but, unfortunately, dismissed their own observations. They interpret the data on intermediate endpoints as demonstrating improved lung function in babies treated with Curosurf and devote much of the discussion to trying to rule out the possibility that Curosurf is the more effective agent. They conclude the paper by stating their belief that Curosurf is more effective at improving oxygenation and reducing ventilatory requirements than Survanta in the first 24 hours and remind readers of the ‘trend’ towards lower incidence of adverse outcomes in the Curosurf group and suggest the need for these results to be confirmed in a large randomised controlled trial. The clear implication of their comments is that they believe that they have demonstrated that Curosurf is more effective than Survanta.

The correct interpretation of the data presented is that the results would be compatible with Curosurf being either considerably better or considerably worse than Survanta at preventing the development of adverse clinical outcomes. The authors specifically mention that patients treated with Curosurf had a lower incidence of pneumothorax, intracerebral haemorrhage grades 3-4, and mortality: in fact, the adjusted 95% confidence limits of the odds ratios for each of these outcomes associated with being treated with Curosurf rather than Survanta are 0.08-2.88, 0.02-1.78, and 0.02-2.54, respectively, which suggests a high probability that these results are due to chance. There is always a temptation to overinterpret data from studies which are too small to provide answers to the questions addressed. In this particular paper there is the added concern that the drug favoured by the authors’ discussion, on the basis of inappropriate interpretation of the data, is manufactured by the company which sponsored the research.

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Dr Gefeller, Professor Speer, et al comment:
A well known dilemma of small-sized clinical trials refers to the repeated observation that they fail to reveal significant differences between the treatment groups, especially when treatment success is assessed using dichotomous outcome variables. For example, a reduction in a treatment complication from 20% to 10% yields a P value of 0.345 (two-tailed Fisher’s exact test) in a comparison of 40 patients randomised to each of two treatment groups (and thus is far away from any claim of a significant difference).

However, when we report our trial of 200 patients in each group, the same difference is highly significant (P = 0.007).

Dr Logan criticises our small-sized randomised clinical trial of two treatment regimens of natural surfactant preparations (Curosurf v Survanta) in neonatal respiratory distress syndrome. He blames us for ‘over-interpreting’ our data, and that we implied that Curosurf was more effective than Survanta.

Dr Logan misunderstood our message. On the contrary, although we agree with his well known general comments on the analysis of randomised trials, we believe that he has chosen an unsuitable example for his crusade against overinterpreted studies.

Specifically, our study was designed as a pilot study for a large scale study (explicitly stated in the paper). Thus, there was no necessity to establish a hierarchy of clinically important endpoints at the design stage. We recorded all relevant complications using standardised criteria and documented measures of oxygenation and ventilatory requirements at fixed time points specified in the study protocol. Our paper presented the results of this pilot trial with respect to all recorded endpoints. Without retracting our final conclusions, we found a consistent picture favouring the Curosurf treatment regimen.

The differences in complications did not reach significance (repeatedly emphasised in the paper), whereas the improvement in oxygenation and the reduction of ventilatory requirements for infants treated with Curosurf did so at several time points during the first 24 hours. We discussed in detail the limitations of our trial and we did not request with the request ‘these findings need to be confirmed in a larger randomised trial’. Consequently, we are puzzled by Dr Logan’s eloquent criticism of our findings.

We take exception to the comments made in the final sentence of Dr Logan’s letter. We openly declared the source of funding of our trial and that it was designed, conducted, and analysed independently by the company according to GCP standards. We assume that since Dr Logan did not declare any financial support in his letter, he has no conflict of interest.

1 Frieman JA, Chalmers TC, Smith H, et al. The importance of beta, the type II error and sample size in the design and interpretation of the

Raised cerebral artery blood flow velocity

EDITOR—Scherjon et al’s findings of raised cerebral artery blood flow velocity (CABFV) in infants who had evidence of fetal brain sparing are very similar to our previously published findings of a raised CABFV at 3 days of age in small for gestational age (SGA) infants weighing less than 1500 g. Two thirds of the patients in our study had evidence of fetal redistribution of blood flow, with absent end diastolic flow in the umbilical artery. Postnatally, we also found persistently lowered coeliac axis, superior mesenteric artery, and renal artery blood flow velocities, indicating that ongoing redistribution of blood flow in SGA babies affects the whole circulation.

Scherjon et al did not find an association between blood pressure and CABFV, whereas we found that for appropriately grown infants a multiple regression analysis indicates a significant correlation between blood pressure, carbon dioxide tension, and CABFV.

We carried out an identical multiple regression analysis, using data from 16 SGA infants weighing less than 1500 g at birth who had invasive monitoring of blood pressure through umbilical or peripheral arterial lines. Measurements of blood flow velocity in the anterior cerebral arterial were taken at 24 hours of life in the same way as for the appropriately grown infants. For the SGA infants there was no effect of blood pressure, although there was an effect of carbon dioxide tension (table).

Correlation coefficients from multiple regression analysis of effects of mean arterial blood pressure (MABP), PaCO2 and PaO2 on anterior cerebral artery blood flow velocity

<table>
<thead>
<tr>
<th>MABP</th>
<th>PaCO2</th>
<th>PaO2</th>
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<tbody>
<tr>
<td>Appropriately grown</td>
<td>0.70***</td>
<td>0.64***</td>
</tr>
<tr>
<td>SGA (n=16)</td>
<td>0.14</td>
<td>0.60*</td>
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*p<0.05, ***p<0.001.

We suggest that Scherjon et al’s failure to find an effect of blood pressure on CABFV may be due to the lack of such effects in the SGA group only. The antenatal experiences of the fetus may modify the responses of many parts of the circulation, producing both baseline changes and differences in the way the circulation responds to factors such as blood pressure. Circulatory data from appropriately grown and SGA babies should probably be analysed separately, and as we understand more, it may be that our management of the circulation should also be different in these babies.

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1 Scherjon SA, Oosting H, Kok JH, Zondervan HA. Effect of fetal brain sparing on the early