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Sander Greenland (Editor)
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II-4.


Woolf's "On Estimating the Relation Between Blood Group and Disease" is most often cited for its introduction of what was probably the first common odds-ratio estimator for a set of two-by-two tables (or studies), and a test for heterogeneity of the odds ratio across the tables; these are now known as Woolf's estimate and Woolf's test of homogeneity (see, e.g., Schlesselman [1982]). Though these statistics have been supplanted by Mantel-Haenszel and maximum-likelihood methods, the paper contains more than just bare mechanics, as it touches on a number of important points which were generally unappreciated in the early literature.

Woolf was motivated by concern about the then-prevalent practice of analyzing and comparing case-control results by computing the difference in exposure proportions between case and control groups. As he notes, this difference would vary with the background frequency of exposure, and artifactual differences between study results could be expected. Instead he chose to work with the ratio of incidence rates, and recognized (independently of Cornfield) that this ratio could be estimated from case-control data, even if the absolute rates could not. (He did not, however, delineate sufficient conditions for the sample odds ratio to estimate the incidence ratio.) He also noted some subtle points that many later users of his methods overlooked, one being that his variance formula for the common odds ratio applied only under the homogeneity assumption.

Reference:
ON ESTIMATING THE RELATION BETWEEN
BLOOD GROUP AND DISEASE

BY BARNET WOOLF

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Following the demonstration of a significant excess of blood group A in patients with cancer of
the stomach (Aird, Bentall & Roberts, 1953) and of group O in sufferers from peptic ulcer (Aird,
Bentall, Mehigan & Roberts, 1954) and from toxaemia of pregnancy (Pike & Dickins, 1954) it
seems certain that many more studies will be made on the relation between blood groups and
disease. It is therefore important that the best possible statistical methods should be used. The
procedure recommended by Aird et al. (1954) is very efficient, but it is open to criticism on one
rather important point. These workers take as criterion the difference in proportion of a given
blood group in the disease and the control series. Denote the two blood types α and β. Suppose
the disease series contains h patients of type α and k of type β, where h + k = n, and the control
series has H of type α and K of type β, where H + K = N. Aird and associates calculate
d = h/n - H/N. This is tested for significance against its sampling variance, combined with
estimates from other bodies of data to give a weighted mean estimate, and compared with these
other estimates in tests for heterogeneity.

Unfortunately, d will differ from one community to another even when the specific attack rate
within any given blood group stays constant. This can be shown by a simple example. Consider
a community of 10,000 people in which H and K are each 5000. Then if h = 100 and k = 50,
d = 100/150 - 0.5, or 0.1667. Now consider another community in which H is 9000 and K is 1000.
In this case h = 180 and k = 10, so d = 180/190 - 0.9, or 0.0474. Even when the essential biolog-
cal conditions are identical, differences in blood-group frequencies in the population will
introduce spurious heterogeneity. This kind of artefact is avoided if one works with incidence
rates in the various blood groups. The data usually do not permit calculation of absolute rates,
nor are they needed. What is wanted and readily obtained is an estimate of the ratio of one rate
to another. The incidence in group α will be h/H x some constant, and that in group β will be
k/K x the same constant. If the ratio is taken as x to 1, an estimate of x will be hK/Hk, and
it may readily be shown that this is the maximum-likelihood estimate. The use of x is recom-
mended instead of d as a criterion of differential incidence of disease in relation to blood group.

In all statistical computations it is best to transform x into its logarithm. This avoids diffi-
culties due to asymmetry. If comparison of α with β gives z = 2 say, comparison of β with α
will give x = 1/z; but log x will retain its numerical value, merely changing in sign. Moreover, the
sampling variance of log x is a very simple expression free of ‘nuisance parameters’. This is
especially true if one transforms into y = loge x. If V is the sampling variance of y, then

\[ V = 1/h + 1/k + 1/H + 1/K, \]

and w, the weight of y, is of course 1/V. If the attack rate is the same for both blood types the
expected value of y will be 0, so the null hypothesis is not to be rejected unless y differs signifi-
cantly from zero. This is tested by \( \chi^2 \) will be \( y^2/V \) or \( wy^2 \) for one degree of freedom. Combination of data
from different communities proceeds as described by Aird et al. (1954). The weighted mean, Y,
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is $\Sigma wy / \Sigma w$, and its antilogarithm, $X$, is taken as the combined estimate of $x$. Significance of $Y$ is tested by $\chi^2 = (\Sigma wy)^2 / \Sigma w$ or $Y^2 \Sigma w$ for one degree of freedom. Heterogeneity is tested by $\chi^2 = \Sigma wy^2 - (\Sigma wy)^2 / \Sigma w$ or $\Sigma wy^2 - Y^2 \Sigma w$ with degrees of freedom one less than the number of sets of data combined. The standard deviation of $y$ is $V^4$, and the approximate fiducial limits at the 95% point are $y \pm 1.96 V^4$. Provided there is no significant heterogeneity the standard deviation of $Y$ is $1/(\Sigma w)^{1/2}$ and the 95% fiducial limits are approximately $Y \pm 1.96(\Sigma w)^{1/2}$. By taking antilogarithms these can be transformed into fiducial limits for $x$ or $X$. This is a 'large-sample' treatment, and the formulae cease to be applicable if any of the observed frequencies is small.

Table 1. Calculation of combined estimate of incidence ratio of peptic ulcer in groups $O$ and $A$

<table>
<thead>
<tr>
<th>City</th>
<th>Peptic ulcer</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$hK$</td>
<td>$y = \log, x$</td>
</tr>
<tr>
<td></td>
<td>$\frac{hK}{k}$</td>
<td>$w = \frac{1}{h + k + 1}$</td>
</tr>
</tbody>
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<th>City</th>
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<th>$\chi^2$ analysis</th>
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$Y = \Sigma wy / \Sigma w = 0.5289.$  
$Y^2 \Sigma w = 62.63.$  
$s.d. of Y = (\Sigma w)^{-1} = 0.0947.$  
95% fiducial limits of $Y = 0.2472 - 0.4106.$  
$X = \text{antilog } Y = 1.39.$  
95% fiducial limits of $X = 1.28 - 1.51.$

Table 2. Incidence ratios of some diseases in relation to blood group

<table>
<thead>
<tr>
<th>Disease</th>
<th>Comparison</th>
<th>$X$ or $x$</th>
<th>95% fiducial limits</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cancer of stomach</td>
<td>Group $A$ with group $O$</td>
<td>1.22</td>
<td>1.12 - 1.32</td>
<td>Aird et al. (1953)</td>
</tr>
<tr>
<td>Peptic ulcer</td>
<td>Group $O$ with group $A$</td>
<td>1.39</td>
<td>1.28 - 1.51</td>
<td>Aird et al. (1954)</td>
</tr>
<tr>
<td>Toxaemia of pregnancy</td>
<td>Group $O$ with all others</td>
<td>1.38</td>
<td>1.15 - 1.66</td>
<td>Pike &amp; Duckins (1954)</td>
</tr>
</tbody>
</table>

Table 1 shows the calculations for comparing incidence of peptic ulcer in groups $O$ and $A$ using combined data from London, Manchester and Newcastle. This is the example worked out by their method by Aird et al. (1954, Table VII). The $\chi^2$ values, 62.63 for significance and 2.99 for heterogeneity, agree closely with 66-21 and 3-91 found by the $d$ method. Similarly, combined data from six centres on cancer of the stomach in groups $O$ and $A$ gave $\chi^2$ values by the $x$ method of 21.28 (1 d.f.) for significance and 2.63 (5 d.f.) for heterogeneity, against 21.49 and 2.69 by the $d$ method. These close concordances are to be expected, since the observations all come from cities in England with very similar population blood-group frequencies. If data from different ethnic groups were combined, the $d$ method would in general be expected to return falsely high $\chi^2$ figures for heterogeneity. This might be of some biological or medical importance as
tending to be confounded with possible genuine heterogeneity arising either from environmental factors or from differential attack rates for the diverse genotypes that may go to make up a single blood group. Even when heterogeneity is not an issue, \( x \) is preferable to \( d \) because it has a direct medical meaning. Table 2 gives some estimated incidence ratios of diseases in relation to blood group, together with fiducial limits. A blood-group difference appears able to increase the risk of disease by as much as 39%.

REFERENCES

