An easier alternative to orthogonal regression for calculation of International Sensitivity Indexes

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Abstract

Aims—To evaluate an easier alternative method to orthogonal regression analysis for calculating International Sensitivity Indexes (ISI).

Methods—ISI for 18 reagents were estimated from reference and test reagent prothrombin times using plasma from 60 stabilised patients undergoing anticoagulation therapy and 20 normal subjects. ISI were also derived for 12 systems (instrument/reagent combination) using lyophilised plasma calibrants. Orthogonal regression and the easier alternative log ratio method were evaluated by comparing resultant International Normalised Ratios (INR) for 58 patients using two test systems.

Results—For the reagent calibrations, the differences in the two methods for the sensitivity slopes were very small. For the system calibrations, slope differences were still of little clinical importance. Parallel observations for INR on the 58 patients confirmed that the bias introduced by easier log ratio derivation of ISI was small and of minor clinical importance, although the bias increases for high INR.

Conclusions—The easier method for ISI determination is a useful alternative to orthogonal regression analysis, particularly when computer assistance is not available, for checking for gross errors in computer computation and for use when calculating ISI from INR calibrants.

Keywords: Oral anticoagulant therapy.

Prothrombin times used for monitoring oral anticoagulant therapy are now expressed as International Normalised Ratio (INR). To derive the INR, the International Sensitivity Index (ISI) of the local reagent needs to be determined. For assessment of the ISI of the local system (reagent/technique combination), a calibration against a reference system is required. Such calibration conventionally requires prothrombin times by local and reference systems on twenty normal subjects and 60 patients stabilised on coumarin treatment. Alternatively, a series of lyophilised plasma samples may be used to calibrate the local reagent and instrument combination system. The calculation of the ISI is achieved by estimating the slope (b) by orthogonal regression analysis of the logarithms of local and reference system prothrombin time results. Such calculation is complex. Computer programs make the calculation less tedious than using logarithm tables or a calculator, but often such programs are not locally available. We have derived a simpler method and in this study have evaluated the ISI obtained. The ISI determined by this method have been compared with those derived from orthogonal regression.

Methods

The usual method for calculating an ISI of a new prothrombin time system involves finding the slope (b) of the line of best fit (by orthogonal regression) of the log prothrombin time of the new system against the reference technique. The ISI of the new system is b x ISI of the reference technique. The easier method for determining b is as follows (see appendix):

\[ \text{Slope } b = \frac{\log \text{ mean reference INR}}{\log \text{ mean (local PR)}^{\text{ISI ref}}} \]

Eighteen calibrations of various reagents were studied. These involved parallel pro-

Figure 1 Eighteen in-house calibrations of various reagents.

* That is, log INR of local system assuming ISI is that of the reference system; PR is the prothrombin ratio.
Three Twelve in-house determinations of system ISI of machines using lyophilised, calibrated plasma samples were also examined. Again slope \( b \) was calculated by the new method and by orthogonal regression. The resulting ISI were plotted and Bland and Altman plots drawn. Using ISI derived by log ratio and regression methods, the resulting INR of 58 patients receiving stable coumarin treatment were compared. This comparison was made for two of the 12 in-house determinations of system ISI.

**Results**

For the 18 calibrations, the mean difference between the slopes determined by the easier method and orthogonal regression was very small (\(-0.0019\)); the limits of agreement being \(-0.0371\) to \(0.0333\). For the 12 in-house machine calibrations using the 20 to 30 lyophilised plasma samples, the mean slope difference was \(-0.0159\) with limits of agreement of \(-0.0521\) to \(0.0203\). The simple plots and Bland and Altman plots for the two types of calibrations are shown in figs 1-4. The parallel observations on INR of 58 patient samples are shown in figs 5 and 6 by means of Bland and Altman plots. The mean INR bias for the system is small when the log mean and orthogonal derived ISI are compared.

**Discussion**

This new method for determining relative sensitivity slopes from thromboplastin calibration exercises, although approximate, appears to give results close to those determined by orthogonal regression. For full calibrations involving observations on 60 patients and 20 normal subjects, the slope \( b \) determined by the new method is unlikely to be more than \(0.0371\) different from the "true" value. In INR terms this would equate to approximately 0.06 under or overestimate for an INR of 2.17 while for an INR of 4.23 it would give approximately 0.20 under or overestimate. When the new method is used for lyophilised plasma samples with smaller numbers of calibrants, calibrations are less close to orthogonal regression analysis. However, in clinical terms, the resultant INR is reasonably close to the estimate determined using the orthogonal regression method. This is confirmed by the parallel observations of INR on 58 patient samples, although the bias increases for high INR values.

The new method does not assess the precision of the calibration or the validity of the assumption that the slope estimate for patients and normal subjects is similar. Graphical examination of the data would permit some qualitative evaluation of such features. This method is therefore a suitable alternative for estimating ISI and may be useful where local facilities do not include orthogonal regression computer assisted analysis. It also offers a simple means of checking for gross errors in the orthogonal regression computation. In addition, it permits
An easier method for calculating ISI

By definition this is to be equal to the reference INR (always the "gold standard").

Thus

\[
\begin{align*}
\text{PT new system}_{\text{INR ref}} & = \text{INR reference} \\
\text{MNPT new system}_{\text{INR ref}} & = \text{INR reference} \\
\text{local PR}_{\text{INR ref}} & = \text{INR reference} \\
\text{"INR new system"}_\text{INR ref} & = \text{local PR}_{\text{INR ref}}
\end{align*}
\]

Take logs of both sides of the equation:

\[
\begin{align*}
\log \text{PT new system}_{\text{INR ref}} & = \log \text{INR reference} \\
\log \text{MNPT new system}_{\text{INR ref}} & = \log \text{INR reference} \\
\log \text{"INR new system"}_\text{INR ref} & = \log \text{local PR}_{\text{INR ref}}
\end{align*}
\]

The best estimate of this is

\[
\frac{\log \text{mean INR reference}}{\log \text{mean "INR new system"}}
\]

Appendix

ALGEBRAIC PROOF

\[
\text{INR new system} = \frac{\text{PT new system}_{\text{INR ref}}}{\text{MNPT new system}_{\text{INR ref}}}
\]

where PT is the prothrombin time and MNPT is the mean normal PT.

determination of local ISI using plasma samples calibrated in terms of INR rather than using plasma calibrated in terms of prothrombin time and orthogonal regression.

