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Evidence at a glance: error matrix approach for overviewing available evidence

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Abstract

Background: Clinical evidence continues to expand and is increasingly difficult to overview. We aimed at conceptualizing a visual assessment tool, i.e., a matrix for overviewing studies and their data in order to assess the clinical evidence at a glance.

Methods: A four-step matrix was constructed using the three dimensions of systematic error, random error, and design error. Matrix step I ranks the identified studies according to the dimensions of systematic errors and random errors. Matrix step II orders the studies according to the design errors. Matrix step III assesses the three dimensions of errors in studies. Matrix step IV assesses the size and direction of the intervention effect.

Results: The application of this four-step matrix is illustrated with two examples: peri-operative beta-blockade initialized in relation to surgery versus placebo for major non-cardiac surgery, and antiarrhythmics for maintaining sinus rhythm after cardioversion of atrial fibrillation. When clinical evidence is deemed both internally and externally valid, the size of the intervention effect is to be assessed.

Conclusion: The error matrix provides an overview of the validity of the available evidence at a glance, and may assist in deciding which interventions to use in clinical practice.

Background

Evidence-based medicine (EBM) was first introduced in 1992 [1], and its increased application is reflected among others by the growth of The Cochrane Library databases as well as implementation of evidence-based guidelines into clinical practice [2]. EBM underpins that information provided from randomized trials, and systematic reviews of randomized trials represent the most reliable evidence regarding intervention effects [3,4]. Thanks to the sustained scientific process (Additional file 1: Table S1), we now know that the reliability of what we observe varies due to a whole array of different factors. There are three dimensions that particularly influence the reliability of our observations in clinical research and they are empirically and theoretically well accepted: the risk of systematic error (‘bias’), the risk of random error (‘play of chance’), and the risk of design error (‘wrong design to answer the posed question’) [4,9].

EBM usually follows a four-phase process starting from a clinical question proceeding to the implementation of new evidence (Figure 1) [3]. Phase 1 is the formulation of a research question and literature search strategy. Phase 2 is the subsequent systematic appraisal and synthesis of the available evidence. Phase 3 covers the initiation of new research. Alternatively, phase 4 is the implementation of all available evidence when statistically and clinically convincing evidence has been obtained.

In daily clinical practice, the question of whether sufficient evidence is available to recommend the implementation of a specific intervention as a treatment arises frequently [3]. Depending on the specific clinical question, often an exhaustive list of references is retrieved when using a sensitive search strategy in multiple databases [3]. After the selection of studies, their data must be interpreted [10-13].

Since results may be contradictory and studies may differ in more than one aspect, to draw a clear, practical conclusion from the publications may be problematic [14].
Figure 1 Overview of the four phases in the process of evidence-based medicine from question to the initiation of new research or implementation of new evidence. PICOT: patients, intervention, control, outcome measure, time.
Objective

We aimed at conceptualizing a visual assessment tool, i.e., a matrix for overviewing studies and their data in order to assess the clinical evidence. The matrix is constructed from the three dimensions of errors: systematic error (‘bias’), random error (‘play of chance’), and design error (‘wrong design to answer the question posed’ or ‘wrong context’). The application of this matrix will be illustrated by two examples: peri-operative beta-blockade initialized in relation to surgery versus placebo for major non-cardiac surgery, and antiarrhythmics for maintaining sinus rhythm after cardioversion of atrial fibrillation.

Methods

The three major error dimensions

The risk of systematic error (‘bias’)

When evaluating a clinical study, one should always try to assess its risk of systematic error [3,4,9-16]. There is increasing agreement on how trials and studies can be placed in a hierarchy when assessing the risk of systematic error [3,4,9-16], depending on the type of research (therapeutic, diagnostic, etiologic, or prognostic) [3,10,11,17]. The risk of systematic error influences the reliability of observed intervention effects [3,10,11,18,19]. A significant association between inadequate or unclear bias protection and overestimation of beneficial effects and underreporting of adverse effects has been demonstrated [16,19-23]. Differences in risk of bias are found both between the different levels of evidence and within each level of evidence [4,16,20].

For randomized trials, there is empirical evidence that at least six components are associated with systematic error: generation of the allocation sequence [24], allocation concealment [25], blinding [26], incomplete outcome measure reporting [4], selective outcome measure reporting [4], and other bias mechanisms (e.g., baseline imbalance, early stopping, vested interests, etc.) [4,16,20,27-29]. The impact of early stopping of trials on bias is largely dependent on how the stopping rules were defined and the level of statistical significance of the interim analysis [30-32]. Trials with one or more systematic error components assessed as inadequate or unclear are considered to be of high risk of bias, while trials with all quality components assessed as adequate are considered to be of low risk of bias [15,27,33]. Trials with a low risk of bias are more likely to estimate the ‘true’ effect of the intervention [16,20,27,33].

The systematic error dimension can be measured by an ordinal variable expressed in the levels of evidence (Table 1).

The risk of random error (‘play of chance’)

The risk of random error is the risk of drawing a false conclusion based on sparse data. There are two types of false conclusions: a false rejection of the null hypothesis (type I error; alpha) or a false acceptance of the null hypothesis (type II error; beta). When data are sparse, then the so called ‘intervention effect’, whether beneficial or harmful, may in fact be caused by randomly skewed variation in prognostic factors between the intervention groups due to sampling error.

The question, however, is how we quantify and compare the risk of random error between different studies with varying numbers of participants. A p-value reflects the risk that the difference in outcome between two interventions has arisen by chance, given the data and the null hypothesis are true. Since random low (and random high) p-values occur, especially during accumulation of data and sequential testing, the p-value does not sufficiently reflect the true risk of random error. Therefore, the p-values of intervention effect estimates certainly are not suitable for comparison of the risk of random error between different studies [32,34-37]. We suggest using the standard error (SE) for the evaluation of the risk of random error. We used the statistical algorithms from the statistical methods group of the Cochrane Collaboration [38]. The SE in a study may be considered a measure of uncertainty. The SE measures the amount of variability in the sample mean; it indicates how closely the population mean is likely to be estimated by the sample mean. The size of the standard error depends both on how much variation there is in the population and on the size of the sample. When two independent proportions \( p_1 = a/n_1 \) and \( p_2 = c/n_2 \) (with \( a \) and \( c \) being the numbers of patients with events, \( b \) and \( d \) being the numbers of patients with no events, and \( n_1 \) and \( n_2 \) being the total numbers of patients in the intervention group and control group, respectively) are considered in an individual study or a trial \( i \), then the relative risk \( (RR_i) \) is defined by:

\[
RR_i = \frac{p_1}{p_2}
\]

The SE of the log risk ratio for an individual study is calculated by the following formula:

| Category | Studies |
|----------|---------|
| Level 1a | Meta-analysis of randomized trials with low risk of bias |
| Level 1b | Randomized trial with low risk of bias |
| Level 1c | Meta-analysis of all randomized trials |
| Level 1d | Randomized trial with high risk of bias |
| Level 2a | Meta-analysis of cohort studies |
| Level 2b | Cohort study |
| Level 3a | Meta-analysis of case-control studies |
| Level 3b | Case-control study |
| Level 4 | Case-series |
| Level 5 | Expert opinion |
The Peto odds ratio ($\text{OR}_{\text{Peto,}i}$) for an individual study or trial $i$ is defined by:

$$\text{OR}_{\text{Peto,}i} = \exp \left[ \frac{Z_i}{V_i} \right]$$

where

$$Z_i = a_i - E[a_i] = a_i - \frac{n_{1i}(a_i + c_i)}{N_i}$$

and $V_i = \frac{n_{2i}(a_i + c_i) \cdot (b_i + d_i)}{N_i} - (N_i - 1)$

The SE of the log Peto odds ratio for an individual study is defined by:

$$\text{SE} \left[ \ln(\text{OR}_{\text{Peto,}i}) \right] = \frac{1}{\sqrt{V_i}}$$

or

$$\text{SE} \left[ \ln(\text{OR}_{\text{Peto,}i}) \right] = \frac{N_i \sqrt{N_i - 1}}{\sqrt{n_{1i} \cdot n_{2i} \cdot (a_i + c_i) \cdot (b_i + d_i)}}$$

In a meta-analysis results of studies or trials are meta-analysed into one intervention effect estimate. For the Mantel-Haenszel pooled risk ratio ($\text{RR}_{\text{MH}}$) the natural logarithm of the $\text{RR}_{\text{MH}}$ has the standard error given by:

$$\text{SE} \left[ \ln(\text{RR}_{\text{MH}}) \right] = \frac{1}{\sqrt{P \cdot S}}$$

where

$$P = \sum_{i} \frac{n_{1i} \cdot n_{2i} \cdot (a_i + c_i) - a_i \cdot c_i \cdot N_i}{N_i}$$

and $R = \sum_{i} \frac{n_{1i} \cdot n_{2i}}{N_i}$ and $S = \sum_{i} \frac{c_i \cdot n_{1i}}{N_i}$

and $N_i$ being the total number of patients in a trial. For the pooled Peto OR ($\text{OR}_{\text{peto}}$) the natural logarithm of the $\text{OR}_{\text{peto}}$ has the standard error given by:

$$\text{SE} \left[ \ln(\text{OR}_{\text{peto}}) \right] = \frac{1}{\sqrt{\sum V_i}}$$

SE depends on the numbers of events and the sample size.

Due to spurious results, incorrect type I error inferences may be drawn. Recent reports indicate that the influence of the ‘play of chance’ may be much larger than generally perceived [39]. In randomized trials, random error may be one reason for the early stopping of trials at interim analyses when benefit or harm appear to be significant [32,40]. Increased random error may also play a role in the repeated analyses of accumulating data in both trials and meta-analyses [36,41-44]. A cumulative meta-analysis subjects accumulating data to repeated testing of the data and is bound to eventually lead to a false rejection of the null hypothesis (‘false positive’ result) [45,46]. The random error phenomenon or ‘multiplicity’ also plays a role in the evaluation of secondary outcome measures [40]. For example, when data on the primary research outcome, on which the sample size calculation was based, may not show statistical significance, while another outcome measure, for which no separate sample size calculation was performed, exhibits statistical significance [47,48].

Random error may be expressed in a continuous variable using the standard error of for example the log of Peto odds ratios or the log of relative risks.

### The risk of design errors (external validity) - the participants included, the outcomes measured, the interventions, etc

When there is sufficient internal validity, i.e., low risks of systematic errors and random errors, it becomes relevant to consider the risks of design errors (external validity). The design (or context) of any piece of research determines its external validity or generalisability (Table 2) [4]. The external validity becomes questionable when a wrong design has been used to answer the question posed. Among the many variables that should be considered, the relevance of different outcome measures are of central importance to clinical research [13]. We, therefore, focus on them from a patient’s perspective.

Outcome measures can be divided into three categories according to the GRADE classifications (Figure 2) [13]. Primary outcome measures are central in deciding the use of one intervention over another. Large differences in the primary outcome measure between groups in a clinical trial may lead to early termination of a trial (following recommendations of a data safety and monitoring committee) [49]. Choice of the primary outcome should concur with the GRADE category of outcomes, ‘critical for decision-making’ [13]. Secondary outcome
measures are additional outcome measures. If they are positively influenced by an intervention, the results may speak for recommending the intervention only if they support a beneficial effect on the primary outcome or if no clinically and statistically significant effect exist on the primary outcomes (e.g., a RR = 1.00 with 95% confidence limits from 0.98 to 1.02). The secondary outcomes should concur with the second and third GRADE categories of ‘important, but not critical outcomes’ [10-13].

GRADE has schematically ordered outcomes according to patients’ perspective on a categorical scale from 1 to 9, with the most critical outcome, mortality, being graded 9 [13]. Depending on the outcomes, this scale should sometimes be considered nominal and in other situations be considered functional. Moreover, the severity of each outcome may differ as well. A stroke can be minor, while a myocardial infarction may involve a substantial worsening of cardiac function. Grading of outcome measures may also vary according to the clinical question. Therefore, outcomes within a category (i.e., critical, important, or not important) may be interchangeable. However, one can hardly argue that outcomes between categories (i.e., critical, important, or not important) are interchangeable (e.g., mortality is always more important than length of stay in hospital).

Eventually, the design error dimension can be expressed by the priority of outcome measures as an ordinal variable according to GRADE [13].

**Conceptualization of the error matrix**

A four-step matrix can be constructed, building upon the three dimensions: systematic error, random error, and design error. Matrix step I ranks the identified studies according to the dimensions of systematic errors and random errors. Matrix step II orders the studies according to the design errors. Matrix step III assesses the three dimensions of errors in studies. Here, a ‘Manhattan-like’ error matrix is constructed where the best evidence is represented by the largest skyscrapers located on the ‘upper-west side’. Matrix step IV assesses the size and direction of the intervention effect, e.g., by calculating the number-needed-to-treat to obtain benefit or to harm one patient.

The principle of the matrix approach can be used in different situations. The overall effort in research should be to minimize all three risks of errors before the size and the direction of the intervention effect can be assessed reliably.
The ‘algorithm’ of the matrix approach is generally applicable to all kinds of interventions, although details may differ according to the specific clinical question. Or, the character of the three dimensions remains the same, while according to the specific question details may differ, like: the preferred hierarchy for levels of evidence, the chosen formula for standard error (RR, ORpeto or any other association metric), and the types of outcomes.

Results

The application of this four-step matrix is illustrated with two examples: peri-operative beta-blockade initialized in relation to surgery versus placebo for major non-cardiac surgery, and antiarrhythmics for maintaining sinus rhythm after cardioversion of atrial fibrillation.

Example 1: Initiating peri-operative beta-blockade for major non-cardiac surgery

A clinical question in PICOT structure illustrates this model. Is initiating peri-operative beta-blockade effective in patients undergoing major non-cardiac surgery?

Patients: patients undergoing major non-cardiac surgery; Intervention: initiating peri-operative beta-blockade; Control: placebo; Outcome measure: mortality, myocardial infarction, and stroke; Time: follow-up of at least 30 days.

We searched in CENTRAL in The Cochrane Library, PubMed, EMBASE, and personal files for all article types up to October 2009, in all languages. Specific searches using the terms ‘beta-blockade’, ‘peri-operative’, ‘placebo’, ‘mortality’, ‘randomised’, and ‘non-cardiac surgery’ were undertaken. The search resulted in multiple publications relevant to our question. References were selected from journals on the basis of importance and relevance [50-58]. We included the publications in our matrix evaluation by extracting information on all-cause mortality, cardiovascular mortality, non-fatal myocardial infarction, and non-fatal stroke. However, the matrix may easily be extended to other outcomes.

In step I, we assessed the systematic error and the random error for the chosen outcomes of each study (Figure 3, Table 3). In step II, we evaluated the design error (Figure 4). In step III, we constructed the three-dimensional matrix (Figure 5). We did not elaborate on the matrix step IV in this example.

From Figure 5 it can be concluded at a glance that peri-operative beta blockade does not reduce mortality in patients undergoing major non-cardiac surgery. Peri-operative beta-blockade in these patients seems to increase all-cause mortality. However, peri-operative beta-blockade does reduce non-fatal myocardial infarction on the expense of an increased cardiovascular mortality and an increased rate of non-fatal stroke.

Example 2: Antiarrhythmics for maintaining sinus rhythm after cardioversion of atrial fibrillation [59]

The conclusion of this Cochrane review focuses on the significant increased mortality associated with use of class 1a antiarrhythmics (odds ratio 2.39; 95% confidence interval (CI) 1.03 to 5.59) [59]. The data of this outcome in class 1a antiarrhythmics in this review as well as in the included randomised trials [60-67] were analysed using the matrix error approach.

In step I, we assessed the risk of systematic error and the risk of random error for the chosen outcome of each study (Figure 6, Table 4). In step II, the design error should be evaluated by assessing multiple outcome measures. However, in this example we only consider the outcome ‘all-cause mortality’, since other outcomes were found to be not statistically significantly different [59]. Therefore, no figure of step II is shown. In step III, we constructed the three-dimensional matrix (Figure 7). We did not elaborate on the matrix step IV in this example, since the available studies are not internally valid (high risks of both systematic and random error).

From Figure 7 it can be concluded at a glance that there is both substantial risk of systematic and random error involved in the evidence available so far considering mortality associated with class 1a antiarrhythmics. The best available level of evidence 1c study shows substantial risk of random error (0.43) and the best available level of evidence 1b study shows high risk of random error (0.78). So, the conclusion in the Cochrane review of a significant increased mortality is based on data with high risks for both systematic and random errors, and should therefore be considered unreliable.

Discussion

The aim of our matrix is to facilitate the overview of evidence in clinical intervention research. The matrix can serve as a tool to provide visual assessment of reliability of observations with respect to systematic error, random error (internal validity), and design error (external validity).

The matrix should not replace the thorough process of systematically reviewing evidence and profound evaluations of data, but could be integrated within these research activities as a tool for overviewing the results. Also, this matrix is not an absolute measure of the risks of errors. The position of studies in relation to each other is relative rather than absolute.

There is a lack of awareness of the importance of the ‘play of chance’ for the reliability of study findings. Ordering the standard errors of the studies might be a tool for ranking studies according to the level of random error. We have used natural logarithm (ln) transformations for calculating standard errors, although the logarithm with the base 10 may be used without producing different conclusions.
As an alternative, the Bayes factor can be considered [37,68]. The Bayes factor is a likelihood ratio comparing one hypothesis versus another, and, therefore, varies with the definition of the possible alternative hypotheses. The Bayes factor is a summary measure that provides an alternative to the p-value for the ranking or the flagging of associations as 'significant' [69]. The Bayes factor:

\[
\text{Bayes factor} = \frac{\text{Probability (Data, given the null hypothesis)}}{\text{Probability (Data, given the alternative hypothesis)}}
\]

Table 3 Ordering of evidence according to levels of evidence (systematic error), standard error (random error), and outcome measures (design error) in peri-operative beta-blockade versus placebo for major non-cardiac surgery (example 1)

| Level of evidence | All-cause mortality | Cardiovascular mortality | Non-fatal myocardial infarction | Non-fatal stroke |
|-------------------|---------------------|--------------------------|--------------------------------|-----------------|
| Bangalore [50]    | 1a                  | 0.12                     | 0.16                           | 0.10            | 0.28            |
| Poise [51]        | 1b                  | 0.13                     | 0.17                           | 0.10            | 0.33            |
| MaV5 [52]         | 1b                  | 1.07                     | Z                              | N               | 0.66            |
| Dipom [53]        | 1b                  | 0.34                     | 0.48                           | 0.91            | Z               |
| Mangano [54]      | 1b                  | 0.85                     | 1.22                           | 1.22            | 1.11            |
| Bangalore [50]    | 1c                  | 0.11                     | 0.15                           | 0.09            | 0.28            |
| Wetterslev [55]   | 1c                  | 0.24                     | N                              | 0.23            | N               |
| Poldermans [56]   | 1d                  | 0.76                     | 0.76                           | Z               | N               |
| Lindenuer [57]    | 2b                  | 0.02                     | N                              | N               | N               |
| AHA Guidelines [58]| 5                   | N                        | N                              | N               | N               |

Z: outcome with zero-events in one or both treatment arms which makes SE incalculable; N: no data.

Some outcome measures may be correlated (e.g. cardiovascular mortality is included in all-cause mortality).

In this example the formulas for SE of lnRR, for individual studies and SE of lnRRcombined for meta-analysis were used.
Figure 4 Matrix step II, ordering of evidence on peri-operative beta-blockade versus placebo for major non-cardiac surgery according to importance of outcome measures (design error) and levels of evidence (systematic error) (example 1). The outcome measures have been adapted to the beta-blockade question.

| Level of evidence | Outcomes | Primary | Secondary |
|-------------------|----------|---------|-----------|
|                   | all-cause mortality | cardio-vascular mortality | non-fatal myocardial infarction | non-fatal stroke | brady-cardia | hype-tension | other |
| Level 1a          | Bangalore 2008 (50) | Bangalore 2008 (50) | Bangalore 2008 (50) | Bangalore 2008 (50) |          |          |       |
| Level 1b          | Mangano 1996 (54); Dipom 2005 (53); Mantis 2006 (52); Poise 2008 (51) | Mangano 1996 (54); Dipom 2005 (53); Mantis 2006 (52); Poise 2008 (51) | Mangano 1996 (54); Dipom 2005 (53); Poise 2008 (51) | Mangano 1996 (54); Dipom 2005 (53); Mantis 2006 (52); Poise 2008 (51) |          |          |       |
| Level 1c          | Bangalore 2008 (50); Wetterslev 2006 (55) | Bangalore 2008 (50); Wetterslev 2006 (55) | Bangalore 2008 (50); Wetterslev 2006 (55) | Bangalore 2008 (50) |          |          |       |
| Level 1d          | Poldermans 1999 (56) | Poldermans 1999 (56) | Poldermans 1999 (56) |          |          |          |       |
| Level 2a          |          |          |          |          |          |          |       |
| Level 2b          | Linderofer 2005 (57) |          |          |          |          |          |       |
| Level 3a          |          |          |          |          |          |          |       |
| Level 3b          |          |          |          |          |          |          |       |
| Level 4           |          |          |          |          |          |          |       |
| Level 5           | AHA Guideline 2007 (58) | AHA Guideline 2007 (58) | AHA Guideline 2007 (58) | AHA Guideline 2007 (58) |          |          |       |
Figure 5 Manhattan-like three-dimensional matrix building upon the risks of systematic error, random error, and design error. The evidence with the lowest systematic, random, and design error is represented by the tallest skyscrapers, located on 'the upper west side'.

a. Outcomes with benefit of peri-operative beta-blockade versus placebo.

b. Outcomes with harm of peri-operative beta-blockade versus placebo.

A 'quick guide' to the perception of the figure: If you want to know what the evidence is for peri-operative beta-blockade to influence myocardial infarction: go to the yellow bars and read 1) Level of evidence (the risk of systematic error) and 2) standard error (the risk of random error). Data with risk of systematic error > level 2b and random error SE > 1.0 were omitted from the figure. The guidelines, which advocate the use of peri-operative beta-blockade, were not included in this figure since the systematic error is level 5 and the random error cannot be calculated (not based on data) [58]. The SE of outcomes with zero events cannot be calculated either. From these 'benefit' and 'harm' Manhattan figures, one can see at a glance that beta-blockers may provide benefit to patients in terms of nonfatal myocardial infarction (yellow bars). However, one can also see that beta-blockers may cause harm to patients in terms of all-cause mortality (red bars), cardiovascular mortality (blue bars), and nonfatal stroke (green bars). Reading the dimension of systematic error it is immediately clear that there is level 1a evidence available for all these four outcome measures. Reading the dimension of random error on this systematic error level of evidence shows that there is a small risk of random error considering all-cause mortality (0.12), cardiovascular mortality (0.16), and nonfatal myocardial infarction (0.10), and a moderate risk of random error considering nonfatal stroke (0.28). It is clear at a glance that the best available evidence does not support peri-operative beta-blockade for major non-cardiac surgery. SE = 0 to 0.10 = ignorable risk of random error. SE = 0.10 to 0.20 = small risk of random error. SE = 0.20 to 0.30 = moderate risk of random error. SE = 0.30 to 0.50 = substantial risk of random error. SE = >0.50 = high risk of random error. A clean version for creating a Manhattan figure can be obtained at the Copenhagen Trial Unit's homepage (http://ctu.rh.dk).
or simple approximations can be very difficult or even impossible to implement for the clinician, since a search for the maximum of the multidimensional posterior may be required for each association [69]. This also includes the asymptotic Bayes factor introduced by Wakefield [69]. In contrast to the Bayes factor, it is possible to calculate the standard error and when available it provides a tool for comparison of the risk of random error between studies of the same intervention.

The aim of minimising error risks according to the three dimensions actually combines the methodological efforts of falsifying any alternative hypothesis in the evaluation of an intervention. Thereby, the matrix concept visualises how far the scientific process has evolved to fulfil Poppers falsification criterion stating that researchers should primarily engage trying to falsify any relevant alternative hypothesis and not only the null hypothesis [5]. The minimisation of systematic errors and random errors, by providing ample room for the null hypothesis, as well as measuring important outcomes is the most audacious attack on any realistic alternative hypothesis.

### Table 4 Ordering of evidence according to levels of evidence (systematic error), standard error (random error), and outcome measures (design error) in antiarrhythmics for maintaining sinus rhythm after cardioversion of atrial fibrillation (example 2)

| Level of evidence | Standard error | All-cause mortality |
|-------------------|----------------|---------------------|
| Byrne-Quinn [60]  | 1d             | 2.02                |
| Hillestad [61]    | 1d             | 2.00                |
| Karlson [62]      | 1d             | 1.42                |
| Lloyd [63]        | 1d             | 1.55                |
| PAFAC [64]        | 1b             | 0.78                |
| Sodermark [65]    | 1d             | 0.73                |
| SOPAT [66]        | 1b             | 1.51                |
| Steinbeck [67]    | 1d             | Z                   |
| Lafuente-Lafuente [59] | 1c   | 0.43                |

Z: outcome with zero-events in both treatment arms which makes SE incalculable.

In this example the formulas for SE of lnORpeto,i for individual studies and SE of lnORpeto for meta-analysis were used.

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**Figure 6** Matrix step I, ordering of evidence according to systematic error (in levels of evidence) and random error (measured by standard error) considering all-cause mortality in antiarrhythmics for maintaining sinus rhythm after cardioversion of atrial fibrillation (example 2). Compare this figure with Figure 3: the studies in this figure are located on the right side of the figure (all SE > 0.40), in contrast with Figure 3 where the studies are concentrated on the upper left side of the figure (six studies with SE < 0.40).
Figure 7 Manhattan-like three-dimensional matrix building upon the risks of systematic error, random error, and design error. The evidence with the lowest systematic, random, and design error is represented by the tallest skyscrapers, located on ‘the upper west side’. 

- a. Outcomes with benefit of antiarrhythmics for maintaining sinus rhythm after cardioversion of atrial fibrillation. 
- b. Outcomes with harm of antiarrhythmics for maintaining sinus rhythm after cardioversion of atrial fibrillation. 

A ‘quick guide’ to the perception of the figure: If you want to know what the evidence is for antiarrhythmics for maintaining sinus rhythm after cardioversion of atrial fibrillation to influence all-cause mortality: go to the red bars and read 1) Level of evidence (the risk of systematic error) and 2) standard error (the risk of random error). Only the Cochrane review and the trials included in this systematic review were considered in this example. Data with risk of random error SE > 1.0 were omitted from the figure. The SE of outcomes with zero events cannot be calculated. From these ‘benefit’ and ‘harm’ Manhattan figures, one can see at a glance that there is no benefit at all and that ‘the upper west side’ is empty. Class 1a antiarrhythmics might increase mortality; however, since high risks for both systematic error and random error are present there is insufficient evidence for reliable conclusions.

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If an array of progressively qualified attacks fails to support the null hypothesis then we can reliably trust the intervention to be either beneficial or harmful.

The conclusion based on an assessment of the evidence using the matrix approach may be implemented into clinical practice or serve as an incentive for new research. The matrix facilitates the identification of lacunae in our knowledge and is likely to benefit the process of developing evidence-based guidelines.

Preference for the highest evidence
One has to be aware of the multiple forms of bias, potentially present in evidence below level 1 (Table 1). Several examples illustrate that large, apparently beneficial intervention effects from lower level evidence, even from randomized trials [54,56,70], may eventually be reversed to harmful effects when new high-quality evidence appears [50,71]. This is where the three dimensions of error are of central importance in providing a tool for reliability assessment.

Limitations
Apart from the three error dimensions influencing the reliability of data, other factors play a role in incomparability and uncertainty of inferences. Many reports of studies appear incomplete, and the lack of details raises questions. Incomplete reporting limits interpretation, but more importantly, this reporting factor should be distinguished from the methodological quality of the trial [72].

Statements like CONSORT [73], PRISMA [74], and MOOSE [75] aim to improve and to maximize the amount and correctness of information to be retrieved from publications. These guidelines also create awareness among researchers about the most important issues to report so that the quality of future research may increase. By following reporting guidelines the yield of the research question is likely to be increased (phase 1 in Figure 1).

Standard error does not consider testing of multiple outcomes and multiple testing on accumulating data, which may also induce risks of random error due to multiplicity as well as correlations.

The division of all outcomes into ‘primary’ and ‘secondary’ outcome measures can be helpful as this division sets the standards for the evaluation of interventions. However, this division is artificial, and outcome measures, situated on the border of primary and secondary outcomes, exist. For example, one can argue that quality of life is a primary outcome rather than a secondary outcome. Further, there is also a quantitative aspect in the artificial division into primary and secondary outcomes. Small significant differences in primary outcome measures (e.g., bile duct injuries in patients undergoing cholecystectomy) may be found favouring one intervention, while large differences in secondary outcome measures (e.g., costs) may favour the comparator. Eventually, one may prefer the larger advantages in secondary outcomes to the smaller disadvantages in a primary outcome measure.

Another limitation in the outcome measure dimension is that often outcome measures are correlated and mostly this correlation is ignored. For example when mortality is an outcome measure and complications is another, which again counts deaths as complications, then there is a correlation between the two outcome measures. Authors usually carry out multiple univariate analyses ignoring correlations between outcome measures.

Step IV of the matrix includes the assessment of the size of the intervention effect, e.g., expressed in numbers-needed-to-treat to obtain benefit or to harm one patient with the intervention. This step is the last one since it is irrelevant to consider effect sizes and their directions if a study does not appear to be internally and externally valid.

Another aspect to consider is heterogeneity [76,77]. Statistical heterogeneity reflects the between trial variance of meta-analytic intervention effect estimates rather than the play of chance [76]. Clinical heterogeneity, however, represents differences in populations, procedures, or interventions in daily practice. All these factors of clinical heterogeneity, together with concordance of in- and exclusion criteria should be considered whenever we want to implement results of available evidence. Assessment and consideration of heterogeneity or diversity, therefore, forms the final step before new evidence is implemented. Assessment of heterogeneity is not included in our matrix.

Conclusions
Assessment of risks of systematic error, random error, and design error are essential factors in evaluating evidence and drawing conclusions. We used the standard error in our matrix to rank studies according to their risk of random error. The risks of these error types were incorporated into a three dimensional matrix to create a schematic overview of the internal and external validity of the evidence, seen at a glance.

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Additional material

Additional file 1: Table S1. Word DOC table showing the sustained scientific process.

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