The future of evidence-based medicine: is the frog still boiling?

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ANALOGY BETWEEN LAW AND MEDICINE

As stated previously, for more than 2,000 years personal experience, anecdotes, and bias have dictated medical practice [1]. This editorial will elaborate further in depth on a paper written in 2013, entitled “Evidence-based medicine: are we boiling the frog?” concluding that in employing an analogy to criminal law, even if the evidence is beyond a reasonable doubt, it is rarely unequivocal or indisputable; thus, evidence is not synonymous with truth [1]. The opinions of medical leaders of successive eras have held sway and have done so from the days of Hippocrates and Galen. Dupuytren caused untold harm by proclaiming that under no circumstances could a structure as insignificant as the appendix be responsible for any abdominal mischief [2]. The traditional hierarchical training structure perpetuated such dogma whereby the consultant’s word was law.

Evidence-based medicine is defined as ‘The conscientious, explicit and judicious use of current best evidence in making decisions about the care of individual patients.’ However, as medicine is not an exact science as such, there may be many different ways to treat and cure our patients. Medicine, therefore, is an uncertain science [1]. Medical practitioners have followed their legal colleagues and nowadays scientific proof, based on the best available evidence, is required to substantiate current practice [1]. In law, decisions in civil cases are made on the basis of “more likely than not” where the balance of probability needs to be only greater than fifty percent. Criminal verdicts, which may incur severe penalties, require the evidence to be beyond reasonable doubt. Scientific proof must reach a higher level of certainty, as compared with the legal system [3]. This search for scientific proof has resulted in the construction of the evidence pyramid with the supposed best level of evidence provided by meta-analysis at the apex, the least reliable being personal opinions at the base (Fig. 1). In between these, in descending order of accuracy, lie systematic reviews, randomized controlled trials, cohort studies, case-control studies, and case reports.

As with legal argument, the evidence provided by each level has been contested, strong opinions being voiced by opposing camps [4, 5]. Paradoxically, personal eye-witness testimony on which a verdict may hinge in a court of law is relegated to the least reliable level of evidence in medicine, namely the individual case report. Strong evidence in clinical research is reportedly underpinned by the randomized controlled trial (RCT) with appropriate numbers of patients, assignment, and blinding. From the compilation of all of these, a meta-analysis is the advocate of the standard of care. Adherence to the requirements for reliable clinical research will ensure, in the majority of studies, that the results are valid [5]. Even when performed within such strict confines, however, bias may confound the results. This is especially true of intervention trials, the studies to which the RCT is most pertinent.

NEGATIVE STUDIES INDUCE PUBLICATION BIAS

A meta-analysis is only as good as the studies selected and inevitably can include only those published. There exists, however, a publishing bias against studies with negative or inconclusive findings in both drug and equipment trials.
[6–8]. Many reasons account for this. Investigators may lose interest in an inconclusive research result; sponsors may be unwilling to release detrimental information concerning their product. Personal opinion may sway reviewers and editors. The exclusion of unpublished data will inevitably skew the findings of any meta-analysis [9]. Clinical trials in which the results show a significant difference are three times more likely to be published than those with insignificant findings. The time to publication is also significantly longer with 80% of positive trials but only 20% of negative trials being published within ten years of completion [6]. An undetected positive bias may also exist in the form of duplicate publication [7]. The same study portraying a different aspect of the data may be published under an alternative sequence of authorship and be included twice in a meta-analysis. In the case of highly significant findings, the treatment effects will be overestimated.

For an unbiased meta-analysis, studies must be carefully scrutinized for potential duplication and a liberal time frame must be allowed for the inclusion of studies containing inconclusive or negative data. Unpublished information is also relevant, and for more than two decades a plea has been made for the registration of all interventional studies, thus allowing access to such data [8–11]. Failure to publish, or allow access to all information has been suggested as being tantamount to scientific misconduct [12]. Recently, the International Committee of Medical Journal Editors (ICMJE) has become insistent on the criteria for acceptance for publication, a move which appears to have succeeded in encouraging the registration of clinical trials [13].

Bias against negative findings goes beyond non-publication where results which are detrimental to the tested product may be deliberately suppressed, the positive aspects alone being published [14–18]. The last 20 years have seen a trend from a time when investigators controlled all aspects of a study from design, data collection, analysis, and publication to an era where the sponsor assumes control and offers financial incentives to enrol patients [11]. A contract between researchers and funders is often agreed upon whereby commercial sponsors control the database, perform the statistical analysis, and after that, present the finished article to the researchers. Unless the researchers have direct access to the entire data and are allowed to conduct an independent review, there remains the opportunity for sponsors to withhold damaging information or skew the data in favour of a positive outcome [18]. Information describing the detrimental effects may only surface at a later date after patient harm has occurred [19].

**EFFECT OF SPONSORSHIP BY FOR-PROFIT ORGANIZATIONS**

Funding of new drug trials is almost invariably, and understandably, provided by a parent company. Fiscal contributions go beyond the costs of delivering the drug and conducting the study, however, and financial incentives are often offered for individual researchers, either personally or for the department to which they belong. In a poll of 459 academic departments, almost two-thirds revealed financial ties with the pharmaceutical or medical equipment industry including acting in a consultative capacity,
on advisory boards, and speakers’ bureaus [20]. Does this have an impact on study outcomes? The answer would appear to be that it does. An analysis of 159 trials involving 12 different specialties concluded that there was a significant finding in favour of the trial drug if the study was funded by for-profit organizations. This could not be explained by methodology, statistical analysis, or the type of study [21]. A similar review of meta-analyses found that 51% of studies funded by for-profit organizations found in favour of the experimental drug compared to only 16% of studies sponsored by non-profit organizations [22]. Trials funded by for-profit organizations were more than 5 times as likely to recommend the trial drug as the treatment of choice. This must cast doubt upon the validity of certain conclusions. As stated by Angell, a former editor of the New England Journal of Medicine for two decades, “Physicians can no longer rely upon the medical literature for valid and reliable information”. She reluctantly concludes that these products are not nearly as effective as the publications suggest [17].

This is not a subjective appraisal and has been substantiated in the literature. An analysis of highly cited trials published in the three journals with the highest impact factor (The New England Journal of Medicine, The Lancet, Journal of the American Medical Association) and those with an impact factor greater than seven, showed that 30% of trials initially reporting highly significant positive findings were found in subsequent studies to either overestimate treatment effects or show no benefit [23], a phenomenon confirmed in more recent studies [24]. Admissible studies covered a broad cross-section of medical and surgical specialties. The danger lies in such initial studies being adopted blindly into clinical practice without confirmation which, due to the time-consuming nature of clinical research, may take years to produce. Scepticism is essential, especially when the initial results show an exceptional outcome. Unfortunately, it is these very studies which are more readily accepted without validation.

The effect of funding affects more than pharmacology or new device trials. Consensus guidelines and statements written by panels of experts are frequently supported by the industry, and the sponsoring company may have financial affiliations with the members of such panels [17, 25]. This has led to specific journals refusing to accept for publication consensus statements, reviews, or commentaries which recommend drugs, equipment or other profitable interventions if authors have declared a financial conflict of interest [25].

**GHOST AUTHORS, GUEST AUTHORS, AND DATA FABRICATION**

Ghost authorship takes two forms. In its benign form, professional medical writers skilled in the use of language may improve a manuscript without altering the scientific content (perhaps they should be regarded as the real “spin doctors”). They may be acknowledged in the text but will not appear on the authors’ list. A more malignant tendency has spread in industry-sponsored studies whereby the initial draft of the manuscript, namely that which shapes the final article, is compiled by company employees. Key opinion leaders that are academically affiliated, are often sourced as principal or second authors without having provided any substantial contribution to the study or article [26]. Articles in high-impact journals under the authorship of respected academics undoubtedly influence clinical practice. Representatives of the companies use them to promote their product while clinicians independently garner evidence from the current literature regarding best practice. In the most adverse scenario, pharmaceutical companies engage the services of contract research organizations to organize the trial, the company then analyses the data, use their employees to write the manuscript, source a respected academic to act as the principal author, and pay communication companies to accelerate publication [27]. The exact incidence of ghost authorship is unknown.

From painted mice to post-operative pain relief, colloids to cardiac protection, instances of trial misconduct and data fabrication have raised their ugly heads. This literary hydra is a cause for serious concern and casts an uncomfortable shadow over medical evidence. A recent analysis reported that 2% of scientists admitted to fabricating or modifying data at least once and one-third confessed to questionable research practices. Interrogation of colleagues revealed more alarming figures of 14% concerning data falsification, and 72% for dubious scientific behaviour [28].

**STATISTICAL VERSUS CLINICAL SIGNIFICANCE**

Statistical significance is the cornerstone of the bridge between clinical trials comparing two or more groups and their conclusions. Simply put, it produces a mathematical probability of whether the results of a study are due to chance, a five percent risk of the effects being falsely positive deemed acceptable. As amusingly described by Hall [29], this is not due to divine intervention but was the learned opinion of the statistician Fisher. In essence, this is, therefore, based on subjective expert opinion, the antithesis of evidence-based medicine.

However, in reality, clinical relevance has little or nothing to do with statistical significance and must not be confused with biological importance [29]. In fact, Luus et al. [30] suggest that clinically relevant differences and statistical significance concur only by coincidence. These authors pointed out that clinicians must have a good understanding of the results (and the statistical methods used), while statisticians need to understand the clinical problem in order to generate analytical results in line with meaningful clinical conclusions.
As the aim of trials is to determine whether the aspect under scrutiny will have an impact on clinical practice, the results should, therefore, be expressed in clinical and not mathematical terms. The latter are largely susceptible to sample size and clinical differences may be statistically significant, albeit meaningless [1]. The reverse also holds true if the sample size is too small. In his book "The Last Well Person", Hadler [31] argues that an absolute reduction in the mortality rate of 1.9% in the West of Scotland statin trial [32], although statistically significant, is clinically irrelevant. The data however are expressed as a relative risk reduction of 29%, a more impressive statistic. He suggests that no study can control for all confounders and an absolute difference of less than 2% should be viewed with caution.

More recently, although first described almost 50 years ago, the concept of intention to treat (ITT) analysis for RCT's has gained widespread advocacy, especially amongst statistical purists [33]. Using this approach, data are analysed according to randomization regardless of dropouts, the crossover between treatment groups, or missing data. They argue that this technique assures homogeneity between randomized groups and minimizes a type I error (the incorrect rejection of the null hypothesis). Clinicians argue that this concept is flawed, increases the risk of a type II error (the false acceptance of the null hypothesis) and propose that two other techniques provide more meaningful clinical information. These are a per protocol (PP) analysis where only compliant patients are included, or in the event of crossovers an as-treated (AT) analysis where subjects who received a particular treatment are included regardless of the group to which they were initially assigned. If the number of patients who drop out or crossover is minimal, the ITT concept holds true, but if higher than 20% the results will be inaccurate [34]. Furthermore, the choice depends on the question to be answered. If this relates to the efficacy of an intervention, then PP or AT analyses are superior. This concept is ably demonstrated by the recent EVAR trial [35] where subjects deemed too unfit for open repair were randomly allocated to either EVAR or no intervention. In the latter group, 34% underwent repair for a variety of reasons. ITT analysis showed a significant reduction in aneurysm-related deaths but no improvement in overall survival at eight years. PP analysis not only showed a much more significant reduction in aneurysm-related fatalities but a marked increase in overall survival. Both effects were even more evident using the AT technique. Hall suggests that in addition to an ITT analysis, authors should declare non-compliance and provide a summary that excludes these patients [29].

The number needed to treat (NNT) is of even greater clinical relevance, the reciprocal of absolute risk reduction, which defines how many patients need to be treated for one to gain a benefit [1, 36], which may also be expressed per 1,000 patients. Of importance, this calculation does not correlate with probability values but assesses clinical impact. Equally, or of perhaps greater concern is the number needed to harm (NNH) which assesses the possible adverse consequences of a particular intervention. The POISE (Post Operative Ischaemic Evaluation) study epitomizes these concepts [37]. This is the largest randomized controlled trial to assess whether peri-operative beta blockade can lower the risks of post-operative cardiovascular events. A highly significant reduction of non-fatal myocardial infarctions was found in the treated group with an NNT of 66. The incidence of stroke, however, doubled with the NNH being 200. For every three patients spared a cardiac event, one would potentially suffer a cerebral insult. In those who suffered a stroke in the treatment group, however, only 15% regained full function while 26% were left severely incapacitated. The choice between the risk and sequelae of a non-fatal myocardial infarct versus a disabling stroke is a matter of clinical judgement and patient preference, not mathematical probability [1].

THE EVIDENCE PYRAMID

Meta-analysis lies at the summit of the evidence pyramid and is regarded as the final arbiter in areas of uncertainty with conflicting or scarce data (Fig. 1). In essence, it is a statistical averaging technique undertaken in the hope that by pooling data, any errors in individual study design or implementation will be minimized. Errors in clinical trials may be random or systematic [38]. The former is unpredictable and may skew data in both positive and negative directions. An increase in sample size reduces its occurrence [1]. Systematic errors are not eliminated by increasing the sample size, and arise when a trend in the data occurs that is actually false [1]. This occurs as a result of three types of bias, namely selection, misclassification, and confounding.

SELECTION BIAS

Selection bias occurs when a test is inadvertently skewed to favour a subset of patients, as in that cited by Tobin and Jubran [38] where patients who failed a weaning test were automatically excluded from a trial of spontaneous breathing.

MISCLASSIFICATION BIAS

Mischclassification bias describes the error of placing patients in an incorrect category, resulting in a heterogeneous rather than homogeneous population being placed under scrutiny [1]. This is especially true where standard therapies are normally titrated against physiological end-points rather than fixed dose regimens [1, 39]. Deans et al. [39] cite the ARDS low tidal volume trial as a prime example where patients were randomized to fixed tidal volumes of either 6 mL kg⁻¹ or 12 mL kg⁻¹ whereas the standard practice would be to titrate
treatment in accordance with airway pressures and compliance. Randomisation to a fixed dose would result is some patients in the low tidal volume group being underventilated and some in the high volume group being overventilated. As emphasized by Vincent [40], the identical scenario pertains to transfusion triggers. Younger patients without coronary artery disease may tolerate a lower haemoglobin level than the elderly cardiopath and conversely, overtransfusion in the young may lead to a detrimental effect [1,40]. Such insufficient or excessive treatment may contribute substantially to differences in the trial results. Thus, ignoring heterogeneity violates sound statistical practice [41]. In truth, there is no such entity as a perfectly matched RCT, for although the control and treatment groups may demonstrate no statistically significant difference between the means of any parameter, each group, even if normally distributed, will inevitably be heterogeneous.

**CONFOUNDING BIAS**

Confounding bias refers to the mistaken relationship found between two variables because of a third unaccounted factor. Even if selected studies are homogeneous, disagreement may arise amongst reviewers who, when presented with identical sets of information, arrive at diametrically opposite conclusions [42]. As with any other scientific publication, meta-analyses must be interpreted with caution. For over fifteen years, there has been an appeal for individual patient data to be used, but even if sourced and included, systematic error, publication, and selection bias may still exist [43]. The most recent example is that of the systematic review and meta-analysis of hydroxyethyl starch for volume resuscitation which purports to find a higher incidence of acute kidney injury with the use of starch solutions [44]. Selected for inclusion in the analysis were such diverse groups as burns, trauma, septic and non-septic ICU patients, organ donors, acute pancreatitis and those resuscitated post cardiac arrest. This study concludes by stating, “Clinical use of hydroxyethyl starch for acute volume resuscitation is not warranted due to serious safety concerns”, a conclusion that may pertain to a sub-group somewhere but cannot be applied to such a diverse patient population. The consequence of such unfounded statements in a high impact publication is that many clinicians will abandon the use of colloids and revert to crystalloids which are known to have serious adverse effects if used as the sole resuscitation fluid.

The precept that systematic reviews should exclude observational studies has recently been challenged, and in situations where randomized trials are either few or of inferior design, non-randomised trials should be eligible for meta-analysis [45]. Given the multitude of problems in conducting randomized controlled trials in the critically ill, Vincent [40] has suggested that these should be abandoned and more attention paid to observational studies. This, he argues, would allow enrolment of almost the entire patient population under investigation with fewer exclusions, making such an approach more clinically relevant.

**ARE WE STILL BOILING THE FROG?**

But what of the boiling frog and its relevance to evidence-based medicine? There is a physiological anecdote (although this may be an urban legend1) that if a frog is placed in boiling water, it will understandably leap out immediately. If the water is initially tepid however, and slowly heated to boiling point, the frog will remain until boiled alive [1]. This example has been used in various scenarios including economics and global warming, to illustrate the concept that slow change may pass unrecognized until harm occurs [1]. The current United Nations Congress on climate change and international fiscal crisis epitomize this concept. Numerous examples exist in medical practice across all specialties; the pulmonary artery catheter in critical care; in surgery abdominal tension sutures when the abdomen is difficult or painful to close; bowel preparation for colonic surgery; the list could be considerable. Proof of benefit of any intervention is an understandable and laudable aim of any clinician and this, coupled with the desire to do no harm, has been the driving force for evidence-based medicine. This concept is not scientifically perfect and must not be viewed as exclusive [45, 46], especially in relation to practice guidelines. Despite claims that these represent the best standard of care, this does not hold true for all patients with the same disease. As observed by Osler, “Variability is the law of life and no two individuals react alike and behave alike under the abnormal conditions which we know as disease. The good physician treats the disease, the great one treats the patient”[47]. Evidence-based guidelines not only ignore patient heterogeneity but depending on the quality of evidence, may not define optimal treatment [48, 49]. Blind acceptance may cause harm, as illustrated by the use of potassium-sparing diuretics as the diuretic of choice by the American Heart Association resulted in a more than four-fold increase in prescriptions. Over the ensuing three years, there was a four-fold increase in hospital admissions for hyperkalaemia and an almost seven-fold increase in death from this complication.

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1The story was based on nineteenth-century experiments in which frogs were shown to stay in heating water as long as it was heated very slowly. The validity of the experiments is, however, disputed. Professor Douglas Melton, Harvard University Biology Department, says: “If you put a frog in boiling water, it won’t jump out. It will die. If you put it in cold water, it will jump before it gets hot — they don’t sit still for you”. (See also: https://www.fastcompany.com/26455/next-time-what-say-we-boil-consultant)
Although whimsically sarcastic, Smith and Pell rightfully conclude there is no evidence that a parachute prevents major harm when jumping from an aircraft, but common sense should prevail. They suggest the same should apply when considering the potential risk and benefit of interventions [51]. In light of the above it is essential that evidence-based medicine be viewed with an open mind. As in criminal law, even if the evidence is beyond reasonable doubt, it is rarely unequivocal or indisputable; evidence is not synonymous with truth [1]. From initially tepid waters the zeal for evidence-based practice has now reached boiling point and if the shortcomings are not appreciated, evidence-based medicine may itself become a boiled frog. Even in modern practice Osler’s aphorism still holds true, namely “Medicine is a science of uncertainty and an art of probability”.

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