Computational modelling of lung injury: is there potential for benefit?

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State-of-the-art medical care of the victims of current conflicts is generating large quantities of quality clinical data as a by-product. Observational research based on these data is beginning to have a profound influence on the clinical management of both military and civilian trauma patients. Computational modelling based on these datasets may offer the ability to investigate clinical treatment strategies that are practically, ethically or scientifically impossible to investigate on the front line. This article reviews the potential of this novel technology to aid development of treatment for blast lung and other unresolved medical scenarios.

Keywords: blast lung; respiratory failure; respiratory distress syndrome; model; computer simulation

1. INTRODUCTION

Many researchers feel that conducting even simple clinical trials can seem a Herculean task. The increasing bureaucracy of research governance, though of critical importance, tends to make the process slow, expensive and time consuming. Even more respect and admiration is therefore due to those who conduct clinical trials during the very real trials of battle. Outstanding efforts have recently been made by clinicians on the front lines in Afghanistan and Iraq into topics such as traumatic coagulopathy [1–2] and damage control surgery [3], and the important insights gained impact not only on future war casualties, but also on the victims of more mundane but equally fatal trauma closer to home [4].

The problems of conducting clinical trials in this environment are multiple. The military train of command does not extend to offering blanket ethical approval, and the concepts of informed consent and ethical oversight might seem very alien on the battlefield. Classical research methodology (for instance, randomization and blinding) can be very difficult or impossible. Patients have heterogeneous patterns of injury, and their numbers, although high in human terms, may offer limited statistical power. Indeed experimentation on the battlefield is prohibited by US federal law [5]. These difficulties lead to the evolution of practice based on retrospective, observational data empowered by the military’s abilities to disseminate best practice protocols rapidly to the front line and ensure that they are followed. These observational and cohort data are frequently used to generate questions that can then be investigated using animal models, the results feeding back into practice protocols amenable to repeated audit. While the evidence base is grade II or III at best [6], there seems no potential for true randomized controlled trials in this context.

2. WHAT DOES MODELLING HAVE TO OFFER?

Much scientific research is conducted using in silico modelling, the fields of climate research and economics being topical examples. While it would be foolish to claim to model reality in every detail, it is widely recognized that carefully developed, empirically based and validated computer models have much to offer. Models are in frequent use throughout medical research but are not always recognized as such; for example, many studies that utilize a correlation between surrogate and clinical outcomes rely on an underlying model, which relates the two. Likewise much statistical analysis used to evaluate the results is reliant on statistical models of data relationships.

Modelling is the reproduction of real-life shape, form or function [7–8]. It can be physical, mental, computational, statistical (probabilistic), mathematical, animal or laboratory. It is an attempt to simulate or recreate the important and interesting aspects of a question or system, while excluding the unpredictable and irrelevant aspects of organisms and populations that add heterogeneity, random variation and pragmatic obstructions. Such modelling may allow us to elucidate complex issues or to formalize or automate decision-making processes with a view to treating patients [9–12]. The valuable efforts of scientists in the battlefield can provide the empirical data on which a model can be built. Treatment strategies can be investigated in an in silico model rapidly and to a level of complexity not possible in animal models [13–14]. Even where there are gaps in the...
knowledge needed to construct a model, it is still possible to gain valid results provided the gaps are known and the resulting limitations acknowledged. A credible in silico model might best be considered akin to an early Petri dish investigation: an efficient, rapid and cheap method of rejecting grossly ineffective strategies and selecting those ripe for further investigation, perhaps in other models or direct to patient cohorts where the validity is strong. Indeed the concept of Monte Carlo testing of many subtly different protocols in an established model is a powerful one [15], and is perhaps the only method of selecting the 'best-fit' algorithm with a high level of fidelity. Furthermore, in silico modelling allows investigation of physiological situations that are impossible or unethical to reproduce in a controlled way in human studies, but which are, nonetheless, credible reflections of existing pathologies [16–17].

Our own respiratory computational model (the Nottingham physiology simulator) integrates respiratory physiology with mechanical ventilation, allowing subtle and precise alteration of more than 50 variables (figures 1–3). For instance, compliance and resistance to both gas and vascular flow to 100 representative alveolar units can be adjusted to mimic known pathological conditions. Downstream tissue perfusion outcome measures (such as arterial gas partial pressures) can be calculated after exposure to a myriad of ventilatory strategies. This model has been successfully deployed in characterizing deterioration in tissue oxygenation during apnoeic episodes, aiding our understanding of management of airway emergencies [16].

What are the questions confronting clinicians at the front that might be illuminated by the application of in silico modelling? The incidence of blast lung appears to be increasing owing to increases in the power of improvised explosive devices and their use in confined spaces [18–19]. The relative infrequency of blast lung [20], its geographical dispersion and the obvious chaos of the initial insult make traditional investigation virtually impossible for the reasons described above. There is an ethical obligation to minimize the number of animals used in research, although animal-derived data are likely to be a very important aspect of model development and subsequent validation. Blast lung's importance to the field is not doubted; it is the commonest cause of death in initial survivors of explosions [18]. Animal models suggest that relatively high pressures are required to trigger blast lung [21] and that victims exposed to real world pressure waves of this magnitude are often sufficiently close to the explosion for fragmentation injuries to predominate [22].

The ability to model individual lung units down to the alveolar level facilitates the study of pathologies whose distribution within the organ are heterogeneous (for instance, adult respiratory distress syndrome—ARDS), and our model has been validated for studying this condition [23]. Heterogeneous lung damage is also seen in blast lung where perihilar infiltrates are virtually pathognomonic [24]. Data extrapolated from ARDS patients have been used to guide therapy in these patients, although the pathophysiology, distribution of injury and progression may be quite different [25]. Thus, most centres have used a low-volume, pressure-limited strategy resorting to rescue therapies such as nitric oxide and ECMO when necessary [18] in blast lung. A validated model based on an empirical dataset gained in the field, and cross-validated against the same patient cohorts, animal models and laboratory models would allow the detailed investigation of unanswered questions; for example, the role of PEEP and the potential advantages of high-frequency oscillatory ventilation. In addition, an unlimited number of clinical scenarios can be investigated to suggest important contributory factors in poor outcome (e.g. massive haemorrhage, periods of hypotension or apnoea).

There is a substantial body of model evidence for physical patterns of blast injury, which has been

![Figure 1. Respiratory physiological variable control panel.](image-url)
undertaken both in the laboratory [26–28] and in silico [29–31] that could inform a preliminary physiological model. Indeed computational modelling was identified as a potentially fruitful avenue for blast lung research by a consensus workshop and in a recent review [22].

Integrated respiratory and cardiovascular models bring with them the possibility of further refining military trauma protocols already being successfully deployed on the battlefield. Such a model would permit detailed investigation of the response of tissue perfusion parameters to almost infinite constellations of pathologies and severity, allowing the development of sophisticated and individualized therapies. For instance, severe burns coupled with lung contusions and major haemorrhage presents a complex challenge, and resuscitation goals in this situation are uncertain. Current available evidence shows that as well as the choice of intervention, its precise timing is critical [32]. The value of the clinical acumen of experienced clinicians cannot be underestimated, but the ability to model responses to different resuscitation strategies may be very helpful (for instance, in rapidly determining the theoretical optimal timing and targets of cardiovascular resuscitation). Decisions regarding timing of risky procedures likely to worsen physiological status (for instance, global repatriation and surgical intervention) can be illuminated by computational models, which also allow for progression of underlying pathological insult, seen in conditions such as blast lung [25].

3. PRINCIPLES OF MODELLING

Modelling can be undertaken in a ‘top-down’ or ‘bottom-up’ methodology. The top-down approach involves construction of a model that appears correct
and then entering a process of validation against datasets with subsequent refinement and repeat investigation, building the ‘machinery’ of the model after the initial credible output is established. Bottom-up construction is scientifically more robust, involving building of a model from first physiological principles, but is slower and requires data that may not exist or may be extremely difficult to collect. The end result of both processes may be a model that looks remarkably similar; in any case, a key concept within modelling is that the model does not represent truth, but is close enough to it for practical purposes (in that no model is valid, merely fit for purpose). Once the model has been developed, validation against existing human and animal datasets, or if possible, prospective comparison against real-time patient cohorts will provide credibility. Investigation of treatment strategies can be begun in earnest at this stage, although the model should be continually developed in the light of emerging knowledge.

4. CONCLUSIONS

Military and civilian victims of war and terrorism have different pathologies and patterns of injury compared with those suffering civilian trauma. These patient cohorts have been very difficult to study historically but there is an increasing quantity and quality of data emanating from Iraq, Afghanistan and other military arenas. The human price that has been paid to make these data available makes the imaginative and
efficient use of these data a moral and practical imperative. The onus is on the scientific community to maximize the benefit derived by future victims, and computational modelling may well have a valuable role to play. The current development of clinical care on the battlefield by meticulous observational data collection and synthesis with current understanding may be considerably quickened by the application of systemic pathophysiological investigation with appropriately validated computational models. Furthermore, the data derived from animal models will accelerate the synthesis of these data into credible models, allowing us to truly translate basic science research into battlefield medicine. Modelling will allow us to open a conduit from human research (with its noisy data), animal research (with its limited mapping to human pathophysiology) to the development of real-world clinical treatment strategies; modelling aggregates our accumulated knowledge, expanding the importance and relevance of previous work, drawing together ideas and data from disparate sources and groups and forging a clear highway from isolated knowledge to human clinical practice.

REFERENCES


