Guidelines for using animal models in blast injury research

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ABSTRACT
Blast injuries are a significant problem for the military and an increasing problem for civilians. This has contributed to an increase in research in blast effects (a PubMed search using the term ‘blast injury’ found over 40 primary research papers for 2016 using animal models vs less than five papers for 2006). Blast injury research is required to enable improvements in body armour and post-injury therapies, for example; this will save lives as well as reduce the impact of such injuries on quality of life. Blast injuries are those caused by an explosive event and result in a wide range of injuries, therefore it is important that those involved in research in this area as well as those involved in the treatment of patients are united with respect to the classification of blast injuries to avoid misunderstanding and inappropriate interpretation of results. This North Atlantic Treaty Organization (NATO) panel has used the classification of blast injuries according to Department of Defense Directive 6025.21E.

INTRODUCTION
Blast injuries are a very complex phenomenon and frequently result in multiple injuries. One method to investigate the consequences of blast injuries is with the use of living systems (animal models). The use of animals allows the examination and evaluation of injury mechanisms in a more controlled manner, allowing variables such as primary or secondary blast injury for example, to be isolated and manipulated as required. To ensure a degree of standardisation across the blast research community a set of guidelines which helps researchers navigate challenges of modelling blast injuries in animals is required.

THE NEED FOR THE GUIDELINES
Blast injury is a very complex phenomenon and frequently results in multiple injuries. While data gathering from those casualties injured by an explosive event is extremely valuable, this does not provide all the answers and due to the sporadic nature of events evaluation of potential therapies in the target population, for example, is not a viable option. One method to investigate the consequences of blast injuries is with the use of living systems (animal models). The use of animals allows the examination and evaluation of injury mechanisms in a more controlled manner, allowing variables such as primary or secondary blast injury, for example, to be isolated and manipulated as required. Animal experiments can control for age, gender and other genetic parameters not possible when examining data from human subjects exposed to blast. The use of animals in blast injury research presents many challenges. Due to the complexity of blast injury it is unlikely that one model will be able to replicate all the relevant injuries and post-injury consequences, therefore it is highly likely that several models will be required. To ensure a degree of standardisation across the blast research community a set of guidelines which helps researchers navigate challenges of modelling blast injuries in animals is required. Existing guidelines for animal studies, for example, traumatic brain injury, are not designed for blast exposure.
Challenges that impact on blast injury modelling in living systems

The injuries sustained from a blast are influenced by a number of factors including the following:

- The physical loads from the explosive event, therefore the impact of any scaling issues needs to be addressed.
- Biological effects from the initial response in tissues as well as secondary effects (the final injury depends on the initial trauma, secondary responses and then any treatment effects).
- Species effects that can result in a failure to replicate the features of interest for human casualties (e.g., post-traumatic stress disorder).

In addition, researchers using animals for blast injury research have an ethical obligation to ensure that the research has scientific and clinical validity and thus ensure that no animals are used unnecessarily.

The complexity of blast injuries and the challenges of modelling such injuries in living systems highlight the importance of the experience of the research team and any research group undertaking blast research using living animals must have the appropriate capability, knowledge, skills and expertise to address the intended research questions.

THE GUIDELINES

The aim of the guidance is to ensure that experiments are validated and replicate the human condition or aspects of the human condition to enable the translation of the results.

The Guidelines are intended to provide a framework for scientifically valid methodological approaches to address the pathological consequences of blast exposures, and assist researchers during all stages of blast trauma animal experiments. The intention is that this will reduce interlaboratory variability and allow valid comparisons of results to be made.

The Guidelines are aimed at research scientists when planning, executing and reporting animal experiments for blast trauma; funding bodies when evaluating a proposed plan of work; and journal editors and reviewers to determine validity and relevance of research presented.

At the heart of these guidelines is good experimental design as this is fundamental to the translation of results from animal studies to clinical practice. The general principals of good experimental design in this context are listed below:

1. Study aim—what problem does the study address?
2. Study hypothesis.
3. Study methodology—how the experiment addresses the hypothesis.
4. Relationship to real-world operational conditions—appropriate levels of ‘blast’ exposure.
5. Choice of model:
   i. Which blast effect is being modelled, for example, primary, secondary, and so on. This will determine the choice of exposure environment, for example, primary blast requires one of the following: open field exposure, shock tube or blast tube.
   ii. Exposure conditions: exposure level and target positioning. Special consideration is needed in terms of positioning of specimen in the shock/blast tubes as well as orientation in relation to incident shock wave. Positioning the animal outside the shock tube results in exposure to a subsonic jet wind, this results in effects that are significantly different from those generated by a shock wave. It has been shown that both the pattern and severity of organ damage caused by blast depend on the orientation of the body towards the shock wave front. In addition, cognitive and behavioural responses in animals to blast are also dependent on the orientation of the animal. The choice of the animal holder is another important component in shock/blast tube experiments.
   iii. Species selection. Consideration needs to be given to the physiological responses to blast injury for the chosen species; the similarity of anatomical properties to humans must be considered; study feasibility must be considered; and model limitations must be acknowledged and discussed to ensure that results are neither misinterpreted nor overinterpreted. All these factors need to be considered to ensure reported results are appropriate and not misleading.
6. Data collection—it is not possible to provide a comprehensive list of parameters to be measured, and the exact data collected will depend on the actual research question. The choice of data collected as well as omitted data will need to be described and justified by the research team.
   i. The method of data collection must be described, for example, frequency of sampling must be appropriate to the parameter being assessed.
   ii. Sample timing must be justified, for example, the time course of disease process is likely to be species dependent and therefore needs to be appropriate.
   iii. Postexperimental analysis must be described and the statistical plan must be appropriate.
7. Limiting variability—not all variability can be eliminated but steps must be taken to reduce and limit its impact. This can be achieved by measuring as many critical parameters as possible and controlling the animal species across laboratories. In addition to experimental design validation is a critically important aspect of animal models and regardless of the research questions to be addressed the criteria every clinically and militarily relevant blast injury model should fulfil are the following:
   i. The injurious component of the blast should be clearly identified and reproduced in a controlled, reproducible and quantifiable manner (see the Guidelines for Reproducing Blast Exposures in the Laboratory).
   ii. The inflicted injury should be reproducible and quantifiable, and mimic components of human blast injury.
   iii. The injury outcome established based on morphological, physiological, biochemical and/or behavioural parameters should be related to the chosen injurious component of the blast.
   iv. The mechanical properties (intensity, complexity of blast signature and/or its duration) of the injurious factor should predict the outcome severity.

One aspect of validation may be dose–response relationships demonstrating that with the increasing intensity of blast exposure the biological responses will be more pronounced and the pathological consequences more severe. Dose–response studies are necessary to determine injury threshold and saturation values. Bowen curves, for example, provide excellent framework for experiments analysing the relationship between primary blast exposure(s) and tissue/organ damage. The Bowen curves have their limitations such as limited usefulness for chronic biological or psychological outcomes, as well as non-primary blast effects. Thus, new dose–response curves are needed based on appropriate scaling laws that would establish the relationship between individual blast components and biological/psychological outcome measures, while taking into account the size, composition and geometry of the exposed body and/or organ.
Studies undertaken with good experimental design using validated animal models will improve the state of the science for blast injury.

CONCLUSION
In an ideal world there would be no requirement for the use of living animals in blast injury research, however there are currently no non-living models that can integrate all the biological (cardiovascular effects, immunological effects, and so on) responses seen after injury. Therefore, animal experiments are necessary and they can generate valuable data regarding blast injury for the disease process and for the investigation of potential therapies. Animal models allow for a more controlled examination of blast injuries. One particular advantage of animal models is that results can be achieved from a relative small number of animals.

No animal model can replicate all the conditions of a blast event and the human’s response to blast exposure and decisions made by the researcher regarding the nature of each experiment has the potential to reduce the relevance of a study.

Limited reproducibility can be a concern in animal studies. This leads to translational problems, both between animal data and real-life blast events but also between different animal studies. Good monitoring of experiments and adherence to guidelines are important ways to decrease such problems. However, multicentre studies could also be an effective way to increase the usefulness of animal studies.

This guidance document provides a framework for the research community with the aim of improving experimental quality. It is anticipated that adherence to this guidance document will help reduce the following: the uncertainly regarding the nature of the blast injury being modelled; the variability and quality in study outcomes; and finally enhance the impact and translation of results such that patient outcomes will be improved.

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