A protocol for a scoping and qualitative study to identify and evaluate indications for damage control surgery and damage control interventions in civilian trauma patients

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ABSTRACT

Introduction: Initial abbreviated surgery with planned reoperation (damage control surgery) is frequently used for major trauma patients to rapidly control haemorrhage while limiting surgical stress. Although damage control surgery may decrease mortality risk among the severely injured, it may also be associated with several complications when inappropriately applied. We seek to scope the literature on trauma damage control surgery, identify its proposed indications, map and clarify their definitions, and examine the content and evidence on which they are based. We also seek to generate a comprehensive list of unique indications to inform an appropriateness rating process.

Methods and analysis: We will search 11 electronic bibliographic databases, included article bibliographies and grey literature sources for citations involving civilian trauma patients that proposed one or more indications for damage control surgery or a damage control intervention. Indications will be classified into a predefined conceptual framework and categorised and described using qualitative content analysis. Constant comparative methodology will be used to create, modify and test codes describing principal findings or injuries (eg, bilobar liver injury) and associated decision variables (eg, coagulopathy) that comprise the reported indications. After a unique list of codes have been developed, we will use the organisational system recommended by the RAND/University of California, Los Angeles (RAND-UCLA) Appropriateness Rating Method to group principal findings or injuries into chapters (subdivided by associated decision variables) according to broader clinical findings encountered during surgical practice (eg, major liver injury).

Ethics and dissemination: This study will constitute the first step in a multistep research programme aimed at developing appropriate, evidence-informed indications for damage control in civilian trauma patients. With use of an integrated knowledge translation intervention that includes collaboration with surgical practice leaders, this research may allow for development of indications that are more likely to be relevant to and used by surgeons. Ethics approval is not required for this study.

Strengths and limitations of this study

- Highly sensitive search strategy covering 11 electronic bibliographic databases, numerous conference proceedings, included article bibliographies, expert files and the grey literature.
- Article selection, data extraction and qualitative coding of the indications for damage control surgery and damage control interventions will be performed independently by two investigators to increase reliability of the study results.
- Although the study will identify candidate indications for systematic review for evidence of validity and reliability, the quality of the included studies will not be assessed in this study given its scoping design.

BACKGROUND

Injury remains a leading international health problem.1 2 Worldwide each year, injury occurs among 700 million people, including 30 million North Americans, and results in 5 million deaths.3 4 In 2010, road traffic (bicycle, motorcycle, motor vehicle and pedestrian) injury was the fifth leading cause of years of life lost due to premature mortality in the USA.1 Injury is also the leading cause of quality years of life lost and preventable morbidity in North America.3 4

As injured patients may lose blood at rates greater than 20 units/h, haemorrhage accounts for at least half of traumatic deaths within 24 h of hospitalisation, many of which are potentially preventable.3 5 Exsanguination, or blood loss exceeding 40% of total blood volume with ongoing bleeding, and contamination secondary to intra-abdominal hollow viscus and/or pancreaticobiliary injuries, are frequently associated with development of a ‘lethal triad’ of hypothermia, acidemia and coagulopathy (defined clinically as the absence of visible
blood clots during surgery or biochemically as an elevated international normalised ratio, prothrombin time or partial thromboplastin time). This triad has been linked with a high risk of mortality despite conventional surgical attempts at controlling haemorrhage and contamination.11 12

**Innovation and current utilisation of trauma damage control surgery**

The steps of a standard trauma laparotomy (also known as single-stage trauma laparotomy) include rapid evacuation of intra-abdominal blood followed by four-quadrant packing with laparotomy pads, complete abdominal exploration and definitive repair of all injuries.13 Although this approach is effective for most patients with abdominal injuries, prolonged or extensive operation in select severely injured patients leads to decreases in body temperature and arterial pH.12–14 Administration of large amounts of resuscitation fluid during single-stage laparotomy also frequently exacerbates or promotes onset of coagulopathy.11 13 15 As such, some major trauma patients have been reported to develop the lethal triad and succumb to their injuries when single-stage laparotomy was utilised.12 14 16

In an attempt to control coagulopathic bleeding in patients with major abdominal injuries, Stone et al17 proposed the abbreviated laparotomy with use of gauze packing and other temporising injury repair techniques, followed by planned relaparotomy for definitive injury repair (usually within 24–72 h). This approach was later named ‘damage control’ in 1993 and represents a paradigmatic change in surgical thinking that focuses more on the physiology of the patient than the need to repair all injuries during the index operation.15 18 Damage control evolved from the use of compressive perihepatic packing for patients with major liver injuries,11 19–25 and is now frequently recommended to manage select severely injured patients with head and neck, thoracic, abdominal and extremity injuries.26–30 Although the suggested number and characteristics of procedural stages vary, they have most frequently been reported to include ‘damage control ground 0’ (preoperative resuscitation and selection of patients appropriate for damage control); damage control 1 (initial abbreviated operation for control of haemorrhage and contamination); damage control 2 (intensive care unit (ICU) resuscitation); damage control 3 (reoperation for definitive repair, which may require multiple operations); and damage control 4 (reconstructive surgery; table 1).31–33

Despite being considered a breakthrough in injury care, there has been limited evaluation of damage control compared with single-stage surgery for management of trauma patients.34 A Cochrane systematic review on damage control laparotomy conducted in 2010 identified only seven relevant observational15 18 25 26 35–38 studies and no randomised controlled trials (RCTs).34 Although this review suggested that ‘good quality RCTs comparing damage control laparotomy with traditional, immediate repair of abdominal injuries [ie, single-stage laparotomy]’ were warranted,34 no trial protocols have appeared in clinical trials registries or peer-reviewed journals. While exact reasons for this are unknown, it may be due to the loss of perceived clinical equipoise among the surgical community regarding the effectiveness of damage control surgery during the adoption phase of the procedure.36 Surgeons have therefore instead focused on defining the effectiveness of individual stages of the damage control process (damage control 0–4).33 40–50

Several studies have recently reported data suggesting that damage control surgery may be overutilised.51–53 This is concerning as the procedure has been associated with substantial complications among survivors.54 55 A recent retrospective cohort study reported that one in...
five patients who received damage control laparotomy at a high-volume trauma centre between 2004 and 2008 failed to meet at least one of the traditional indications. In this study, only 33% were acidotic, 43% hypothermic and 48% coagulopathic on arrival to the ICU from the operating room. Although it could be argued that these patients may have simply been selected for the procedure before they developed the lethal triad, another retrospective cohort study suggested that applying damage control to patients who are not in physiological extremis could potentially lead to harm. In this study, use of damage control versus single-stage surgery in less-acute ill trauma patients was associated with an increased risk of bowel ischaemia/perforation, sepsis and multiorgan failure; a prolonged hospital stay and an elevated risk of death.

**Study rationale**

As no single set of appropriateness indications for damage control surgery exists, a large number of heterogeneous, sometimes non-specific, and even contradicting indications for the procedure have been proposed. In addition to the lethal triad (or its component parts), indications have been suggested based on specific patient injuries, characteristics of the surgeon or healthcare team (eg, limited surgeon experience with major trauma), and even various trauma care structural or environmental factors (eg, a non-level I trauma centre with little surgical or intensive perioperative monitoring capabilities). Those based on biochemical or laboratory measurements (eg, pH) or the temperature or fluid resuscitation requirements of the patient (eg, the number of units of packed red blood cells administered) have also been reported to have a large number of cut-offs or decision thresholds. Moreover, while indications have been proposed across all phases of trauma surgical decision-making (prehospital, emergency department and intraoperative), it is unclear when the decision to perform damage control over single-stage surgery should best be made, with some authors suggesting that this should occur preoperatively and others intraoperatively. It also remains unknown whether indications differ according to the type of damage control procedure (eg, thoracic vs abdominal), whether indications for damage control surgery are dynamic (ie, an intraoperative conversion to a single-stage procedure could be performed if haemorrhage is rapidly controlled and patient physiology improves during the procedure), and which indications for the procedure may be valid and/or reliable. The above lack of consensus regarding damage control indications has frequently resulted in the inclusion of heterogeneous populations of patients with unbalanced determinants of outcomes in damage control studies, resulting in difficulties in comparing outcome data across investigations. It also likely contributes to the aforementioned damage control practice variation.

**Study objectives**

The objectives of this mixed methods study are to systematically scope the literature on damage control surgery in civilian trauma patients, identify its proposed indications, map and clarify their definitions and examine the content and evidence on which they are based. We also seek to generate a comprehensive and well-defined list of unique indications to inform a subsequent appropriateness rating process. To improve understanding of the findings on which the decision to perform damage control is based, indications will be classified into a predefined conceptual framework and categorised and described using qualitative content analysis. Constant comparative methodology will be used to create, modify and test codes describing the principal findings or injuries (eg, bilobar liver injury) and associated decision variables (eg, coagulopathy) that comprise the reported indications for damage control. After a unique list of codes have been developed, we will use the organisational system recommended by the RAND/University of California, Los Angeles (RAND-UCLA) Appropriateness Method (RAM) to group those describing findings or injuries and associated decision variables into chapters according to broader clinical findings encountered during surgical practice (eg, major liver injury). This work will constitute the first step in a multistep research programme aimed at development of evidence-informed indications for the appropriate use of damage control in civilian trauma patients.

**METHODS AND ANALYSIS**

**Protocol design**

Methods for this study were developed following suggestions for designing and performing scoping studies conducting qualitative content analyses of textual data and for creating a list of indications for a surgical procedure.

**Conceptual framework**

Our scoping study will utilise a conceptual framework for damage control surgery indications (table 2). According to this framework, the decision to perform the procedure may occur in either the preoperative (prehospital or emergency department) or intraoperative phase of trauma surgical decision-making. This decision may be influenced by characteristics of the patient, providers, patient response to care and/or healthcare environment. Patient-based indications may be further subclassified according to those that are physiology based (eg, arterial pH<X), injury based (eg, pulmonary hilum injury) or resuscitation based (eg, administration of >X units of packed red blood cells). Injury-based indications can be further subcategorised by affected anatomical region (eg, neck, thorax or abdomen). These influencing factors are not meant to be mutually exclusive, as indications for damage control (eg, penetrating injury to the femoral artery) may be
dependent on other factors (eg, concomitant haemorrhagic shock).

Identifying relevant citations
With the assistance of an information scientist/medical librarian (HLR), we used the COre Standard Ideal (COSI) model\(^\text{72}\) to develop an ‘ideal’ (highly sensitive) search strategy to identify indications for damage control surgery and damage control interventions in civilian trauma patients. We will search the following electronic bibliographic databases from their earliest available dates without language, publication date or other restrictions: Ovid MEDLINE and EMBASE, PubMed, Web of Science, Scopus and the six databases contained within the Cochrane Library (see online supplementary table S1 for details of our planned electronic bibliographic database search strategies). Additional citations will be located by contacting several damage control experts and by searching reference lists of included citations. In an attempt to find studies about to be published, we will also review abstracts from selected conferences held between 2009 and 2013, including meetings of the American Association for the Surgery of Trauma (AAST), Australasian Trauma Society, American College of Surgeons, Canadian Association of General Surgeons (CAGS), Eastern Association for the Surgery of Trauma (EAST), International Association for Trauma Surgery and Intensive Care (IATSIC), Trauma Association of Canada (TAC) and the Western Trauma Association (WTA).

In order to further increase the sensitivity and coverage of the search, one investigator (DJR) will also search the grey literature for additional indications not reported in the peer-reviewed literature. This will involve searching relevant organisational websites (AAST, American College of Surgeons, American Trauma Society, Australasian Trauma Society, British Trauma Society, CAGS, Critical Care Society, EAST, International Trauma Anaesthesia, National Trauma Research Institute, the Society of Trauma Nurses, TAC, http://www.trauma.org, and the WTA), Google Scholar (the first 10 web pages) and two clinical trial registries (http://www.clinicaltrials.gov and http://www.controlled-trials.com) using various combinations of the following key terms: trauma, injury, damage control, damage control surgery, bailout surgery, abbreviated surgery, planned reoperation, indication and predictor. We will also manually search several trauma and surgery textbooks. Textbooks of interest will be identified by searching for relevant books listed in Access Medicine (http://accessmedicine.com), the University of Calgary Ebrary (which contains more than 30 000 book titles, and is located at http://site.ebrary.com.ezproxy.lib.ucalgary.ca/lib/ucalgary/home.action), and within the surgery category of Books@Ovid (http://www.ovid.com).

Citation selection
Two investigators (DJR and NB) will independently screen all identified citations in duplicate and select those that mention damage control, related terms (eg, staged laparotomy, planned relaparotomy or open abdominal management) or damage control interventions (eg, solid organ or intracavitary packing, temporary intravascular shunting or balloon catheter tamponade) in their title or abstract. These two investigators will then independently review the full text of the articles for these abstracts and include all citations (original or unoriginal) that involve civilian trauma patients and explicitly report one or more indications for damage control surgery or a damage control intervention. We will define an indication as an objective or subjective reason (or hypothetical clinical scenario) that the authors provided in order to guide surgeons

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<th>Phase of trauma surgical decision-making</th>
<th>Influencing characteristics or factors</th>
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<td>Preoperative Prehospital</td>
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<td>Emergency department</td>
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<td>Intraoperative</td>
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*Examples of indications that are dependent on both a principal clinical finding (eg, penetrating abdominal injury) and an associated decision variable (eg, the presence of haemodynamic instability). INR, international normalised ratio; PT, prothrombin time; PTT, partial thromboplastin time.
towards the use of damage control (or a specific damage control intervention) over single-stage surgery (or a definitive surgical procedure). 57–59 As no standardised or consensus definition yet exists, damage control will be broadly defined as a multistep operative intervention, which includes an abbreviated initial surgical procedure (or set of procedures) that aims to control obvious mechanical bleeding or contamination as compared to definitively repairing all injuries. Potentially relevant non-English language original articles will be identified for review by interpreters to assess whether they satisfy inclusion criteria. We will exclude animal studies and data sources involving exclusively non-civilian or burn patients and non-trauma patients with general (eg, intra-abdominal sepsis) or cardiovascular surgical emergencies. We will also exclude articles focusing solely on damage control for orthopaedic or neurological injuries, including spinal trauma. Eligibility disagreements will be resolved by consensus. Interinvestigator agreement regarding inclusion of titles/abstracts and full-text articles will be quantified using the \( \kappa \) statistic 73 and the ordinal \( \kappa \)-statistic agreement categories suggested by Altman. 74

**Collating, synthesising and summarising the scoping study results**

Characteristics of included articles and indications for damage control surgery and damage control interventions will be summarised using counts and proportions. Indications will be broadly classified into non-mutually exclusive categories of the above conceptual framework. We will also group indications according to patient age (adult vs paediatric depending on whether the included patients were \( \geq 16 \) years old vs \(< 16 \) years old, respectively), country and year of article publication in order to assess whether differences appear to exist across regions, practice types or time periods. Statistical analyses will be performed using Stata MP V 13.1 (Stata Corp., College Station, Texas, USA).

**Qualitative content analysis of indications for damage control surgery and damage control interventions**

Two investigators (DJR and NB) will conduct an in-depth analysis of the identified indications for damage control surgery and damage control interventions using qualitative content analysis. 67–70 While the initial stages of descriptive coding will begin as data are extracted from included manuscripts (basic unit of analysis=the text describing each indication), we anticipate that the process of coding and refinement of codes based on similarities and differences between indications will continue to emerge into the analysis phase of the study. This constant comparison method of analysis will be used to guide basic, open coding of indications independently between investigators. As the simultaneous extraction and coding of damage control indications is undertaken, our initial set of independently created codes will be reviewed, compared and then discussed by both investigators in order to test, build up and break down categories and subcategories describing the content on which the indications for damage control are based. Analysis will continue as relational similarities emerge in the form of natural groupings. Although it is possible that the resulting categories and subcategories could be similar to those outlined by the conceptual framework, our analysis will be carried out without any preconceived assumptions, given that existing theory on damage control indications is limited. 68 Once the coding and analysis phase is complete, the full code list will be reorganised into two groupings: principal findings or injuries (eg, bilobar liver injury) and associated decision variables (eg, coagulopathy). Finally, in order to identify common dichotomous decision thresholds for measured indications or associated variables with explicitly reported cut-offs (eg, intraoperative pH-X), we will calculate medians or means summarising these thresholds as appropriate. Ranges for these data will also be reported in order to display the variation in reported cut-offs across the literature.

**Organising the coded list of indications into RAM chapters**

Using the organisational system recommended by the RAM, and for ease of subsequent use, we will organise

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**Charting the data**

Two investigators (DJR and NB) will independently extract data on included articles as well as their reported indications using an electronic data extraction spreadsheet. This spreadsheet will be pilot tested using a randomly selected sample of 50 relevant English-language articles identified for inclusion in the scoping study. During pilot testing, the spreadsheet will be serially revised until consistent data collection can be demonstrated (\( \kappa \) statistic \( \geq 0.75 \)). 75

We will extract data regarding: (1) characteristics of included citations; (2) author’s institution(s); (3) number (where applicable) and characteristics of the patient population that received (or would be considered for) damage control surgery or a damage control intervention; (4) definitions of damage control surgery; (5) definitions of damage control indications and whether they were meant to be applied in the preoperative (prehospital or emergency department) or intraoperative phase of trauma surgical decision-making; (6) interventions suggested by authors to constitute damage control (in order to link the identified indications with suggested damage control techniques) and (7) outcomes associated with use of damage control surgery or a damage control intervention (see online supplementary table S2 for details regarding planned data items for collection). To identify candidate indications for systematic review for their evidence of validity and/or reliability, the type of study design for original research articles (abstract or full text) will be classified using the scheme developed by Oleckno. 75 As cohort studies and case series are frequently confused, we will use the tool recommended by Dekker et al 66 to distinguish between these study designs.

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codes into ‘chapters’ according to the broader clinical findings that may be encountered during surgical practice. Codes for principal findings or injuries within these chapters will be subdivided by those for associated decision variables, therefore allowing for later creation of a series of RAM tables (see Table 3 for examples of principal clinical findings and associated decision variables that may be used to create an interlinked RAM table).

ETHICS AND DISSEMINATION

This study will constitute the first step in a multistep research programme aimed at developing appropriate, evidence-informed indications for damage control surgery and damage control interventions in civilian trauma patients. As all data will be collected from publicly available materials, this study does not require ethics approval. To facilitate knowledge translation efforts, our team will utilise an integrated approach to dissemination and translation of study findings that enhances interactivity and clinician engagement. This will be carried out to ensure that knowledge user participation is meaningful, productive and mutually beneficial across all phases of the research process.

Several end-of-synthesis outputs are anticipated from this study that will subsequently be tailored for dissemination. Outputs will include a summative list of unique indications, an evidence map with validity/reliability assessments, a detailed evaluation of the content underlying indications and a RAM-based sorting of indications into chapters subdivided by associated decision variables.

Table 3 Example indication subdivided into its principal clinical finding and associated decision variables

| Example principal intraoperative clinical finding | Major liver injury |
| Example associated decision variables |
| Associated injuries | Inaccessible major venous injury |
| Patient physiology at the beginning of laparotomy | pH ≤ X |
| Patient physiology during laparotomy | pH > X |
| Extent of fluid resuscitation | pH improves to > X |
| Transfusion of > X units of packed red blood cells since presentation | pH does not improve to > X |
| Length of the operative procedure | ≤ X min |
| Hospital level of care designation/ability to provide comprehensive perioperative care | > X min |

To advance both awareness and trauma surgeon dialogue around damage control surgery indications and our research, we plan to disseminate the above outputs in user-friendly formats to knowledge users and their professional associations (TAC, AAST and the Australasian Trauma Society) at the national and international levels. When study results become available, presentations will be given tailored for academic and clinical audiences.

As the opinions of surgical practice leaders may be more likely to change surgical practice than clinical practice audit or guidelines, trauma surgeon involvement is likely to be essential for building a successful research programme on damage control surgery indications. A structured effort to purposively build relationships with and among several surgical practice knowledge users and practice leaders will therefore be undertaken. Engaging these individuals will help to ensure that study findings are driven by the primary change agents in the field. By involving knowledge users in the research team (DJR, AWK and CGB) and deliberately connecting with surgical practice leaders through established collegial and professional networks, we will work to ensure the research plan and emerging results are aligned with the needs of practicing trauma surgeons.

This research will identify and evaluate the proposed indications for damage control surgery and damage control interventions in civilian trauma patients. Our findings will be required to inform the subsequent creation of consensus indications among stakeholders using the RAM, and may ultimately allow for development of appropriate, evidence-informed consensus indications. They may also identify areas of uncertainty regarding whether damage control or single-stage surgery should be applied, which could guide the design of future RCTs comparing the two approaches. Development of evidence-informed indications for damage control surgery will therefore help standardise surgical practice, guide future clinical trials and improve care for seriously injured patients.

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Length of the operative procedure
| Hotizontal level of care designation/ability to provide comprehensive perioperative care |
| ACS COT level I trauma centre |
| ACS COT level II or III trauma centre |
| ACS COT level IV trauma centre |

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