A core outcome set for damage control laparotomy via modified Delphi method

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ABSTRACT

Objectives Damage control laparotomy (DCL) remains an important tool in the trauma surgeon’s armamentarium. Inconsistency in reporting standards have hindered careful scrutiny of DCL outcomes. We sought to develop a core outcome set (COS) for DCL clinical studies to facilitate future pooling of data via meta-analysis and Bayesian statistics while minimizing reporting bias.

Methods A modified Delphi study was performed using DCL content experts identified through Eastern Association for the Surgery of Trauma (EAST) ‘landmark’ DCL papers and EAST ad hoc COS task force consensus.

Results Of 28 content experts identified, 20 (71%) participated in round 1, 20/20 (100%) in round 2, and 19/20 (95%) in round 3. Round 1 identified 36 potential COS. Round 2 achieved consensus on 10 core outcomes: mortality, 30-day mortality, fascial closure, days to fascial closure, abdominal complications, major complications requiring reoperation or unplanned re-exploration following closure, gastrointestinal anastomotic leak, secondary intra-abdominal sepsis (including anastomotic leak), enterocutaneous fistula, and 12-month functional outcome. Despite feedback provided between rounds, round 3 achieved no further consensus.

Conclusions Through an electronic survey-based consensus method, content experts agreed on a core outcome set for damage control laparotomy, which is recommended for future trials in DCL clinical research. Further work is necessary to delineate specific tools and methods for measuring specific outcomes.

Level of evidence V, criteria

INTRODUCTION

Damage control laparotomy (DCL) is a well-established tool in the surgical armamentarium since its original 1908 description in liver trauma by Dr J. Hogarth Pringle.1 It is subsequently achieved renewed interest and acceptance in the 1970s2 and 1980s.3 The term ‘damage control’ was inspired by the naval military experience and captured the concept of containment without definitive repair, specifically in reference to ‘the capacity of a ship to absorb damage and maintain mission integrity’.4 Damage control in abdominal surgery is initial control of hemorrhage and contamination by followed rapid and temporary closure, further resuscitation to normal physiology in the intensive care unit and subsequent definitive re-exploration.5 DCL generally involves a temporary abdominal closure, frequently with pending bowel anastomoses and/or vascular reconstructions to perform. This description is also often referred to as an abbreviated laparotomy. The key elements are rapid control of bleeding and contamination, and in the unstable, acidotic, coagulopathic cold patient, resuscitation prior to performing definitive injury repairs. Adoption of an abbreviated laparotomy has since spread to other surgical fields including emergency general surgery and obstetrics. DCL is associated with decreased mortality5 in patients with the lethal triad (ie, hypothermia, acidosis, and coagulopathy), yet it also has known or posited associated complications including failure to achieve fascial closure, surgical site infection, fascial dehiscence, and enterocutaneous and enteratmospheric fistula formation.6–10 This has subsequently created elective surgical practices to address such surgical complications. As the DCL experience and number of survivors continue to grow, studies to refine current practice are needed. The National Academies of Medicine suggest that we should approach these problems using an evidence based, data-driven approach.11 However, this can be challenging when reported outcomes in the literature are inconsistent and differ between studies.

One way to approach this challenge is by developing a core outcome set (COS)12–14 for all future studies. A COS is a minimum set of outcomes reported that does not preclude investigators from reporting data on additional outcomes.15 This allows for future comparisons via meta-analysis or Bayesian statistics. Furthermore, it minimizes bias since these measures were prespecified and disallow selective reporting of major outcomes. The Core Outcome Measures in Effectiveness Trials (COMET) Initiative16 seeks to develop relevant resources, facilitate the exchange of ideas and information, and foster methodological research to avoid unnecessary duplication of effort, raise awareness of current problems with outcomes in clinical...
trials, and encourage evidence-based COS development. The Eastern Association for the Surgery of Trauma (EAST) clinical practice management guideline committee has brought attention to the limitations caused by lack of standardized outcomes across published manuscripts in trauma. Therefore, the purpose of this study was to develop a COS for future studies regarding DCL via an ad hoc task force of experts.

**METHODS**

The Damage Control Laparotomy Core Outcomes Set (DCL COS) study was developed following the COMET tool and conducted in accordance with recommendations from the Core Outcome Set-Standards for Development and Reporting (COS-STAD). Additionally, the Core Outcome Set Standards for Reporting were used to ensure transparency and clarity in reporting outcomes. The study was registered with the COMET database.

Delphi methodology was selected as the preferred consensus method when compared with nominal process and consensus development. Delphi is time-efficient and cost-efficient and was popularized by the RAND Corporation in the 1940s. Advantages to Delphi technique include elimination of bias by maintaining anonymity, avoiding compromised decision versus an actual consensus, and obviating specific meeting times, which allows respondents to make thoughtful decisions at a convenient time. Disadvantages include participant burnout as rounds increase, and judgment may be influenced by researcher feedback.

Panel members were identified based on contribution to the peer-reviewed DCL literature. All EAST Landmark Papers for DCL were initially evaluated by a scoping review (MZ and MB performed). All papers that were not primary literature (eg, reviews) were removed, as were papers with a topic-focus other than open-abdomen decision-making or outcomes. The first and last authors of the EAST landmark papers were considered content experts; additional content experts were selected by members of the EAST ad hoc COS task force.

Twenty-eight experts were identified based on this method. In the first round, DCL content experts were queried for suggested COS without any limitations. These were collated and grouped by the DCL COS task force with duplicates removed. In the second round, experts were asked to rank each variable using the Grading of Recommendations Assessment, Development and Evaluation scale of 1–9, with scores of 1–3 signifying a lesser important variable, 4–6 important but not critical and 7–9 a critically important variable. Consensus was defined as >70% of scores ranging from 7 to 9 and <15% of scores ranging from 1 to 3 as in previous clinical Delphi studies. Each panelist was provided with their answers as well as the distribution of the group and histograms along with the intraclass correlation (ICC) and current consensus COS of the group. We allowed panelists to include additional COS candidates as well as anonymously give feedback that they felt the other panelists should consider. This de-identified feedback was given to the group and a third round commenced including the voting on the non-consensus COS from the prior round, as well as two yes/no comments related to semantics and definitions brought up previously by the panelists that were prespecified as requiring supermajority of 70% to take effect. The flow diagram is viewable in figure 1. At no time were the panelists given information regarding the identities of the other panelists. Two rounds were required and this has previously been determined to be optimal, with more rounds requiring explanation.

**RESULTS**

Twenty-eight content experts were identified and were invited to participate. Round 1 was conducted between December 15 and December 31, 2020 with a response rate of 20/28 (71%). Each of the 20 participants (online supplemental file 1) proposed between 3 and 10 outcomes by free response, resulting in 36 unique proposed outcomes (box 1). Several of the proposed entries were demographics and not actual outcomes (study population, ‘primary diagnosis’, ‘indication for abbreviation of laparotomy’) and were removed in subsequent rounds. There were four variables included for consideration for mortality (in-hospital, 28-day, 30-day and 90-day) and these were not considered mutually exclusive. Round 2 was conducted between January 1 and January 31, 2021, with 20/20 (100%) participants. Ten outcomes achieved consensus: in-hospital mortality, 30-day mortality, fascial closure, days to fascial closure, abdominal complications, major complications requiring reoperation or unplanned re-exploration following closure, gastrointestinal anastomotic leak, secondary intra-abdominal sepsis (including anastomotic leak), enterocutaneous fistula, and 12-month functional outcome. Results were shared with the group along with individual feedback. ICC (two-way mixed-effects model with the average of k=20 raters) was 0.87, 95% CI (0.81 to 0.92), p<0.001. Email feedback was encouraged from participants to identify any additional outcomes and to provide additional feedback supporting or opposing any particular candidate outcome. A definition of intra-abdominal infection (IAI) was proposed to combine terms of gastrointestinal anastomotic leak, abscess/ deep surgical site infection, secondary intra-abdominal sepsis, and abdominal complication. Feedback also challenged the
contention that abdominal closure was synonymous with fascial closure. Two additional proposed COS (IAI and maximum lactate in first 24 hours) were submitted along with feedback regarding six existing proposed outcomes. These were provided to the group as well in preparation for round 3 voting.

The Delphi process for round 3 was conducted between February 15 and April 1, 2021. No new COS were identified through consensus and neither the semantics challenge nor new definition met the supermajority (70%) requirement to change the language of the study. The overall response rate was 95% (19 of 20), with one expert not responding in the format requested. ICC (two-way mixed-effects model with the average of k=19 raters) was 0.83, 95% CI (0.73 to 0.90), p<0.001. With no further improvement in consensus, the Delphi study was concluded (box 2).

**DISCUSSION**

DCL is an established tactic in emergency surgery for trauma, and its long-term consequences and outcomes are not entirely well-defined. The EAST ad hoc COS task force identified DCL as an important topic that would benefit from a COS to improve future research. Using an accepted consensus method, 19 content experts participated in this study to define a 10 item COS (box 2), which represent the minimum outcomes any future DCL study should include. This is the first COS developed for DCL with the intention of facilitating higher quality studies,27 with more easily combined results for pooled analysis. We recommend DCL studies measure these outcomes at minimum and also encourage investigators to include other outcomes relevant to their hypotheses.

The goal of DCL is to stop hemorrhage, re-establish critical vascular perfusion, and control contamination while delaying definitive surgery once the patient’s physiology is optimized. Given the ultimate goal of DCL is survival, it is not surprising that two core outcomes are in-hospital mortality and 30-day mortality. Mortality was one of five outcomes reported in a recent DCL systematic review and meta-analysis along with days to fascial closure and abdominal complications which were also identified as core outcomes by our expert panel.28 Fascial closure at index hospitalization, major complications requiring reoperation or unplanned re-exploration following closure, gastrointestinal anastomotic leak, secondary intra-abdominal sepsis (including anastomotic leak), and enterocutaneous fistula were all identified as core outcomes. Functional outcome at 12 months was also identified as a COS variable and was the only outcome (besides 30-day mortality, potentially) that usually extends beyond the patient’s hospital course. Patient-reported outcomes have recently been identified as an important future area of study.28

**LIMITATIONS**

Factors such as the participant panel, number of survey rounds, feedback between rounds, and the ability of panelists to add their own views must be considered according to COS-STAR.18 While we attempted to follow best practices outlined via COMET and Delphi guidelines, it should be noted that there are several
limitations to this study. First, there are limitations specific to Delphi compared with other consensus methods. Expert panels are ideally 6–11 participants; our panel was larger, with 20 participants. Prior studies identify panel sizes of 6–11 as ideal. Dagenais reported a monotonic increase in reliability of Delphi as the size of the panel increases. His study ended at 11 participants on the panel with reliability index of 0.76. Nair et al suggested that a number of experts on a panel ‘can be hundreds, but at least 10–30’, and that <6 had limited reliability and groups over 12 had insignificant reliability. This larger cohort makes consensus harder to achieve but alternatively, smaller groups usually have fewer novel suggestions.

Unfortunately, we were unable to include DCL survivors due to recruitment and organizational restraints. These would be important stakeholders to include in future COS development as the importance of patient-centered outcomes is paramount to the next evolution of surgical care. In the future, formalized community programs using national organizations such as the Coalition for National Trauma Research may facilitate patient participation in this important effort. Clear definitions of our trauma outcomes are not well-defined based on the several definition clarifications we had to make within the modified Delphi process among our panel experts and while the group defined functional outcome as a consensus outcome, we did not strictly define which measure should be reported in the literature. Finally, the authors acknowledge that the technique of damage control is not standardized within the trauma community. Although achieving consensus about the ideal technique is outside the scope of this core outcome study, this represents an area ripe for future research.

CONCLUSION

Our Delphi process using 20 content experts of damage control laparotomy achieved consensus on 10 core outcomes. This study is the first to standardize important outcomes for future DCL research and facilitate pooling of results via systematic review, meta-analysis, and/or Bayesian analysis.

SYNTHESIS

The study was deemed exempt according to University of Tennessee Health Science Center Institutional Review Board, Reference 908142, and all involvement with human participants were in accordance with the ethical standards of the institutional committee and with the 1964 Helsinki Declaration and its later amendments. Errors’. EH was a paid speaker for the Vizient Hospital Improvement Innovation Network (VHIIN) VTE Prevention Acceleration Network. TL was supported by the National Institute of General Medical Sciences of the National Institutes of Health under Award Number K23 GM140268.

Patient consent for publication Not applicable.

Ethics approval The study was deemed exempt according to University of Tennessee Health Science Center Institutional Review Board, Reference 908142, and all involvement with human participants were in accordance with the ethical standards of the institutional committee and with the 1964 Helsinki Declaration and its later amendments.

Provenance and peer review Not commissioned; internally peer reviewed.

Data availability statement All data relevant to the study are included in the article or uploaded as supplementary information.

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