Effect of triclosan-coated sutures for abdominal wound closure on the incidence of abdominal wound dehiscence: a protocol for an individual participant data meta-analysis

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ABSTRACT

Introduction Acute abdominal wound dehiscence (AWD) or burst abdomen is a severe complication after abdominal surgery with an incidence up to 3.8%. Surgical site infection (SSI) is the biggest risk factor for the development of AWD. It is strongly suggested that the use of triclosan-coated sutures (TCS) for wound closure reduces the risk of SSI. We hypothesise that the use of TCS for abdominal wound closure may reduce the risk of AWD. Current randomised controlled trials (RCTs) lack power to investigate this. Therefore, the purpose of this individual participant data meta-analysis is to evaluate the effect of TCS for abdominal wound closure on the incidence of AWD.

Methods and analysis We will conduct a systematic review of Medline, Embase and Cochrane Central Register of Controlled Trials for RCTs investigating the effect of TCS compared with non-coated sutures for abdominal wound closure in adult participants scheduled for open abdominal surgery. Two independent reviewers will assess eligible studies for inclusion and methodological quality. Authors of eligible studies will be invited to collaborate and share individual participant data. The primary outcome will be AWD within 30 days after surgery requiring reoperation. Secondary outcomes include SSI, all-cause reoperations, length of hospital stay and all-cause mortality within 30 days after surgery. Data will be analysed with a one-step approach, followed by a two-step approach. In the one-step approach, treatment effects will be estimated as a risk ratio with corresponding 95% CI in a generalised linear mixed model framework with a log link and binomial distribution assumption. The quality of evidence will be judged using the Grading of Recommendations Assessment Development and Evaluation approach.

Ethics and dissemination The medical ethics committee of the Amsterdam UMC, location AMC in the Netherlands waived the necessity for a formal approval of this study, as this research does not fall under the Medical Research involving Human Subjects Act. Collaborating investigators will deidentify data before sharing. The results will be submitted to a peer-reviewed journal.

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INTRODUCTION

Rationale

Abdominal wound dehiscence (AWD), also known as acute fascial dehiscence or burst abdomen, is a severe complication after abdominal surgery with a reported incidence of up to 3.8%. AWD frequently requires reoperation and is associated with prolonged hospital stay, lower quality of life, increased healthcare costs and mortality rates as high as 45%. In the USA, the Nationwide Inpatient Sample demonstrated that AWD results in US$40323 additional hospital costs per...
patient. The most important risk factor for the development of AWD is surgical site infection (SSI), increasing the odds more than six times. Several recently published meta-analyses investigate the effect of the use of triclosan-coated suture (TCS) for wound closure; they all report that TCS reduces the risk of SSI. One meta-analysis investigates the effect of TCS on the risk of AWD as a secondary aim, but found that current published trial data provide insufficient information to draw conclusions. To date, cumulative information of the effect of TCS on the risk of AWD is lacking. Although there are multiple randomised controlled trials (RCTs) investigating the use of TCS for abdominal wound closure, only two describe its effect on the incidence of AWD. The largest trial reports a statistically significant decrease in AWD, but concludes this to be clinically irrelevant as rates of deep SSI are comparable among treatment arms. Also, the study was not powered to detect a difference in AWD. In the second largest trial, AWD was an exclusion criteria. An individual participant data meta-analysis (IPDMA) is a meta-analysis of the original trial data and provides the opportunity to include unpublished trial data, standardise inclusion criteria and statistical analysis, check the raw data for integrity and missing data and identify baseline effect modifiers. To be able to detect the relative risk that is found in the largest trial (RR 0.42), a study would need 1436 participants. Prior the start of this study, the principle investigators of the two largest trials confirmed that individual participant data (IPD) could be made available. A pooled analysis of just these two trials would contain 2152 participants and therewith easily be able to detect the expected risk difference.

**Objectives**

The purpose of this IPDMA is to evaluate the effect of using TCS for abdominal wound closure on the incidence of AWD within 30 days after surgery in patients undergoing open abdominal surgery. Subgroup analyses will be performed according to the specific type of suture that is used for wound closure (polyglactin 910 or polydioxanone) and the level of contamination. We hypothesise that wound closure with TCS reduces the risk of AWD. This may occur through reduction of deep SSI by the use of TCS at the fascial level, or by the use of TCS at more superficial tissue layers reducing superficial SSI and its potential spread to the fascia.

**METHODS**

This study consists of a systematic review and a consecutive IPDMA. We will contact authors of studies that meet the inclusion criteria and invite them to contribute to the IPDMA. This study is registered with the International prospective register of systematic reviews (PROSPERO) (registration number CRD42019121173). This protocol is reported according to the Preferred Reporting Items for Systematic Review and Meta-Analysis Protocols (PRISMA-P) statement. Description and date of all amendments will be reported. The final manuscript will be reported according to PRISMA-Individual Participant Data (PRISMA-IPD) Statement.

**Systematic review**

**Eligibility criteria**

Randomised trials that investigate the use of TCS, compared with the exact same but non-coated sutures, in patients that underwent open abdominal surgery are potentially eligible. Studies investigating the effect of TCS for abdominal skin closure, and/or abdominal fascia closure will both be eligible. If studies only report the SSI incidence but not the AWD incidence, authors will be asked if AWD incidence is registered (either for the trial or in the medical record for regular care) and available. Trials will only be included if they can share either IPD or aggregated data on the incidence of AWD within 30 days after surgery. If AWD incidence is not available, the study will not be included. We will exclude studies if TCS is part of a bundle of interventions, and studies that investigate the use of TCS after right lower quadrant incision for appendectomy. There will be no restrictions on publication date, language or publication status.

**Literature search**

The PubMed (Medline), Embase online databases (Ovid) and Cochrane Central Register of Controlled Trials will be searched. To identify potential unpublished evidence or any ongoing trials, the International Clinical Trials Registry Platform will be searched. References of included studies will be hand searched for any additional relevant studies. In addition, meta-analyses investigating the effect of TCS on the incidence of SSI will be searched for possibly missed eligible studies. The corresponding authors from the collaborating studies will be contacted to review the list of identified studies for omission of potentially relevant studies.

A professional clinical librarian will be consulted to develop the search strategy. The search includes the free text and index terms: sutures, polyglactin 910, vicryl, polydioxanone, PDS, triclosan, wound infection, surgical wound dehiscence, fascial dehiscence and burst abdomen. These terms will be combined with the Cochrane highly sensitive search strategy for identifying randomised trials. The final search strategy is presented in online supplemental appendix 1.

**Study selection**

All studies, identified by the search strategy, will be handled through a free web app (http://rayyan.qcri.org). Duplicates will be removed. Two reviewers (AST and NW) will independently assess the studies based on previously described eligibility criteria. After screening title and abstract, full text of potentially eligible studies will be retrieved and assessed. When it is not possible to retrieve the manuscript or study eligibility is not clear, the authors will be contacted to provide further information. Any discrepancies in study selection will be resolved.
through discussion or, when necessary, by consultation with the principle investigator. We will keep a list with reasons for exclusions for all articles that pass title and abstract screening but are deemed ineligible for inclusion. Only trials that can provide either IPD or aggregated data on AWD inclusion will meet the criteria for final inclusion in the IPDMA.

**Individual participant data meta-analysis**

**Study collaboration invitation**

Authors from potentially eligible studies will be contacted and invited to contribute if their study indeed meets the inclusion criteria. An email invitation letter will be sent to the corresponding authors, outlining the IPDMA goals. If no reply is received within 2 weeks, a second email request will be sent to both the corresponding and first author. If again no response is received, we will try to contact all individual authors by email and/or telephone. IPD and/or aggregated data on AWD will be considered unavailable if numerous times (at least five) no reply is received, if authors no longer have access to the study data or authors do not consent for collaboration. Collaborating investigators will be asked to critically appraise the study protocol, provide feedback, approve the finalised version and will be offered coauthorship on the publication of the study protocol. By sharing their IPD, collaborators will be offered one coauthorship on the IPDMA manuscript, with one additional coauthorship if data of more than 300 participants are shared.

**Risk of bias**

Two reviewers (AST and NW) will independently assess the quality of the included studies using the revised tool for assessing Risk of Bias in randomised trials (Rob V.2). Studies will be judged as ‘low risk’, ‘some concerns’ or ‘high risk of bias’. Only data from the original manuscripts and study protocols will be used to ensure consistent and uniform assessments of studies that do and studies that do not provide IPD. Presence of publication bias will be assessed with the construction of a contour-enhanced funnel plot.

**Data collection process**

The collaborating investigators will be requested to sign a data transfer agreement before deidentified IPD is shared. The agreement describes the purposes of the IPDMA, the ownership of the IPD and confirms that the IPD is stored on a secure location. A researcher (AST) will conduct data collection, an interview on the study protocol and a formal handoff of the data codebook, if possible, in person. The primary objective will be to collect IPD for all outcomes. Aggregated data will only be collected if IPD is not available. If aggregated study data are not reported in the publication, this will be requested from the study authors.

**Data items**

We will propose a selection of data items of interest (with definitions and measures). All collaborating investigators will be asked to criticise and supplement this list. To ease the process of data handover, collaborating investigators can opt to share the complete data set of their study. We will select and clean only those data items that were selected collaboratively. After repeated consultation with the collaborating investigators, we selected data items on study level and data items on participant level. The list of data items with definitions is presented in online supplemental appendix 2. Study-level data include: study design (number of participating centres, blinding, randomised tissue layer, TCS specification, sample size), inclusion and exclusion criteria, and primary and secondary outcomes. Participant-level data include: baseline characteristics (age, gender, ASA score, body mass index, chronic obstructive pulmonary disease (COPD), smoking status, and previous midline incisions), and procedural characteristics (received suture, procedural status, target organ, wound classification, duration of surgery and incision type).

**Outcomes**

The primary outcome is the incidence of AWD requiring reoperation. AWD is defined as spontaneous dehiscence of the abdominal fascia within 30 days postoperatively. Reoperation, for any indication other than AWD, is not regarded as AWD.

Secondary outcomes are incisional SSI within 30 days after surgery according to the Center for Disease Control and Prevention (CDC) criteria (specified as superficial and/or deep), skin wound dehiscence, length of hospital stay, all-cause reoperations within 30 days after surgery and all-cause mortality within 30 days after surgery.

**Data integrity**

IPD will be checked for missing, invalid, out of range and inconsistent outcomes and for discrepancies with the published aggregated data. When detected, we will seek to resolve the issues with the collaborating investigators to improve data quality and ensure that trials are represented accurately. To ensure all randomised participant are included, IPD will be compared with the aggregated data from the original studies. In the case of any concerns on IPD integrity that cannot be resolved with the collaborating investigators, the data of the concerning study will not be included in the primary analysis. Checking baseline imbalances will be used to assess randomisation and allocation concealment. Pattern and extent of follow-up will be checked.

**Missing data**

For the primary analysis, we will not perform imputation of the complete variable for a study if variables are systematically missing in one or multiple trials. Missing data at participant level will be assumed to be at random. Multiple imputation by chained equations (MICE) will be used to handle missing data. Multiple rounds of imputation will be used to estimate the missing value. Percentage
of missing data will determine the number of imputation sets. MICE will be done for each individual trial before merged in the aggregated database.

**Data synthesis**

The raw IPD from each study will be copied to a separate database and recoded according to the predefined IPDMA settings. The recoded IPD will then be combined into one database containing the IPD from all studies. Dichotomous data will be expressed using risk ratios (RR) with corresponding 95% CI. Continuous data will be expressed using weighted mean differences with corresponding 95% CI. Data will be analysed according to the intention-to-treat principle, meaning that the original randomisation allocation is used to define treatment groups, regardless of the treatment that is actually received.

The primary analysis will be performed in a one-step approach using only IPD. Because the availability of IPD is not an inclusion criterion, it might occur that some trials can only share aggregated data for one or more outcomes. In the additional two-step analysis, aggregated data of outcomes for which IPD are not available will be added and analysed. For the one-step approach, we will use a generalised linear mixed model framework and an appropriate statistical model for the type of outcome. We will use a linear regression model for continuous outcome data and a log-binomial model for binary outcome data. If the log-binomial model fails to converge, we will use a log-binomial generalised estimating equation (GEE) or a log Poisson GEE model. A random intercept and, if appropriate, a random slope will be added to account for clustering of patients within studies. Potential confounding variables that, despite randomisation, show baseline imbalances across treatment arms will be added to the appropriate model. Variable selection will be based on VanderWeele principles of confounder selection. In short, we will control for each variable that is considered a cause of the intervention, the outcome, or both and for any proxy of unmeasured variable that is considered a cause of the intervention and outcome. We will limit the number of variables included in the model by the number of observed events in the dataset with a factor of 1:10. Only variables that are available in all trials are eligible for confounder selection. Additionally, we will perform a two-step approach. In this analysis, IPD from all studies will be reanalysed separately in a similar fashion as the one-step approach but without the term for trial clustering. Aggregated study data of outcomes for which no IPD is available will be added in the two-step approach. The aggregated data of each study will then be summarised, synthesising an overall estimate using DerSimonian and Laird method assuming random effects.

Statistical heterogeneity among studies will be evaluated using the $\chi^2$ test and expressed using the $I^2$ statistic. The between-study variance will be assessed using the $\tau^2$ statistic. As all tests are prespecified and effects follow from our hypothesis, no correction for multiple testing will be performed.

**Additional analyses**

All additional analyses will be performed using the one-step approach. Besides the intention-to-treat analysis, we will perform an as-treated analysis in which participants are analysed according to the type of suture that was actually used rather than the randomisation allocation. When a patient is reoperated, the study-suture is removed and the effect of the suture used on future AWD is diminished if not completely absent. As a result, inclusion of participants that underwent a reoperation might affect the observed treatment effect. We will investigate this in a per-protocol analysis in which participants that underwent a reoperation for any indication other than AWD are excluded. This analysis was added during the peer review process.

Subgroup analyses will be performed according to the specific type of suture that is used for wound closure (polyglactin 910 or polydioxanone), and the level of contamination (according to the CDC criteria).

The risk to develop an incisional hernia is higher after a midline incision than after a non-midline incision. As such, different incision types may also have different risks for AWD. Inclusion of participant with a non-midline incision introduces some degree of clinical heterogeneity and may affect the observed treatment effect. Therefore, we will perform a sensitivity analysis specifically investigating midline incisions. Additional sensitivity analyses will be performed to assess the effect of the additional use of TCS for skin closure and the effect of adding confounders that pass criteria for confounder selection but are not included in the former model as the variables are not reported in all included studies. Potential bias will further be explored in sensitivity analyses specifically investigating trials that blinded participants and personnel and through exclusion of trials assessed at high risk of bias. A complete case analysis will be performed to investigate the effect of imputation of missing data.

**Confidence in cumulative estimate**

The quality of evidence will be judged using the Grading of Recommendations Assessment Development and Evaluation working group methodology for the following domains: risk of bias, unexplained inconsistency, indirectness, imprecision, publication bias, magnitude of effect and residual confounding. The level of evidence will be downgraded for imprecision based on the optimal information size and the CI. If the optimal information size is met and the CI fails to excluded important benefit or harm, we will rate down for imprecision. We set a default threshold for appreciable benefit and harm that warrants rating down (relative risk reduction or RR of 25% or more). The level of evidence will be upgraded for a large magnitude of effect (RR>2 or <0.5) or very large magnitude of effect (RR<5 or <0.02). The overall quality will be
classified using four levels: high, moderate, low and very low.

Software
Statistical analysis will be done using R V.4.0.3.

Patient and public involvement
No patients or patient federations are involved in the design of this study protocol nor the IPDMA. Yet, the disastrous consequences of AWD are well described, underlining the need for (surgical) interventions that reduce the risk of AWD.

Study status
Currently, we have executed the systematic review. We are in contact with the authors from the original studies. We have not collected any data from the original manuscripts nor received IPD from any of the collaborators.

DISCUSSION
We designed an IPDMA with the aim to evaluate the effect of using TCS for abdominal wound closure on the incidence of AWD. This protocol describes intended methodology and statistical analysis ahead of analysis to provide transparency and receive timely feedback.

Based on the observed risk difference in the largest published trial, a new RCT investigating the effect TCS on AWD should include around 1500 participants. Such trial would be very time consuming and expose numerous patients to random assignment of two treatments while sufficient information to assess comparative effectiveness may already be available. Moreover, the effect of TCS for wound closure on the risk of SSI is well-documented, and SSI and subsequent AWD risk are closely related. A new RCT is therefore not ethical before the already available information has been optimally analysed.

IPDMA is considered the ‘gold standard’ in meta-analysis. At the core of its strength is the use of individual participant data of available trials that allows standardisation of inclusion criteria, definitions and statistical methods to reduce both clinical and statistical heterogeneity. Individual participant data also allow testing of interaction effects to assess subgroup differences and permits exploration of data that was not included in the original publications. Importantly, IPDMA requires intensive collaboration with all trialists on a certain topic, and consequently contributes to consensus on the interpretation of the available data among subject matter experts.

Despite these advantages, an IPDMA has some potential limitations. Its quality depends on the quality, size and number of available studies, the number of included participants, the availability of high-quality data and, most importantly, the willingness to collaborate among the original trialists. We have been incredibly fortunate to find so many of the original researchers willing to collaborate and contribute to the project. The expert input of all involved trialists has greatly contributed to the completion of the study protocol. In consensus meetings, we discussed the differences in data collection and variable definition between the studies. Consequently, we selected a primary outcome for which all studies would be able to uniformly provide data, being AWD requiring reoperation. Despite being a universally available outcome definition, it remains limited by the absence of a strict criteria on when to reoperate. Variation between clinicians exists and the consideration on whether or not to reoperate is hard if not impossible to retrieve. As selective reoperation by biased investigators may affect the results, we will perform a sensitivity analysis only including trials that blinded both participants and personnel making selective reoperation near impossible. Blinding for allocation is easily performed because the sutures look identical.

In conclusion, this study protocol describes an individual participant data meta-analysis in which we aim to investigate if the use of TCS for abdominal wound closure reduces the risk of AWD. If a lower incidence of AWD is observed, this may have considerable consequences for daily practice.

ETHICS AND DISSEMINATION

Ethical approval
All individual trials were approved by a medical ethics committee according to national legislation. The medical ethics committee of the Amsterdam UMC, location AMC in the Netherlands waived the necessity for a formal approval of this study, as this research does not fall under the Medical Research involving Human Subjects Act.

Dissemination
The results of this study will be submitted to peer-reviewed journals regardless of the outcome. The protocol will be submitted before the data are gathered and analysed.

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**REFERENCES**


