Faculty of Health Science
Department of Clinical Medicine

**Identifying and measuring patient harms**

* A study of measuring adverse events in hospitalised patients by the Global Trigger Tool record review method

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**Kjersti Mevik**

*A dissertation for the degree of Philosophiae Doctor – February 2019*
For all future patients
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for your humor and support and that you take so good care of the girls and me (and the house and the boat and the cabin). I love you and I need you.

Bodø 01.02.19
SUMMARY

Patient harms, or adverse events (AEs) which is the term used in this PhD thesis, is a major global health problem. They cause suffering for patients, are stressful for involved health personnel and costly for the healthcare services. Acknowledging that such events happen is necessary in order to improve patient safety. The Global Trigger Tool (GTT) has been used to track AEs over time in Norwegian hospitals from 2011. The method involves a review team who screens randomly selected patient records for predefined triggers (situations) that could indicate that an AE has happened. A trigger can be use of blood products, an infection, abrupt medication stop or a readmission. If one or more of such triggers are present, a more in-depth review is performed to decide if the trigger represent an AE. The GTT method has demonstrated high sensitivity in comparison to other methods, such as voluntary incident reporting, quality indicators from administrative data and claims for compensation. However, the GTT method is criticized because of the sampling strategy, low agreement between review teams and that the method is time consuming to perform.

This PhD evaluated if increasing the number of records to be reviewed (Paper I), changes of reviewers (Paper II) and automatically identification of triggers (Paper III) improved the reliability and validity of the GTT method.

The results showed that increasing the number of reviewed records seven times increased the rate of identified AEs by 45%. The confidence interval was narrower in a large sample compared to a small sample. Review teams with at least one identical reviewer demonstrated substantial agreement compared to moderate agreement between review teams with no identical reviewers. Automatic identification of triggers saved review time and use of this tool identified equal rates of AEs comparable to the original GTT method with manual trigger identification.

In conclusion, these studies showed that if the number of reviewed records is increased, at least one reviewer is consistent and automatic trigger identification is used, the method’s reliability and validity are improved and the review time reduced.
SAMMENDRAG (summary in Norwegian)

Pasientskader, eller uønskede hendelser som er begrepet brukt i denne ph.d. avhandlingen, er et betydelig globalt helseproblem. De forårsaker lidelse hos pasienter, er belastende for involvert helsepersonell og kostbare for helsevesenet. Anerkjenning av at slike hendelser skjer er nødvendig for å kunne bedre pasientsikkerheten. Metoden Global Trigger Tool (GTT) ble derfor innført ved alle norske sykehus fra 2011 med det formål å følge antall uønskede hendelser over tid. Metoden går ut på at ett granskningsteam gransker et tilfeldig utvalg av pasientopphold etter forhåndsdefinerte triggere (situasjoner) som kan indikere at en uønsket hendelse kan ha skjedd. En trigger kan være bruk av blodprodukter, en infeksjon, plutselig seponering av ett medikament eller en reinnleggelse. Hvis en eller flere slike triggere er tilstede, gjøres en mer grundig gjennomgang for å finne ut om triggeren er assosiert med en uønsket hendelse. GTT metoden har høy sensitivitet i forhold til andre metoder som avviksmeldinger, kvalitetsindikatorer basert på administrative data og klagesaker. Imidlertid er GTT metoden kritisert fordi de baseres på granskning av små utvalg av pasientopphold, har dårlig samsvar mellom forskjellige granskningsteam og at metoden er tidskrevende å gjennomføre.

Denne doktorgradsavhandlingen evaluerte om økning av antall pasientopphold som granskes (Artikkel I), utskifting av granskere (Artikkel II) og automatisk identifisering av triggere (Artikkel III) bedret metodens reliabilitet (pålitelighet) og validitet (gyldighet).

Resultatene viste at ved å øke utvalget av granskede pasientopphold sju ganger, økte raten av antall identifiserte uønskede hendelser med 45 %. Konfidensintervallet var smalere i et stort utvalg sammenlignet med ett lite utvalg. Granskningsteam som hadde minst en lik gransker viste godt samsvar sammenlignet med team som ikke hadde noen like granskere. Automatisk identifisering av triggere sparer granskningstid, og bruk av dette verktøyet identifiserte samme rate av uønskede hendelser som ved bruk av den original GTT metoden med manuell trigger identifisering. Oppsummert viser studien at hvis man gransker større utvalg av pasientopphold, beholder minst en gransker stabil i granskningsteamet og bruker automatisk identifisering av triggere, vil metodens reliabilitet og validitet forbedres og tidsbruken reduseres.
LISTS OF PAPERS

This thesis is based upon three publications, referenced in the text by their respective roman numerals:

   Does increasing the size of bi-weekly samples of records influence results when using the Global Trigger Tool? An observational study of retrospective record reviews.
   *BMJ open*, 2016, 6.4: e010700.

II. **Mevik K**, Griffin FA, Hansen TE, et al.
   Is inter-rater reliability of Global Trigger Tool results altered when members of the review team are replaced?
   *International Journal for Quality in Health Care* 28.4 2016: 492-496

III. **Mevik K**, Hansen TE, Deilkás EC, et al.
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GTT</td>
<td>Global Trigger Tool</td>
</tr>
<tr>
<td>AE</td>
<td>Adverse events</td>
</tr>
<tr>
<td>IHI</td>
<td>Institute for Healthcare Improvement</td>
</tr>
<tr>
<td>COSMIN</td>
<td>Consensus-based Standards for the selection of health status Measurement Instruments</td>
</tr>
<tr>
<td>NPE</td>
<td>Norsk pasientskadeerstatning</td>
</tr>
<tr>
<td>WHO</td>
<td>World Health Organization</td>
</tr>
<tr>
<td>HMPS</td>
<td>Harvard Medical Practice Study</td>
</tr>
<tr>
<td>SPC</td>
<td>Statistical Process Control</td>
</tr>
<tr>
<td>EHR</td>
<td>Electronic health records</td>
</tr>
<tr>
<td>INR</td>
<td>International normalized ratio</td>
</tr>
<tr>
<td>QI</td>
<td>Quality indicator</td>
</tr>
<tr>
<td>PSI</td>
<td>Patient safety indicator</td>
</tr>
<tr>
<td>PROM</td>
<td>Patient reported outcome measure</td>
</tr>
<tr>
<td>CI</td>
<td>Confidence interval</td>
</tr>
<tr>
<td>SE</td>
<td>Standard error</td>
</tr>
<tr>
<td>RR</td>
<td>Risk ratio</td>
</tr>
</tbody>
</table>
“To err is human; to cover up is unforgivable; and to fail to learn is inexcusable.”

Sir Liam Donaldson at the launch of the World Alliance for Patient Safety Oct 2004

1 INTRODUCTION

1.1 Background

Patient harms, or adverse events due to medical care, is a major global health problem as they cause suffering for patients and are stressful for involved healthcare professionals [1]. In addition they are costly for the healthcare services [2]. Acknowledging that such events happen and measuring them, are necessary for improving health care and increasing patient safety [3].

The common methods (i.e.; incident reporting, quality indicators, processes for dealing with complaints and mortality & morbidity conferences) of reporting and analysing adverse events are unfortunately inappropriate for measuring adverse events mostly due to reporting bias [3]. These systems depend on either the patients, their relatives or health personnel voluntary reporting the adverse events.

Review of patient records for specific triggers (situations) such as use of blood products, abrupt stop in medication or readmissions, is an alternative method to identify and measure adverse events. Such method has demonstrated high sensitivity in comparison to the referred methods above [4]. The widely used method for identifying and measuring adverse events is the Global Trigger Tool (GTT), developed by the Institute of Healthcare Improvement (IHI) in Cambridge, USA [5]. Frequent use of the GTT method has demonstrated that adverse events are far more common than first assumed [6], [7]. Estimates show that adverse events happen as frequent as up to 30 % of the inpatient population [6].

However, the GTT has some practical disadvantages. It is rather resource intensive due to time and personnel required. The sampling approach, reviewing only small samples of records, together with frequent replacement of reviewers question the reliability and validity of the method [8], [9]. This thesis examined the effect on the results of identified adverse
events by increasing the number of reviewed records and changing the reviewers. Use of automatic identification of triggers was also evaluated. As the GTT is used in all Norwegian hospitals the aim of the thesis was to make the GTT method a more efficient, valid and reliable strategy to identify and measure adverse events in hospitalised patients.
1.2 Adverse events

1.2.1 Definitions

Several different terms describing adverse outcomes of medical care are used (table 1). Inconsistent use of terms, which appear both in the literature and in the clinical settings, complicates the understanding of adverse outcomes due to medical care [10].

Table 1 Terms describing adverse outcomes

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
<th>Pros</th>
<th>Cons</th>
</tr>
</thead>
<tbody>
<tr>
<td>Errors</td>
<td>a failure to carry out a planned action as intended or application of an incorrect plan [11]</td>
<td>Identify failures</td>
<td>Promotes blaming</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Inhibit system approach</td>
</tr>
<tr>
<td>Injuries</td>
<td>damage to tissues caused by an agent or event [11]</td>
<td></td>
<td>Only severe events</td>
</tr>
<tr>
<td>Patient harms</td>
<td>an outcome that negatively affects a patient’s health and/or quality of life [12]</td>
<td>Already in use</td>
<td>Used differently whatever</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>considered a severe event, a claim or adverse outcomes</td>
</tr>
<tr>
<td>Adverse events</td>
<td>unintended physical injury resulting from or contributed to by medical care that requires additional monitoring, treatment or hospitalisation, or that results in death [5]</td>
<td>System approach</td>
<td>New term</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Promotes a no blame culture</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Promotes interventions to reduce them</td>
</tr>
<tr>
<td>Complications</td>
<td>an unfavourable evolution or consequence of a disease, a health condition or a therapy [13]</td>
<td>Already in use</td>
<td>Acceptance of the incidence of the events</td>
</tr>
<tr>
<td>Healthcare-associated harm</td>
<td>harm arising from or associated with plans or actions taken during the provision of healthcare, rather than from an underlying disease or injury [10]</td>
<td>No doubt that the harm is due to the healthcare given</td>
<td>Too complicated for everyday use</td>
</tr>
</tbody>
</table>
Identification and measurement of adverse outcomes from medical care depend on a common definition of what constitutes this term, in order to increase the understanding of such events [14]. Consistent use of patient safety terms is also necessary for making comparison between facilities possible and to track the trends over time [10]. A group, initiated from the World Health Organisation (WHO), agreed upon 48 concepts aiming for that this agreement could pave the way for a common understanding of the concepts of patient safety [10]. Common definitions would probably increase the focus on these events promoting implementation of interventions to prevent them. However, deciding the contribution of medical intervention in regard to the underlying disease to an event, is often difficult. For example; an unplanned unit of blood is infused to an anaemic patient after an operation. It is not always obvious if the anaemia is due to the medical condition or due to the operation. The type of medical condition is important to consider when deciding if the event was due to the condition. A definition including criteria for defining it as an adverse outcome due to medical care, would make it easier to decide. A discussion concerning when to use and not to use the different terms follow, as well as their suitability as measures of adverse outcomes.

Using the term error for the adverse outcome often brings up the question of whom is to blame. The blame perspective makes the culture for analysis the event difficult. A “just” culture promotes a system approach, rather than blaming and shaming on individuals [3], [15], [16]. Most errors are committed by good hardworking people and identifying who’s to blame is a distraction. It is far more productive to identify the situations that caused the error and implementing systems that will prevent them from happening again [17]. However, the fact that all errors do not result in adverse outcomes and all adverse outcomes are not necessary a result of errors, makes measuring errors not suitable as a measure [18].

The terms injury or harm do not distinguish between injuries as adverse outcomes due to medical care or due to injuries caused by the patients’ disease or by an accident. In the clinical setting the term patient harm has traditionally been used when a patient suffers a harm due to a severe and highly unexpected event caused by the medical care given. This unresolved understanding of the term patient harm was not considered when the Norwegian Patient Safety Campaign (later defined as program) “In safe hands” (“I trygge hender”) was launched
in 2011. They chose to use the term *patient harm* (pasientskade) for all events when implementing the GTT to measure adverse outcomes due to medical care [19]. The manual of the original GTT define such events as adverse events and do not use the term patient harm. “Patient harm” used in the Norwegian campaign included both minor events, such as catheter based urinary infections, and more severe events, such as injury to the ureter during a laparotomy. This “new” use of the term *patient harm* was not immediately adapted by the clinical health personnel in Norwegian hospitals as they have reserved this term for the severe events and events that could qualify for compensation through the Norwegian System of Patient Injury Compensation (NPE) [20]. According to the Act on Patient Injury Compensation [21] three criteria must be fulfilled before a claim for compensation is accepted. It must have been a failure in treatment (with some exceptions), economic loss of more than 10000 NOK and the injury could not be more than three years old when applying. The patient harms measured by the GTT method is mostly less severe than the events traditionally defined as patient harm by the clinical health personnel.

The term *complication* does neither distinguish between events caused by the patients’ underlying disease, or by medical care. However, complications are often agreed as foreseeable unintended events due to medical care. If an event is considered foreseeable it is often a silent acceptance that they happen from time to time. Accepting that such events happen could act as an obstacle to identify, measure and prevent them. The Norwegian Patient Safety Program wanted to include events that were defined as complications as well as events that were previously not considered a patient harm (i.e.: urinary tract infection due to catheter), addressing all these events as patient harms.

The original GTT defined the adverse outcomes due to medical care as *adverse events* (uønskede hendelser) with the definition described in table 1. As described previously, unplanned and unintended events have traditionally been defined as complications, if acknowledged at all by the clinical health personnel. The authors of the GTT focused on the events that harm the patients rather than errors that easily promote a perspective of whom to blame. *Adverse events* has been used in the literature for decades, but first used in relation to patient harms in the mid 80’s [22].
In this thesis we will investigate how the GTT’s ability to identify and measure adverse outcomes could be improved. We therefore decided to use the term *adverse event* in this thesis. We argue that this term includes most of the relevant events due to medical care; whether considered a complication, a preventable event, an error or a failure of systems.
1.2.2 Identification

Table 2 shows the different systems that are used for reporting or measuring adverse events in hospitals [23]. These are unlike the methods that are used for dealing with adverse events such as root cause analysis, mortality & morbidity conferences, malpractice claims and compensation systems which all are inappropriate to use as measurement methods due to reporting bias. Also, selection bias, confounding bias, information bias or hindsight bias could influence the reporting of adverse events in the different measurement methods referred to in table 2. Selection bias could occur when patients are seemingly selected non-randomly, but for whatever reason still selected due to a specific variable such as their age, sex, department admitted to or selected because of the adverse event. Confounding bias can occur if an alternative explanation of the adverse event (which is not accounted for) is present, such as age. For example, if age is not adjusted for, the adverse event rates could be explained by that the selected patients are mainly above a certain age. If there is an error concerning the measurement method, it is defined as information bias. This could be present if there is something wrong with the measurement method. Hindsight bias could be due to that the outcome is known for the reviewer when determining if adverse events are present.

Voluntary incident reports and patient reported outcome measures, rely on the commitment of health personnel and patients to report adverse events. These systems are therefore subject to reporting bias. The patient voice is an emerging part in the patient safety field, but there is so far no tradition to include patient reports in the measurement of adverse events [24]. Patients mostly identify problems related to doctor-patient-relationship (lack of respect, time pressure, rudeness, break of confidence), coordination, access (long waiting time, no appointments available) and communication (between doctor and patient, among health care professionals) [24], [25]. Medical record review is the method with highest correlation with patient reported events, in contrast to incident reporting by staff with no or low concordance with patient reported events [26]–[29]. The few studies performed suggest that patient reported outcomes can be included in the hospitals measurement of adverse events, but the risk of both overestimating and underestimating due to inconsistent use of terms must be accounted for [28].
### Table 2: Strengths and limitations of common methods to identify adverse events

<table>
<thead>
<tr>
<th>Methods</th>
<th>Strengths</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Administrative data</td>
<td>Few resources required</td>
<td>Low sensitivity - many false positives</td>
</tr>
<tr>
<td>(e.g.: ICD 10 codes)</td>
<td>Inexpensive</td>
<td>Requires correct diagnosis, procedures</td>
</tr>
<tr>
<td></td>
<td>Utilize readily available data</td>
<td></td>
</tr>
<tr>
<td>Quality indicators (QI)</td>
<td>No clinical resources needed for computerized systems</td>
<td>Low sensitivity - many false positives</td>
</tr>
<tr>
<td>(e.g.: readmission after 30 days)</td>
<td>Objective measure</td>
<td>Requires correct documentation of the data</td>
</tr>
<tr>
<td></td>
<td>Inexpensive to run when first developed</td>
<td></td>
</tr>
<tr>
<td>Patient safety indicators (PSIs)</td>
<td>Do not rely on clinical judgment</td>
<td>Requires technology development</td>
</tr>
<tr>
<td>(e.g.: decubitus ulcer)</td>
<td>Identifies adverse events directly</td>
<td>Depend on the accuracy of the ICD-10 coding</td>
</tr>
<tr>
<td></td>
<td>Comprehensive</td>
<td>Some indicators are just indicators of adverse events, and not just an adverse event by itself</td>
</tr>
<tr>
<td></td>
<td>Screening tool</td>
<td>Narrow range of adverse events</td>
</tr>
<tr>
<td></td>
<td>Inexpensive to run when first developed</td>
<td>Administrative data lack information about the severity</td>
</tr>
<tr>
<td>Voluntary reporting</td>
<td>Inexpensive</td>
<td>Relies on awareness and willingness of staff to volunteer submit event notification</td>
</tr>
<tr>
<td>(e.g.: incident reporting)</td>
<td>Can detect latent events (near-misses)</td>
<td>Requires a no blame culture, Reporting bias</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Hindsight bias</td>
</tr>
<tr>
<td>Trigger tools</td>
<td>Sample based</td>
<td>Rely on documentation in the health record</td>
</tr>
<tr>
<td>Manual</td>
<td>Commonly used</td>
<td>Requires extensive clinical resources</td>
</tr>
<tr>
<td>(e.g.: GTT, HPMS)</td>
<td>No technology development required</td>
<td>Inter-rater reliability can vary</td>
</tr>
<tr>
<td></td>
<td>Works in paper records</td>
<td>Hindsight bias</td>
</tr>
<tr>
<td>Automatic</td>
<td>Inexpensive when first developed</td>
<td>Technology development required</td>
</tr>
<tr>
<td>(e.g.: automatic trigger identification)</td>
<td>Efficient</td>
<td>Manual review required of the triggered records</td>
</tr>
<tr>
<td></td>
<td>Objective identification of triggers</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Integrates multiple data sources</td>
<td></td>
</tr>
<tr>
<td>Full chart review</td>
<td>Works in paper records</td>
<td>Incomplete medical records</td>
</tr>
<tr>
<td></td>
<td>Commonly used</td>
<td>Judgment of adverse events are subject to reviewers decision</td>
</tr>
<tr>
<td></td>
<td>Gold standard?</td>
<td>Expensive</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Resource intensive</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Hindsight bias</td>
</tr>
<tr>
<td>Patient reported outcome measure (PROM)</td>
<td>Reflects the patients view of adverse events</td>
<td>Inconsistent reporting routine</td>
</tr>
<tr>
<td>(e.g.: EKG of all post-operative patients)</td>
<td>No technology development required</td>
<td>No standard definition of an adverse events</td>
</tr>
<tr>
<td>Clinical surveillance</td>
<td>Accurate and precise</td>
<td>Costly as all patient in a cohort are screened</td>
</tr>
<tr>
<td>(e.g.: EKG of all post-operative patients)</td>
<td>Limited to specific interventions</td>
<td></td>
</tr>
<tr>
<td>Observation of patient care</td>
<td>Direct observation</td>
<td>Confidentiality concerns (punishments)</td>
</tr>
<tr>
<td>(e.g.: videotaping or observation)</td>
<td></td>
<td>Hawthorne effect (people do not act “normal” when observed)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Evaluates a specific situation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Resource intensive training of observers</td>
</tr>
</tbody>
</table>
1.2.3 Evaluations of measures

Measures should be of high precision and with high accuracy. Precision refers to if the measure consistently provides the same results if it is repeated. The accuracy refers to whether the measure measures exactly what it is supposed to measure [30]. The precision describes the difference between repeated measures of the same value and the accuracy reflects the difference between the measured and the true value (figure 1).

Figure 1 Precision and accuracy (Illustration by Laila Bjølgerud)

<table>
<thead>
<tr>
<th>PRECISE</th>
<th>ACCURATE</th>
<th>NOT ACCURATE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><img src="image1.png" alt="Diagram" /></td>
<td><img src="image2.png" alt="Diagram" /></td>
</tr>
<tr>
<td>PRECISE AND ACCURATE</td>
<td><img src="image3.png" alt="Diagram" /></td>
<td><img src="image4.png" alt="Diagram" /></td>
</tr>
<tr>
<td>NOT PRECISE</td>
<td><img src="image5.png" alt="Diagram" /></td>
<td><img src="image6.png" alt="Diagram" /></td>
</tr>
<tr>
<td>ACCURATE BUT NOT PRECISE</td>
<td><img src="image7.png" alt="Diagram" /></td>
<td><img src="image8.png" alt="Diagram" /></td>
</tr>
<tr>
<td>NEITHER PRECISE NOR ACCURATE</td>
<td><img src="image9.png" alt="Diagram" /></td>
<td><img src="image10.png" alt="Diagram" /></td>
</tr>
</tbody>
</table>

The confidence interval (CI) is calculated from the observed data based on the standard error (SE). The confidence level is usually set to 95%. The accuracy regarding the CI defines if the interval contains the true population mean while the precision refers to the width of the CI. To increase accuracy the confidence level is increased which will widens the CI. But if the width of the CI increases the precision goes down.
Methodological quality in studies on measurement properties can be assessed by using the Consensus-based Standards for the selection of health status Measurement Instruments (COSMIN) checklist [31]. The checklist include the measurement properties internal consistency, reliability, measurement error, content validity, structural validity, hypotheses testing, cross-cultural validity, criterion validity, responsiveness and interpretability [31]. The measurement properties used in this thesis is further discussed.

For academic use the term reliability describes how reliable and precise the results from a measure are. Reliability refers to the consistency of a measure with the types: test-retest reliability, internal consistency and inter-rater reliability. Test-retest reliability is administering a test to a group of individuals, re-administering the same test to the same group at some later time, correlating the first set of scores with the second in a scatterplot computing Pearson’s r [14]. Inter-rater reliability is the correlation of scores between two or more reviewers who scores the same item. This is typically measured by the Cohen’s Kappa coefficient where kappa is the “true” agreement when accounting for agreement by chance [32]. This method could also be used to evaluate the agreement of repeated administration of a test performed by one rater (intra-rater reliability). Internal consistency is the correlation between different items on the same test measured by Cronbach’s alpha [33].

Validity is not defined by one definition [34]. It could be explained as the degree of which a concept measure what it is supposed to measure and how valid and accurate the results from the measure are. It could be evaluated by comparing the results of the measure to the results of another measure (referred to as gold standard) [35]. Content validity evaluate if the content of an instrument is an adequate reflection of the item to be measured. If this is obtained by expert opinions as a descriptive evaluation without any statistically analysis, it is called face validity. Construct validity evaluates if the measure measures what it is supposed to measure [36]. Criterion validity is how good the measure correlates with or predicts another valid and observable variable at the same time (concurrent validity) or later (predictive validity). For example, if the adverse event urinary tract infection is related to the rate of indwelling urine catheter used [37]. Validity is also divided in internal and external validity. Internal validity refers to whether the findings relate or are caused by the phenomena under investigation. For
example, if the adverse event identified, really is caused by the intervention given in the actual admission. External validity is the extent to which the results can be generalized for other patient groups [38].

A measure needs to have high reliability and high validity, but low validity is considered more critical than low reliability. If the measure measures some other variable and not the one we think it measures or if the measure is systematically wrong, a larger sample will not help, it will rather do more harm [36]. For example, if the method used for measuring adverse events have low validity, the events measured might not be true adverse events. Low reliability could be improved by increasing the sample size.

### 1.2.4 Types

A brief description, prevalence and source of the main types of adverse events referred to in the literature are presented in table 3.

#### Table 3 Overview of the common types of adverse events

<table>
<thead>
<tr>
<th>Type</th>
<th>Including</th>
<th>Incidence in hospitalised patients</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infections</td>
<td>Healthcare associated infections, hospital acquired infections, iatrogenic infections and nosocomial infections such as Ventilator associated pneumonia, Pneumonia, Central line associated bloodstream infections, Catheter associated urinary tract infections, Surgical site infections, Gastrointestinal illness, Blood stream infections</td>
<td>5 %</td>
<td>Surveillance</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Prevalence study, Trigger tools, Record tools, QIs, Administrative data, Chart review</td>
</tr>
<tr>
<td>Surgical</td>
<td>Surgical site infections, Hematoma/Bleeding, Postoperative thromboembolism, Wrong site surgery, Retained foreign objects</td>
<td>2 %</td>
<td>Surveillance</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>PSIs, Chart review</td>
</tr>
</tbody>
</table>
- Medical device related harms (gas/air embolism, burning, stent thrombosis)

<table>
<thead>
<tr>
<th>Obstetric/perinatal</th>
<th>0.3 %</th>
<th>Surveillance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Foetal asphyxia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anal sphincter tear</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infections</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shoulder dystocia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Injury of intestines or urinary tract</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Uterine rupture</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thromboembolism</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Falls</th>
<th>20 %</th>
<th>Surveillance</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Voluntary reporting</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Chart review</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Pressure ulcer</th>
<th>14 %</th>
<th>Surveillance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bedsores</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Decubitus ulcer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pressure sores</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Medications</th>
<th>20 %</th>
<th>Surveillance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adverse drug event</td>
<td></td>
<td>Trigger tools</td>
</tr>
<tr>
<td>Adverse blood infusion event</td>
<td></td>
<td>Chart review</td>
</tr>
<tr>
<td>Adverse infusions events (vaccines)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Diagnostics</th>
<th>Unknown</th>
<th>PROMs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Misdiagnosis</td>
<td></td>
<td>Malpractice</td>
</tr>
<tr>
<td>Missed diagnosis</td>
<td></td>
<td>claims/Compensation system</td>
</tr>
<tr>
<td>Delayed diagnosis</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Infections**

Infections associated with medical care has been named healthcare associated infections, hospital acquired infections, iatrogenic infections or nosocomial infections as opposed to community-acquired infections. The terms are mostly used interchangeably, but “healthcare associated infection” are recommended to use when the patient recently has been hospitalised, had haemodialysis, received intravenous chemotherapy or resided in a long-term care facility in contrast to “hospital acquired” infection where the patient received the infection diagnose within 72 hours of admittance to hospital or developed the infection within 10 days of discharge from the hospital [39]. The percentage of patients experiencing at least one healthcare associated infection is approximately 4 % in the US [40], 5.7 % in Europe and 4.9 % in Norway [41] making this one of the most common type of adverse event.
Adverse events following surgery

According to the WHO almost half of the identified adverse events (48%) are related to surgical procedures [42]. The most frequent adverse surgical events are blood loss, surgical site infections and postoperative venous thromboembolism. Surgical site infections increase mortality, length of stay, readmissions and use of health-care services [43]. Postoperative venous thromboembolism is a common adverse event, occurring in 7% of hospitalised patients [44] and is associated with reduced survival and substantial health-care costs [45].

Wrong site surgery could be defined as surgery on the wrong person, on the wrong body part or at the wrong side of the patient body [46]. Wrong site surgery and retained foreign objects are rare but receive major attention if they occur. Risk factors are emergency operations, unusual time pressures to start or complete a procedure or the involvement of different surgeons [47].

Manufacturer-related errors, user errors and design errors of medical devices can cause adverse events such as gas emboli after laparoscopy/hysteroscopy, air embolism after infusions, stent thrombosis and burning scar after diathermic procedures [43]. In some cases it is difficult to identify these as the cause of the adverse event [48].

Obstetric and perinatal adverse events

Worldwide the maternal and infant mortality rates are high mostly due to lack of access to medical facilities and adequate medical care [43]. The rate of obstetric related adverse events has been reported to less than 1% in developed countries [49]. However, despite their infrequencies, obstetric events are one of the ten most common cases for claims for compensation in the Norwegian System of Compensation to patients (e.g.: fetal asphyxia, anal sphincter tear, infections, shoulder dystocia, injury of intestines or urinary tract, uterine rupture and thromboembolism) [50].

Fall with injury and pressure ulcer

Patient fall is the most common reported adverse event in the voluntary reporting systems [43]. The overall rate of patient fall is estimated to 5-9 per 1000 patient days and 30% of the
events lead to harms. Negative outcome of a fall frequently includes hip fractures with prolonged hospitalisation. The prevalence of pressure ulcer in hospitals is estimated from 10% to 15% of admitted patients and the risk factors includes immobility, friction, incontinence, cognitive impairment and poor nutritional status [42].

**Adverse drug events**

An adverse drug event can be caused both by drugs, blood products or fluid infusion. The adverse events related to drug treatment are one of the most common adverse events in developed countries. The adverse events relate mostly to prescribing, monitoring and administering medicines with look-alike labelling, wrong use of medication or failure to recognize drug interactions [43]. The consequences of an adverse drug event could be substantial, and it is estimated that it occurs in 1 of 16 hospitalised patients, with huge financial impacts [51]. Injections are one of the most common healthcare procedures with 16 billion injections annually in developed countries including immunizations, local anaesthetics and contraceptives. Adverse events concerning injections are mostly related to devices that could transmit infections and not to the drug itself [43].

**Diagnostics challenges**

Diagnostics challenges include missed diagnosis, misdiagnosis and delayed diagnosis. This is an unexplored perspective of patient safety but is rarely registered as a type of adverse events on its own. This could be due to the difficulty studying the problem and the complex causes of it [43]. Many of the claims in the Norwegian compensation system for patient harm are related to delays in diagnosis or delayed or missed follow-ups. Andreasen et al found considerable variations of experts’ evaluations regarding the claims after alleged birth complications demonstrating the difficulty of studying the issues related to diagnostics challenges [52].

**1.2.5 Incidence**

Measuring number of patients being harmed while hospitalised was first referred by the Tort system of medical malpractice in the U.S [53], [54]. Later, the Harvard Medical Practical
Study (HMPS) measured adverse events and negligence in hospitalised patients by reviewing patient records [22], [55]. The definition of an adverse event “as an injury that was caused by medical management (rather than the underlying disease) that prolonged the hospitalisation, produced a disability at the time of discharge, or both”, was applied. They estimated that adverse events occurred in 3.7 % of the hospitalised patients. The Institute of Medicine’s report “To Err is Human” brought the issue of measuring adverse events to national and international attention as they estimated that 98,000 Americans died as a results of medical errors every year [56]. This made measuring adverse events in hospitalised patients a growing focus for quality and safety in healthcare worldwide [57].

Several studies followed, demonstrating that the level of adverse events was higher than first estimated [58]–[62]. However, comparing the results between the studies were challenging as the studies applied different definitions of what they had measured [63]. Although no gold standard to identify the true level of adverse events exists, it is a common agreement that adverse events is a major global health problem [1], [63]. Valid and reliable methods that measure adverse events are demanded. The existing systems, such as the GTT, are inadequate to count the actual number of events, [38] but are used for estimating the rate of adverse events.

The WHO estimated a total of 47.7 million events when including seven different types of adverse events occurring annually in patients across the world [64]. In Norway, commissioning documents from the Ministry of Health have instructed the hospitals since 2011 to perform the GTT to measure adverse events yearly. The most common identified adverse events in Norwegian hospitals during the period 2010-2015 were hospital-acquired infections and medication related harms [65]. Interventions to reduce adverse events were initiated and implemented in the hospitals as part of the Norwegian Patient Safety Program (“I trygge hender”). In the period from 2010 to 2017 the rate of adverse events have slowly decreased from 16 % to 14 % of the admissions (figure 2) [65], [66]. This rate is below the rate of adverse events reported in international studies [6]. The reduction of the rate in Norway could reflect a true reduction of rate, or it could be due to random variability. Even though the total rate remains unchanged, the rate of the different types of adverse events could
have changed [67]. Many have argued that the rate of adverse events is still persistently high, despite the many different interventions implemented to reduce the rate of adverse events [67]. [68].

Figure 2 Percent of admissions with adverse events in Norwegian hospitals measured by the GTT

As described previously, the results from the systems for dealing with and reporting adverse events can only estimate the number of adverse events. However, when reporting systems are used for estimating how many patients who are harmed, the results of this are often misleading. The Norwegian claims for compensation system are based on patients’ own claims, voluntary reports rely on health personnel to report, and severe events are investigated by the health supervision only if someone report the events. To illustrate how many events the different systems handle, we compared the reported adverse events per 100 admissions between the existing systems in our trust for 2013. Unfortunately, patient reported outcome measures are not included. The results are illustrated in figure 3. The data were collected from
the NPE, the trust’s system for voluntary reporting of adverse events, the Norwegian Board of Health supervision and the GTT results. The GTT identified four times more adverse events than the other systems. We argue that this demonstrates that the GTT is the most appropriate system to quantify the number of adverse events. However, in most cases the events were reported only by one of the methods. Others have found similar results with no overlap of the identified events between the methods [69]. According to these findings, different methods might be used to reveal as many adverse events as possible.

**Figure 3** Adverse events/reported events in 2013 per 100 admissions by the different systems in Nordland Hospital
1.3 The Global Trigger Tool (GTT)

1.3.1 Background

Identification of triggers in patient records to measure adverse events was first introduced by Jick in 1974 [70]. Classen et al developed the method further to be used for identifying adverse drug events [71]. Later, these trigger tools were introduced to measure adverse events in surgical departments, intensive care departments and children’s departments [4], [71]–[74]. The trigger tools represented an alternative approach to measure adverse events [55]. The IHI developed the GTT initially for reviewing randomly selected paper patient records to identify triggers that could represent that an adverse event had occurred [5]. The GTT has successfully been advocated with the aim to monitor adverse events in adult inpatients demonstrated by widespread adoption [61], [75]–[78].

The intention of the GTT was to develop an easy-to-use approach for the hospitals to identify and measure adverse events [5]. The results were not intended for benchmarking between hospitals as they have different demographic background of the patients, they treat different conditions, the number of inpatients differ, and the functions of the hospitals differ. These issues make comparing GTT results between different hospitals challenging. The developers of the GTT argued that the results should be used within the hospital to acknowledge the rate and severity of the adverse events. Once the adverse events are identified, interventions that can prevent them from happening should be implemented. The effect of the interventions can be evaluated by the use of the GTT following the rate trends over time [5].

Reviewing all inpatient records manually is impossible except in very small hospitals, hence the sample strategy. To obtain consistent results regarding the rate of adverse events, the sampling methodology needs to be truly random as the numbers of records selected must be identical in every sampling period from the same discharge lists. The recommended sample size in the GTT is ten closed inpatient records for every bi-weekly period. The patients eligible for selection must be 18 years or older, admitted for more than 24 hours and not be admitted for rehabilitation or psychiatric care since the triggers are not developed for these
areas of care. The triggers in the GTT are neither developed for children and teenagers or for outpatients.

1.3.2 Implementation

The GTT is a two-step method with manual retrospective review of records: Two primary reviewers individually review the records for 53 specific triggers (see appendices) and determine if the triggers represent any adverse events, before reaching consensus (step 1). A secondary reviewer, a physician, authenticates their findings (step 2) [5]. The two primary reviewers, either nurses or other health personnel with clinical background, review the records independently in a predefined order; discharge codes (particularly infections, complications, or certain diagnoses), discharge summary, medications administration record, laboratory results, prescriber orders, operative record, nursing notes, physician progress note and last if time permits; history, consult notes and emergency department notes. The reviewers look for any of the triggers and possible concurrent adverse events within a maximum 20-minute review time limit per record. The intention of reviewing for triggers is that this provides a more efficient and focused review of the records to identify adverse events instead of reviewing the records in their entirety. This approach help select the records in the sample that are more likely to have documented an adverse event. The triggers are classified according to the care that is provided in addition a medication module:

- General Patient care
- Surgical care
- Perinatal care
- Intensive care
- Care given in the emergency department

If a trigger is identified, the reviewer checks the relevant documentation to determine if the trigger is related to an adverse event according to the GTT definition: “unintended physical injury resulting from or contributed to by medical care that requires additional monitoring, treatment or hospitalization, or that results in death” [5]. For example, a venous thrombosis in the leg after a hip replacement is an unintended outcome, while the permanent scar from the
surgery is an intended outcome. The former is an adverse event and the latter is not. With this approach, all unintended events presented as signs, symptoms and diseases and that requires intervention, are considered an adverse event. To help the reviewers to determine whether an event is an adverse event, the following questions should be asked [5]:

- “Would I be happy if it happened to me?”
- “Was it a natural progression of the underlying disease?”
- “Was it an intended result of care?”

If the answers are no in all three questions, it is likely an adverse event. With these questions the method focus on how the patient perceives the event and stress that the patient’s perspective should be emphasized when deciding if the event is an adverse event or not.

In some cases, it can be difficult to distinguish between consequences of medical care and the natural progression of the underlying disease as referred earlier. For example, if the patient suffers from a brain tumour and is treated with an operation and the patient receives blood transfusion after the operation- is the blood transfusion a result of an adverse event (e.g. the patient experienced unexpected or excessive blood loss) or was it due to the disease? In this case there was no reaction to the transfusion, but the transfusion was not a planned event. In such cases the event could be defined as an adverse event. Another example of an adverse event is if a patient develops a urinary tract infection while or after having an indwelling urine catheter. In the last case the infection is obviously due to the use of the catheter. Determining that this is an adverse event should be straightforward. The former described case with the blood transfusion is more difficult. Hence, the determination is to some extent a matter of the subjectivity of the reviewers although the common definition and guidelines should be used.

After an adverse event is identified, the reviewer determines the severity level of the event. The grading of the severity is based on a modification from the National Coordinating Council for Medication Error Reporting and Prevention Index with categories ranging from A to I (NCC MERP) [79]. The categories A-D concern events that do not reach or cause any harm to the patient (near-misses): Category A is circumstances or events that have the capacity to cause adverse events, while category B is adverse events that do not reach the patient. Category C is adverse events that reach the patient but do not cause harm, and
category D is adverse events that reach the patients and monitoring to confirm that no harm occurred is required. The few events reported through the voluntary reporting system are often near-misses. The category A-D is not included in the GTT definition as only events that cause harm to the patient are classified as adverse events in the GTT:

- Temporary harm to the patient that required intervention (Category E)
- Temporary harm to the patient that required initial or prolonged hospitalisation (Category F)
- Permanent patient harm (Category G)
- Intervention required to sustain life (Category H)
- Patient death (Category I)

The adverse events are often classified according to their type. Classification of types it not a part of the original GTT, but included in the Norwegian translation of the GTT [19] (see appendices).

The results of the reviewed bi-weekly data are then presented in three ways:

- Adverse events per 1,000 patient days
- Adverse events per 100 admissions
- Percent of admissions with an adverse event

“Adverse events per 1,000 patient days” is the recommended measure to apply when evaluating the rate of adverse events, since this measure accounts for the different length of stay in the records. Longer length of stay is associated with adverse events [80]. The “Percent of admissions with adverse events” is more easily understood by non-clinical staff and is recommended to use when the results are shared public [5]. This measure does not include that some patients experience more than one adverse event or the variability of length of stay [5].

To visualize how the rate of adverse events change over time, continual data plotting in a run chart enables to uncover either upwards or downwards trends. The data series are plotted in a time sequence. Special cause variations are identified by looking for trends (six consecutive
jumps above or over the mean/median), shifts (eight or more point above/over the central line), patterns (pattern that reoccur) and last looking for outliers that lie far from the central line. A more advanced version is the control chart in Statistical process control (SPC) which includes the upper and lower control limits which detect special cause variation quicker and more accurate [81]. Random variations are synonym with common causes that are causes that cannot be eliminated or determined. If sample size increases, random variation decreases. The SPC is used to identify special causes, or systematic errors, that might influence the process [82].

When identifying adverse events according to the definition given in the GTT, the preventability of the events is not considered. The authors of the GTT explain that this is not included as the definition of what is preventable constantly change. Events considered unpreventable today can quickly change to preventable when new innovations are introduced. When evaluating the adverse events over time, categorization of preventable versus unpreventable adverse events will be meaningless over time [5]. In Sweden, the assessment of preventability of adverse events has been evaluated by a grading system from 1-6; where 1-3 are considered non-preventable and 4-6 are considered preventable [9]. Schildmeijer et al found great differences in the assessments of preventability and doubt the benefit of including this aspect as there are no standard of how to decide preventability. They argue, as other also have [6], that all adverse events should be considered preventable.

Also, when using the GTT to identify adverse events, events due to omission is excluded as the definition only includes events due to medical care given. For example, if the patient does not receive his antithrombotic medication when indicated, and a cardiac attack occur, this type of adverse event is not included in the GTT. Such cases are often due to missed diagnoses which is difficult to reveal as discussed in 1.2.4.

Hanskamp-Sebregts et al reviewed the literature concerning validity and reliability of the record review methods using the COSMIN checklist [83]. They evaluated the studies in regards to face validity and concurrent validity and they found no reference that the validity of the GTT were evaluated [38]. The inter-rater reliability between different review teams
have been reported moderate to substantial [38]. However, the face validity of the GTT is evaluated to some extent by Schildmeijer et al [84]. They found that the GTT was a useful method to identify adverse events.

Further discussions regarding challenges with identifying and measuring adverse events with the GTT method will be described in the next chapter.

1.3.3 Challenges

There are some issues to consider when using the GTT as a measure of adverse events. First, critics argue that the GTT is too resource intensive due to time and labour required [7], [85]–[87]. The GTT is based on a 20-minute maximum review time per record per primary reviewer which equates a maximum of six hours per reviewer per month if 10 records are reviewed bi-weekly. In addition, the time used of the authenticator is estimated to one to two hours per month [5]. Also, the method requires trained personnel to perform the review. The training is a recurring event every time a reviewer or authenticator is replaced.

Second, the results of the GTT are used to make estimates of the rate of adverse events which are based on reviewing a small sample of records. The authors of the GTT explain that if the same sampling strategy is used, the method is reliable for evaluating if the rate of adverse events is reducing or increasing [74]. The results are less accurate when a small number of records is used for estimating the rate, and make it less valid as a measure of the total number of adverse events [88]. The number of identified adverse events are used to estimate the total incidence of adverse events by extrapolation. Extrapolation is a statistical method estimating a value (e.g.: expected rate of adverse events) based on extending a known sequence of values beyond the area that is certainly known [89].

Third, identification of the individual triggers varies between reviewers as triggers based on indexed variables (i.e.; blood transfusion and dialysis) have higher agreement than triggers based on free text (i.e.; pressure ulcers, patient fall) [87]. The results are to some extent subject to the reviewers subjectivity as inter-rater reliability between reviewers and review teams have been reported from low to moderate [74], [90].
Schildmeijer et al addressed strength and limitations from the GTT reviewer’s perspective. They interviewed the reviewers concerning the usefulness and application of the GTT, preventability of the adverse events, review teams and dependence of the documentation provided in the health records [84]. They concluded that changing the approach of the method could influence the GTT results. They also meant that the reviewers should be more focused at looking at the patient’s perspective when deciding if an adverse event had happened.

These issues are further discussed in a review of the GTT which found widespread adoption with different modification demonstrating its flexibility [91]. With these concerns Hibbert et all proposed that “the GTT should be reframed as an opportunity to identify adverse events, raise awareness of these within hospitals and to describe the most frequent type of adverse events to prioritize quality improvement”, rather than an exclusively measuring method [91]. This demonstrate that the GTT could be modified in order to act as a method both for acknowledging and measuring adverse events.

Forster et al demonstrated that triggers were identified in 19-56 % of the records suggesting that half of the records are excessively reviewed when manual review for triggers are performed [92]. With automatic identification of triggers, manual reviews are only needed in records where triggers are identified in order to determine if the trigger is associated with any adverse events [87]. This reduce the number of records needed to be reviewed as the first part of the review (trigger identification) is done automatically. Such approach has been demonstrated to identify adverse drug events and adverse paediatric events with promising results [93], [94].

Accounting for these challenges we initiated our studies to evaluate the GTT method regarding sample size, inter-rater reliability and automatic identification of triggers.
2 AIMS OF THE THESIS

Overall aim
The general aim of this thesis was to evaluate the GTT method regarding sample size, changes of reviewers and automatic trigger identification to improve the method’s reliability and validity and to reduce the resources required.

Specific aims:

Paper I
To investigate the influence on the results of increasing the sample of reviewed records by the GTT.

Paper II
To evaluate the inter-rater reliability when reviewers are replaced when identifying adverse events by the GTT.

Paper III
To evaluate a modified GTT method with automatic trigger identification to the original GTT method with manual trigger identification.
3 MATERIAL AND METHODS

3.1 Setting

The GTT was implemented in Norwegian hospitals in 2011 as a part of the National Patient Safety Program “In safe hands” launched in 2010. All hospitals were required to review ten closed inpatients records randomly selected every bi-weekly period. Our trust, Nordland Hospital trust, chose to multiply the recommended sample size times seven. This was done partly because we wanted to measure adverse events separately for our seven main units, but we also thought that ten records reviewed bi-weekly were too small for reliable results. The trust implemented seven different GTT review teams corresponding to the seven different units. The seven review teams reviewed records discharged from their department respectively. The reviewers in the studies were recruited from the GTT review teams in the trust and had the same basic training with the GTT.

The electronic health record (EHR) system was implemented in the trust in 1992 (DIPS, ASA). The EHRs include both free text (i.e.: discharge summaries, operative reports, pathology reports, radiology results, transfer of service notes, admission notes, medical progress notes and notes from other healthcare professionals) and indexed variables (i.e.: laboratory results, admissions and discharge data, diagnosis and procedure codes). In Norwegian hospitals medication administration, prescriber orders and vital parameters are still hand-written and scanned into the EHRs but are currently being digitalized and indexed.

The first national Norwegian GTT results from all Norwegian hospital were used to estimate the number of deaths and harms caused by medical treatment. These calculations were made by extrapolations from the rate of the identified adverse events and contributed to major resistance and objections from health personnel against the GTT when published [20], [95]. The critics from the health personnel were mainly concerning the small sample size. Also, the definition of the adverse event defined as patient harm were not necessarily acknowledged by the clinical staff. Last, the GTT required resources which were considered unmanageable by the clinical staff. We designed these studies to examine the arguments from the critics.
3.2 Study design

All records included are selected from the discharge lists in Nordland Hospital Trust (figure 4 and figure 5). A total of 3153 different admissions were included altogether. Exclusion criteria were; patients aged 17 years or younger, patients admitted primarily for psychiatric or rehabilitation care, or patients with a length of stay less than 24 hours. The exclusion criteria were adapted from the GTT as the triggers are developed for adult somatic inpatients only [5].

Anonymous bi-weekly discharge lists were obtained from the hospital administrative system. Included records were randomly selected as described in the Norwegian GTT [19]. The discharge lists included information regarding type of admission (acute or planned), diagnoses, services which the patient was admitted to, case mix index (the value is dependent on diagnosis and the allocation of resources to care for and/or treatment included in the admission), wherever the patient underwent surgery, sex, age and length of stay.
Figure 4 Flowchart of the study populations in Paper I and Paper II

14267 records eligible for inclusion in 2010

Paper I
1680 records (10 records from each bi-weekly period in seven units from January 1th to December 31th)

Paper I
240 records (10 records from each bi-weekly period from January 1th to December 31th)

Paper II
120 records (10 records from bi-weekly periods from July to December)
In all three papers collection of data was done retrospectively. All three studies are observational studies. Observational studies are either prospective, retrospective or cross-sectional studies. In prospective studies a sample of study objects are classified in some way and then followed over a period to see if they develop a condition. In retrospective studies the cases of the condition have already occurred at study initiation, and the study investigates if the subjects were exposed to any risk factors. In cross-sectional studies the samples are randomly selected at a specific point in time and cross-classified if they have the condition or not. Strengthening the Reporting of Observational Studies in Epidemiology (STROBE)-checklist [96] was used to include relevant information in the papers.

In Paper I, the study was designed as an observational retrospective study. We chose this design as the data were collected retrospectively and the rate of identified adverse events in two different samples were compared. In the paper we have defined it as a cross-sectional study. This is not correct, as patient days were accounted for. We used the appropriate
statistical measurements and evaluated the study as a retrospective cohort study. The hypothesis was: will increasing number of records reviewed affect the results of identified adverse events? This was examined by comparing the rate, type and severity of adverse events identified by the GTT in two different sample sizes; one small and one large by obtaining the risk ratio (RR). Altogether 1920 records selected from the bi-weekly discharge lists were included. The large sample included records selected as 10 records bi-weekly from the seven units discharge lists (n=1680) while the small sample included ten records selected bi-weekly from the trust´s discharge lists (n=240) (figure 4). The manual review to identify triggers and adverse events differed in some way between the two samples. The records in the large sample were reviewed by one of three primary reviewers (two physicians and one nurse) and all three reached consensuses regarding the adverse events they all had identified separately. The records in the small sample were reviewed as described in the GTT; two primary reviewers (nurses) individually reviewed the records and reached consensus regarding the adverse events before a secondary reviewer (a physician) authenticated their common findings [5]. The reviewers of both samples were the same, except that one of the physicians from the small sample was replaced by a nurse in the large sample.

Paper II was designed as an observational cross-sectional study including 120 records (figure 4). We did not consider an alternative design as the study compares agreement between different reviewers regarding the prevalence of identified adverse events within a sample. The study evaluated the reproducibility of the method. The length of patient stay in the different records do not affect the results as the review teams review the same 120 records. Three review teams review the records as described in the GTT with two primary reviewers and one secondary reviewer [5]; Team I (three consistent reviewers- two primary reviewers and one secondary reviewer), Team II (one of the two primary reviewers or/and the secondary reviewer from Team I are replaced for different review periods) and Team III (no identical reviewers with Team I or Team II). The presence, type and severity of the adverse events identified by the three review teams were compared to assess the inter-rater reliability between the teams.
Paper III describes an observational cross-sectional study including 1233 records. The study evaluates two different methods to identify and measure adverse events, the modified GTT method versus the original GTT method. As in Paper II we did not consider an alternative design. 70 records were selected bi-weekly from the discharge lists from March 1th to December 31th 2013 (figure 5), but 167 records were excluded as data for these records were missing in the automatic trigger system. A modified GTT method, including automatic identification of triggers with manual review of the triggered records performed by a physician, was compared to the original GTT method [5]. The original GTT method included manual review of triggers and possible corresponding adverse events by two primary reviewers and authentication of their findings regarding adverse events was performed by a secondary reviewer. The identified adverse events by the modified GTT method were compared to the adverse events identified by the original GTT method. The concurrent validity of the modified GTT was evaluated by obtaining sensitivity, specificity, precision and reliability.

3.3 Intervention

In all three papers we evaluated the use of the GTT. The definition of an adverse event adopted from the GTT was applied in all three papers [5]. The training of the reviewers included the following understanding of how to determine if an adverse event was present: If the patient had experienced an unplanned event that led to either treatment, prolonged stay, permanent injury, immediate treatment to sustain life or death, the event was defined as an adverse event. The perspective of the patient was assessed by asking the questions as addressed in the GTT, mentioned in the chapter 1.3.2. [5]. Near misses that did not lead to the above criteria were not counted as adverse events and preventability of the adverse events was not evaluated.

The reviewers followed the approach as described in 1.3.2 except from the review team who reviewed the records of the large sample in Paper I. The triggers identified by the review teams and by the automatic trigger identification system was recorded in the databases. If the review team identified an adverse event, the type and severity was decided and recorded in
the databases. The categories of type of adverse events was adopted from the Norwegian GTT manual and sub classified in these main categories [19]:

- Surgical complications
- Bleeding/thrombosis
- Medication harm
- Patient fall
- Pressure ulcers
- Obstetric harm
- Other

### 3.4 Methodological consideration

Possible bias, presented in 1.2.2, could be present in all three studies. In Paper I selection and confounding bias are most likely to occur as the selection of records differed between the two samples. In Paper II hindsight bias could occur as the reviewers reviewed the records with different reviewers. The subjectivity of the reviewers could influence the results as the reviewers decided to some extent by themselves if the event was an adverse event or not. Selection bias is less likely in Paper II and Paper III as the reviewers reviewed the same records. In all three papers information bias could be present, as the findings of the adverse events rely on documentation in the records. Also, identification of triggers relies on that the information needed to identify a trigger is documented in the patient records. We consider that the results could be generalized for patient populations elsewhere.

### 3.5 Statistical analyses

The size of the small sample size in Paper I and Paper II are equal to the recommended sample size in the GTT with ten records selected bi-weekly. The size of the large sample in Paper I and Paper III with 70 records selected bi-weekly is the same as the total sample size used in our trust for the GTT. Power estimates of the sample sizes in Paper I was done with 80 % power. We assumed that the incidence of adverse events was 20 %. We then needed at least 7 % difference in the rate of identified adverse events between the samples for
significant results. In Paper II and Paper III kappa statistic was used. Power estimates was not performed, but if the CI is narrow then the power is considered good. The primary endpoint in all three papers was the rate of identified adverse events. Secondary endpoints were severity and types of adverse events.

To examine the means of the adverse event rates in the different samples, SPC charts were applied using QI Macros for Excel. The SPC charts include control limits, which are calculated based on the values presented and are ± 3 standard deviations from the central line (average). Any variation between the control limits are common causes of variation, while variation above or under the control limits are due to special cause variation.

Poisson regressions in Generalized linear models were applied to compare the rates of adverse events, severity level and categories of types of adverse events between the different sample sizes. Poisson regression was selected as it accounts for variation of number of cases and length of stay. Adjustments of demographical variables were done by including these as covariates. RR was obtained.

We used Cohen’s kappa to determine the inter-rater reliability. For nominal data (value of 0 if no agreement and value of 1 if perfect agreement) kappa statistic was applied. For ordinal data (values from 1 to usual not more than 4 or 5 is applied for the different categories) weighted kappa was applied. Weighted kappa was applied when comparing severity level as we decided that it was less agreement if the event was rated category E by one reviewer and category H by another reviewer than rated category E versus category F. If no adverse events were identified the value of 1 was applied, if the adverse event was severity category E value 2 was applied, severity category F value 3 was applied, severity category G value 4 was applied, severity category H value 5 was applied and severity category I value 6 was applied. The interpretations from Landis and Koch was used for the Cohen kappa coefficient: poor (<0.0), slight (0.00-0.20), fair (0.21-0.40), moderate (0.41-0.60), substantial (0.61-0.80) and almost perfect (0.81-1.00) [97]. The inter-rater reliability has been used to evaluate measures of adverse events by many [98].
Statistical association of categorical variables was assessed by Chi-square test while continuous data were compared using independent t-tests. To compare the number of adverse events identified by different methods or by different teams, Paired t-tests were used. When evaluating the performance of the modified GTT method, the original GTT method was set as gold standard. We calculated sensitivity (recall as used in Paper I), positive predictive value (or precision) (PPV) and specificity with their respective 95 % confidence intervals (CI) to evaluate the validity of the modified GTT method [99]:

\[
\text{Sensitivity} = \frac{\text{No. of correct positive records identified by the modified GTT method}}{\text{No. of positive records identified by gold standard}}
\]

\[
PPV = \frac{\text{No. of correct positive records identified by the modified GTT method}}{\text{Total no. of positive records identified by the modified GTT method}}
\]

\[
\text{Specificity} = \frac{\text{No. of correct negative records identified by the modified GTT method}}{\text{No. of negative records identified by gold standard}}
\]

The CI for sensitivity, PPV and specificity was calculated using the Wilson score method [100]. CI for Cohen’s Kappa was calculated as κ± 1.96*SE.

For all analyses, we used two-sided tests. The significance level was set at 5 % and 95 % CI were reported if relevant. The statistical analyses were performed using the SPSS statistical package, version 22.0 (SPSS Chicago, IL, USA).

When performing a statistical test there is always a chance of committing Type I error (incorrectly rejecting a true null hypothesis). The maximum probability of Type I error equals the specified significance level (5%) and we reject the null hypothesis whenever the p-value is lower than 0.05. In situations where we have multiple tests the Type I error probability in at least one test would be higher than 5 %. One way of reducing this error probability is to reduce the significance level proportionally to the number of tests (Bonferroni adjustments). Regardless of significance level, adjustment for other variables (age, length of stay etc.) increase the validity of our results.
3.6 Ethical consideration

The studies were performed in accordance to the Helsinki Declaration of 1975, and approved by the Norwegian Regional Committee for Medical and Health Research Ethics (Protocol ID: 2012/1691) and the Data Protection Office at the Nordland Hospital trust.

The information from the patient records were anonymised when extracted from the hospital administrative system and included in databases. The databases were hosted within an encrypted environment restricting the access to granted personnel only.

Information from medical records can be obtained for the purpose of internal control and quality assurance according to the Norwegian Health Professional Act. The trusts are required to obtain information and develop statistics about unintended events involving patients according to the regulations of the Norwegian Health Specialized Service Act. The health information can be obtained without consent in such cases. According to the same act the health services are also obligated to report severe adverse events to the Norwegian Board of Health Supervision.

The need for patient consent was waived for the records included in Paper III and for the largest sample in Paper I on the basis of: 1) the records had already been selected and reviewed as a part of the trust’s measurement of adverse events; 2) retrospectively collecting informed consent from patients or relatives of deceased patients would be costly with respect to time and money and might be considered a burden or inconvenience for the patients/relatives; 3) that the risk of being included and disadvantages of not being informed are considered minimal. This is in accordance to the criteria for waiving consent by Baker et al [101]. In Paper I we included a sample of records (n=240) that were not already selected for the trust’s measurements of adverse events. 120 of the 240 records used in Paper I were also used in Paper II. We argued that these patients should be contacted and asked for consent when we applied for approval of the study. 26 denied consent or did not respond to the consent letter. To include the correct amount of records (ten records bi-weekly), replacements of records were performed with random selection from the same discharge lists where the
patients had denied consent or not responded. The “new” included patients were also asked for consent. We included information in the consent letter that the patients could contact the study leader upon questions and that they at any time could withdraw their consent. Retrospectively we considered that asking these patients for consent was not necessary in concordance with the referred criteria.

There is great variability in the interpretation of research issues related to patient safety and quality. Research committees and national legislations practice consent waiving differently [102]. The WHO recommend that when in doubt, all projects should be submitted to the ethic committees before study start, to determine if consent is needed [103]. The ethic committees can waive the usual requirement of individual informed consent when the research involves minimal risks and obtaining consent would be impracticable [104]. An alternative when formal consent is waived, is to provide information of the studies being performed either by posters, leaflets or as a part of the general patient information [103]. In our trust we provide information in the lobby regarding that patient records are being reviewed to identify adverse events, along with the identified rate of adverse events (figure 6).

**Figure 6** The lobby in Nordland Hospital Trust, Bodø
4. RESULTS

The thesis examined different methodological aspects of the GTT record review method. In the first part of the study, the results suggested that increasing the sample sizes narrowed the CI thereby giving more precise results that can be extrapolated to institutional levels. The rate of identified adverse events was higher in a large sample compared to a small sample. The second part of the study evaluated how a larger sample of records could be efficient reviewed with valid results. The review of the triggered records should be done with consistent reviewers and automatic trigger identification enabling increasing the sample size without increasing the resources needed.

4.1 Patient characteristics

Demographic variables for the included 3153 patients compared to all records eligible for selection in 2010 and 2013, are presented in table 4. Median age was 64 (range 18-84) years. Most of the patients were women (60 %). Adverse events were identified in 655 (21 %) of the records included in the studies.
Table 4 Demographic characteristic

<table>
<thead>
<tr>
<th>Hospital</th>
<th>Included records (n=3153)</th>
<th>Mean (SD)</th>
<th>All records (n=25938)</th>
<th>Mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N (%)</td>
<td></td>
<td>N (%)</td>
<td></td>
</tr>
<tr>
<td>Bodø</td>
<td>2301 (73)</td>
<td></td>
<td>17153 (66)</td>
<td></td>
</tr>
<tr>
<td>Lofoten</td>
<td>395 (13)</td>
<td></td>
<td>4201 (16)</td>
<td></td>
</tr>
<tr>
<td>Vesterålen</td>
<td>457 (15)</td>
<td></td>
<td>4584 (18)</td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤ 65</td>
<td>1632 (52)</td>
<td>60.1 (21.3)</td>
<td>12335 (48)</td>
<td>62.1 (20.7)</td>
</tr>
<tr>
<td>&gt; 65</td>
<td>1521 (48)</td>
<td>(21.3)</td>
<td>13603 (52)</td>
<td></td>
</tr>
<tr>
<td>Type of admission</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acute</td>
<td>2240 (71)</td>
<td></td>
<td>19092 (74)</td>
<td></td>
</tr>
<tr>
<td>Planned</td>
<td>913 (29)</td>
<td></td>
<td>6846 (27)</td>
<td></td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>1258 (40)</td>
<td></td>
<td>11189 (43)</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>1895 (60)</td>
<td></td>
<td>14749 (57)</td>
<td></td>
</tr>
<tr>
<td>Number of patient days</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3≤</td>
<td>1421 (45)</td>
<td>6.1 (6.3)</td>
<td>11082 (43)</td>
<td>6.1 (6.5)</td>
</tr>
<tr>
<td>3&gt;</td>
<td>1732 (55)</td>
<td>(7.3)</td>
<td>14856 (57)</td>
<td></td>
</tr>
<tr>
<td>Adverse event present*</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>655 (21)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>2498 (79)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Adverse events identified in at least one of the studies
4.2 Paper I

We found that a large sample size of 70 records selected bi-weekly identified 45% (RR: 1.45 CI: 1.07-1.97) more adverse events per 1000 patient days, than a smaller sample size of ten records selected bi-weekly. In the large sample 39.3 adverse events per 1000 patient days (CI: 35.8-43.1, SE: 1.86) were identified while in the small sample 27.2 adverse events per 1000 patient days (CI: 20.3-36.4, SE: 4.05) were identified. The difference was significant (p=0.02, CI: 1.04-1.93). As expected, the CI was narrower and the SE was lower in the large sample than in the small sample. However, there was no difference regarding variation over time between the samples. This is in accordance with the main purpose of the GTT; to monitor the rate of adverse events over time. There was no significant difference between the samples regarding length of stay, average age or sex. When adjusting for services, diagnosis, case mix index, surgical treatment, acute or planned admission and numbers of transfers related to the index hospitalisation, the overall results were not altered.

SPC charts were applied to compare the mean rate of adverse events over time to examine if any of the tests of special causes were positive. In the small sample test 1 was positive (i.e.; data points outside the control limits). None of the tests were positive for the large sample.

**Figure 7** Number of adverse events per 1000 patient days in SPC U-chart
Hospital acquired infection was the most frequent type of adverse event in both samples followed by surgical related harms, medication harms, bleeding/thromboembolism, patient falls, pressure ulcer and obstetric harms (figure 8). No significant difference between the samples regarding the types of adverse events or the severity level was identified. 57% of the adverse events identified in the large sample were defined as category E (harms requiring interventions) compared to 56% in the small sample (RR: 1.5 p= 0.054, CI: 0.99-2.26). Respectively 39% and 33% of the adverse events were category F (RR: 1.69 p=0.051, CI: 1.00-2.86) and 3% and 11% were defined as severe adverse events (category G, H or I) (RR: 0.47 p=0.14, CI: 0.17-1.27).

**Figure 8** Types of adverse events identified
4.3 Paper II

120 records were reviewed by three review teams; Team I, Team II and Team III. Team I and Team II had one or two identical reviewers throughout the review of the 120 records. Team III had none identical reviewers with Team I and Team II. Team I identified 23 adverse events, Team II identified 20 adverse events and Team III identified 18 adverse events (figure 9). Team I and Team II identified six identical adverse events. The same six adverse events were not identified by Team III. In seven records Team III disagreed with Team I in regard of type of adverse event, while Team II disagreed to Team I in three records.

Figure 9 Number of identified adverse events by the three teams

We found that the agreement in regards of presence of adverse events was substantial ($\kappa=0.64$) when one or two of the reviewers were identical (Team I versus Team II) compared to moderate ($\kappa=0.47$) when none reviewers were identical (Team I versus Team III).

Regarding the number of adverse events and categorizing of the severity level, the agreement
was substantial between Team I and Team II compared to between Team I and Team III (table 5).

**Table 5** The level of agreement between Team I and Team II and between Team I and Team III in terms of adverse events and severity level

<table>
<thead>
<tr>
<th></th>
<th>Team I vs Team II (kappa coefficient, 95 % CI)</th>
<th>Team I vs Team III (kappa coefficient, 95 % CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Presence of adverse events*</td>
<td>0.640 (0.434-0.846)</td>
<td>0.468 (0.232-0.703)</td>
</tr>
<tr>
<td>Number of adverse events**</td>
<td>0.661 (0.479-0.842)</td>
<td>0.468 (0.278-0.694)</td>
</tr>
<tr>
<td>Severity level**</td>
<td>0.652 (0.469-0.836)</td>
<td>0.442 (0.260-0.624)</td>
</tr>
</tbody>
</table>

*Unweighted kappa analysis, **Weighted kappa analysis
4.4 Paper III

We evaluated the performance of a modified GTT method (figure 10). The modified GTT method included manual reviews for adverse events in 658 records identified with triggers by an automatic trigger identification system. The automatic trigger system screened 1233 records. The results were compared to the original GTT method which included manual review of all 1233 records to identify both triggers and adverse events.

Figure 10 The modified GTT method (Illustrated by Laila Bjølgerud)

The modified GTT method identified the same rate of adverse events as the original GTT method; 35 adverse events per 1000 patient days. Sensitivity, PPV, specificity and reliability for records identified with adverse events were respectively 0.59, 0.58, 0.92 and 0.51 for the modified GTT method in respect to the original GTT method as gold standard. The total manual review time in the modified GTT method was 23 hours, while the manual review time using the original GTT method was 411 hours.

Number of records identified with adverse events (15.3 % versus 15.1 % of the total number of records, p=0.81, CI; -0.02-0.02) and number of identified adverse events (p=0.90, CI; -0.03-0.03) did not differ significantly between the modified GTT method and the original
The modified GTT method reduced the number of records needed to be manual reviewed by 50% (figure 11).

**Figure 11** Number of records identified with triggers and adverse events by the modified GTT method and the original GTT method

- Records with adverse events identified by both methods, n=110
- Records with adverse events identified by the modified GTT method, n=189
- Records with adverse events identified by the original GTT method, n=186
- Records with triggers, n=762
- Records without triggers, n=471
- Records in sample, n=1233
- Records with triggers identified by the modified GTT method, n=658
- Records with triggers identified by both methods, n=522
  \((257+69+45+109)\)
- Records with triggers identified by the original GTT method, n=626
  \((104+522)\)
5. DISCUSSION

5.1 Summary of strength and weaknesses

A summary of the strengths and weaknesses of the studies included in this thesis are presented in Table 6.

Table 6 A summary of strength and weaknesses

<table>
<thead>
<tr>
<th>Weaknesses</th>
<th>Strengths</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Study design</strong></td>
<td></td>
</tr>
<tr>
<td>Only two sample size (10 vs 70 records bi-weekly) compared</td>
<td>Large number of records reviewed from bi-weekly periods</td>
</tr>
<tr>
<td>Different sampling methods</td>
<td>Adjusted for different sample methods</td>
</tr>
<tr>
<td>No power estimates were performed</td>
<td>Records were randomly selected</td>
</tr>
<tr>
<td>Some of the triggers were not possible to identify automatically</td>
<td>The sample size recommended in the GTT was applied</td>
</tr>
<tr>
<td>Small samples increase the risk of type 2 error</td>
<td>Automatic identification of 42 triggers</td>
</tr>
<tr>
<td>Information bias due to retrospective collection of data</td>
<td></td>
</tr>
<tr>
<td>No validation to an external patient cohort</td>
<td>The results correspond with other studies</td>
</tr>
<tr>
<td>Cross-sectional studies require large samples</td>
<td>Observational study design allows for different variables</td>
</tr>
<tr>
<td><strong>Reviews</strong></td>
<td></td>
</tr>
<tr>
<td>All reviews were performed by expert reviewers</td>
<td></td>
</tr>
<tr>
<td>Reviewers underwent the same training program</td>
<td></td>
</tr>
<tr>
<td>Data rely on documentation in the EHR</td>
<td></td>
</tr>
<tr>
<td>Identification of adverse events rely on triggers identified</td>
<td>A common definition of adverse events was applied in all paper</td>
</tr>
<tr>
<td>Manual review demanded time and methodical skills to assembly</td>
<td></td>
</tr>
<tr>
<td><strong>Data analysis</strong></td>
<td></td>
</tr>
<tr>
<td>Generalized method was used accounting for different length of stay and different sample sizes</td>
<td></td>
</tr>
<tr>
<td>Performance of a modified GTT method demonstrated a valid method to measure adverse events</td>
<td></td>
</tr>
<tr>
<td>Manual review is difficult to reproduce and compare between studies</td>
<td>Inter-rater reliability between reviewers was obtained when applicable and was substantial</td>
</tr>
<tr>
<td>A minimum P-value has increased type 1 error (false positive) and difficult to compare across studies</td>
<td>A minimum P-value approach is appropriate for exploratory studies (reducing type 2)</td>
</tr>
</tbody>
</table>
5.2 Paper I

Our study demonstrated that increasing the sample size affected the rate of adverse events, while the type and severity of identified adverse events were not influenced. 1.45 more adverse events per 1000 patient days were identified in the large sample (n=1680 records) than in the small sample (n=240). We argue that the rate of adverse events identified in the large sample is more representative and precise than the rate of adverse events identified in the small sample due to the narrow CI in the large sample.

A narrow CI makes the results more precise, and extrapolation with such results makes estimates of the total number of adverse events less uncertain. Many have debated the difficulty with estimating rates of adverse events based on small samples, and our results demonstrate this challenge [63], [105], [106]. Also the infrequent severe adverse events are often missed when sampling approaches are used [107]. No severe adverse events (category I) were identified in the small sample. Due to the infrequently occurrence of severe adverse events, other methods should be used to monitor these specific types of events, for example, investigating all hospital deaths [108], [109].

In the small sample there were an outlier with excessive patient days. We tested if exclusion of 10% of the patient days form each sample altered the results. In the large sample 24 records with a total length of stay of 1150 patient days were excluded. In the small sample two records were excluded with a length of stay of 197 patient days. The result was not altered as the RR was 1.55 (CI: 1.1-2.1). In the large sample the rate was 39.2 adverse events per 1000 patient days while in the small sample the rate was 25.3 adverse events per 1000 patient days. As an explanation of the difference in rate, we therefore argue that this is most likely due to random variation as other explanations were adjusted for.

Several studies refer to high sensitivity [6] and acceptable reliability [7], [110] of the GTT, but the impact of the sample size has been discussed only by few. To our knowledge this study is the first attempt to assess the impact of the sample size on the results identified with the GTT. The large sample in our study represented approximately 12% of the overall patient
population in our trust. Kennerly et al, among others, proposed that the sample size of records to be reviewed should be adjusted to the hospital size [6], [75], [111].

It is important to consider the Simpson’s paradox when evaluating the results. This is implying that statistical results from aggregated data could give a different result when extracting the results to a group-level analysis [112]. This is important to be aware of when using the statistics for causal interpretations. We therefore adjusted for the variations such as case mix, if surgery was performed, case mix index, hospital locations, units and type of admissions, which were correlated to the index discharges (the sources of the selection of the records). When adjusting for these variations the results were not changed; the rate of adverse events was still significantly different between the two samples while the type and severity were not. The results did also not differ when adjusting for demographic variables such as gender, age and total length of stay. We do therefore not consider that the Simpson’s paradox is a relevant problem in this study.

Another factor is that we do not have any information of the patient records in the small sample where patients had denied consent or refused to answer. These patients could have experienced an adverse event and would not participate because of a bad experience with the trust. This could bias the result from the small sample as the aim was to compare the rate of adverse events between the samples.

The SE of the mean represents the degree of the variability of the mean. The SE is low in the large sample while the small sample has a higher SE. The means of the rate of adverse events identified in the large sample has less random variability. With these assumptions we consider that a larger sample include more trustworthy results. The smaller sample is less resilient for outliers as there are too few records included.

The review process of the two samples differed slightly as described in the study design. To adjust for this possible bias, we assessed the agreement between the different authentication processes and found the agreement to be substantial. Zegers et al concluded that different
authentication processes did not impact the results [113]. We argue that the difference in review process between the samples did not influence the results in our study.

The power estimate of the large sample size was based on a difference of 7 % between the samples with 80 % power. The difference in the identified adverse events rate between the samples was 45 %. We therefore assume that the sample size was large enough. A larger sample size could reflect the population more accurate than a smaller sample and the rate of adverse events that were identified in a larger sample could be more reliable [114]. Further, we could have included more records by enhancing the study length period. Variation of number of patients and medical care given differ more between the different parts of the year than between two different years. We assumed that inclusion of records from one year was enough to obtain reliable results.

Other limitations when interpreting the results, is the categorization of types of adverse events which are not mutually exclusive. The determination of type of adverse event is based on the subjectivity of the reviewers as no common definitions of which type of adverse events to include in the different categories exists.

The length of stay in the records, which is the denominator in the estimated rate of adverse events per 1000 patient days, must be accounted for when comparing the means of the rate of adverse events. We therefore applied the Poisson regression in the generalized linear models as it is appropriate for rate data when the dependent variable is a count of events divided by some measure of that unit’s exposure, i.e. number of adverse events per 1,000 patient days. The difference in number of records included in the two samples is also being accounted for when using this statistical test. The RR could then be obtained. The wide control limits in the SPC chart of the mean rates in the small sample demonstrated that these rates did vary more than the rates identified in the large sample.

Our results imply that the recommend sample size of ten records reviewed bi-weekly is too uncertain. Hence, further studies are needed to determine whether there is an optimal sample size. For example, if the sample size should be based on hospital size, especially as reviewing
larger sample sizes requires more resources. Until further studies, we have suggested using a relative increase in sample size to 8–10% of total number of discharges when using the GTT to achieve a narrow CI and hence more precise results. The increase in sample size requires a more effective strategy to review the records which is evaluated in Paper II and Paper III.
5.3 Paper II

In this study we evaluated the inter-rater reliability as we compared the results from different review teams who reviewed the same records. Others have examined the inter-rater reliability and have found at best a moderate to substantial agreement [98]. However, in this study we demonstrated substantial inter-rater reliability between review teams where at least one of the reviewers were identical. Moderate inter-rater reliability was found between review teams with no identical reviewers.

Members in the review teams performing the GTT are often replaced due to practical issues such as relocation of work place, sick leave and maternity/paternity leave. To our knowledge, this is the first attempt to assess inter-rater reliability between review teams experiencing replacement of reviewers to varying degrees. Evaluating the inter-rater reliability between all different teams, as replacement of all reviewers, have been described previously and reported to be poor [68]. We therefore evaluated how the results are affected when review members are changed except from one of the primary reviewers. We chose to keep one of the primary reviewers consistent as the GTT recommend that the primary reviewers are the ones who conduct the first screening of the records and therefore most important to keep consistent [5]. We considered replacement of both primary reviewers as equal to replacement of all review members as the primary reviewers perform the initial review. The secondary reviewer only authenticates the findings without accessing the records routinely. Unlike our assumptions O’Leary et al highlighted that the variation was higher between confirmation of adverse events than for identification of potentials adverse events [115].

The variables concerning the reviewers such as review experience, clinical background and years of experience could influence the results. The mean years of clinical experience was 18.3 years (range 7-29) of the reviewers and the total mean years of review experience of the three teams were 2 years. To evaluate the agreement of identified adverse events between the teams, the kappa statistics was used. This analysis is not able to adjust for clinical experience between the teams. We have therefore not discussed any influence such as psychology or social influence of the consistent reviewer. This viewpoint would be more of a study of group
dynamics which was not the intention of this study. We intended to evaluate a practical solution with a pool of reviewers performing the GTT at different times without influencing the results.

Our findings indicate that hospitals can rely on rotating reviewers from a consistent pool of reviewers in order to optimize resources. With this approach hospitals are encouraged to perform the GTT even if they experience frequent replacement of reviewers. However, the CI is wide which indicate that the sample size might not be large enough. Our results must therefore be interpreted with some caution.
5.4 Paper III

Identifying and measuring adverse events in hospitalised patients is challenging. So far, we consider the GTT the most robust method to measure adverse events in comparison to most other existing methods. The practical disadvantage with the method being resource intensive, can to a certain extent be addressed by automating the trigger identification. We developed an automatic trigger identification system to automate 42 of the GTT triggers. The study demonstrated that the modified GTT method using automatic trigger identification is a valid measure in respect to the original GTT method. To our best knowledge such study has not been performed previously.

Since the late 90’s Classen et al along with others, have demonstrated computerized surveillance of adverse drug event by automated detection of triggers that could represent possible adverse events [116]–[121]. When triggers are automatically identified, only the records with triggers are reviewed manually to determine if the trigger represents an adverse event. This approach has showed promising results [93], [94].

The “gold standard” of determination of an adverse event has traditionally been the judgment of clinicians [122]. Automatic identification of adverse events based on administrative data have showed disappointing results [123]. With such approach the positive predictive value are reported to be low, ranging from 12-30 % [124]–[136]. We consider that a manual review is still needed to determine if the triggers automatically identified, represent an adverse event according to the GTT definition applied in the study. However, machine learning is slowly integrated in medical decisions, such as radiology imagination and treatment outcomes [137], [138]. In the future it is therefore possible that adverse events can be identified automatically. We consider that the automatic trigger identification system could be further developed to a system that can predict which patients who are at risk to experience an adverse event enabling the clinicians to act in real-time to prevent adverse event.

The modified GTT method, with manual review of only records with automatic identified triggers, demonstrated a more resource efficient method than the original GTT method. The
number of manually reviewed records were reduced by 50 % with the modified GTT method (n=658) compared to the records manually reviewed in the original GTT method (n=1233). This is because the original GTT method demands the reviewers to screen all records to identify any triggers and then do a more in-depth review when triggers are identified to find any possible corresponding adverse events. In the modified GTT method the automatic trigger identification system performed the screening of triggers and manual review was only performed in the triggered records. The time used with the modified GTT method was only 6 % of the time used with the original GTT method. We consider this an exceptional result. Others have showed that the time using computerized strategies is 20 % compared with the time used with manual strategies [107], [139].

We found good agreement between the two methods with regards to the records identified with adverse events (κ=0.51 CI: 0.44–0.57). Our results demonstrated better agreement between the automated method versus all manual methods, compared to other studies, who have found only up to 12 % agreement [115], [139]. The modified GTT method identified 59 % of the records identified with adverse events by the original GTT method (110 of 186 records). The variation between the methods concerning the difference of number of records identified with adverse events could be explained by using different review team. The automatic triggered records were reviewed by one physician. The original GTT method was performed as described in the GTT manual [5]. We have argued that using different reviewers may affect the results as demonstrated by for example O’Leary et al [115]. However, we concluded that this did not bias the results in this study.

A recent review by Hibbert et al found that the GTT identified adverse events in 7-40 % with a cluster around 20-29 % of the reviewed records [91]. O’Leary et al found that 22- 26 % of records identified by automatic system were confirmed with adverse event [115]. The modified GTT method confirmed adverse events in 33 % of the triggered records. To examine how many of the total number of records are triggered, we ran the automatic trigger identification system for all admissions eligible for inclusion in the GTT in 2017. The automatic trigger identification system identified at least one trigger in 62 % (n=10807 records) of the records. The modified GTT method identified adverse events approximately in
30 % of the triggered records in the study, constituting 15 % of the original record sample. If we apply this result to the aggregated numbers of 2017, the estimated number of records with adverse events would be 3242 records; or one of five hospitalised patients are harmed due to medical care. This emphasize the modified GTT method as a valid method to measure adverse events [6].

We have not considered the financial aspect of the automatic tool as this was beyond the scope of the study. The GTT method is criticized because it is resource intensive due to time and personnel required. We have therefore evaluated how to reduce resources in regard to personnel and time needed for reviewing the records.

Our results recommend that the modified GTT should be preferred rather than the original GTT method, as the modified GTT method is less resource intensive. The resources saved by using the modified GTT method is considerable, enabling increasing the sample size as proposed in Paper I and reviewing the records with consistent reviewers as demonstrated in Paper II.
6. CONCLUSION

In Paper I we found that increasing the sample size provides a narrower CI, reduce the random variation and increase the precision of the results. The rate of identified adverse events was higher in a large sample than in a small sample. We argue that a large sample should be preferred as this is a more reliable source for extrapolation of rates when calculating the total number of adverse events. In Paper II we demonstrated that keeping one reviewer consistent provide more reliable results. Using the modified GTT method, as demonstrated in Paper III to identify and measure adverse events, is a time-effective strategy. We suggest that our findings can guide hospitals to identify and measure adverse events more effectively and that using such approaches would gain valid and reliable results.
7. IMPLICATION FOR FUTURE RESEARCH

The results of these studies lead to suggestions of some changes in the practical use of the GTT. First, we suggest that the sample size should be increased, second, we argue the importance of keeping one of the primary reviewers consistent and finally we introduced automatic trigger identification as a successful alternative approach to the manual GTT. These implications could facilitate more widespread adoption of the GTT as a method to identify and measure adverse events.

Future research could include a comparison of review of the automated triggered records by two different review teams or by two different reviewers. This could be done as a cross-sectional study comparing the rate of identified adverse events but also comparing the findings in each record by the two review teams/reviewers. If the agreement is substantial it could demonstrate that automatic trigger identification increase the agreement with an objective screening of the records.

Also, an automatic trigger identification system could be developed further to a prospective approach. Sammer et al presented a system allowing for real-time bedside intervention, real-time trend analysis and continued learning about harm measurement using a sociotechnical approach of people, process and technology [107]. Their framework emphasizes the framework of Donabedian [140] to assess quality of care; structure, process and outcome. We believe that moving to a prospective system, to identify patient at risk, would be beneficial for the clinical health personnel as it allows them to prevent adverse events from happening to the actual patient. Novel technologies such as identifying risk factor for developing adverse events must be integrated in the EHR. This could be performed either as a cohort study or as a cross-sectional study depending on the study questions. Such prospective system could be used to improve clinical outcome, optimize treatment, reduce the financial burden of patient harm, reduce the burden for involved health personnel and most importantly; reduce the suffering for patients due to adverse events.
8. REFERENCES


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[77] P. Davis and New Zealand. Ministry of Health, “Adverse events in New Zealand


[98] K. Walshe, “Adverse events in health care: issues in measurement,” *JAMA*, vol. 265,


APPENDICES
Norwegian GTT Trigger sheet

Norwegian GTT Categories of harm and severity

Approval from the Regional Committee for Medical and Health and Research Ethics

Letter from the Norwegian Social Science Data Service

Approval from the Data Protection Office at the Nordland Hospital trust

Trial invitation letter

Informed consent form

Paper I-III
## Non-Automatic Triggers

<table>
<thead>
<tr>
<th>Trigger</th>
<th>Care module Triggers</th>
<th>Medication Module Triggers</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>Transfusion or use of blood products</td>
<td>M1</td>
</tr>
<tr>
<td>C2</td>
<td>Code/arrest/rapid response team</td>
<td>M3</td>
</tr>
<tr>
<td>C3</td>
<td>Acute dialysis</td>
<td>M4</td>
</tr>
<tr>
<td>C4</td>
<td>Positive blood culture</td>
<td>M5</td>
</tr>
<tr>
<td>C5</td>
<td>X-ray or Doppler studies for emboli or DVT</td>
<td>M6*</td>
</tr>
<tr>
<td>C6</td>
<td>Decrease of greater than 25% in hemoglobin or hematocrit</td>
<td>M7*</td>
</tr>
<tr>
<td>C7</td>
<td>Patient fall</td>
<td>M8*</td>
</tr>
<tr>
<td>C8</td>
<td>Pressure ulcers</td>
<td>M9*</td>
</tr>
<tr>
<td>C9</td>
<td>Readmission within 30 days</td>
<td>M10*</td>
</tr>
<tr>
<td>C10*</td>
<td>Restraint use</td>
<td>M11*</td>
</tr>
<tr>
<td>C11</td>
<td>Healthcare-associated infection</td>
<td>M12*</td>
</tr>
<tr>
<td>C12</td>
<td>In-hospital stroke</td>
<td>M13*</td>
</tr>
<tr>
<td>C13</td>
<td>Transfer to higher level of care</td>
<td><strong>Intensive Care Module Triggers</strong></td>
</tr>
<tr>
<td>C14</td>
<td>Any procedure complication</td>
<td>I1</td>
</tr>
<tr>
<td>C15*</td>
<td>Other</td>
<td>I2</td>
</tr>
<tr>
<td></td>
<td><strong>Surgical Module Triggers</strong></td>
<td>I3</td>
</tr>
<tr>
<td>S1</td>
<td>Return to surgery</td>
<td>I4</td>
</tr>
<tr>
<td>S2</td>
<td>Change in procedure</td>
<td>P1*</td>
</tr>
<tr>
<td>S3</td>
<td>Admission to intensive care post-op</td>
<td>P2</td>
</tr>
<tr>
<td>S4</td>
<td>Intubation/reintubation/BiPap in PACU</td>
<td>P3</td>
</tr>
<tr>
<td>S5*</td>
<td>X-ray intra-op or in PACU</td>
<td>P4</td>
</tr>
<tr>
<td>S6</td>
<td>Intra-op or post-op death</td>
<td>P5</td>
</tr>
<tr>
<td>S7</td>
<td>Mechanical ventilation greater than 24 hours post-op</td>
<td>P6*</td>
</tr>
<tr>
<td>S8*</td>
<td>Intra-op epinephrine, norepinephrine, naloxone, or romazicon</td>
<td>P7</td>
</tr>
<tr>
<td>S9</td>
<td>Post-op troponin level greater than 40 ng/l</td>
<td>P8</td>
</tr>
<tr>
<td>S10</td>
<td>Injury, repair, or removal of organ because of accidental injury</td>
<td>P9</td>
</tr>
<tr>
<td>S11</td>
<td>Change in anesthesia procedure</td>
<td>P10</td>
</tr>
<tr>
<td>S12</td>
<td>Insertion of artery catheter or central venous catheter</td>
<td><strong>Emergency Department Module Triggers</strong></td>
</tr>
<tr>
<td>S13</td>
<td>Surgery more than 6 hours</td>
<td>E1</td>
</tr>
<tr>
<td>S14*</td>
<td>Any operative complication</td>
<td>E2</td>
</tr>
</tbody>
</table>

*Non-Automatic Triggers
**Severity of adverse event**

- Category E: Temporary harm to the patient and required intervention
- Category F: Temporary harm to the patient and required initial or prolonged hospitalisation
- Category G: Permanent patient harm
- Category H: Intervention required to sustain life
- Category I: Patient death

**Type of adverse event**

<table>
<thead>
<tr>
<th>Hospital acquired infections</th>
<th>Bleeding/thrombosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urinary tract infection</td>
<td>Thrombosis/Embolism</td>
</tr>
<tr>
<td>CVC infection</td>
<td>Bleeding</td>
</tr>
<tr>
<td>Ventilator associated pneumonia</td>
<td>Bleeding after surgery</td>
</tr>
<tr>
<td>Other infection</td>
<td></td>
</tr>
<tr>
<td>Lower respiratory infection</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Surgical complications</th>
<th>Patient fall /fracture</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infection after surgery</td>
<td>Patient fall</td>
</tr>
<tr>
<td>Respiratory complications after surgery</td>
<td>Fracture</td>
</tr>
<tr>
<td>Return to surgery</td>
<td></td>
</tr>
<tr>
<td>Injury, repair or removal of organ</td>
<td></td>
</tr>
<tr>
<td>Occurrence of any operative complication</td>
<td></td>
</tr>
<tr>
<td>Switch in surgery</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Other</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Other</td>
<td>Allergy</td>
</tr>
<tr>
<td></td>
<td>Medical technical harm</td>
</tr>
<tr>
<td></td>
<td>Deterioration and chronic illness</td>
</tr>
<tr>
<td></td>
<td>Medication harm</td>
</tr>
<tr>
<td></td>
<td>Obstetric harm</td>
</tr>
<tr>
<td></td>
<td>Pressure ulcer</td>
</tr>
</tbody>
</table>
Barthold Vonen  
Nordlandssykehuset HF  
Prinsens gt 164  
8000 Bodø

2012/1691 Validering av Global Trigger Tool som målemetode for kartlegging av pasientskader

**Forskningsansvarlig:** Nordlandssykehuset HF  
**Prosjektleder:** Barthold Vonen

Vi viser til søknad om forhåndsgodkjenning av ovennevnte forskningsprosjekt. Søknaden ble behandlet av Regional komité for medisinsk og helsefaglig forskningsetikk (REK vest) i møtet 18.10.2012. Vurderingen er gjort med hjemmel i helseforskningsloven § 10, jf. forskningsetikklovens § 4.

**Prosjektomtale**


**Vurdering**

**Søknad/protokoll**

Det hersker stor usikkerhet omkring spørsmålet om både omfang og alvorlighetsgrad av pasientskader ved norske sykehus. Både for pasientenes egen del og for samfunnets evne til å foreta nødvendige prioriteringer innenfor helsevesenet, er det svært viktig at det finnes gode og sammenlignbare oversikter over pasientskadene. Som politisk tema er dette også høyaktuelt. Derfor er dette en søknad REK Vest mener er svært viktig. Komiteen mener også at protokollen er egnet til å besvare de spørsmål en reiser.

**Rekruttering/samtykke**

Datamateriale hentes fra to kilder:

forskningsdatabase hvor koblingsnøkkel er fjernet. Videre opplyses det at arbeidet med å etablere forskningsdatabasen gjøres av personell utenfor selve forskningsprosjektet og i regi av forskningsansvarlig.

Adgang til bruk av helseopplysninger som er innsamlet i helsetjenesten til forskning er regulert i helseforskningslovens § 35. Vilkårene for å kunne tillate dette uten innhenting av samtykke, er at forskningen skal være av vesentlig interesse for samfunnet og at hensynet til deltakernes velferd og integritet er ivaretatt.

REK Vest mener at samfunnsnytten er godt dokumentert. Slik en har lagt opp anonymiseringsprosessen, mener komiteen at hensynet til deltakernes velferd og integritet også er godt ivaretatt. REK Vest vil godkjenne søknaden på dette punkt.


Det vedlagte utkast til forespørsel er imidlertid av dårlig kvalitet. En må bestrebe seg på å benytte et mer allment tilgjengelig språk hvor det er på en enklere måte beskrives hva deltakelse innebærer. REK Vest ønsker å få det reviderte skrivet tilsendt, før endelig vedtak fattes.

Informasjonssikkerhet

Det opplyses at koblingsnøkkel oppbevares ved egen institusjon og at personidentifiserbare opplysninger oppbevares på institusjonens server. REK Vest forutsetter at koblingsnøkkel og personidentifiserbare opplysninger oppbevares separat.

Vedtak

Søken utsettes i påvente av tilbakemelding på ovennevnte merknad.

Vennligst benytt skjema for tilbakemelding som sendes inn via saksportalen til REK http://helseforskning.etikkom.no.

Med vennlig hilsen

Jon Lekven
komitéleder, dr.med.

Kopi til: kso@nlsh.no
Barthold Vonen

2012/1691 Validering av Global Trigger Tool som målemetode for kartlegging av pasientskader

Forskningsansvarlig: Nordlandssykehuset HF
Prosjektleder: Barthold Vonen

Vi viser til tilbakemelding om forhåndsgodkjenning av ovennevnte forskningsprosjekt. Tilbakemeldingen ble behandlet av leder av REK Vest på fullmakt. Vurderingen er gjort med hjemmel i helseforskningsloven § 10, jf. forskningsetikkloven § 4.

Vurdering:

Tilbakemelding
REK Vest krevede at informasjonsskrivet ble forfattet i et mer allment tilgjengelig språk. Et revidert skriv foreligger nå.

Nytt vurdering i REK
REK Vest finner det nye informasjonsskrivet tilfredsstillende og har ingen ytterligere innvendinger til prosjektsøknad.

Vedtak:

REK Vest godkjener prosjektet i samsvar med søknad og tilbakemelding.

Sluttmelding og søknad om prosjektendring
Prosjektleder skal sende sluttmelding til REK vest på eget skjema senest 30.06.2016, jf. hfl. §12. Prosjektleder skal sende søknad om prosjektendring til REK vest dersom det skal gjøres vesentlige endringer i forhold til de opplysninger som er gitt i søknaden, jf. hfl. §11.

Klageadgang

Med vennlig hilsen

Jon Lekven
komitéleder

Rolf Fredriksen
prosjektleder

Kopi til: postmottak@nlsb.no

Jon Lekven
komitéleder

Øyvind Straume
seniorkonsulent
Barthold Vonen  
SKDE  

2012/1691 Validering av Global Trigger Tool som målemetode for kartlegging av pasientskader  

Forskningsansvarlig: Nordlandssykehuset HF, Nordlandssykehuset HF  
Prosjektleder: Barthold Vonen  

Vi viser til søknad om prosjektendring datert 05.02.2018 for ovennevnte forskningsprosjekt. Søknaden er behandlet av leder for REK vest på fullmakt, med hjemmel i helseforskningsloven § 11.  

Vurdering  
REK vest omfatter det slik at prosjektendringen innebærer ikke innsamling av nye data. Det er testing av validiteten til GTT som er formålet med prosjektendringen. Videre søker prosjektlederen om forlengelse av prosjektet til 05.07.2018.  

Vurdering:  
REK vest merker seg at prosjektet er gått ut på dato 31.08.2017. Vi gjør oppmerksom på at søknad om forlengelse av prosjektet skal sendes inn før prosjektsluttdato.  

Vedtak  
REK vest godkjenner prosjektendringen i samsvar med forelagt søknad.  

Klageadgang  

Med vennlig hilsen  
Marit Grønning  
dr.med.  
Avdelingsdirektør, professor  

Anna Stephansen  
sekretariatsleder  

Kopi til: postmottak@nlsh.no; postmottak@nlsh.no
AVSLUTTET SAKSBEHANDLING

Vi viser til meldeskjema mottatt 23.10.2014 for prosjektet:

40442 Validering av Global trigger tool som metode for kartlegging av patientskader

Vi viser også til REK-godkjenningen som var vedlagt meldeskjema, og telefonsamtale 28.11.14 med bekreftelse på at REK-godkjenningen dekker hele prosjektet.

REK har vedtatt at prosjektet faller inn under helseforskningslovens bestemmelser. REK sin godkjenning er tilstrekkelig for behandling av personopplysninger i prosjektet.

Personvernombudet avslutter dermed saksbehandlingen av meldingen uten å realitetsbehandle denne. Vi avslutter også all videre oppfølgning av prosjektet.

Ta gjerne kontakt dersom noe er uklart.

Vennlig hilsen

Katrine Utaaker Segadal

Kontaktperson: Inga Brautaset inga.brautaset@nsd.uib.no

Kopi: Institutt for klinisk medisin, UiT Norges arktiske universitet
ANBEFALING AV BEHANDLING AV PERSONOPPLYSNINGER

Viser til melding om behandling av personopplysninger, mottatt 21.06.

Tittel: Validering av GTT som målemetode for kartlegging av pasientskader

Formål med prosjektet: Å teste verktøyet GTT som brukes til kartlegge pasientskader. Metoden går ut på å screene pasientjournaler etter utvalgte triggere (lab verdi, fall, infeksjoner som kan oppstå i pasientforløpet) som kan indikere at en pasientskade har skjedd. Målet med studien er å finne den optimale utvalgsstørrelsen som trengs for å estimere antall skader, om utskifting av de som screener påvirker resultatet og om ett automatisk verktøy kan erstatte den manuelle granskningen


Forskningsprosjektet krever forhåndsgodkjenning av REK. Personvernombudets (PVO) rolle er å ha oversikt over forskningsprosjekter samt se til at informasjonssikkerheten og personvernet blir ivaretatt.

Det forutsettes at prosjektet gjennomføres i tråd med de opplysningene som er gitt i selve meldingen samt i øvrig korrespondanse og samtaler. Videre forutsettes det at bestemmelsene i lov om behandling av personopplysninger og lov om helseregistre og behandling av helseopplysninger med forskrifter følges. Prosjektet må videre gjennomføres i henhold til annet relevant regelverk, herunder de alminnelige regler om taushetsplikt.

- Dersom registeret skal brukes til annet formål enn det som er nevnt i meldingen må det meldes særskilt i hvert enkelt tilfelle.
- Dersom prosjektet har varighet på mer enn tre år skal prosjektansvarlig hvert tredje år sende bekreftelse til personvernombud på at behandlingen skjer i overensstemmelse med søknaden og vilkårene som er nevnt i denne godkjenningen.
- Det skal gis tilbakemelding til personvernombudet når registret er slettet.

Med hjemmel i personopplysningslovens forskrift § 7-12 godkjennes det at behandlingen av personopplysningene kan gjennomføres med de vilkårene som nevnt ovenfor.
Med hilsen
NORDLANDSSYKEHUSET HF

Alisa Larsen
Informasjonssikkerhetsrådgiver/Personvernombud

Vedlegg 1
Vedlegg – forskningsprosjekt

Helseforskningsloven
§ 10. Søknad om forhåndsgodkjenning
Søknad om forhåndsgodkjenning av et forskningsprosjekt skal sammen med forskningsprotokollen sendes til den regionale komiteen for medisinsk og helsefaglig forskningsetikk.

Den regionale komiteen for medisinsk og helsefaglig forskningsetikk skal foreta en alminnelig forskningsetisk vurdering av prosjektet, og vurdere om prosjektet oppfyller kravene stilt i denne loven eller i medhold av denne loven. Den regionale komiteen for medisinsk og helsefaglig forskningsetikk kan sette vilkår for godkjenning.

Vedtak vedrørende forhåndsgodkjenning kan påklages til Den nasjonale forskningsetiske komité for medisin og helsefag, jf. lov 30. juni 2006 nr. 56 om behandling av etikk og redelighet i forskning § 4.

Departementet kan gi forskrifter om krav til søknaden, om saksbehandlingsfrister for den regionale komiteen for medisinsk og helsefaglig forskningsetikk, og om de nærmere vilkårene for forhåndsgodkjenning

Forskrift om behandling av personopplysninger
§ 7-12. Personvernombud
Datatilsynet kan samtykke i at det gjøres unntak fra meldeplikt etter personopplysningsloven § 31 første ledd, dersom den behandlingsansvarlige utpeker et uavhengig personvernombud som har i oppgave å sikre at den behandlingsansvarlige følger personopplysningsloven med forskrift.
Personvernombudet skal også føre en oversikt over opplysningene som nevnt i personopplysningsloven § 32.
Forespørsel om deltakelse i forskningsprosjektet:

Validering av Global Trigger Tool som målemetode for kartlegging av pasientskader

Bakgrunn og hensikt


Hva innebærer studien?


Mulige fordeler og ulemper

For deg som pasient innebærer studien ingen ulemper eller direkte fordeler. Hvis du samtykker til denne undersøkelsen vil helsepersonell som deltar i dette forskningsprosjektet få innsyn i din pasientjournal. Finner vi at du har opplevd en alvorlig uønsket hendelse eller blitt skadet som følge av behandlingen, vil du bli kontaktet og informert om dette og du vil få tilbud om samtale med en av de som har gjennomgått journalen din. Vi vil ikke lete etter eventuelle nye lidelser/diagnoser, men kun vurdere om det forelå en pasientskade eller uønsket hendelse som følge av behandlingen du mottok i løpet av det aktuelle oppholdet.

Hva skjer med informasjonen om deg?

Informasjon som registreres om deg skal kun brukes slik som beskrevet ovenfor. Når alle data fra sykehusoppholdet er gjennomgått, blir eventuelle skader eller uønskede hendelser registrert og lagret atskilt fra journalen din uten ditt navn, fødselsnummer eller andre direkte eller indirekte identifiserbare opplysninger (anonymisert). Det vil heller ikke være mulig å identifisere de enkelte deltagerne i de publiserte resultatene av studien. Dersom vi senere ønsker å bruke de opplysningene vi har samlet inn til et annet forskningsprosjekt, vil du bli forespurt og videre bruk forutsetter at du samtykker også til det.

1.1 Personvern

Opplysninger som ønskes registrert om deg skal hentes fra Nordlandssykehuset elektroniske journalsystem. I vår studie skal dette ikke koples til andre lokale/nasjonale registre eller bli overlatt til andre forskere. Nordlandssykehuset ved administrerende direktør Paul Martin Strand er ansvarlig for håndtering og lagring av data.
1.2   **Rett til innsyn og sletting av opplysninger om deg**

Hvis du sier ja til å delta i studien, har du rett til å få innsyn i hvilke opplysninger som er registrert om deg. Du har videre rett til å få korrigert eventuelle feil i de opplysningene vi har registrert. Dersom du trekker deg fra studien, kan du kreve å få slettet alle innsamlede data, med mindre opplysningene allerede er benyttet i analyser eller i vitenskapelige publikasjoner. Studien er finansiert gjennom forskningsmidler fra Helse Nord og resultatene fra studien blir publisert i nasjonale og internasjonale fagtidsskrifter.

**Frivillig deltakelse**

Det er frivillig å delta i studien. Du kan når som helst og uten å oppgi noen grunn trekke ditt samtykke til deltagelse uten at dette vil noen konsekvenser for deg i din fremtidige kontakt med Nordlandssykehuset. Dersom du ønsker å delta, undertegner du samtykkeerklæringen på siste side snarest mulig og returnerer dette i vedlagt konvolutt. Dersom du senere ønsker å trekke deg eller har spørsmål til studien, kan du kontakte:

Tittel: LIS lege kir avdeling Kjersti Mevik, kjersti.mevik@nlsh.no

Telefon:75534000

Mvh

Kjersti Mevik

Barthold Vonen

Stipendiat phD/LIS lege kir avd

Prosjektleder/medisinsk direktør
Samtykke til deltakelse i studien Validering av Global Trigger Tool som målemetode for kartlegging av pasientskader

Jeg har lest prosjektbeskrivelsen ovenfor og gir samtykke til at journaldata fra min eller en annens person (i de tilfeller hvor samtykke gis av pårørende) brukes i studien:

(navn, dato, sted)

Bruk vedlagt ferdig frankert svarkonvolutt.
Paper I
Paper II
Paper III