Comparison of the prevalence and characteristics of inpatient adverse events using medical records review and incident reporting

W M Macharia, MB ChB, MMed (Paeds), MSc; C M Muteshi, MB ChB, MMed (O&G); S Z Wanyonyi, MB ChB, MMed (O&G); A M Mukaindo, MMed (O&G); A Ismail, MB ChB, MMed (Paeds); H Ekea, MB ChB; A Abdallah, MB ChB, MMed (Surg); J M Tole, MB ChB, MMed (Paeds); A K Ngugi, BSc, MSc, PhD

1 Department of Paediatrics, Faculty of Health Sciences, Aga Khan University, Nairobi, Kenya
2 Department of Obstetrics and Gynaecology, Faculty of Health Sciences, Aga Khan University, Nairobi, Kenya
3 Department of Medicine, Faculty of Health Sciences, Aga Khan University, Nairobi, Kenya
4 Department of Surgery, Faculty of Health Sciences, Aga Khan University, Nairobi, Kenya
5 Population Health Sciences, Faculty of Health Sciences, Aga Khan University, Nairobi, Kenya

Corresponding author: W M Macharia (william.macharia@aku.edu)

The incidence of medical adverse events (AEs) in hospitalised patients was estimated at 9.2% (interquartile range 4.6 - 12.4) by de Vries et al. in 2008. An event was defined as an incident resulting in death, a prolonged hospital stay or disability but not attributable to the underlying medical condition. The Canadian AEs study found a rate of 7.5% from review of 3 745 medical records in four hospitals. Public hospitals in a multicentre study conducted in Africa and the Middle East had a range of 2.5 - 18.4%. Other methods used to assess AEs include incident reports, interviewing of healthcare providers, direct observation, external audits, confidential inquiries and complaints. Use of a hybrid electronic medical record system has demonstrated some benefit in combining prospective and retrospective approaches to uncover surgical AEs. Laing et al. identified 71.4% errors prospectively and 28.6% retrospectively. They found that a tick-box system improved the quality of documentation, which could help solve the disadvantage inherent in retrospective reviews. Prospective approaches involve voluntary written documentation or reporting of incidents, with concerns of under-reporting when a culture of quality monitoring is not entrenched.

Objective
The practice of AE reporting has not gained ground in many institutions in low-income countries. This may be due to inadequate awareness of its importance in informing service quality improvement to minimise harm to patients. This study was carried out with the objective of comparing the prevalence and patterns of AEs in hospitalised patients, using review of medical records and incident reporting.

Methods
The study was carried out in a 254-bed tertiary hospital in Nairobi, Kenya. The hospital has ~20 000 admissions per annum, comprising medical, surgical, paediatric and obstetric patients. A retrospective review of randomly selected medical records of patients admitted in 2010 was undertaken to identify the presence or absence of AEs documented in the index year. All inpatient medical records and reported clinical incidents for the index year were eligible for inclusion. When more than one incident occurred in a patient in the index year, all were included as separate occurrences provided they were not related.

Results
The study identified 53 UEs from 2 000 randomly selected medical records and 33 reported UEs from 23 026 admissions in the index year. The prevalences of AEs from medical records review and incident reports were 1.4% (95% confidence interval (CI) 0.9 - 2.0) and 0.03% (95% CI 0.012 - 0.063), respectively. Compared with incident reporting, review of medical records identified more disability (13.2% v. 0%; p=0.03) and prolonged hospital stays (43.4% v. 18.2%; p=0.02).

Conclusions
Review of medical records is preferable to incident reporting in determining the prevalence of AEs in health facilities with limited inpatient quality improvement experience. Further research is needed to determine whether staff education and a positive culture change through promotion of non-punitive UE reporting or a combination of approaches would improve the comprehensiveness of AE reporting.

opted to increase the number of randomly sampled files to 2 000. All incidents recorded in the year were reviewed for fulfillment of predefined criteria for unintended or adverse clinical events. All the 1 665 incidents reported in the index year were included in the review.

Unexpected clinical outcomes were defined as unexpected clinical events (UEs) to distinguish them from AEs. Events had to have: (i) occurred at any time before the index admission and been detected during, or been responsible for, the index admission; (ii) occurred and been detected during the index admission; or (iii) occurred during the index admission but been detected on a subsequent admission. An AE was defined as an incident that resulted in death, disability or a prolonged hospital stay and was not explainable by an underlying medical condition. Other occurrences were defined as UEs.

Two experienced nurses used RF1 forms (Appendix 1) to screen for unexpected events using 18 criteria comprising conditions or circumstances commonly associated with AEs. Four physicians then reviewed selected files using RF2 forms (Appendix 2). A panel of content experts in the surgical and medical fields was identified to act as a resource for technical input, but not to determine the occurrence of incidents. All reviewers underwent 2-day training. Pilot testing was conducted using a convenience sample of 20 records to ensure comprehension of the study techniques.

The nurse reviewers ensured that the physicians were not assigned to review medical records of patients in whose care they could previously have participated.

The physician reviewers then scrutinised all the records that screened positive using RF2 forms for the presence of AEs. They then categorised incidents according to severity, location where the event occurred, attribution and preventability.

Incident reports were identified by accessing both computerised records and paper files to determine the number of events reported in the year, the nature of the incidents and the circumstances surrounding them. Initial screening was done to exclude non-clinical incidents. Information on remedial measures taken following root-cause analysis was abstracted and entered into data collection forms. A random sample of 10% of the medical records was subjected to a second review by a reviewer with longer experience in similar studies, and who was not involved in the earlier reviews, for the purpose of validation.

The primary outcome measure was the prevalence of AEs, while secondary outcomes included UEs, circumstances leading to the UEs, site of incident occurrence and preventability. The prevalence of AEs from medical records review was computed as a percentage using the total number of AEs as numerator and the total number of sampled admissions as denominator. Similarly, for incident reporting the numerator was the number of AEs and the denominator the total number of hospital admissions in the index year. A p-value of <0.05 was considered significant, and 95% confidence intervals (CIs) were determined around primary outcome estimates. The χ² test was used to compare independent categorical variables. Prevalence and bias-adjusted kappa statistics were used to estimate inter-rater agreement, with a value of >0.4 considered poor agreement, 0.4 - 0.6 moderate agreement and >0.6 good agreement.¹²

The study was approved by the World Health Organization (Ref: PS090004) and the Aga Khan University Hospital Ethics Review Committees (AKU/REC-06052011). Anonymous record keeping was used to delink case records from study data, and patient confidentiality was maintained.

Results

There were 23 026 hospital admissions in the year 2010, from which 2 000 records were randomly selected for review. Of these, 317 were screening criteria-positive, with 53 (2.7%; 95% CI 2.0 - 3.5) having documented occurrence of a UE, but only 28/2 000 cases were associated with disabilities, a prolonged hospital stay or disabilities not attributable to the primary medical conditions, giving a prevalence of 1.4% (95% CI 0.9 - 2.0). During the same study period, 233/1 665 (14.0%) of the reported incidents were of a clinical nature. From the total of 23 026 admissions for 2010, 33 (0.14%, 95% CI 0.10 - 0.20) had UEs but only 7 fulfilled the criteria for an AE, giving a prevalence of 0.03% (95% CI 0.01 - 0.06).

Forty-three out of 428 (10.0%) of the combined medical review and incident reports were reviewed by the ‘expert reviewer’ for validation. The inter-reviewer agreement for the reviewers was moderate at κ=0.45 (95% CI 0.38 - 0.74) after adjusting for prevalence and bias. Evidence of healthcare causation was identified in 31/53 cases (58.5%) from medical records and in 20/33 cases (60.6%) by incident reporting. Comparing review of medical records with incident reporting, disability (13.2% v. 0%) and prolonged hospital stay (43.4% v. 18.8%) were more likely to be detected from review of records (Table 1).

The majority of UEs took place in the study hospital, predominantly in the patient’s ward or room (Table 2).

The most frequently observed consequence of UEs from both data sources was prolonged hospital stay. Of the UEs detected by medical review, 11/15 (73.3%) occurred while the patient was hospitalised and 4/15 (26.7%) outside the facility.

Thirty-one of 53 medical records (58.5%) attributed UE causation to healthcare management. Another 77.4% (95% CI 52.0 - 87.8) of the events were non-procedure-related. Drugs were associated with UEs in 86.9% and 80.0% according to medical records and incident reports, respectively. Other health products that contributed to UEs, but less commonly, were blood products, medical devices and medical equipment. Clinical circumstances associated with UEs as determined by medical records and incident reports are shown in Table 3.

Table 1. Nature of UEs identified from the medical records and incident report reviews

<table>
<thead>
<tr>
<th>Description of AE</th>
<th>Medical record (N=53), n (%)</th>
<th>Incident report (N=33), n (%)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Events associated with death</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>0 (0.0)</td>
<td>1 (3.0)</td>
<td>0.38</td>
</tr>
<tr>
<td>No</td>
<td>53 (100.0)</td>
<td>32 (97.0)</td>
<td></td>
</tr>
<tr>
<td>Events associated with disability</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>7 (13.2)</td>
<td>0 (0.0)</td>
<td>0.03</td>
</tr>
<tr>
<td>No</td>
<td>46 (86.8)</td>
<td>33 (100.0)</td>
<td></td>
</tr>
<tr>
<td>Events associated with prolonged stay</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>23 (43.4)</td>
<td>6 (18.2)</td>
<td>0.02</td>
</tr>
<tr>
<td>No</td>
<td>30 (56.6)</td>
<td>27 (81.8)</td>
<td></td>
</tr>
<tr>
<td>Healthcare causation/preventability</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>31 (58.5)</td>
<td>20 (60.6)</td>
<td>0.85</td>
</tr>
<tr>
<td>No</td>
<td>22 (41.5)</td>
<td>13 (39.4)</td>
<td></td>
</tr>
</tbody>
</table>
According to medical records, the underlying health status of the patient alone could have contributed to the occurrence of UEs in 64.7% of patients (95% CI 46.5 - 80.3). Eight patients experienced UEs that were not associated with prolonged hospital stay, disability or death so were considered not to be serious, in keeping with the a priori definition. Incident reports identified 5 cases related to a therapeutic intervention, in 2 of which there was an error during administration, while the rest were considered unavoidable or due to undeterminable circumstances. Three cases were rather complex to manage, so it was not possible to attribute their cause to any specific care omission or commission.

Of 19 cases categorised by degree of urgency, 4 were of moderate urgency while 15 were of low or no urgency. Four of 6 events were related to inadequate communication or reporting by the healthcare team. Seventy-five percent of percent of cases (9/12) could have benefited to a great extent or moderately if appropriate management had been given. The risk that a UE or AE would have resulted from management provided was moderate to high in 57.9% of the patients with UEs (95% CI 33.5 - 79.7%). Reviewers considered that 81% of competent healthcare professionals would have managed the patients with UEs in a similar manner.

### Discussion

Our previous work in a multicentre study involving review of medical records in public hospitals in Africa and the Middle East found a prevalence of AEs of 2.5 - 18.4%. The mean in the two hospitals in Kenya was 14.5%, which is much higher than the 1.4% in the medical records review in this study. A similar methodology was used for the medical records review, and some reviewers who had participated in the previous study were involved. This difference may be attributed to the higher standards of care in the current study site, which is a tertiary not-for-profit academic medical centre that at the time of the study was in the preparatory stages of Joint Commission International accreditation, which was eventually granted in 2013. Whereas AEs that are not associated with serious complications could fail to be documented, leading to underestimation, we only sought to document the severe ones that would readily be picked up from records, since death, disabilities and prolonged hospital stay would be evident.

Review of medical records in this study identified many more AEs than incident reporting (1.4% v. 0.03%), clearly indicating that many important events are never reported. This very low reporting may be a result of fear of being held responsible for omissions or commissions. It could also be due to inadequate staff education on the importance of reporting.

The majority of AEs in this study occurred in the rooms in which patients were receiving medical care. There can be major variations in the quality of care provided to hospitalised patients at different levels of care, especially for trauma patients.[7] Medical records review was better than incident reporting at identifying events that resulted in disability (13.2% v. 0%; \( p = 0.03 \)) and prolonged hospitalisation (43.4% v. 18.2%; \( p = 0.02 \)). This is not really surprising considering that these factors were triggers for AE scrutiny in medical records reviews, while incident reporting is expected to be spontaneous. An important drawback of the medical records approach is inability to fully reveal circumstances surrounding the event. Complementing review of records with structured morbidity and mortality meetings may be valuable.

Clarke et al.[8] demonstrated the usefulness of such audit meetings in dissecting out human error contributions. Incident reporting requires a change in institutional culture so that it is not punitive to those who disclose UEs. Where such a culture is not entrenched,
non-reporting would be even higher for patients on treatment for chronic and complex conditions, as true AEs in such patients may erroneously be assumed to be inevitable. There is also a tendency for health professionals to interpret AEs as expected complications of medical procedures even when they are not, and therefore failing to report them. This problem can, however, be overcome through comprehensive medical record review.13,14

Christians-Dingelhoff et al.13 found that only 3.6% of AEs identified by record review were identified by other reporting methods. As in our study, others also found different methods to be complementary with little overlap in reporting of AEs.13,14 This is especially important because some AEs not captured by incident reports would not have been detected using alternative methods, thus concealing opportunities for service improvement. Complementing incident reports with medical records review and other reporting modalities would ensure a more comprehensive assessment of AEs. Clarke et al.15 demonstrated how modern error theory used in commercial aviation could be exploited in health to identify missed injuries in trauma. However, we did not find any literature on how this could be applied for missed AEs in facility incidence or prevalence estimation.

By its nature, incident reporting included a large number of non-clinical reports that tended to mask non-reporting of important clinical incidents. A good reporting system should be able to clearly separate clinical and non-clinical incidents, as users are different. Our review was made possible by robust medical record keeping and an electronic incident reporting system. Lack of appropriate systems for data retrieval and analysis would be a major constraint for resource-deprived facilities. Nurse and physician reviewers with ample experience and training, as we had in our study, help to minimise observer variability.

Management flaws were responsible for nearly 60% of the AEs, with management itself likely to have contributed to AEs in 58% of cases. Irrespective of the data collection method used, >80% of AEs related to medications. Our reviewers estimated that in 75% of AEs cases. Irrespective of the data collection method used, >80% of AEs related to medications. Our reviewers estimated that in 75% of AEs the patient could have benefited from more appropriate care than had been provided.

Level of agreement among physicians completing RF2 forms was estimated using the prevalence-adjusted kappa, as described by Bennet et al. in 1954 and elaborated by Nam15 to give a more reliable degree of agreement. This reflected an acceptable chance-corrected agreement of 0.45, representing moderately good agreement.15

We observed that review of medical records also exposed more disabilities and prolonged hospital stay associated with the events. Laing et al.,16 using a hybrid electronic method, found almost three times more events prospectively than by scrutiny of records. Although the settings may not be comparable, given that the studies were carried out in different continents serving different patient populations, the difference in findings suggests major under-reporting of incidents in our study. Institutional cultural change through staff education on the purpose of reporting and assurance that reporting will not result in punishment may be needed to alter perceptions and practice. There is little overlap in the events, emphasising the need for combining different approaches to be comprehensive. The search for innovative approaches to identify the many factors that impact on quality of care to hospitalised patients continues, even as known combined approaches demonstrate promise.

**Conclusions**

Review of medical records is preferable to incident reporting in health facilities with limited inpatient quality improvement experience. The approach identifies more AEs and exposes more factors associated with events. Further research is needed to determine whether staff education and positive culture change through promotion of non-punitive reporting, or a combination of approaches, would improve comprehensiveness of AEs reporting.

**Authors’ contributions.** WMM was involved in conceptualisation, proposal writing, analysis and drafting of the final manuscript. CMM participated in study implementation, analysis and drafting of the final manuscript. SZW was involved in conceptualisation, proposal writing and drafting of the final manuscript. AMM, AI, HE, AA and JMT all participated in data collection and drafting of this manuscript. AKN participated in data analysis and drafting of the manuscript. All the authors reviewed the manuscript and approved submission.

**Acknowledgements.** This work was supported by the World Health Organization Small Grants Program (WHO Ref: 2011/167789-0). We wish to acknowledge the enormous amount of guidance we received from Nittita Prasopa-Plaizer (WHO, Geneva) during the proposal writing and manuscript preparation stages. Georgeh Banica (WHO, Geneva) similarly provided us with requisite administrative support for completion of this work. Janet Musia made invaluable input in data and project management.

---


Accepted 7 February 2016.
Appendix 1. RF1 form: Adverse event detection questionnaire

CONFIDENTIAL

RF1: adverse event detection questionnaire

REVIEWER

Reviewer ID Number: 

Clinical department n° 

Date of data collection: D D M M Y Y

Time interview commenced: (use 24 hour clock)

Time interview finished: 

PATIENT NAME: 

(Surname) (Given Names)

CASE Number 

Birth date (at least the year of birth) D D M M Y Y Y Y

Gender (1 male / 2 female) 

Admission Status (1 elective / 2 acute / 3 do not know) 

Date of Admission: D D M M Y Y

Date of Discharge (if known): D D M M Y Y

<table>
<thead>
<tr>
<th>SCREENING CRITERIA</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Unplanned admission in the 12 months prior to the index admission as a result of any health care management.</td>
</tr>
<tr>
<td>2. Hospital-incurred patient accident or injury.</td>
</tr>
<tr>
<td>3. Adverse drug reaction / drug error</td>
</tr>
<tr>
<td>4. Hospital acquired infection/sepsis.</td>
</tr>
<tr>
<td>5. Unplanned removal, injury or repair of organ or structure during surgery, invasive procedure or vaginal delivery.</td>
</tr>
<tr>
<td>6A. Unplanned return to the operating theatre during this admission</td>
</tr>
<tr>
<td>6B. Unplanned visit to the operating theatre during this admission</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td>7</td>
</tr>
<tr>
<td>8</td>
</tr>
<tr>
<td>9</td>
</tr>
<tr>
<td>10</td>
</tr>
<tr>
<td>11</td>
</tr>
<tr>
<td>12</td>
</tr>
</tbody>
</table>
13. Unplanned transfer from general care to intensive care/higher dependency.  

14. Unplanned transfer to another acute care hospital  

15. Unexpected death (i.e. not an expected outcome of the disease during hospitalisation)  

16. Any other undesirable outcomes (not covered by any of the other criteria).  

Are any Criteria present?  

If Yes, total number of criteria  

Appendix 2. RF2 form: Second questionnaire

Second questionnaire (RF2)

Take account of all adverse events identified in RF1, not only the most serious. Complete one form for each adverse event (AE).
If the patient has more than one AE, use two RF2 questionnaires and quote “1 out of 2” and “2 out of 2.”

Case number (same number as in RF1).
Reviewer ID Number:
Date of data collection: D D M M Y Y
Time when interview commenced: H H M M (use 24 hour clock)
Time interview finished:

Q1 & Q2  AE No □ out of a total of □ AE

Information sources used
Q3 Physician □ I = Yes 2 = No
Q4 Head Nurse □ I = Yes 2 = No
Q5 Nurse □ I = Yes 2 = No
Q6 Medical record □ I = Yes 2 = No
Q7 Other source □ I = Yes 2 = No
Q8 Did the patient experience an adverse event (injuries or complications)? [ ] 1 = Yes 2 = No

Q9 When did this event occur? [ ] [ ] [ ] [ ] [ ] [ ]

Q10 Clinical summary of the case and description of the adverse event

Main disease

Known comorbidities

History of disease (in particular specify if the disease was known before admission)

Cause for hospital admission

Main events during hospitalisation

Adverse event: (for example, answer briefly the following “what, who, when, where, how” questions) (Give any relevant laboratory/imaging results) continue on back if needed
ADVERSE EVENT DETERMINATION

SEVERITY (answer : 1 = Yes 2 = No)
Q11 Did the injury or complication caused the hospitalisation? ☐
Q12 Was the injury or complication associated with death of the patient ☐
Q13 Was the injury or complication associated with disability/deficit at the time of discharge? ☐
Q14 Was the injury or complication associated with prolonged hospital stay? ☐
(including readmission)

CAUSATION
Q15 In your best judgement, is there evidence that healthcare management caused the adverse event?
In answering this question, consider, when relevant, the following questions and complete the appropriate boxes.
Q151 Could the event be expected, giving the disease or the health status of the patient? 1 = Yes 2 = No 3 = Don’t know ☐
Q152 Are there indications that health care management caused the injury? 1 = Yes 2 = No 3 = Don’t know ☐
Q153 Does the timing of events suggest that the injury was related to the treatment or lack of treatment? 1 = Yes 2 = No 3 = Don’t know ☐
Q154 Are there other reasonable explanations for the event? 1 = Yes 2 = No 3 = Don’t know ☐
Q155 Is there general recognition that the intervention or lack/delay of intervention or diagnosis (depending in the instance) causes this kind of adverse event? 1 = Widely recognised by scientific community 2 = Recognised by some specialists only 3 = No recognition 4 = Don’t know ☐
Q156 Was there an opportunity prior to the occurrence of the injury for intervention which might have prevented it? 1 = Yes 2 = No 3 = Don’t know ☐
Q157 Was the AE recognised during the hospital stay? 1 = Yes 2 = No ☐
Q157a Was appropriate action taken? 1 = Yes 2 = No 3 = not applicable ☐
Q157b Did the AE respond to the appropriate action? 1 = Yes 2 = probably 3 = too early to know 4 No 5 Don’t know 6 Not applicable ☐

Consider all of the above questions above before continuing

Q16 After due consideration of the clinical details of the patient's management, irrespective of preventability what level of confidence do you have that the HEALTH CARE MANAGEMENT caused the injury?

Confidence Score:
1 = Virtually no evidence for management causation (Then STOP, no AE)
2 = Slight to modest evidence for management causation (Then STOP, no AE)
3 = Management causation not likely; less than 50-50 (Then STOP, no AE)
4 = Management causation more likely than not, more than 50-50
5 = Moderate/strong evidence for management causation
6 = Virtually certain evidence for management causation

Score ☐

The questionnaire is complete if your score is three or less

Q17 Location of occurrence

Q171 Where did the healthcare management causing the AE occur? (choose one)
1 outside this hospital   2 inside this hospital

Q172 If outside this hospital
01 = Public hospital
02 = Private hospital
03 = Home with professional healthcare management
04 = Home without professional healthcare management
05 = Nursing home
06 = GP office
07 = other

Q173 If inside hospital, in the clinical unit in which the patient was hospitalised??
Yes 2 = No
1 =

Q174 If not in patient’s clinical unit, in which one?

SURGERY
01 = Cardiac Surgery
02 = Colon/Rectal Surgery
03 = General Surgery
04 = Orthopaedic Surgery
05 = Paediatric Surgery
06 = Plastic Surgery
07 = Thoracic Surgery
08 = Urological Surgery
09 = Vascular Surgery
10 = Neurosurgery
11 = Obstetrics
12 = Ophthalmology
13 = ENT
14 = Stomatology
15 = Cardiology
16 = Dermatology
17 = Endocrinology
18 = Gastroenterology
19 = Geriatrics
20 = Gynaecology
21 = Haematology
22 = Immunology and Allergy
23 = Infectious Disease
24 = Internal Medicine
25 = Physical Medicine
26 = Neonatology
27 = Nephrology
28 = Neurology
29 = Medical Oncology
30 = Paediatrics
31 = Pulmonary Disease
32 = Psychiatry
33 = Medical Intensive Care Unit
34 = Rheumatology
35 = A&E
36 = Other

Q18 If inside hospital, where exactly?

INSIDE HOSPITAL
01 = Theatres
02 = Recovery Room
03 = ICU
04 = Catheterisation, endoscopic unit
05 = Consultation, out-patients clinic
06 = Therapy/Rehabilitation
07 = Patient’s room
08 = Labour and Delivery
09 = Radiology
10 = A&E
11 = Service Area (stairs, halls, elevator)
12 = Other site in hospital
13 = Don’t know
### Q19 CLASSIFICATION OF ADVERSE EVENT

**Q191** To which type of care management was the adverse event mainly related?  
1 prevention 2 diagnosis 3 therapeutic 4 rehabilitation

**Q192** What was the main cause of AE (the most important one)?  
1 = Error in the choice of management  
2 = Delay for its implementation  
3 = Error during its implementation.  
4 = Other (mainly unavoidable events)  
5 = Don’t know

**Q193** Was the AE related to a procedure?  
1 = Yes 2 = No

**Q193a** If yes, which procedure?  
1 = Surgery  
2 = Anesthesiology  
3 = Surgical intervention during radiology.  
4 = Radiology using contrast product  
5 = Endoscopy  
6 = Biopsy  
7 = Puncture or tapping  
8 = Catheter, perfusion or injection  
9 = Urinary catheter  
10 = Gastric  
11 = Intubation  
12 = Dialysis  
13 = Radiotherapy  
14 = Instrument assisted delivery  
15 = Physiotherapy  
16 = Other

**Q194** Was the AE related to a substance or health product?  
1 = Yes 2 = No

**Q194a** If yes, which product?  
1 = drug  
2 = blood product  
3 = medical device.  
4 = equipment (laser, electric bistoury..)  
5 = Dietetic product  
6 = Local preparation (e.g. chemotherapy product…)  
7 = Other

### Q20 Patient-related contributory factors?

**Q201** Patient’s global health status and disease  
1 = Yes 2 = No

**Q202** Patient’s behaviour  
1 = Yes 2 = No

**Q203** Family’s behaviour  
1 = Yes 2 = No

**Q204** Other  
Specify: ........................................
**Q21 System-related contributory factors?** 1 = Yes 2 = No

(For all 16 next items, answers are 1 = Yes 2 = No)

Q211 Inadequate or defective premises
Q212 equipment or supplies not available or defective
Q213 inadequate staffing at the time of the AE (not merely in terms of numbers, take account of balance among different competences and experience, in particular at weekend and during holidays)
Q214 recent organizational changes inside the unit
Q215 defective coordination inside the unit
Q216 inadequate reporting or communication
Q217 inadequate training or supervision of doctors or other personnel
Q218 delay in the provision or scheduling of services (e.g. lab tests, x-rays or follow-up visits)
Q219 failure to implement protocol or plan
Q220 inadequate monitoring of patient
Q221 inadequate discharge procedure
Q222 defective coordination between the unit and other units (e.g. pharmacy, blood bank or catering)
Q223 No protocol/healthcare policy
Q224 Other

Describe the most important contributing factor to the adverse event

**Q23 inappropriate adaptation to an unexpected event**

Describe inappropriate adaptation
### Q24 PREVENTABILITY

Consider and evaluate the following questions before making a judgement on preventability.

<table>
<thead>
<tr>
<th>Q241</th>
<th>How serious was this case PREVIOUS to the occurrence of an AE?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Very serious</td>
</tr>
<tr>
<td>2</td>
<td>Moderately serious</td>
</tr>
<tr>
<td>3</td>
<td>not very serious</td>
</tr>
<tr>
<td>4</td>
<td>Not serious</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q242</th>
<th>How complex was this case? (co-morbidity, global health status)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Very complex</td>
</tr>
<tr>
<td>2</td>
<td>Moderately complex</td>
</tr>
<tr>
<td>3</td>
<td>not very complex</td>
</tr>
<tr>
<td>4</td>
<td>Uncomplicated</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q243</th>
<th>What was the degree of emergency in management of the case prior to the occurrence of adverse event?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Critical and very urgent</td>
</tr>
<tr>
<td>2</td>
<td>Moderate</td>
</tr>
<tr>
<td>3</td>
<td>low</td>
</tr>
<tr>
<td>4</td>
<td>Not urgent</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q244</th>
<th>Was the management of the illness appropriate?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Large consensus</td>
</tr>
<tr>
<td>2</td>
<td>Consensus moderate</td>
</tr>
<tr>
<td>3</td>
<td>No consensus</td>
</tr>
<tr>
<td>4</td>
<td>Management non-indicated or contra-indicated</td>
</tr>
<tr>
<td>5</td>
<td>Don’t know</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q245</th>
<th>What was the degree of deviation of management from recommendations?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>None</td>
</tr>
<tr>
<td>2</td>
<td>Slight</td>
</tr>
<tr>
<td>3</td>
<td>Moderate</td>
</tr>
<tr>
<td>4</td>
<td>Marked</td>
</tr>
<tr>
<td>5</td>
<td>Don’t know</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q246</th>
<th>What was the chance of benefit associated with the management of the illness which led to the AE?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>High</td>
</tr>
<tr>
<td>2</td>
<td>Moderate</td>
</tr>
<tr>
<td>3</td>
<td>Low</td>
</tr>
<tr>
<td>4</td>
<td>absent</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q247</th>
<th>What was the risk of an adverse event related to the management?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>virtually absent</td>
</tr>
<tr>
<td>2</td>
<td>low</td>
</tr>
<tr>
<td>3</td>
<td>Moderate</td>
</tr>
<tr>
<td>4</td>
<td>High</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Q248</th>
<th>On reflection, would a reasonable doctor or health professional have managed the care in a similar manner?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Definitely would</td>
</tr>
<tr>
<td>2</td>
<td>Probably would</td>
</tr>
<tr>
<td>3</td>
<td>Probably would not</td>
</tr>
<tr>
<td>4</td>
<td>Definitely would not</td>
</tr>
</tbody>
</table>
Consider all the questions 241-248 above before continuing

25 Rate on a 6 point scale your confidence in the evidence for preventability.

   Confidence score:
   1 = Virtually no evidence for preventability
   2 = Slight to modest evidence for preventability
   3 = Preventability not really likely; less than 50-50
   4 = Preventability more likely than not; more than 50-50
   5 = Strong evidence for preventability
   6 = Virtually certain evidence for preventability

   Score

26 Please describe in which way the adverse event may have been prevented