Impact of standardization of in-ward urine output measurement in post-operative patients: a clinical audit

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Keywords: Postoperative care; urine outputa audit

Abstract
Urine output (UOP) is a key parameter used in post-operative patients to monitor the recovery. In our setup, this is done by patients or caregivers. An audit cycle was carried out to develop and evaluate a standardized protocol to improve the accuracy of UOP measurement. The pre-audit assessment was done evaluating: individual's ability in measuring and record keeping, characteristics of the measuring container and error of measurement. Subsequently, corrections were done in these areas. The cycle was completed comparing the error of measurement as an outcome. The intervention group had fewer errors compared to the pre-audit group. Errors can be overcome by a standardized measuring protocol.

Introduction
Variation of urine output is a valuable surrogate marker of impending acute kidney injury (AKI) and tissue perfusion (1). Accurate measurement of urine output is also important in maintaining proper fluid balance in critically ill patients (2).

In our setup, most of the patients after major surgery are managed in general wards. In our ward setup, the urine output (UOP) is measured usually by caretakers. No standardized methods are followed in the process of measurement. The usual practice is to visually read the recording in the urine-collecting bag and record it. Such readings are usually inaccurate. Errors can happen during urine measuring container manipulation, visual assessment and manual data recording. Errors that happen during measuring, monitoring and recording UOP can end up in delays of identification and early intervention of episodes of oliguria (3) hence clinical diagnosis and decision making.

We observed these drawbacks in our practice as well. An audit cycle was carried out focusing on standardizing the measurement of urine output of inward post-surgical patients.

Methodology

Study sample
This clinical audit was carried out for 03 months. Twenty-five post-operative patients were included in the pre-audit assessment and another 25 post-operative patients were included in the re-audit cycle. Based on the first cycle results, intervention protocol was prepared to focus on standardizing the process. Accuracy of the urine output measurements was prospectively audited.

Inclusion and exclusion criteria
The primary inclusion criterion was patients undergoing major surgeries needing postoperative measurement of urine output. Patients younger than 18 years, the patient who can't read numbers, patients with visual impairments and patients who didn't give the consent to participate were excluded.

First audit cycle
In 25 (n=25) post-operative patients (group A) as the first step, using an interviewer-administered questionnaire data was collected on, individuals (patient’s or bystander’s) knowledge on volume reading from the measuring cup (figure 1), writing numbers, record keeping of volume. An interviewer observed the device used for measurements (whether it is a calibrated measuring container or not). Further interviewer assessed the volume reading ability and knowledge on numbers by giving pre-measured water sample and asking them to measure the volume and record it.

To measure the accuracy of measurement, patient or bystander was asked to measure the UOP for next 24 hours duration. Urine was collected to a separate large container given by the interviewer. After 24 hours, interviewer assessed the true volume of urine. As the outcome, the difference between true volume and patient’s measurement was recorded. The difference over 50ml was considered significant.

Intervention protocol
Based on the findings three interventions were done. An information sheet was given to the patient/caregiver as a step by step guide on the measuring process. Separate volume
The intervention we made was a simple change that can make a significant difference in accurate monitoring of post-operative patients. Adapting this strategy in our clinical setup in different units needs to be considered.

All authors disclose no conflict of interest. The study was conducted in accordance with the ethical standards of the relevant institutional or national ethics committee and the Helsinki Declaration of 1975, as revised in 2000.

References