Limitations of Gravimetric Quantitative Blood Loss during Cesarean Delivery

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Abstract

Objective This study examined the accuracy, sources of error, and limitations of gravimetric quantification of blood loss (QBL) during cesarean delivery.

Study Design Blood loss determined by assays of the hemoglobin content on surgical sponges and in suction canisters was compared with QBL in 50 parturients.

Results QBL was moderately correlated to the actual blood loss ($r = 0.564; p < 0.001$). Compared with the reference assay, QBL overestimated blood loss for 44 patients (88%). QBL deviated from the assayed blood loss by more than 250 mL in 34 patients (68%) and by more than 500 mL in 16 cases (32%). Assayed blood loss was more than 1,000 mL in four patients. For three of these patients, QBL was more than 1,000 mL (sensitivity = 75%). QBL was more than 1,000 mL in 12 patients. While three of these had an assayed blood loss of more than 1,000 mL, 9 of the 46 patients with blood losses of less than 1,000 mL by the assay (20%) were incorrectly identified as having postpartum hemorrhage by QBL (false positives). The specificity of quantitative QBL for detection of blood loss more than or equal to 1,000 mL was 80.4%.

Conclusion QBL was only moderately correlated with the reference assay. While overestimation was more common than underestimation, both occurred. Moreover, QBL was particularly inaccurate when substantial bleeding occurred.

Keywords ► cesarean delivery ► blood loss quantification ► postpartum hemorrhage ► quality improvement

Key Points
- QBL is inaccurate in cesarean delivery.
- QBL deviated from the assay result by more than 500 mL in 32% of cases.
- QBL sensitivity and specificity for hemorrhage is 75.0% (95% confidence interval [CI]: 0.19–0.93) and 80.4% (95% CI: 0.69–0.92), respectively.

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Quantification of blood loss (QBL) is recommended over visual estimation of blood loss for women having cesarean delivery.\(^1\)\(^,\)\(^2\) QBL is thought to enhance the accuracy of blood loss measurements, improve hemorrhage recognition, and allow for appropriate activation of treatment stage-based protocols which may consequently improve patient outcomes.\(^3\)

A common way of measuring QBL is the gravimetric method that involves weighing of soaked sponges and measurement of fluid in calibrated suction canisters.\(^4\) After adjusting for the dry weight of soaked materials, the estimated amount of amniotic fluid and the amount of surgical irrigation are deducted from the total weight, presumably resulting in an accurate measure of blood loss. While intuitively sensible, technical problems obtaining accurate measurements, difficulty in estimating amniotic fluid, and retention of irrigation fluid in the abdomen can lead to inaccurate results.\(^5\)\(^,\)\(^6\) Moreover, some blood loss may not be captured on surgical sponges and in suction canisters leading to additional error.

In a previous publication that compared visually estimated blood loss, gravimetric QBL, and a colorimetric method of estimating blood loss to a reference hemoglobin assay measurement of the amount of blood on surgical sponges and in suction canisters during scheduled cesarean delivery, we found that gravimetric QBL measurements were poorly correlated to the more accurate reference hemoglobin assay results.\(^7\) The aim of this secondary analysis was to determine the sources of error and limitations of the gravimetric QBL method in greater detail, using both correlations and Bland–Altman measures of agreement.

**Materials and Methods**

The original protocol was approved by the Santa Clara Valley Medical Center Institutional Review Board (San Jose, CA; reference no.: 12–003; August 12, 2013). The details of the study methodology have been previously described.\(^7\) Briefly, all sponges and calibrated surgical canisters from 50 patients having scheduled cesarean deliveries were evaluated, and relevant patient and procedural information was collected. The original study compared visually estimated blood loss, gravimetric QBL, and a colorimetric blood loss estimation to a reference hemoglobin extraction assay during scheduled cesarean delivery. This secondary analysis is focused on the relationship between gravimetric QBL and the reference hemoglobin assay.

Quantitative gravimetric measurement methods were adopted from published guidelines.\(^4\) At the time of the uterine incision, the surgical technician or circulating nurse recorded the canister volume using the graduated markings. After aspiration of amniotic fluid, a second measurement was made, and the difference was recorded as the estimated amniotic fluid volume. At the conclusion of the surgery, the technician recorded the total amount of any irrigation fluid used. Additionally, any blood loss expressed by uterine massage was collected and transferred to one or more sponges. Immediately following the case, all sponges and suction canisters were individually weighed using a calibrated digital scale, and dry sponge weights and canister weights were subtracted. To determine the total QBL estimate, all individual sponge and canister measurements were tallied, and the measured amount of amniotic fluid and irrigation fluid used was subtracted from the total weight.

For the reference hemoglobin extraction assay method, all soaked laparotomy sponges and calibrated suction canisters were transferred to an on-site benchtop facility for hemoglobin measurement at the completion of each procedure.\(^7\) Sponges were individually soaked in 400 mL of normal saline, compressed by hand for 60 seconds to a mean weight of 50 g. This process was repeated four times and the hemoglobin concentration of the final extraction fluid was measured using the plasma/low spectrophotometer (Hemo-Cue AB, Ängelholm, Sweden). The yield of the extraction process was independently characterized by depositing banked blood on sponges in known quantities and performing the same mechanical extraction. A linear regression analysis revealed mean hemoglobin recovery rates of 89.5% (95% confidence interval [CI]: 86.8–92.1%) for individual sponges (\(n = 116\)) and this value along with the patient’s preoperative hemoglobin was used to convert the extracted amount of hemoglobin (\(g\)) to the amount of blood in the sponges (\(mL\)).

The hemoglobin concentration in the canisters was separately assayed by using either a whole blood or low-concentration hemoglobin analyzer and converted to a canister blood volume based on the patient’s preoperative hemoglobin concentration (in \(g/dL\)).\(^7\) The blood loss in the canisters was then combined with the blood loss from the sponges to give a total assayed blood loss. The blood and fluid that remained on the surgical drapes, gowns, towels, and other materials was not accounted for by either the quantitative gravimetric assessment or the hemoglobin assay.

**Statistical Analysis**

The sample size was based on the previous study. The distributions of the blood loss measurements and other analysis quantities were assessed using descriptive univariate statistics: the mean and standard deviation (±), as well as the median and the first and third quartiles. Since the primary outcome of this paper is the accuracy of the gravimetric method compared with the hemoglobin assay method (i.e., considered to be the accurate reference value), the absolute difference between the hemoglobin assay and gravimetric methods was assessed instead of the signed difference between the measures.

Estimates of sensitivity and specificity used a binomial model. Since only four cases were observed with assayed blood loss greater than 1,000 mL, CIs for sensitivity were computed using the exact small-sample formula. CIs for specificity used the standard large sample formula for the binomial distribution.

The strength of the relationship between the gravimetric and assay methods was assessed by correlation coefficients; Fisher’s \(Z\)-transformation was used to compute CIs for correlation coefficients. A quantitative relationship between the gravimetric and assay methods was derived using the
nonparametric regression method of Passing and Bablok. The Bland–Bablock regression line provides a nonparametric estimate of the linear relationship between two measures that assumes both are subject to error, without making further assumptions about the statistical distributions of the measures. The strength of correlations was assessed as follows: $r < 0.5$ (low); $r = 0.5$ to $0.7$ (moderate), $r = 0.7$ to $0.9$ (high); and $r > 0.9$ (very high). For comparative analysis, a $p$-value of 0.05 was considered significant.

In addition, agreement between the extraction assay and gravimetric QBL was evaluated using the Bland–Altman method. The Bland–Altman bias (mean the difference between the two measures) and upper and lower limits of agreement (mean $\pm 1.96 \times SD$) with the respective 95% CIs was computed. All statistical computations were computed using SAS, Version 9.4 (SAS Institute, Cary, NC).

**Results**

Data were successfully collected from all 50 patients. The mean preoperative hemoglobin level was $12.2 \pm 1.0$ g/dL. Additional patient characteristics were described in the previous publication.

The gravimetric QBL method was moderately correlated to the actual (hemoglobin assayed) blood loss ($r = 0.564$; mean absolute difference $= 422 \pm 330$ mL; median absolute difference $= 365; [204–538]$ mL; $p < 0.001$). The 95% CI for the correlation coefficient was $[0.334, 0.725]$. As previously reported, the assessment of agreement between gravimetric QBL and the extraction assay revealed a bias (95% CI) of $353 \pm 405$ mL. The upper limit of agreement (95% CI) was $1,145$ mL and the lower limit of agreement (95% CI) was $-441$ mL. The Bland–Altman plot is represented in Fig. 1.

Compared with the hemoglobin assay result, gravimetric QBL underestimated blood loss for 44 of the 50 patients (88%). In 32 patients (64%) the underestimate was greater than 250 mL and for 14 patients (28%), the underestimate was greater than 500 mL. In two instances, there was an underestimation of more than 500 mL when compared with the assay (528 and 784 mL, respectively). Collectively, gravimetric QBL was only within 250 mL of the actual amount in 16 (32%) patients and all but three of these patients had an assayed blood loss of less than 500 mL (mean $= 378 \pm 179$ mL). The Bland–Bablock regression line (Fig. 2) demonstrates that gravimetric QBL is typically about twice the blood loss determined by the hemoglobin assay method.

Assayed blood loss was greater than 1,000 mL in four patients. For three of the four patients gravimetric QBL recognized that the blood loss was greater than 1,000 mL (sensitivity = 75%; 95% CI: 0.194–0.932) but underestimated it by 617, 533 and 255 mL, respectively. For the fourth patient the assayed blood loss was 1,041 mL but the gravimetric QBL was only 513 mL. Seventeen patients (34%) had an assayed blood loss of greater than 500 mL. For those 17 patients, the mean absolute difference between the QBL measurement and the assay was $549 \pm 420$ mL (median $= 528; [275–696]$ mL). Gravimetric QBL overestimated the assayed blood loss in 13 of the 17 patients (mean overestimation $= 597 \pm 445$ mL) and underestimated it in 4 (mean underestimation $= 394 \pm 326$ mL). Only 8 of the 17 (47%) values were within 500 mL and quantitative QBL was within 250 mL of the assayed result in only 3 (18%) cases.

Gravimetric QBL was greater than 1,000 mL (the American College of Obstetricians and Gynecologists [ACOG] criteria for hemorrhage recognition) in 12 patients. While three of these had an assayed blood loss of greater than 1,000 mL, 9 of the 46 patients (20%) with blood losses of less than 1,000 mL by the hemoglobin assay were incorrectly identified as having postpartum hemorrhage (false positives). The specificity of gravimetric QBL was 80.4% (95% CI: 0.690–0.919). In 37 cases (74%), there was concordance among both methods that the blood loss was less than 1,000 mL. These relationships are illustrated in Table 1. Interestingly, if the criteria for hemorrhage
recognition was 900 mL rather than 1,000 mL, gravimetric QBL would have recognized hemorrhage in 16 patients, only 3 of whom had an assay of greater than 900 mL. Thus, 13 of the 46 patients (28%) with blood losses of less than 900 mL would have been incorrectly identified as having postpartum hemorrhage. Conversely, if the criteria for hemorrhage recognition were increased to 1,100 mL, the false positive rate would be 8 of 47 patients (17%).

Corrections for amniotic fluid and irrigation were collectively applied to the sponge and calibrated canister quantities. The measured amount of amniotic fluid varied from 100 to 2,100 mL (mean = 632 ± 507 mL; median = 500 mL; [250–850]). The measured amount of irrigation varied from 50 to 2500 mL (mean = 760 ± 437 mL; median = 775 mL; [400–1,000]).

An average of 15 surgical sponges were used per case (mean = 15.1 ± 4.9; median = 14.5; [12–17]). The gravimetric QBL on the sponges was substantially higher than the hemoglobin assayed amount of blood on the sponges in every case (mean absolute difference = 480 ± 182 mL). The number of sponges used was highly correlated with the total assayed blood loss on the sponges (r = 0.748; 95% CI: [0.588, 0.847]; p < 0.0001; i.e., the use of more sponges was associated with a higher assayed blood loss).

A single-calibrated suction canister was utilized in 49 cases; the remaining case required use of two canisters. The assayed amount of blood in the canisters was substantially less than the total volume of fluid collected in the canisters (mean absolute difference = 1,264 ± 627 mL). The correlation between the total amount of fluid in the canisters and the actual amount of blood in the canister was low to moderate (r² = 0.466; 95% CI: [0.211, 0.656]). Irrigation and amniotic fluids affected both the sponge and canister values. Assuming that all of the amniotic fluid was aspirated into the canisters, 88% of the irrigation fluid went into the canisters and only 12% was absorbed by the sponges (► Fig. 3).

**Discussion**

Our study found that despite the use of trained technicians to carefully perform gravimetric QBL using recommended guidelines, the results deviated substantially from the actual (hemoglobin assayed) amounts of blood on the soiled sponges and in the surgical canisters by greater than 250 mL in 68% of cases and by greater than 500 mL in 32% of cases. While overestimation was more common than underestimation, both occurred. Detection of hemorrhage (blood loss greater than 1,000 mL)⁹ was aligned in 40 patients (80%); however, in 10 cases (20%), the results were not aligned. If quantitative QBL were used to implement a hemorrhage protocol,⁴ the protocol would have been inappropriately applied in the unaligned group.

Postpartum hemorrhage is the leading preventable cause of maternal mortality both in the United States and worldwide.¹¹,¹² Since treatment is often complicated by delayed or imprecise recognition, accurate and timely measurement of the actual amount of bleeding is recommended to identify hemorrhage early and institute protocol-based treatment plans that have led to improved outcomes.⁴,¹³

During vaginal delivery, blood loss is captured by calibrated under-buttocks drapes, surgical sponges and other substrates such as towels and pads that can be accurately weighed. Estimation of the volume of amniotic fluid is relatively simple since the collection of amniotic fluid precedes most of postpartum bleeding. During cesarean delivery, however, there is mixing of amniotic and irrigation fluid with maternal blood loss, further complicating the measurement of maternal blood loss.

**Strengths and Limitations**

The strength of our analysis is that the actual amount of blood loss was diligently assayed creating a blood loss “gold standard” for comparison. The major limitation was that only four of the patients studied (8%) had a hemoglobin assayed blood loss of greater than 1,000 mL, fulfilling the criteria for postpartum hemorrhage.¹³ The large differences in patients whose hemoglobin assayed blood loss was greater than 500 mL and greater than 1,000 mL demonstrate that the inaccuracy of quantitative QBL is substantial at higher amounts of blood loss.

Visual estimation of blood loss is known to be inaccurate.¹⁴,¹⁵ Therefore objective measures, such as gravimetric QBL, are recommended and promoted.¹² This analysis demonstrates significant inaccuracies in the gravimetric QBL method, suggesting alternative objective measures to

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**Table 1** Distribution of blood loss by quantitative QBL and the reference assay (mL)

<table>
<thead>
<tr>
<th>QBL ≤ 1,000 mL</th>
<th>Assay ≤ 1,000 mL</th>
<th>Assay &gt; 1,000 mL</th>
</tr>
</thead>
<tbody>
<tr>
<td>QBL &gt; 1,000 mL</td>
<td>37</td>
<td>1</td>
</tr>
<tr>
<td>QBL &gt; 1,000 mL</td>
<td>9</td>
<td>3</td>
</tr>
</tbody>
</table>

Abbreviation: QBL, quantification of blood loss.
accurately measure blood loss during cesarean delivery are needed. Several studies have shown that the colorimetric method is a practical and reliable way of accurately estimating maternal blood loss in the operating room.\textsuperscript{7,16} Using the change in hemoglobin from the preoperative level to the value following delivery, Saoud et al confirmed the accuracy of the colorimetric method.\textsuperscript{17} Katz et al recently demonstrated that quantifying blood loss using the colorimetric method resulted in improved identification of postpartum hemorrhage, changes in patient management and cost savings.\textsuperscript{18} Further studies regarding the utility of accurate QBL using the colorimetric method and its effect on reduction of maternal morbidity and mortality are needed.

\section*{Conclusion}

In summary, this study demonstrates that quantitative QBL is only moderately correlated with actual blood loss and is potentially inaccurate when bleeding is substantial. These inaccuracies were apparent despite a dedicated assessor of gravimetric QBL and optimal methodology within a study. Further studies evaluating the quantitative method in patients with high-risk deliveries and in emergency cases with higher amounts of blood loss, as well as comparisons with alternative methodologies and effects on clinical outcomes, are needed.

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\section*{Conflict of Interest}

None declared.

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\section*{References}

2. Obstet Gynecol 2019;134(06):e150–e156