RESEARCH ARTICLE

The decision to delivery interval in emergency caesarean sections: Impact of anaesthetic technique and work shift
[version 1; peer review: 2 approved]

Anette Hein, David Thalen, Ylva Eriksson, Jan G. Jakobsson

Department of Anaesthesia & Intensive Care, Institution for Clinical Science, Karolinska Institutet, Danderyds University Hospital, Stockholm, Sweden

Abstract

Background: One important task of the emergency anaesthesia service is to provide rapid, safe and effective anaesthesia for emergency caesarean sections (ECS). A Decision to Delivery Interval (DDI) <30 minutes for ECS is a quality indicator for this service. The aim of this study was to assess the DDI and the impact of chosen anaesthetic technique (general anaesthesia (GA), spinal anaesthesia (SPA) with opioid supplementation, or “top-up” of labour epidural analgesia (tEDA) with local anaesthesia and fentanyl mixture) and work shift for ECS at Danderyds Hospital, Sweden.

Methods: A retrospective chart review of ECS at Danderyds Hospital was performed between January and October 2016. Time between decision for CS, start of anaesthesia, time for incision and delivery, type of anaesthetic technique, and time of day, working hours or on call and day of week, Monday – Friday, and weekend was compiled and analysed. Time events are presented as mean ± standard deviation. Non-parametric tests were used.

Results: In total, 135 ECS were analysed: 92% of the cases were delivered within 30 minutes and mean DDI for all cases was 17.3±8.1 minutes. GA shortened the DDI by 10 and 13 minutes compared to SPA and tEDA (p<0.0005). DDI for SPA and tEDA did not differ. There was no difference in DDI regarding time of day or weekday. Apgar <7 at 5’ was more commonly seen in ECS having GA (11 out of 64) compared to SPA (2/30) and tEDA (1/41) (p<0.05).

Conclusion: GA shortens the DDI for ECS, but the use of SPA as well as tEDA with opioid supplementation maintains a short DDI and should be considered when time allows. Top-up epidural did not prolong the DDI compared to SPA. The day of week or time of ECS had no influence on the anaesthesia service as measured by the DDI.

Keywords

Caesarean section, anaesthesia, time factors
Introduction

There are around 110,000 births annually in Sweden, and the national statistics shows a trend for an increasing number of caesarean sections (CS). In 2014, 17% of all births in Sweden were CS. CS may be divided into emergency and elective procedures. Emergency CS (ECS) are commonly defined as follows: to be performed within an adequate time frame to avoid negative effects on neonate and/or mother, while elective are performed where there is no time constrain. Lucas four graded scale categorize CS by degree of urgency, as follows: 1) immediate threat to life of woman or foetus; 2) maternal or foetal compromise that is not immediately life-threatening; 3) needing early delivery but no maternal or foetal compromise; and 4) at a time to suite patient and maternity team. Dupuis suggested a coloured system to distinguish grade of emergency, to facilitate the communication, thus facilitating the process and shorten the DDF.

Need for an urgent CS is among the most dramatic anaesthetic events, requiring effective and vigilant services. It has been suggested that neonates should be delivered within 20 to 30 minutes after the decision of an urgent CS has been made. The time interval is, however, an extrapolation around time for the development of serious, life threatening acid base compromise. Various logistical programs aiming at improving the service have shown that time between decision and delivery can be reduced. The Swedish Society for Anaesthesia & Intensive Care has set recommendations that anaesthetic services should include an anaesthesiologist available within 5 minutes from the decision of an obstetric emergency and that an emergency CS incision should be possible to start within 15 minutes from the decision. The explicit evidence to support a clear medical benefit of the 30 minute decision to delivery interval (DDI) limit is sparse and this may be more of a tool to use for auditing of anaesthesia services. The NICE guidelines extend category 2: Perform category 2 CS in most situations within 75 minutes of making the decision. It should also be acknowledged that the DDI is a composite end-point, including delay between obstetric decision, press the alarm button, time until start of anaesthesia, anaesthesia ready for surgery, and surgery time.

The primary aim of the present study was to assess the impact of anaesthetic technique and work shift on the DDI in emergency CS, with a decided DDI <30 minutes at our hospital, Danderyds Hospital (category 1 and emergency category 2 CS).

Methods

The study was reviewed and approved by the Stockholm Ethical Review Board (reference number: 2016/825-31).

This is a retrospective chart review study using a proforma protocol; alarm logs and patient records for ECS at Danderyds Hospital from 1st of January to October 31st 2016 were collected and analysed. From the alarm logs, we assessed the time of the event and we then matched this information with the performed CS in our electronic surgical registration system (Orbit 5.7), from where we retrieved start time of anaesthesia, start and end of operation and type of anaesthesia performed. The follow up regarding foetal status and need of treatment was collected from the patient journal.

Routines at the department

Danderyds Hospital is an emergency hospital with about 530 beds for general and gynaecological surgery, medicine and cardiac clinic, and includes two delivery departments with a total of 10800 deliveries/year. Two anaesthesia specialists and one anaesthesia registrar are in house on call for all anaesthesia services, including intensive care. There are at minimum three surgical teams, each including one anaesthesia nurse, one surgical nurse and one or two nursing assistants. One surgical team is located in and reserved for the women’s surgical department. In case of obstetric emergency collisions, a team from the general surgical department (located in the same building) reach the women’s department in 1–2 minutes to assist.

When the attending obstetrician decides on ECS in the most severe cases, needing immediate delivery, the alarm is pressed gathering the surgical team together with anaesthesia specialist, anaesthesia registrar and neonatologist. When the obstetrician estimates the ECS need a DDI within 30 minutes, the obstetrician first calls the anaesthesia specialist by phone to give a short report, including whether there is a well working epidural to top-up and then presses the alarm to gather the team. The obstetrician follows the patient to facilitate the process.

The OR is located central of the largest delivery department on the same floor and close to the women’s surgical department – within reach in 30–60 seconds. When occupied, an OR in the women’s surgical department is used.

When an ECS with an estimated DDI time >30 minutes is decided, the surgical team is gathered by phone and no alarm is used.

The anaesthesiologist specialist on call, responsible for the obstetric anaesthesia services decides on anaesthetic technique per set routines at the department. When the obstetrician urges for an immediate delivery general anaesthesia (GA) is recommended in most cases by our routines and with an explicit need for delivery within 30 minutes’ regional anaesthesia (RA) is recommended.

In the present study, only ECS needing the alarm, delivery immediately and up to 30 minutes, were studied, with the most common cause being sign of foetal distress.

No intervention or change of routine was initiated by or associated with the present study. GA was based on a rapid sequence induction with propofol and suxamethonium following
pre-oxygenation and sevoflurane until umbilical cord is divided. Neuroaxial anaesthesia, spinal and top-up epidural followed standard routines. Spinal anaesthesia (SPA) with bupivacaine (approximately 2.4 ml 5 mg/ml), morphine (100 µg) and fentanyl (10 µg); top-up epidural (tEDA) with ropivacaine (7.5 mg/ml) and fentanyl (100 µg/20 ml ropivacaine) 15 – 20 ml.

Statistical analysis
Descriptive statistics regarding the ECS and the variables in this study was made using mean, standard deviation (SD) and range, as well as median and interquartile range (IQR), as appropriate. Mann-Whitney U test was used for comparing means between two variables and Kruskal-Wallis H test was used when comparing two or more groups. Chi-square-test was used for test of differences between category data. A p-value <0.05 was considered significant and all data were analysed in IBM SPSS Statistics 23.

Results
During the study period, 150 ECS were identified from the record systems. Data for analysis was not retrieved in 15 cases, thus 135 ECS were included in the present analysis (Figure 1).

The median DDI for the 135 ECS studied was 17 minutes (range, 5–41). The median DDI was significantly shorter for GA (10 minutes) compared to RA (SPA 20 minutes, tEDA 23 minutes) (p<0.0005; Table 1).

There was no significant difference in DDI between different working shifts: daytime, on call during the week and weekends (Table 2).

---

**Figure 1. Patient inclusion.**

**Table 1. Time events for different anaesthetic techniques.** Data are presented in minutes as median (range).

<table>
<thead>
<tr>
<th></th>
<th>Call to start anaesthesia</th>
<th>Start anaesthesia to ready for surgery</th>
<th>Surgery to delivery</th>
<th>DDI</th>
</tr>
</thead>
<tbody>
<tr>
<td>GA (n = 64)</td>
<td>6 (1–17)</td>
<td>2 (1–8)</td>
<td>2 (1–4)</td>
<td>10 (5–21)***</td>
</tr>
<tr>
<td>SPA (n = 30)</td>
<td>8 (1–23)</td>
<td>9 (1–16)</td>
<td>3 (1–7)</td>
<td>20 (13–33) ns.</td>
</tr>
<tr>
<td>tEDA (n = 41)</td>
<td>8 (1–25)</td>
<td>10 (1–23)</td>
<td>3 (1–8)</td>
<td>23 (12–41) ns.</td>
</tr>
<tr>
<td><strong>ALL (n = 135)</strong></td>
<td><strong>6 (1–25)</strong></td>
<td><strong>5 (1–23)</strong></td>
<td><strong>2 (1–8)</strong></td>
<td><strong>17 (5–41)</strong></td>
</tr>
</tbody>
</table>

*** P < 0.0005 compared to reginal anaesthesia, ns. No significant difference between SPA and tEDA. GA, general anaesthesia; SPA, spinal anaesthesia; tEDA, top-up epidural anaesthesia; DDI, decision to delivery interval.
Table 2. Time events for the different work shifts: daytime, on-call during the week and at weekends. Data are presented in minutes as median (range).

<table>
<thead>
<tr>
<th></th>
<th>Call to start Anaesthesia</th>
<th>Start anaesthesia to ready for surgery</th>
<th>Surgery to delivery</th>
<th>DDI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Daytime (n = 37)</td>
<td>6 (1–25)</td>
<td>7 (1–23)</td>
<td>3 (1–6)</td>
<td>21 (6–41)</td>
</tr>
<tr>
<td>On-call (n = 60)</td>
<td>6 (1–23)</td>
<td>4 (1–19)</td>
<td>2 (1–7)</td>
<td>14.5 (6–36)</td>
</tr>
<tr>
<td>Weekend (n = 38)</td>
<td>9 (1–17)</td>
<td>6 (1–17)</td>
<td>2 (1–8)</td>
<td>18 (5–39)</td>
</tr>
<tr>
<td>All (n = 135)</td>
<td>6 (1–25)</td>
<td>5 (1–23)</td>
<td>2 (1–8)</td>
<td>17 (5–41)</td>
</tr>
</tbody>
</table>

DDI, decision to delivery interval.

Figure 2. Decision to delivery interval for emergency caesarean sections studied (n=135) in relation to time of day. GA, general anaesthesia; SPA, spinal anaesthesia; tEDA, top-up epidural anaesthesia; DDI, decision to delivery interval.

Fourteen neonates had an Apgar score of <7 at 5 minutes: 11 out of the 64 mothers that received GA, 2 out of the 30 that received SA and 1 out of the 42 that received tEDA (p < 0.05). In all, 39 neonates were transferred to the neonatal intensive care for further observation and treatment: 22, 10 and 7 had GA, SPA and tEDA, respectively (ns.; Table 3).
Discussion

Our study was designed as a quality audit of an important part of our anaesthesia service, providing effective anaesthesia for ECS. Our anaesthesia service was seemingly effective: work shift and day of the week did not impact the DDI. General anaesthesia was expectedly associated with the shortest time for anaesthesia, as well as the lowest DDI; however the DDI was kept within 30 minutes in a clear majority of cases also when spinal anaesthesia and top-up epidural anaesthesia were chosen. The conversion of an established labour epidural, increased time for anaesthesia and DDI, but only marginally. We did not find that the use of spinal anaesthesia or top-up epidural worsened neonate outcome. Thus, we do consider that our anaesthetic service is in line with national and local guidelines, since time to establish surgical anaesthesia was achieved in a timely fashion 24/7.

Time recommendations, such as a 30 minute DDI for ECS is more of a general recommendation than based on firm evidence. Anaesthesia for CS should always be managed on a benefit vs. risk basis. The degree of foetal and or maternal distress should form the basis for management and haste of delivery. One of our primary aims was to assess how our emergency obstetric anaesthesia service performed and thus an analysis of time lines was found to be a reasonable indicator. In 2006, Blom et al. published the results from a study assessing DDI in the US. Of the included 11,481 CS, 2,808 were performed for an emergency indication. Of these, 1,814 (65%) began within 30 minutes of the decision to operate, thus a lower figure than ours. Likewise, in a more recent meta-analyses, Tolcher et al. found that 79% of category 1 deliveries and 36% of category 2 deliveries were achieved within 30 minutes, with significantly shorter time in category 1 compared to category 2 deliveries. Thus, our service was found effective and “superior” to the results found in that study.

The conversion of a labour epidural to regional anaesthesia suitable for CS has been debated, but is today seemingly well-accepted practice. The success rate for conversion is high; however prolonged duration of labour analgesia, repeated need of clinician administered bolus doses and obesity are factors suggested to increase the risk of failure. Lidocaine with adrenalin with fentanyl supplementation is commonly used for conversion. Allam et al. showed carbonated lidocaine with adrenaline to be twice as fast as sole levo-bupivacaine to achieve a T5 touch/T4 cold block, when used for conversion. Carbonated local anaesthetics are not available in Sweden. A previous study comparing lidocaine/ adrenaline/ fentanyl to plain bupivacaine did not show significant difference in time to be ready for surgery. Sng et al. compared 2% lignocaine with adrenaline and fentanyl, 0.75% ropivacaine and 0.5% levo-bupivacaine for extension of low dose epidural analgesia for urgent CS and did not find any significant difference in time to reach surgical anaesthesia. We used ropivacaine and fentanyl mixture and achieved a rapid conversion.

Table 3. Comparison of anaesthetic technique and neonatal outcome.

<table>
<thead>
<tr>
<th>Type of anaesthesia</th>
<th>GA (n = 64)</th>
<th>SPA (n = 30)</th>
<th>tEDA (n = 41)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apgar 5*, median (IQR)</td>
<td>9 (3)</td>
<td>10 (2)</td>
<td>10 (1)</td>
</tr>
<tr>
<td>Apgar 5’&lt;7, n (%)</td>
<td>11 (17)*</td>
<td>2 (7)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>Umbilical cord arterial, mean</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>7.13</td>
<td>7.21</td>
<td>7.23</td>
</tr>
<tr>
<td>pCO₂</td>
<td>9.07</td>
<td>7.40</td>
<td>7.28</td>
</tr>
<tr>
<td>Base excess</td>
<td>-9.53</td>
<td>-6.56</td>
<td>-6.55</td>
</tr>
<tr>
<td>CPAP, n (%)</td>
<td>31 (48)</td>
<td>12 (40)</td>
<td>13 (31)</td>
</tr>
<tr>
<td>Ventilation, n (%)</td>
<td>26 (40)</td>
<td>13 (43)</td>
<td>6 (14)</td>
</tr>
<tr>
<td>Intubation, n (%)</td>
<td>5 (8)</td>
<td>1 (3)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>Neonatal unit, n (%)</td>
<td>22 (34)</td>
<td>10 (33)</td>
<td>7 (17)</td>
</tr>
</tbody>
</table>

* p<0.05 Chi-square test, ns none significant between anaesthetic techniques

GA, general anaesthesia; SPA, spinal anaesthesia; tEDA, top-up epidural anaesthesia; Apgar 5:, Apgar score at five minutes; CPAP, continuous positive airway pressure.

We did not find any major difference in time delay between spinal anaesthesia, combining bupivacaine, fentanyl and morphine, and top-up epidural combining ropivacaine and fentanyl regarding time or neonatal outcome. Strouch et al. studied neonatal acid-base status and did not find any further acidosis associated to conversion epidural compared to spinal anaesthesia. We used a bupivacaine, morphine and fentanyl combination for the spinal anaesthesia, and ropivacaine and fentanyl for the epidural top-up. We used the 100-µg intrathecal dose morphine since it has been suggested to be an adequate balance between its benefits and side effects, pruritus and nausea/vomiting. Fentanyl facilitates onset and improves intraoperative analgesia. The addition of fentanyl for epidural anaesthesia has also been shown to improve quality of anaesthesia. However, the intraoperative effect has been discussed for elective CS.

The conversion of a labour epidural to regional anaesthesia for CS has been debated, but is today seemingly well-accepted practice. The success rate for conversion is high; however prolonged duration of labour analgesia, repeated need of clinician administered bolus doses and obesity are factors suggested to increase the risk of failure. Lidocaine with adrenalin with fentanyl supplementation is commonly used for conversion. Allam et al. showed carbonated lidocaine with adrenaline to be twice as fast as sole levo-bupivacaine to achieve a T5 touch/T4 cold block, when used for conversion. Carbonated local anaesthetics are not available in Sweden. A previous study comparing lidocaine/ adrenaline/ fentanyl to plain bupivacaine did not show significant difference in time to be ready for surgery. Sng et al. compared 2% lignocaine with adrenaline and fentanyl, 0.75% ropivacaine and 0.5% levo-bupivacaine for extension of low dose epidural analgesia for urgent CS and did not find any significant difference in time to reach surgical anaesthesia. We used ropivacaine and fentanyl mixture and achieved a rapid conversion.
The mother is exposed to increased anaesthetic risk when ECS is performed under general anaesthesia\textsuperscript{25}. Endler \textit{et al} suggested, following their review of maternal mortality in 1988, regional anaesthesia to be used when possible, avoiding the risk for serious airway complications\textsuperscript{26}. However, regional anaesthesia is not without risk\textsuperscript{27,28} and the Cochrane meta-analysis published in 2012 could not show any significant difference between general and regional anaesthesia in terms of risks\textsuperscript{29}.

The present results must be put into the perspective of the routine at our institution. When a push-button call is made by the obstetrician for a category 1 CS, we have always at least one experienced anaesthesiologist who is called together with anaesthesia and scrub nurses and a neonatologist. We have been working with the communication and process to facilitate regional anaesthesia when that is an option. The obstetrician informs the anaesthesiologist immediately if there is a labour epidural to top-up. The routine is to start the surgery activating dose outside, but close to, the operation theatre when it seems appropriate, with continued supervision by the anaesthesiologist, when a well working labour epidural is in place and mother and child’s status allow. If a well working labour epidural is not in place a rapid spinal is chosen, if there is time. Warm Ringers lactate (1000 ml) is used as co-load and phenylephrine as first line to stabilize blood pressure to be combined with ephedrine where pulse is < 75 min\textsuperscript{-1}. During day time when the regular operation program is on-going there is always one room spared for emergent CS.

There are several limitations with our study. We have unfortunately not been able to discriminate absolute grade of emergency apart from the attending obstetricians’ decision of a DDI less than 30 minutes in the performed CS, due to our record keeping. The decision for anaesthetic technique may of course have been influenced by the degree of foetal and/or mother compromise. It is common practice to choose GA for the most urgent category 1 ECS, and to reserve spinal and epidural top-up for cases with less maternal or foetal compromise. We found a tendency of less pCO\textsubscript{2}, lower need of CPAP, ventilation and admission to neonatal unit in favour for EDA vs. GA and even vs. SA. Indeed, epidural top-up might have been chosen for the healthiest foetus. We did furthermore not explicitly study maternal effects, e.g. need for supplementation analgesia during regional anaesthesia. Consequently, further studies are warranted.

Terbutaline administration i.v. is common practice in our intrapartum intra-uterine resuscitation routine in case of asphyxia ECS, but we did not analyse the number of patients receiving tocolytics. Other possible factors are the occurrence of maternal hypotension events and amount consumed vasopressor, ephedrine and phenylephrine, which was not analysed in our study. Foetal heart rate monitoring is continued during the top-up epidural procedure and as foetal heart rate pattern is improved the need of urgency decreases and this might influence the DDI time.

In conclusion, we found our emergency obstetric anaesthesia service effective and adherent to guidelines for DDI. Anaesthesia for ECS must, however, always be based on an individual assessment, benefit vs. risk for mother and child. General anaesthesia was as expected associated with a more rapid DDI, but spinal anaesthesia with bupivacaine, morphine and fentanyl mixture, as well as top-up labour epidural provided similar rapid time to delivery and seems a reasonable benefit vs. risk option for category 1 ECS with acceptable DDI within 20–30 minutes. Further studies assessing effect on neonatal outcome associated with the choice of anaesthetic technique are warranted.

### Data availability
Dataset 1: Raw data for the present study. doi, 10.5256/f1000research.13058.d183533\textsuperscript{30}

### Competing interests
No competing interests were disclosed.

### Grant information
The author(s) declared that no grants were involved in supporting this work.

### Acknowledgements
The study has been supported by the department of Anaesthesia, with no external funding.

### References

5. Weiner E, Bar J, Fainstein N, \textit{et al}.: The effect of a program to shorten the...


24. Sng BL, Pay LL, Sia AT: Comparison of 2% lignocaine with adrenaline and fentanyl, 0.75% ropivacaine and 0.5% levobupivacaine for extension of epidural analgesia for urgent caesarean section after low dose epidural infusion during labour. Anaesthesia Intensive Care. 2008; 36(5): 659–64. 


Data Source
Open Peer Review

Current Peer Review Status: ✓ ✓

Version 1

Reviewer Report 30 November 2017

https://doi.org/10.5256/f1000research.14157.r28440

© 2017 Özkan Seyhan T. This is an open access peer review report distributed under the terms of the Creative Commons Attribution Licence, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Tülay Özkan Seyhan
Department of Anesthesiology, Istanbul Faculty of Medicine, Istanbul University, Istanbul, Turkey

The manuscript titled “The decision to delivery interval in emergency caesarean sections: Impact of anaesthetic technique and work shift” is a well written study with limitations due to the retrospective nature.

I have three comments for possible revision:

1. Please specify the top-up timing: The most interesting result as the similar onset of epidural top-up to spinal anesthesia without any given information about the used local anesthetic volume. When were top-ups injected? (in the delivery unit just following c-section decision or in OR?)

2. Please specify the type of spinal bupivacaine (hyper-/ iso-baric?)

3. The main limitation is the lack of details about cesarean delivery indications. It is not surprising that general anesthesia allowed to quickest possible operation start. But particularly general anesthesia group includes possibly more emergent cases leading to lower Apgar scores. It may give more insight to the reader, if the authors can give data about the number of neonates with pH<7.1 at birth.

Is the work clearly and accurately presented and does it cite the current literature?
Partly

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Partly

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

---

Reviewer Report 16 November 2017

https://doi.org/10.5256/f1000research.14157.r27757

© 2017 Vaananen A. This is an open access peer review report distributed under the terms of the Creative Commons Attribution Licence, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Antti J. Vaananen
Department of Anesthesiology and Intensive Care, Helsinki University Central Hospital, Helsinki, Finland

This is a sound piece of work. Some points to consider regarding the article:

Methods:
The detailed description of the institutional procedures is a merit to this paper as it facilitates comparison of the results to other institutions.
Were the parturients allocated into the different anesthesia groups according to the intended anesthesia type or final anesthesia type (is failed epidural top-up resulting in GA analyzed as epidural top-up or GA)?
Was the rate of regional anesthesia failures addressed?

Regarding epidural top-ups:
The "start anesthesia to ready for surgery" is surprisingly fast for the epidural group given that ropivacaine (7.5 mg/ml) is used as the epidural anesthetic. It is stated in the paper that the top-up was initiated at the labour ward before transfer to the operating room which can be considered safe even in the absence of haemodynamic monitoring during transfer IF: a) the attending anesthesiologist is following the parturient to the operating room in these cases and b) the distance (=transfer time) from the labour room to the operating room is not long.
It would be helpful for the reader to see a brief description about the underlying labour analgesia system used (continuous infusion vs boluses on demand with or without background infusion) for the parturients with epidural catheters as this may have a major effect on the onset time of epidural top-up.

General comments:
As the authors note in the discussion, it is evident that GA is more likely to be chosen in the more urgent cases (evident also as a 2 minutes lower median "call to start of anesthesia -time" and worse infant outcome parameters). This underlying difference in the obstetric urgency affects direct comparison of the anesthesia modes on outcome DDI times and could warrant presentation of the data in Table 2 separately for the GA cases as well as for the regional cases (even separately for all three anesthesia subgroups).
Ultimately this study provides important aspects in relation to the very similar overall DDI time when either
spinal anesthesia or epidural anesthesia is employed (Table 1 and Figure 2) while showing that GA results in superior outcome DDI times - at least when obstetric emergency requires.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Partly

Are the conclusions drawn adequately supported by the results?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Anesthesia for cesarean delivery

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response (Member of the F1000 Faculty and F1000Research Advisory Board Member) 06 Dec 2017

Jan Jakobsson, Danderyds University Hospital, Stockholm, Sweden

Dear Referees. Thank you for effective read and constructive and important comments. We have tried to amend the paper accordingly below are explicit responses to your comments. Responses in Italics

Methods:
The detailed description of the institutional procedures is a merit to this paper as it facilitates comparison of the results to other institutions.
Were the parturients allocated into the different anesthesia groups according to the intended anesthesia type or final anesthesia type (is failed epidural top-up resulting in GA analyzed as epidural top-up or GA)? Was the rate of regional anesthesia failures addressed?
We did not have failure of blocks explicitly registered, thus we can’t give any exact data/details.

Regarding epidural top-ups:
The "start anesthesia to ready for surgery" is surprisingly fast for the epidural group given that ropivacaine (7.5 mg/ml) is used as the epidural anesthetic. It is stated in the paper that the top-up was initiated at the labour ward before transfer to the operating room which can be considered safe even in the absence of haemodynamic monitoring during transfer IF: a) the attending
anesthesiologist is following the parturient to the operating room in these cases Added to discussion, 
and b) the distance (=transfer time) from the labour room to the operating room is not long. Added in discussion, 
It would be helpful for the reader to see a brief description about the underlying labour analgesia system used Added in the discussion 
(continuous infusion vs boluses on demand with or without background infusion) for the parturients with epidural catheters as this may have a major effect on the onset time of epidural top-up.

General comments: 
As the authors note in the discussion, it is evident that GA is more likely to be chosen in the more urgent cases (evident also as a 2 minutes lower median "call to start of anesthesia -time" and worse infant outcome parameters). This underlying difference in the obstetric urgency affects direct comparison of the anesthesia modes on outcome DDI times and could warrant presentation of the data in Table 2 separately for the GA cases as well as for the regional cases (even separately for all three anesthesia subgroups). Ultimately this study provides important aspects in relation to the very similar overall DDI time when either spinal anesthesia or epidural anesthesia is employed (Table 1 and Figure 2) while showing that GA results in superior outcome DDI times - at least when obstetric emergency requires. Thank you for the adequate comment, the outcome is in our eyes overall positive and we the importance of no difference between spinal and top-up EDA 
The manuscript titled “The decision to delivery interval in emergency caesarean sections: Impact of anaesthetic technique and work shift” is a well written study with limitations due to the retrospective nature. 
I have three comments for possible revision: 
● Please specify the top-up timing: The most interesting result as the similar onset of epidural top-up to spinal anesthesia without any given information about the used local anesthetic volume. The mixture and volumes used are addressed in the method section, we can't provide single patient data about volume/time. 
When were top-ups injected? (in the delivery unit just following c-section decision or in OR?) This is explicitly addressed and added to discussion 
● Please specify the type of spinal bupivacaine (hyper-/ iso-baric?) Hypebaric added 
◆ The main limitation is the lack of details about cesarean delivery indications. It is not surprising that general anesthesia allowed to quickest possible operation start. But particularly general anesthesia group includes possibly more emergent cases leading to lower Apgar scores. It may give more insight to the reader, if the authors can give data about the number of neonates with pH<7.1 at birth. The aim was not to explicitly address neonatal outcome. We found that 14 out of the 64 neonates GA mothers had a Ph < 7.1, there was 2 neonates in spinal group of mothers (DDI 15 and 17 minutes), and only 1 out of the 41 mothers having top-up EDA (DDI 24 minutes). Added to the results.

Competing Interests: No competing interests were disclosed.
The benefits of publishing with F1000Research:

- Your article is published within days, with no editorial bias
- You can publish traditional articles, null/negative results, case reports, data notes and more
- The peer review process is transparent and collaborative
- Your article is indexed in PubMed after passing peer review
- Dedicated customer support at every stage

For pre-submission enquiries, contact research@f1000.com