Is adequate pain relief and time to analgesia associated with emergency department length of stay?  
A retrospective study

Catalina Sokoloff,1,2 Raoul Daoust,1,2 Jean Paquet,1,3 Jean-Marc Chauny1,2

ABSTRACT
Objectives: Evaluate the association of adequate analgesia and time to analgesia with emergency department (ED) length of stay (LOS).
Setting and Design: Post hoc analysis of real-time archived data.
Participants: We included all consecutive ED patients ≥18 years with pain intensity >6 (verbal numerical scale from 0 to 10), assigned to an ED bed, and whose pain was re-evaluated less than 1 h after receiving analgesic treatment.
Outcome measures: The main outcome was ED-LOS in patients who had adequate pain relief (AR=50% pain intensity) compared with those who did not have such relief (NR).
Results: A total of 2033 patients (mean age 49.5 years; 51% men) met our inclusion criteria; 58.3% were discharged, and 41.7% were admitted. Among patients discharged or admitted, there was no significant difference in ED-LOS between those with AR (median (25th–75th centile): 9.6 h (6.3–14.8) and 18.2 h (11.6–25.7), respectively) and NR (median (25th–75th centile): 9.6 h (6.6–16.0) and 17.4 h (11.3–26.5), respectively). After controlling for confounding factors, rapid time to analgesia (not AR) was associated with shorter ED-LOS of discharged and admitted patients (p<0.001 and <0.05, respectively). When adjusting for confounding variables, ED-LOS is shortened by 2 h (95% CI 1.1 to 2.8) when delay to receive analgesic is <90 min compared with >90 min for discharged and by 2.3 h (95% CI 0.17 to 4.4) for admitted patients.
Conclusions: In our study, AR was not linked with short ED-LOS. However, rapid administration of analgesia was associated with short ED-LOS.

INTRODUCTION
Emergency department (ED) overcrowding has been a concern for many years, and Canada is no exception, with nearly 60% of EDs reporting that problem in 2007.1 The phenomenon of ‘boarding’ is one of the principal factors identified as its cause.2–3 ‘Boarding’ (or ‘access block’) refers to situations where bedridden emergency patients wait for the allocation of a bed in the ward for an unreasonably long time period (prolonged ED length of stay (LOS)) with consequent patient overflow in EDs. However, a recent retrospective study revealed that among patients waiting more than 6 h in the ED, 50% were finally admitted while the other 50% were discharged,1 indicating that non-boarding patients’ LOS also contributes to overcrowding. It has been shown to be a strong predictor of low satisfaction among patients5 and healthcare workers.1 It is also associated with long hospital LOS,6 7 and high short-term and medium-term mortality rates.8–10 Furthermore, overcrowding is linked with reduced timelines and quality of interventions and treatments,11–15 including delayed analgesic administration,14 15 particularly when pain is severe,16 all of which contributes to the snowball effect of cumulating waits.

Pain represents more than 40% of consultations in EDs.17 In large studies of patients with moderate-to-severe pain, only 21–68%18–27 received analgesics, and 50–74% still had moderate-to-severe pain at discharge.17 Severe, persistent pain may also lead to unwanted physiological responses, namely, increased adrenergic tone, augmented oxygen consumption, predisposition to hypercoagulability, decreased immune function and heightened risk of delirium.28 29 Moreover, adequate and
timely treatment of acute pain could reduce the risk of chronic pain. The relationship between pain management and LOS has not been studied as a primary outcome. However, a study of intermittent injection versus patient-controlled analgesia (PCA) for sickle cell crisis pain in the ED, established that PCA was associated with a significant reduction in the ED-LOS, although there was no difference in initial or final pain intensity score.

Recent studies have attempted to identify the factors contributing to prolonged ED-LOS. Many of them have already been recognised, namely, number of laboratory examinations required, having to undergo X-ray or scan, the need for more than three medications, and number of consultants. To the best of our knowledge, the adequacy and effectiveness of pain management have never been investigated in this regard. We sought to evaluate which component of initial pain management was associated with ED-LOS reduction. We hypothesised that ED-LOS would be lessened in patients with significant pain relief.

MATERIALS AND METHODS
Study design
We conducted a post hoc analysis of real-time archived data on all consecutive patients presenting with severe pain at our ED between March 2008 and February 2011. The aim of our study was to assess if pain relief was associated with ED-LOS reduction. As a secondary objective, we evaluated if time to receiving analgesic treatment was linked with lessened ED-LOS.

Setting
Hôpital du Sacré-Cœur de Montréal is an urban, adult, level I trauma centre with 540 inpatient beds and 60 000 ED visits/year. It sustains 22 000 hospitalisations annually, of which 51% are admitted through the ED. The study was approved by the institutional review board.

Selection of participants
Patients 18 years or older were included if they were assigned to an ED treatment bed, had severe pain at triage (defined as >6 on an 11-point verbal numerical scale from 0 to 10), received an analgesic, and had their pain intensity re-evaluated in less than 1 h after such medication. Patients were excluded if they died during their ED stay, were pregnant or had been transferred from another hospital. We also excluded patients with altered mental status, intoxicated subjects and anyone with chest pain necessitating emergent percutaneous coronary intervention, because their LOS would be determined by treatments other than pain management.

Data collection
Data were extracted from computerised information and nursing records in our ED (MedUrge, MediaMed Technologies, Mont-Saint-Hilaire, Québec, Canada). This system is an integrated and mandatory working tool for all physicians, nursing staff, and any employee involved in the ED healthcare process. It contains all demographic data, triage information (including vital signs, purpose of consultation, and pain level when relevant) as well as any pertinent data collected in real time by nurses during their re-evaluation rounds, including medication administration, and pain intensity.

Data processing
The cut-off of >6 on 10 was chosen because it was felt that lower pain intensity is less likely to warrant observation in itself. ED-LOS was measured in hours from the time of arrival at the ED to discharge or admission to a ward. We defined adequate pain relief (AR) as reduction of 50% or more of the initial pain level scored on the numerical scale within 1 h after receiving the first analgesic. The 50% reduction and the 1 h threshold are based on previous literature suggesting that they represent a meaningful decline and acceptable delay in managing severe pain.

Initial pain was the one reported on the triage form. Time between arrival and analgesic administration was dichotomised into ≤90 vs >90 min and also analysed by three categories (<1; between 1 and 2; and >2 h). We selected a 90 min threshold because it is the median time to analgesia reported in many EDs with a pain scale integrated in their triage assessment.

Our primary outcome was ED-LOS of patients with AR and without adequate pain relief (NR). Our secondary outcome was ED-LOS of patients who received their medication in ≤90 min compared with those who received it after a longer time period.

Data analysis
Median LOS (25th–75th centile) between groups of patients was compared by the Mann-Whitney U test and relationship among LOS and continuous predictors by Spearman rank-order correlations. Median differences and their 95% CI are also reported. All LOS are presented in hours and separately for patients with intravenous versus patients with other than intravenous route of analgesia administration. To examine the relative influence of AR and time to analgesia on LOS, generalised linear model regressions with γ-distribution and a log link function were undertaken for patients discharged from the ED and those admitted to a ward, controlling for age, gender, route of analgesia administration (intravenous vs other), number of doses of analgesia, type of arrival at ED (ambulance or walk-in), triage priority (high vs low), crowding defined as number of patients in ED beds at the time of arrival, time of day of arrival with high or low LOS (calculated from a database of 162,000 patients of 18 years or older assigned to a bed between March 2008 and February 2011 from the same ED and selecting hours of arrival with high LOS and hours of arrival with low LOS), time between arrival and physician’s first assessment, number of examinations,
number of specialty consultations, baseline pain intensity score, trauma versus non-trauma, abdominal pain versus other, need for oxygen and for isolation. Generalised linear model was chosen because LOS is largely skewed and tends to produce less prediction errors than traditional linear regression.41 Mean LOS difference and Wald 95% CI adjusted at mean covariates were produced from estimated marginal means. The Canadian healthcare system being public and free, the presence or absence of insurance was not analysed.

**RESULTS**

A total of 2033 patients met our inclusion and exclusion criteria. Of these patients, about half (51%) were male, more than two-third arrived at ED alone, 1186 (58.3%) criteria. Of these patients, about half (51%) were male, more than two-third arrived at ED alone, 1186 (58.3%) were analysed with SPSS V.20 (IBM, Somers, New York, USA).

Among patients who were discharged from the ED, 45.7% had AR compared with 40.3% of admitted patients. There was no significant difference in ED-LOS between patients with AR compared with those NR (p=0.41 for discharged patients and p=0.87 for admitted patients; table 2).

Among patients who were discharged from the ED, 533 (45%) received analgesia in ≤90 min, with unadjusted ED-LOS reduction of 2.2 h (95% CI 1.4 to 3.0; p<0.001) compared with those with >90min. The same analysis was applied to patients being admitted: only 265 (31%) received their medication in that interval, and their median unadjusted ED-LOS reduction was 3.9 h (95% CI 2.0 to 5.7; p<0.001; table 2). Median ED-LOS for three different times to receive analgesia is displayed in table 3.

Tables 4 and 5 show the bivariate relations between LOS and all confounding variables controlled for route of analgesia administration for discharged and admitted patients, respectively. For discharged patients with intravenous route of analgesia administration—only type of arrival, crowding and baseline pain intensity score were not related to LOS; while for other than intravenous route of analgesia administration—type of arrival, triage priority, oxygen support, time of day LOS, and time to patient care by physician were not associated with LOS. For admitted patients with intravenous route of analgesia administration—tria priority, type of arrival, blood testing, time of day LOS, crowding, number of doses and baseline pain intensity score were not associated with LOS; while for other than intravenous route of analgesia administration—gender, triage priority, type of arrival, trauma injury, abdominal pain, time of day LOS and time to patient care by physician were not associated with LOS.

Multivariate analysis showed that when controlling for confounding variables, a brief time period (≤90 min) before analgesic administration (not AR) is associated with shortened ED-LOS for discharged and admitted patients (β=0.16; 95% CI 0.10 to 0.22; p<0.001 and β=0.09; 95% CI 0.006 to 0.18; p<0.05, respectively). When adjusting for confounding variables, ED-LOS is shortened by 2 h (95% CI 1.1 to 2.8; p<0.001) when time to receive analgesic is ≤90 compared with >90 min for discharged and by 2.3 h (95% CI 0.17 to 4.4; p<0.05) for admitted patients.

**LIMITATIONS**

The main limitation of our study is its post hoc design and preformed database. Potential confounding variables, such as ethnicity and linguistic barrier, which are not recorded in demographic charts of our computerised system, could not be taken into consideration. Time from pain onset, component of chronic pain and pharmacological or non-pharmacological analgesia prior to arrival at the ED were also unknown. Case complexity assessment was difficult, although we controlled for number of examinations, number of consultants, need for oxygen and for isolation, which are markers of complexity. Likewise, we do not know if some patients did...
only in Canada, but also around the world. This goal is far from being achieved in many EDs, not when Wilson and Pendleton 44 first de- fined the term ‘oligoanalgesia’. Recently, the Pain and Emergency Medicine Initiative study demonstrated that patient satisfaction was associated more with the way ED physicians responded to their symptoms of pain than to the actual result of pain treatment.19 Which components of this response to pain were significant was not specified, but a possible part of it was the promptness with which pain was addressed. Patients with severe pain probably associate receiving pain medication quickly with quality of care and are more inclined to accept a medical treatment plan, even if they do not get relief. This might explain why we observed improved ED-LOS with prompt analgesic administration in patients being discharged or admitted.

In our study, the adjusted ED-LOS was 2 h shorter in discharged patients who received their medication in ≤90 min than in those treated in >90 min. The rapid administration of analgesia, associated with shorter ED-LOS, could have a significant impact on ED overcrowding. For example, our centre received an average of 5000 patients/year with severe pain on an ED bed. If we extrapolate the proportion of patients who received analgesia in >90 min after arrival and the time saved if received in less than 90 min from our study of this population, a bed could be available during 16 h every day. Such economy of beds would contribute to better throughput of patients and render our EDs more efficient, as espoused by Asplin et al2 with their conceptual model of overcrowding in 2003.

A recent consensus of the Canadian Association of Emergency Physicians has ranked ‘ED-LOS’ and ‘time to not receive an analgesic nor had suboptimal pain management because of refusal. However, it is doubtful that any of these confounding variables would cause significant differential bias. Finally, our single-centre study in an academic hospital might limit the generalisation of our results.

### DISCUSSION

As far as we know, this is the first investigation to evaluate the impact of pain relief on ED-LOS, and our results demonstrated that rapid administration of analgesia (not AR), is associated with shorter ED-LOS. It has been reported that patients expect to receive pain medication 25–30 min after their arrival,42 which coincides with the guidelines of our triage system (Canadian Emergency Department Triage and Acuity Scale).43 Unfortunately, this goal is far from being achieved in many EDs, not only in Canada, but also around the world.19 27 42 This is a persistent problem that dates back to the late 1980s when Wilson and Pendleton44 first defined the term ‘oligoanalgesia’.

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| Disposition after ED | IV LOS in hour median (25th–75th centile) | Other than IV LOS in hour median (25th–75th centile) |
|----------------------|------------------------------------------|-----------------------------------------------|
|                      | (N=698)                                  | (N=481)                                        |
| Discharged patients  |                                          |                                               |
| Time to receive analgesia |                                          |                                               |
| <1 h                 | 8.6 (6.0–11.8)                           | 6.6 (4.4–9.5)                                 |
| From 1 to 2 h        | 10.5 (6.9–15.9)                          | 8.2 (5.4–12.2)                                |
| >2 h                 | 12.9 (8.9–18.0)                          | 10.1 (6.3–19.2)                               |
| Admitted patients    | (N=556)                                  | (N=289)                                       |
| Time to receive analgesia |                                          |                                               |
| <1 h                 | 16.4 (10.8–23.8)                         | 17.2 (10.7–24.8)                              |
| From 1 to 2 h        | 14.9 (10.4–22.6)                         | 18.1 (10.0–26.4)                              |
| >2 h                 | 18.7 (11.6–27.4)                         | 19.6 (12.9–28.7)                              |

ED, emergency department; IV, intravenous route of analgesia administration; LOS, length of stay.
first dose of analgesic’ in the top 12 priority indicators of quality care. In the USA, the Joint Commission on Accreditation of Healthcare Organizations mentions ‘early intervention’ as the first goal in the treatment of acute pain. Similarly, the Australian National Institute of Clinical Studies ranked ‘reduced time to analgesia’ as the top priority and is currently working on improving their numbers. New solutions are being proposed to improve the initial approach to pain management. For example, the simple act of making pain scoring mandatory at triage has been shown to reduce time to analgesia by 45 min. Extension of this practice could also integrate pain treatment as early as triage to limit further delays. Such measures have been introduced in Australia where nurse-initiated pain protocols are currently being evaluated. A paediatric ED study has shown 50% reduction of time to analgesia with such a protocol. Early administration of analgesics has been investigated in prehospital settings, and appears to be safe and effective, particularly with the use of intranasal fentanyl. Even if no study has yet shown a benefit of this practice in LOS, it certainly has promising advantages, and further investigations should be considered.

In summary, we found that shorter time to analgesia administration is associated with ED-LOS reduction. This observation supports recent interest in analgesia implementation as early as triage or in prehospital settings to improve the throughput component of the overcrowding phenomenon seen in EDs around the world.

### Table 4 Relationship between LOS and all confounding variables for discharged patients

| Confounding variables | IV LOS in hour (N=702) | Other than IV LOS in hour (N=484) |
|-----------------------|------------------------|-----------------------------------|
| **Categorical confounders** |                       |                                    |
| Gender                |                        |                                    |
| Male                  | 9.5 (6.7–15.0)**       | 8.0 (5.5–12.3)**                   |
| Female                | 11.1 (7.9–16.9)        | 9.8 (6.1–18.4)                    |
| Triage priority       |                        |                                    |
| High (1–2)            | 9.7 (6.7–15.3)**       | 8.8 (5.8–13.1)                    |
| Low (3–5)             | 10.8 (7.9–17.0)        | 8.8 (5.8–15.0)                    |
| Arrival               |                        |                                    |
| Ambulance             | 10.8 (7.3–16.3)        | 8.8 (5.7–15.5)                    |
| Walk-in               | 10.0 (7.2–15.7)        | 8.7 (5.8–13.5)                    |
| Trauma injury         |                        |                                    |
| Yes                   | 7.1 (4.1–13.0)**       | 6.4 (4.6–12.7)**                  |
| No                    | 10.4 (7.3–15.9)        | 9.0 (5.9–14.8)                    |
| Abdominal pain        |                        |                                    |
| Yes                   | 11.3 (7.8–17.5)**      | 10.5 (5.7–13.4)**                 |
| No                    | 9.6 (6.9–14.6)         | 8.3 (6.9–14.6)                    |
| Blood test            |                        |                                    |
| Yes                   | 16.4 (10.2–23.6)**     | 18.5 (11.6–25.0)**                |
| No                    | 10.0 (7.1–15.5)        | 8.6 (5.8–13.8)                    |
| Heart-rate monitoring |                        |                                    |
| Yes                   | 14.6 (10.6–24.4)**     | 12.7 (9.2–27.8)**                 |
| No                    | 9.8 (6.9–15.2)         | 8.2 (5.7–13.4)                    |
| Oxygen support        |                        |                                    |
| Yes                   | 13.5 (8.7–23.0)**      | 10.4 (8.7–23.0)                   |
| No                    | 10.0 (7.1–15.4)        | 8.6 (5.7–14.0)                    |
| Isolation             |                        |                                    |
| Yes                   | 22.7 (11.3–36.7)**     | 22.6 (11.7–44.3)**                |
| No                    | 10.0 (7.1–15.7)        | 8.6 (5.8–13.5)                    |
| Time of day of arrival with |                  |                                    |
| Low LOS               | 9.0 (7.1–10.7)**       | 8.6 (6.1–10.4)                    |
| High LOS              | 11.1 (7.2–16.9)        | 8.8 (5.7–15.2)                    |
| **Continuous confounders** | Spearman rank-order correlation | Spearman rank-order correlation |
| Age                   | 0.18**                 | 0.09*                             |
| Crowding              | 0.06                   | 0.06                              |
| Time to patient care by physician | 0.21**                 | 0.19**                           |
| Number of examinations (range 0–15) | 0.33**                 | 0.29**                           |
| Number of specialist consultations (range 0–8) | 0.41**                 | 0.44**                           |
| Number of doses (range 1–7) | −0.13**                | −0.10*                            |
| Baseline pain intensity score | −0.06                  | −0.01                            |

IV, intravenous route of analgesia administration; LOS, length of stay.

*p<0.05; **p<0.01.
Table 5  Relationship between LOS and all confounding variables for admitted patients

| Confounding variables                      | IV LOS in hour (N=558) Median (25th–75th centile) | Other than IV LOS in hour (N=289) Median (25th–75th centile) |
|-------------------------------------------|---------------------------------------------------|----------------------------------------------------------|
| **Categorical confounders**               |                                                   |                                                          |
| Gender                                    |                                                   |                                                          |
| Male                                      | 15.4 (10.5–23.6)*                                 | 18.9 (11.4–27.5)                                         |
| Female                                    | 18.0 (11.7–25.4)                                  | 19.5 (12.7–27.9)                                         |
| Triage priority                           |                                                   |                                                          |
| High (1–2)                                | 16.4 (10.4–24.5)                                  | 18.4 (11.1–25.8)                                         |
| Low (3–5)                                 | 17.2 (11.7–25.0)                                  | 19.5 (12.9–27.6)                                         |
| Arrival                                   |                                                   |                                                          |
| Ambulance                                 | 17.0 (10.7–25.3)                                  | 20.7 (13.5–28.4)                                         |
| Walk-in                                    | 16.6 (11.4–24.5)                                  | 18.4 (12.0–26.9)                                         |
| Trauma injury                             |                                                   |                                                          |
| Yes                                       | 12.5 (8.7–21.1)*                                  | 15.7 (8.2–27.6)                                          |
| No                                        | 17.0 (11.4–24.8)                                  | 19.4 (12.6–27.6)                                         |
| Abdominal pain                            |                                                   |                                                          |
| Yes                                       | 16.0 (10.9–23.3)*                                 | 19.4 (12.9–27.6)                                         |
| No                                        | 18.0 (11.5–27.6)                                  | 19.3 (11.6–28.2)                                         |
| Blood test                                |                                                   |                                                          |
| Yes                                       | 19.8 (11.8–28.5)                                  | 27.9 (16.4–35.8)**                                       |
| No                                        | 16.4 (10.9–24.5)                                  | 18.8 (11.6–26.8)                                         |
| Heart-rate monitoring                     |                                                   |                                                          |
| Yes                                       | 23.1 (16.2–38.6)**                                | 30.0 (25.3–45.2)**                                       |
| No                                        | 15.6 (10.7–23.1)                                  | 17.8 (11.3–25.7)                                         |
| Oxygen support                            |                                                   |                                                          |
| Yes                                       | 20.9 (12.3–34.4)**                                | 26.8 (15.1–45.0)**                                       |
| No                                        | 15.9 (10.8–24.1)                                  | 18.7 (11.7–26.7)                                         |
| Isolation                                 |                                                   |                                                          |
| Yes                                       | 30.1 (21.8–54.3)**                                | 35.5 (22.6–57.8)**                                       |
| No                                        | 15.9 (10.8–23.8)                                  | 18.5 (11.6–25.7)                                         |
| Time of day of arrival with               |                                                   |                                                          |
| Low LOS                                   | 15.7 (9.7–29.1)                                  | 16.2 (10.6–24.2)                                         |
| High LOS                                  | 17.0 (11.4–24.5)                                  | 19.8 (12.7–27.8)                                         |
| Continuous confounders                    | Spearman rank-order correlation                    | Spearman rank-order correlation                          |
| Age                                       | 0.15**                                            | 0.22**                                                   |
| Crowding                                  | 0.03                                              | 0.17**                                                   |
| Time to patient care by physician          | 0.16**                                            | 0.06                                                     |
| Number of examinations (range 0–15)       | 0.34**                                            | 0.36**                                                   |
| Number of specialist consultations (range 0–8) | 0.31**                                      | 0.37**                                                   |
| Number of doses (range 1–7)               | −0.03                                             | −0.15**                                                  |
| Baseline pain intensity score              | 0.003                                             | −0.12*                                                   |

IV, intravenous route of analgesia administration; LOS, length of stay.
*p<0.05; **p<0.01.

Author affiliations
1Department of Emergency Medicine, Research Centre, Hôpital du Sacré-Cœur de Montréal, Montréal, Québec, Canada
2Faculty of Medicine, Université de Montréal, Montréal, Québec, Canada
3Department of Surgery, Centre for Advanced Research in Sleep Medicine, Hôpital du Sacré-Cœur de Montréal, Montréal, Quebec, Canada

Contributors CS, RD and J-MC conceived the study and obtained the research funding. JP mined and analysed the data. CS drafted the manuscript, and all authors contributed substantially to its revision. CS is the guarantor. All coauthors have had the opportunity to review the final manuscript and have provided their permission to publish the manuscript.

Funding This study was supported by the Emergency Department Research Fund of Hôpital du Sacré-Cœur de Montréal.

Competing interests None.

Ethics approval The study was approved by the institutional review board and Scientific and Ethics committee of Sacré-Cœur Hospital.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement No additional data are available.

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