Incidence of mortality due to rebound toxicity after ‘treat and release’ practices inprehospital opioid overdose care: a systematic review

Jennifer Anne Greene,1,2 Brent J Deveau,1,3 Justine S Dol,1 Michael B Butler1

ABSTRACT

Introduction  Death due to opioid overdose was declared a public health crisis in Canada in 2015. Traditionally, patients who have overdosed on opioids that are managed by emergency medical services (EMS) are treated with the opioid antagonist naloxone, provided ventilatory support and subsequently transported to hospital. However, certain EMS agencies have permitted patients who have been reversed from opioid overdose to refuse transport, if the patient exhibits capacity to do so. Evidence on the safety of this practice is limited. Therefore, our intent was to examine the available literature to determine mortality and serious adverse events within 48 hours of EMS treat and release due to suspected rebound opioid toxicity after naloxone administration.

Methods  A systematic search was performed on 11 May 2017 in PubMed, Cochrane Central, Embase and CINHAL. Studies that reported on the outcome of patients treated with prehospital naloxone and released at the scene were included. Analyses for incidence of mortality and adverse events at the scene were conducted. Risk of bias and assessment of publication bias was also done.

Results  1401 records were screened after duplicate removal. Eighteen full-text studies were reviewed with seven selected for inclusion. None were found to be high risk of bias. In most studies, heroin was the source of the overdose. Mortality within 48 hours was infrequent with only four deaths among 4912 patients (0.081%) in the seven studies. Only one study reported on adverse events and found no incidence of adverse events from their sample of 71 released patients.

Conclusion  Mortality or serious adverse events due to suspected rebound toxicity in patients released on scene post-EMS treatment with naloxone were rare. However, studies involving longer-acting opioids were rare and no study involved fentanyl.

Key messages

What is already known on this subject  
▸ We knew opioid overdoses were on the rise. Illicit drug deaths have increased 6.87 times that of 2010 since the introduction of synthetic opioids.
▸ We knew that use of emergency medical services (EMS) for care of opioid overdose was increasing; in some services estimated at over 75% increase since 2012.
▸ Given the overall rise in emergency services use, we reviewed the literature to determine whether treat and release at the scene is safe.

What this study adds  
▸ In this systematic review of seven studies that reported mortality after naloxone administration by EMS and release at the scene, we found very low incidence of mortality (0.081%), suggesting the safety of this practice.
▸ The data are limited as to largely involving overdoses from heroin, a short-acting agent. It is not clear if this data can generalise to methadone. Our findings may be more generalisable to the more recent epidemic of fentanyl abuse, because of the similar-short acting nature.

INTRODUCTION

Drug-induced mortality has reached a public health crisis status1,2; overdose deaths now exceed motor vehicle collisions as a preventable cause of death in the USA.3 In Canada, there has been a 79.2% increase in illicit drug-related death in British Columbia (BC) alone between 2015 and 2016.4 The recent introduction of more powerful opioids, such as fentanyl, to recreational use is an acute cultural shift compounding the existing problem of opioid overdoses.4 Death from fentanyl overdose was declared a public health crisis in Canada in September 2015.5 The BC Coroner’s office reported 374 fentanyl-related deaths in 2016.6 This is over double that which was reported in 2015.7,8 The increase in opioid use and overdoses has led to increased demands on prehospital services and emergency departments (EDs). Emergency medical services (EMS) routinely stock naloxone, an opioid antagonist which can reverse the effects of opioids by competitively binding to receptor sites. Naloxone itself is chemically similar to an opioid with superior binding affinity to opioid receptors but little to no other pharmacological effect.9 There are several observational studies on opioid overdose treatment approaches by EMS.6–10 Currently, most, approximately 90.6%, opioid overdose patients are transported by EMS.11 The rationale for these conservative ‘support and transport’ approaches are to limit violent behaviour or instantaneous withdrawal symptoms that may ensue on delivery of naloxone12 or to limit risk of rebound toxicity if naloxone used to reverse in the field.13,14,15 The ‘support and transport’ approach may include a
titrated dose of naloxone if the patient with apnoea is difficult to ventilate making oxygenation inadequate.

An approach that warrants further exploration based on emerging evidence is for EMS professionals to use a ‘treat and release’ approach, in which paramedics fully reverse overdoses with naloxone and monitor the patient for a short observation period before the patient makes an informed decision about transport to an ED or home.20–14 However, the risks and benefits of this approach are uncertain. Studies have reported a very low incidence of rebound toxicity related to prehospital release post-naloxone administration for accidental overdose.9–14 A recent pooled analysis by Kolinsky found that 3/3875 (0.8%) of released patients had mortality related to rebound toxicity.32 However, a rigorous systematic review, with an exhaustive methodological search of the literature and assessment of the quality of the studies provides a more reliable source of information for EMS decision-making.

We therefore conducted a systematic review to assess the incidence of mortality due to suspected rebound toxicity within 48 hours after ‘treat and release’ practices by EMS professionals for patients with suspected opioid overdose.

METHODS
Data sources and search strategy
A systematic search of the literature was conducted on 11 May 2017 to identify relevant articles on the incidence of mortality and adverse events associated with EMS treatment and release of opioid overdose patients. We followed Cochrane methodology. The protocol was registered with PROSPERO: registration number CRD42017067898. We searched PubMed, Cochrane Central, Embase and CINAHL using search strategies developed with the aid of a health sciences librarian and in conjunction with an experienced advanced care paramedic (ACP). We searched using the following terms using appropriate synonyms, MeSH headings and wildcards: ‘emergency medical services’, ‘ambulances’ ‘emergencies’, ‘emergency medical technicians’, ‘paramedic’, ‘prehospital’ ‘naloxone’ and ‘narcotic antagonists’. No limits by date, language or age were applied. The bibliographies of the included studies were screened for additional relevant articles. A Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) tracking tool was used to aid in organising the search results.23 The full search strategy can be found in the online supplementary appendix A.

Data selection
Primary studies were included if they were conducted in a prehospital setting that investigated mortality in adult (>16 years) opioid overdose patients that were reversed with naloxone by EMS and subsequently refused transport to the ED or were released on scene. We included any primary study design in this review. This included but was not limited to abstracts, case studies, case series, cohorts, trials and randomised controlled trials. Secondary analysis of data, such as systematic reviews or narrative reviews, was excluded as well as were editorials, commentary or expert opinion pieces. Data from paediatrics, intentional overdoses, non-opioid overdoses, opioid overdoses treated exclusively by non-EMS practitioners or by EDs were not analysed. Our primary outcome of interest was mortality due to suspected rebound toxicity at or before 48 hours. Our secondary outcome was any adverse event due to suspected rebound opioid toxicity within 48 hours. This included any call for EMS or any ED presentation within 48 hours of the index EMS encounter.

We chose the time frame of 48 hours for two reasons: prior familiarity with the literature on this topic suggested that this was commonly studied time frame and second, considering the half-life of even the longest acting opioids (eg, methadone), 48 hours offered a clinically representative time frame that would capture every rebound due to any possible residual opioid.

We conducted title and abstract screening using Covidence systematic review software (Veritas Health Innovation, Melbourne, Australia; available at www.covidence.org), independently with two reviewers (JAG and BJD) between 8 June and 11 July 2017. Disagreements were resolved by discussion. The same reviewers similarly assessed full texts for inclusion.

Data extraction
Reviewers JAG and BJD independently extracted data from the included full texts using a predetermined extraction form. This form was designed to collect information on study design, population demographics, EMS setting/provider, intervention, outcomes and results. Our primary outcome was ‘mortality within 48 hours and secondary outcome was ‘adverse event within 48 hours’. Data on route, dose and culprit opioid were also extracted. Abstraction discrepancies were resolved via discussion. Risk of bias was evaluated using a modified Quality in Prognostic Studies (QUIPS) tool.24 25 We selected this tool a priori due to its ability to address prognostic/retrospective studies with no comparisons. Studies with high risk of bias were to be excluded from analysis.

RESULTS
Search yield
The searches yielded 1649 studies. After duplicate removal, 1401 studies remained for screening at the title and abstract phase. The Covidence programme combines these stages into one. Thirty-three studies were discussed after title and abstract screening for inclusion, resulting in 18 articles reviewed as full texts (figure 1). From these, 11 items did not meet criteria for inclusion (online supplementary appendix B). One item was removed because it was an abstract of a published study and therefore was duplicate information. One study presented data as a Cox regression and HR of death after 48 hours. Several attempts were made to contact the author in order to request data before 48 hours, but no response was received;thus, we excluded this study from analysis. Two studies were in non-English languages. Both were translated as full texts and found to not assess the outcome of interest. Therefore, these two articles were also excluded. The other studies failed to meet the inclusion criteria related to design, outcome or setting. Seven studies met criteria and were included in the analysis (online supplementary appendix B). Inter-rater agreement was unity at both title/abstract and full-text review stages.

Risk of bias
Using the modified QUIPs tool, no studies were found to be high risk of bias, and thus all were included in the analysis (table 1). Despite the tool revealing no high risk of bias, we note that all but one of the studies are retrospective. The study by Heyerdahl was prospective.26 Because of this, we consider the overall quality of evidence is limited.

Population
A total of 4912 patients were included in this review.6,7,10 12–14,16 Mean age was similar in all studies with a combined mean age
of 36.0 years (SD=5.48) (in some cases mean was estimated from the median). Females made up 20.4% of the combined patient population. All seven studies reported on prehospital opioid overdose patients treated with naloxone by a crew with at least one EMT/paramedic present (table 2). Three studies were conducted in Scandinavia and the other four were in California and Texas. Two studies were set in a fire-based system. Two other studies had a paramedic/doctor crew configuration: one in Oslo and the other Copenhagen.10 26 All studies had ACPs involved in the service: five had exclusively ACPs, two had a basic and ACP combination.

Control groups and interventions
Our intervention of interest was treatment with naloxone by EMS with subsequent release, regardless if by refusal or meeting criteria for a protocol driven release on scene. All studies investigated this intervention by retrospective chart review. Two studies were conducted within services with policies delegating specific criteria for discharge on scene post-naloxone treatments. Both of these studies were Scandinavian with a paramedic/doctor configuration. Five other studies reported on patients that had refused transport against medical advice.6 7 12–14

Three studies had comparisons to patients transported to hospital. These transported patients may be inherently different: these patients may not have responded well to the naloxone, may be suicidal or may be suffering from a multisubstance overdose. We did not aim to address these comparisons in this current review.

Several naloxone dosing regimens were used for treatment. Most dosing was at the clinician’s discretion within the parameters of 0.8 mg–2.0 mg with additional doses when required. One study reported a lesser median dose of 0.4 mg.6 Route of administration was most commonly intravenous and/or intramuscular with two studies including the intranasal route.7 10 The primary culprit opioid was heroin. In two studies, methadone and possible morphine use were reported.10 14

Outcomes
Of the 4912 patients who were included in this review, four (0.081%) died within 48 hours. The medical examiner’s office (MEO) in all cases determined the official cause of death. The time frame in which the outcome follow-up occurred varied from 12 hours to 1 week, although the most commonly used time frame

<table>
<thead>
<tr>
<th>Author (year)</th>
<th>Goal: to judge the risk of selection bias</th>
<th>Goal: to judge the risk of attrition bias</th>
<th>Goal: to judge the risk of measurement bias related to how PF was measured</th>
<th>Goal: to judge the risk of bias related to the measurement of outcome</th>
<th>Goal: to judge the risk of bias due to confounding</th>
<th>Goal: to judge the risk of bias related to the statistical analysis and presentation of results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Levine et al7</td>
<td>Low</td>
<td>N/A</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Boyd et al6</td>
<td>Low</td>
<td>N/A</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Rudolph et al10</td>
<td>Low</td>
<td>N/A</td>
<td>Low</td>
<td>Low</td>
<td>Moderate</td>
<td>Low</td>
</tr>
<tr>
<td>Vilke et al12</td>
<td>Low</td>
<td>N/A</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Vilke et al13</td>
<td>Low</td>
<td>N/A</td>
<td>Low</td>
<td>Low</td>
<td>Moderate</td>
<td>Low</td>
</tr>
<tr>
<td>Wampler et al14</td>
<td>Low</td>
<td>N/A</td>
<td>Low</td>
<td>Low</td>
<td>Moderate</td>
<td>Low</td>
</tr>
<tr>
<td>Heyerdahl et al26</td>
<td>Low</td>
<td>Low</td>
<td>Moderate</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
</tr>
</tbody>
</table>

N/A, Not applicable.
to assess mortality was 12 hours (table 1). Levine observed one death within 24 hours with a MEO reported cause of death as ‘coronary artery disease and heroin’. Authors reported that it was not definitively determined if this death was suspect to rebound toxicity or a possible new overdose. This finding was conservatively included in the four patients with the outcome.

Boyd also investigated altered level of consciousness suspect to rebound postrelease, finding no incidence of that outcome. This was the only study that reported on our secondary outcome of adverse events due to suspected rebound toxicity within 48 hours.

Statistical analysis

The incidence of the mortality outcome was so low that the assumptions necessary to perform a weighted meta-analysis could not be met (eg, normality of the observed effect sizes and weighting of the effect sizes inversely proportional to their variance). When assessing for heterogeneity, a 0.00% I² statistic was calculated and the Wald’s test statistic showed a p value of 0.9740; thus, the null hypothesis of homogeneity was not rejected. When compiling the forest plot for effect size, a non-zero constant (0.05) had to be artificially introduced to the zero incident cases in order to obtain a meaningful estimate (figure 2). There were zero events in most studies, thus not permitting for an overall proportion to be reported.

We assessed for publication bias using a funnel plot. None of the studies fall outside of the 95% CI. This plot does not indicate any evidence of publication bias (figure 3).

DISCUSSION

We uncovered seven studies (six retrospective and one prospective design) with no high risk of bias. The findings suggest very low risk of mortality when treat and release practices are employed by EMS providers in accidental opioid overdose. One challenge to generalising this existing evidence into practice is concerning the opioid used and its corresponding half-life. Our included studies found heroin to be the primary culprit in 48 of 5779 opioid toxic events in Canada in 2016–2017 causing hospitalisation, 473 were due to methadone. This is concerning for rebound toxicity, after giving a short-acting antidote such as naloxone. Naloxone has a duration of action of 30–120 min with a half-life estimated at 28.2 min, depending on the route of administration. However, in the studies with predominant heroin overdose, naloxone treat and release appeared safe. Synthetic opioids, such as fentanyl, are significant contributors to the current crisis and have varying similar pharmacokinetics to heroin. Heroin is less potent than fentanyl and this has led many healthcare providers to believe that a larger dose of naloxone is necessary when treating fentanyl overdose, yet this is generally not seen in practice and animal and case reports suggest otherwise.

The secondary outcome concerned adverse events postrelease related to suspected rebound toxicity. The intention was to investigate suspected risk factors associated with opioid reversal that may happen after the patient has been released from care

---

**Table 2** Summary of characteristics for included studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Setting</th>
<th>Mean age of EMS patients (years)</th>
<th>n, EMS treatment group</th>
<th>% female EMS patients</th>
<th>Culprit opioid</th>
<th>Intervention</th>
<th>Primary outcome</th>
<th>Secondary outcome</th>
<th>Deaths within 48 hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Levine et al⁷</td>
<td>US EMS/LAFD</td>
<td>41</td>
<td>205</td>
<td>13</td>
<td>Heroin</td>
<td>Naloxone with release</td>
<td>Death within 24 hours</td>
<td>Death with in 30 days</td>
<td>1</td>
</tr>
<tr>
<td>Boyd et al³⁶</td>
<td>Helsinki EMS</td>
<td>26</td>
<td>71</td>
<td>17.2</td>
<td>Heroin</td>
<td>Naloxone with release</td>
<td>Death within 12 hours</td>
<td>Rebound toxicity</td>
<td>0</td>
</tr>
<tr>
<td>Rudolph et al⁹⁸</td>
<td>Copenhagen MECU</td>
<td>–</td>
<td>2241</td>
<td>–</td>
<td>Heroin/morphine</td>
<td>Naloxone with release</td>
<td>Death within 12 hours</td>
<td>Death likely to rebound toxicity</td>
<td>3</td>
</tr>
<tr>
<td>Vilke et al²⁴</td>
<td>US EMS/SDFD</td>
<td>39.8</td>
<td>317</td>
<td>16.3</td>
<td>Heroin</td>
<td>Naloxone with release</td>
<td>Death within 12 hours</td>
<td>N/A</td>
<td>0</td>
</tr>
<tr>
<td>Vilke et al¹³</td>
<td>US EMS/SDFD</td>
<td>37.7</td>
<td>998</td>
<td>16.7</td>
<td>Heroin</td>
<td>Naloxone with release</td>
<td>Death within 12 hours</td>
<td>N/A</td>
<td>0</td>
</tr>
<tr>
<td>Wampler et al¹⁴</td>
<td>US EMS/SADF</td>
<td>38</td>
<td>552</td>
<td>28</td>
<td>Heroin/methadone</td>
<td>Naloxone with release</td>
<td>Death within 12 hours</td>
<td>N/A</td>
<td>0</td>
</tr>
<tr>
<td>Heyerdahl et al⁸²</td>
<td>Oslo EMS, outpatient clinic and ED</td>
<td>34</td>
<td>528</td>
<td>31</td>
<td>Heroin</td>
<td>Naloxone with release</td>
<td>Death due to rebound in 1 week</td>
<td>N/A</td>
<td>0</td>
</tr>
</tbody>
</table>

ED, emergency department; EMS, emergency medical service; LAFD, Los Angeles Fire Department; MECU, Mobile Emergency Care Unit; SAFD, San Antonio Fire Department; SDFD, San Diego Fire Department. N/A, not applicable.

---

*Figure 2* Plot of effect size for mortality.
they are safe to be discharged after 1 hour of observation if they meet the following criteria: (1) can mobilise as usual; (2) have oxygen saturation on room air of >92%; (3) have a respiratory rate >10 breaths/min and <20 breaths/min; (4) have a temperature of >35.0°C and <37.5°C; (5) have a heart rate >50 beats/min and <100 beats/min and (6) have a Glasgow Coma Scale score of 15. This length of time may not be feasible in all EMS settings but it suggests safety of the release practice. Further research is required to answer that question.

Limitations

While there are many strengths of this review, there are limitations, mainly related to the design and reporting of the included studies. All but one of the included studies were retrospective chart reviews lending to the biases and charting inconsistencies inherent to this design. Most of the studies state that they could not account for a death or adverse event outside the catchment area of their protocol. The potentially illegal and stigmatised nature of this clinical situation may lead to patients providing aliases or inaccurate information. However, measures were taken in all cases to mitigate this potential. There were also some missing data, most of which were from the Rudolph study in which 1517 of 4762 patients remained unidentified at follow-up. There was only one study that reported data on the second outcome of adverse events. However, this may be due to the fact that this outcome would be difficult to capture after the patient was released from care, particularly in a patient population that is notoriously difficult to track. Many of these studies predate the current fentanyl-related crisis and thus have limited generalisability. Prospective enrolment and rigorous follow-up would be the ideal design for this question.

CONCLUSIONS

Mortality or serious adverse events in the included studies due to suspected rebound toxicity in patients released on scene post-EMS treatment with naloxone was rare. There was very limited evidence available reporting on adverse events.

Acknowledgements

The authors would like to thank Dr Jillian Hayden for supervising this systematic review; Robin Parker for her assistance in the development of our search strategy and support during the search process and Ida Zavattin and Erik Palmén for their contribution in translating the non-English studies.

Contributors

JAG was the main author and initiated the idea and protocol, managed all key actions and created all first drafts. BJD was the second screener/abstractor and contributed significantly to the pharmacology content. JSD is the senior researcher who provided guidance and recommendations throughout and aided in the data organisation and cleaning. MBB was the statistician for this project who analysed all data and created all figures. All authors contributed to drafting and editing content for the manuscript.

Funding

The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests

None declared.

Patient consent for publication

Not required.

Provenance and peer review

Not commissioned; externally peer reviewed.

Data sharing statement

Additional data can be provided upon request to corresponding author.

REFERENCES


4. Zavattin and Erik Palmen for their contribution in translating the non-English studies.

Figure 3 Funnel plot assessing publication bias of included studies.
Original article


