A mobile device application to reduce medication errors and time to drug delivery during simulated paediatric cardiopulmonary resuscitation: a multicentre, randomised, controlled, crossover trial

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Summary
Background Vasoactive drug preparation for continuous infusion in children is both complex and time consuming and places the paediatric population at higher risk than adults for medication errors. We developed a mobile device application (app) as a step-by-step guide for the preparation to delivery of drugs requiring continuous infusion. The app has been previously tested during simulation-based resuscitations in a previous single-centre trial. In this trial, our aim was to assess this app in various hospital settings.

Methods We did a prospective, multicentre, randomised, controlled, crossover trial to compare this app with an internationally used drug-infusion-rates table for the preparation of continuous drug infusion during standardised, simulation-based, paediatric post-cardiac arrest scenarios using a high-fidelity manikin. The scenarios were split into two study periods to assess the two preparation methods consecutively, separated by a washout distraction manoeuvre. Nurses in six paediatric emergency centres in Switzerland were randomly assigned (1:1) to start the scenario with either the app or the infusion-rates table and then complete the scenario using the other preparation method. The primary endpoint was the proportion of participants committing a medication error, which was defined as a deviation from the correct weight dose of more than 10%, miscalculation of the infusion rate, misprogramming of the infusion pump, or the inability to calculate drug dosage without calculation and guidance help from the study team. The medication error proportions observed with both preparation methods were compared by pooling both study periods, with paired data analysed using the unconditional exact McNemar test for dependent groups with a two-sided α level of 0·05. We did sensitivity analyses to investigate the carryover effect. This trial is registered with ClinicalTrials.gov, number NCT03021122.

Findings From March 1 to Dec 31, 2017, we randomly assigned 128 nurses to start the scenario using the app (n=64) or the infusion-rates table (n=64). Among the 128 drug preparations associated with each of the two methods, 96 (75%, 95% CI 67–82) delivered using the infusion-rates table were associated with medication errors compared with nine (7%, 3–13) delivered using the mobile app. Medication errors were reduced by 68% (95% CI 59–76%; p=0·0001) with the app compared with the table, as was the mean time to drug preparation (difference 148·2 s [95% CI 124·2–172·1], a 45% reduction; p=0·0001) and mean time to drug delivery (168·5 s [146·1–190·8], a 40% reduction; p=0·0001). Hospital size and nurses’ experience did not modify the intervention effect. We detected no carryover effect.

Interpretation Critically ill children are particularly vulnerable to medication errors. A mobile app designed to help paediatric drug preparation during resuscitation with the aim to significantly reduce the occurrence of medication errors, drug preparation time, and delivery time could have the potential to change paediatric clinical practice in the area of emergency medicine.

Funding Swiss National Science Foundation.

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Introduction

Fast, accurate, and safe preparation and administration of continuous infusion is both complex and time consuming in paediatric critical situations, such as septic shock, cardiogenic shock, and return of spontaneous circulation (ROSC) following cardiopulmonary resuscitation from cardiac arrest.12 Most drugs given intravenously to children are provided in vials originally prepared for the adult population, which leads to the need for a specific individual, weight-based drug dose calculation and preparation for each child that varies widely across age groups. This error-prone process and the lower dosing-error tolerance of children place them at a high risk for life-threatening medication errors.13 Medication errors have been reported in up to 41% of cases during simulated paediatric resuscitations, 65% of which were incorrect.
articles were then submitted to our inclusion criteria for eligibility: articles had to describe time to preparation or delivery of vasoactive drugs for continuous infusion during cardiopulmonary resuscitation in children, or related medication errors. We only identified our previous single-centre trial as meeting these criteria.

**Added value of this study**

We developed a mobile device application—the paediatric accurate medication in emergency situations (PedAMINES) app—as a step-by-step guide for preparation to delivery of drugs requiring continuous infusion. The key finding is that medication errors with current tools were frequent and that time to preparation and delivery of vasoactive drugs for continuous infusion was dramatically reduced with the use of the app. We add new evidence of the benefit of a mobile app to improve the management of paediatric life-threatening situations by quickly delivering expertise in vasoactive drug administration, compared with an internationally used drug-infusion-rates table.

**Implications of all the available evidence**

We consider that this app has the potential to change critical care clinical practice when vasoactive continuous infusions have to be prepared and to improve quality of care in the paediatric vulnerable population. Its development also contributes to the goals of the WHO 3rd Global Patient Safety Challenge, which has the aim to reduce severe, avoidable medication-associated harm by 50% in all countries over the next 5 years.

**Methods**

**Study design and participants**

We did a prospective, multicentre, randomised, controlled, crossover trial at three tertiary and three regional paediatric emergency departments in Switzerland with a total of approximately 150 000 visits per year. The trial protocol has been published previously.11 The trial was approved by the Geneva institutional ethics committee and conducted according to the principles of the Declaration of Helsinki and Good Clinical Practice guidelines. The study was

**Research in context**

**Evidence before this study**

Medication errors are among the most common medical errors. Most drugs given intravenously to children are provided in vials prepared for the adult population, which leads to the need for a specific individual, weight-based drug dose calculation and preparation. This error-prone process places children at a high risk for life-threatening medication errors, particularly in critical situations where the preparation and administration of continuous infusions are both complex and time consuming. Efforts have been undertaken to develop cognitive tools such as tables, calculators, colour-coded resuscitation tapes, or prefilled syringes to reduce medication errors. However, these tools were intended for drugs requiring a direct intravenous bolus route and not continuous infusion. Although increasingly used in some countries, there is mixed evidence that smart infusion pumps prevent medication errors and adverse drug events. The evaluation of new methods to reduce medication errors and time to deliver vasoactive drugs as continuous infusions during paediatric resuscitation is of paramount importance but research in this area is scarce. We searched PubMed for all available studies referring to continuous infusion of vasoactive drugs during cardiopulmonary resuscitation. The search extended arbitrarily from Jan 1, 1990, until Dec 11, 2018, with no language restriction. The following medical search terms were explored and connected by Boolean operators: “resuscitation” “cardiopulmonary resuscitation”, “CPR”, “drug”, “medication errors”, “risk”, “effect”, “infusion”, “time”, and “delay”. The search was restricted to paediatric ages (from birth to age 18 years). We searched for all clinical trials, reviews, case reports and meta-analyses. 39 articles whose titles or abstracts included the search terms were reviewed. All identified articles were then submitted to our inclusion criteria for eligibility: articles had to describe time to preparation or delivery of vasoactive drugs for continuous infusion during cardiopulmonary resuscitation in children, or related medication errors. We only identified our previous single-centre trial as meeting these criteria.

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Randomisation and masking
Nurses were randomised using a stratified, single, constant 1:1 allocation ratio determined with web-based software. Written informed consent was obtained from each participant after full information disclosure prior to participation in the study. Blinding to the vasoactive drugs and doses intended for use was maintained during recruitment to minimise preparation bias. Allocation concealment was ensured with the allocation software and was not released until the nurses started the scenario. Study team members were revealed to the participants just before the scenario started. All scenarios were video-recorded for later analysis. Post-scenario video review was done without blinding by two reviewers (JNS and FL) who reviewed footage independently and were blinded to each other’s reviews. The data analyst (CC) was not blinded to treatment allocation.

The PedAMINES app
The app was developed at Geneva University Hospitals (Geneva, Switzerland) following a user-centred and evidence-based approach with emergency department caregivers, software developers, and ergonomists. On the basis of paediatric resuscitation observations and focus groups, the team worked closely together to identify the key functionalities and processes to be implemented.14

The app lists all the available resuscitation drugs with doses automatically adapted to the weight or age of the patient based on information entered when starting the app. At the time of the study, 15 drugs for continuous infusion and 19 drugs for direct intravenous injection were listed in the app and were at the participating nurse’s disposal. With one touch, any of the listed drugs can be selected and shown with a detailed preparation according to a standardised and simplified pathway. In the case of a continuous infusion, this pathway is composed of three steps: (1) drug selection, (2) dilution of the initial drug concentration, and (3) conversion of the prescribed dose rate in μg/kg per min into an infusion pump rate in mL/h.

For each drug, the exact amount to prepare is clearly displayed and thus avoids the necessity for calculations (see appendix for an example screenshot). This is based on the app’s ability to automatically calculate the optimal weight-based final infusion-pump rate and describe the preparation sequence required to achieve it, independently of the user’s competency in this domain. When using the app, the user can interact with it to start, pause, stop, increase, or decrease the perfusion rate at any time.

Multiple drugs can be prepared and run in parallel. All actions done by the user are sequentially saved locally on the device in historic files to preserve information that can be retrieved at any time for debriefing or medicolegal purposes. Historic files can also be erased or safely exported and saved in institutional electronic health records. We experienced no technical issues with the app during the study.

Procedures
On the day of participation after random allocation, each participating nurse received a standardised 5-min training session on how to use PedAMINES. The participants were then asked to perform a 15-min highly realistic cardiopulmonary resuscitation scenario on a high-fidelity manikin (SimJunior; Laerdal Medical, Stavanger, Norway), including post ROSC. The procedure was standardised across all sites using the same manikin. The scenario was conducted in situ in paediatric shock rooms to increase realism and was filmed with three video cameras (GoPro; San Matteo, CA, USA), worn by the participating nurse and placed within the room.

The untimed portion of the simulation involved a resuscitation team comprising two study team members—a doctor (JNS) and a nurse (KH)—and the participating nurse. The untimed portion of the simulation started by turning on the three video cameras, with the participant and the doctor waiting outside the shock room. Both were invited to enter the shock room by the nurse investigator. When entering the room, a clinical statement to recognise the life-threatening condition of the patient, including his weight and age, was given by the nurse investigator as follows: “Here is Junior, a 16 kg, 3-year-old boy who drowned 8 min ago in a pool and was brought to the emergency department by his parents. He is unconscious, pale, and not breathing.” At this moment, the doctor asked the participant to take a central pulse. Because of the invariable absence of a pulse, the participant was asked to assist the doctor in doing a 2-min full course massage and ventilation (15:2 ratio) manoeuvre, with the massage carried out by the participant, to increase the participant’s stress level. During this time, the doctor asked the nurse investigator to place a 3-derivation electrocardiogram, an upper-arm blood pressure monitor, and a digital pulse oximeter on the manikin. Monitoring alarms were activated to increase the realism. The doctor then asked the nurse investigator to place a peripheral vascular access on the
manikin’s right hand. At this time, an asystole rhythm was recognised and verbalised by the doctor. On the basis of the American Heart Association paediatric cardiac arrest algorithm for asystole, a bolus of 0·01 mg/kg epinephrine (0·1 mL/kg of 0·1 mg/mL concentration) was ordered by the doctor and administered by the nurse investigator. ROSC ensued with arterial hypotension. The doctor said, “He now has a return of spontaneous circulation with a pulse but with low blood pressure. It’s hypotensive shock! This patient needs a vasoactive drug, right now!” The participant was then asked to prepare and inject a 5 μg/kg per min continuous infusion of dopamine, using a syringe pump already in place, for a 16 kg boy, either with the help of the app or with the Shann infusion-rates table depending on their randomised allocation, and the timed scenario began.

During the timed scenario, both the doctor and the nurse investigator maintained a stressful resuscitation atmosphere by frequently reporting vital sounds aloud and asking the participant to promptly provide the drugs. The nurse investigator was asked to administer a 20 mL/kg sodium chloride 0·9% intravenous bolus. The nurse investigator then had to evaluate and repeat the primary assessment (ABCDE approach) according to the Pediatric Advanced Life Support recommendations. The participant was then asked to do a washout distraction manoeuvre consisting of aspirating secretions in the throat when the manikin emitted a retching sound. After this task had been completed, the crossover occurred. The doctor said, “OK, airways are now clear. But despite the volume expansion with sodium chloride and dopamine infusion, the patient is still in hypotensive shock! He needs a second vasoactive drug, right now!” The participant was then asked to prepare and inject a 0·1 μg/kg per min continuous infusion of norepinephrine by crossing over the procedure—ie, by using the app if previously they had used the infusion-rates table or vice versa. To render the task uniform between both groups, the final volume of norepinephrine required a decimal point-dependent calculation with both the infusion-rates table and mobile device app preparation methods. During this time, both the doctor and the nurse investigator maintained a stressful resuscitation atmosphere as described above. When the drug was ready to be injected, the participant was asked to deliver it to the patient using a second syringe pump already in place. The beginning of the injection corresponded to the end of the scenario. The GoPro cameras were turned off 1 min later. Before leaving the shock room, the participant was asked to recall and describe precisely how they had prepared both drugs and was asked to complete a questionnaire about the scenario immediately afterwards.

The delivery of both drugs required programming the same pump in a similar manner among all participants. The time elapsed after drug preparation until its delivery—ie, time needed to set up the pump—was assessed for all participants to ensure uniformity. The measured deviation between the amount of drug delivered and the actual prescribed dose were measured by the amount of drug in the syringe and video recorded. To ensure that participants had heard and understood the prescription orders correctly, they had to confirm the orders verbally and written transcriptions were checked and video recorded.

**Outcomes**

The primary outcome was the proportion of medication dosages containing errors that occurred during the sequence from drug preparation to drug injection. We defined an emergency medication dose administration error as a deviation from the correct weight dose of more than 10%. These errors were measured both as the percentage deviation from the amount of delivered drug compared with the correct weight dose as prescribed by the doctor and the absolute deviations from that dose. Miscalculation of the infusion rate, misprogramming of the infusion pump, and the inability to calculate drug dosage without calculation and guidance help from the nurse investigator were also considered medication errors. The accumulation of some or all of these errors was defined as a cumulative error.

Secondary outcomes were the elapsed time in seconds between the oral prescription by the doctor and time to drug preparation completion by the participant, the elapsed time in seconds between the oral prescription by the doctor and time to drug delivery by the participant, analysis of the type of medication errors (ie, error in transcription of the doctor’s order into the medication dose, wrong choice of drug, wrong vial’s initial concentration), and perceived stress and satisfaction scores after completion of the scenario, as measured in the questionnaire using ten-point Likert scales (appendix).

Other secondary outcomes were measured in this trial and they will be analysed and discussed in a separate article. These outcomes were (1) the participants’ stress level assessed by measuring continuously their heart rate using a smartwatch during the resuscitation scenario and (2) the acceptability and usability testing of the app assessed using a 52-item questionnaire based on the unified theory of acceptance and use of technology model.

**Statistical analysis**

Power calculations were based on the detection of a minimum difference of 30% in the proportion of nurses committing a medication error, which we considered to be a sufficient difference to modify the practice (appendix). Assuming that 15% of nurses would commit a medication error with the app and 45% without the app, eight participants per group had to be recruited in each participating centre to provide the trial with 90% power at a two-sided α level of 0·05. To prevent a potential loss of power due to mis-specification of assumptions, ten participants were recruited per group and per centre, giving a total sample size of
120 participants. Further information regarding the
time to drug delivery. Second, an effect size modification
of the app due to these factors was also tested using
logistic GEE models for the analysis of the error
proportions and t tests for independent groups, as well as
ANOVA tests for analyses of time to drug preparation
and delivery. Finally, means and SDs were determined
for perceived stress and satisfaction scores of individuals
from the Likert-scale questionnaire for each preparation
method and compared using a t test for paired data.

All videos were reviewed by the first reviewer (JNS). To
assess the reproducibility of the video review procedure,
a second reviewer (FL) independently duplicated the
review in a randomly selected 10% of all videos. Details
regarding the statistical analysis of the inter-rater
reliability scores are available in the appendix.

In the case of missing data, a complete case analysis
was done. No multiple imputation was planned. All
statistical tests were two-sided with a type one error risk
of 0·05. We used GraphPad Prism version 7 for graph
figures, Stata/IC version 14 for descriptive analyses,
R version 2.15.2 for GEE models and statistical tests, and
StatXact version 11.1.0 for exact statistical tests and exact
95% CIs. This trial is registered with ClinicalTrials.gov,
number NCT03021122.

Role of the funding source
The funder of the study had no role in study design, data
collection, data analysis, data interpretation, or writing of
the report. The corresponding author had full access to
all the data in the study and had final responsibility for
the decision to submit for publication.

Figure 1: Trial profile
Results

From March 1 to Dec 31, 2017, 130 nurses were assessed for eligibility, of whom 128 were randomly assigned to either the mobile app preparation method first (n=64) or the infusion-rates table preparation method first (n=64), without any dropouts or missing data (figure 1). Baseline characteristics seemed balanced in the two groups (table 1). We observed good to excellent inter-rater agreement for video reviewing (appendix).

Because data did not support a carryover effect in the proportion of participants committing a medication error (appendix), both study periods were pooled. Of the 256 drug doses delivered, 96 (75%) of 128 delivered using the infusion-rates table method were associated with medication errors compared with nine (7%) of 128 delivered using the mobile app (table 2). Medication errors were therefore reduced by 88% (95% CI 91–94%; appendix). Five (6%) of the remaining errors were due to an inappropriate amount of drug drawn from the vial whereas three (3%) were due to inappropriate dilution with sodium chloride. Three (3%) of the remaining errors were due to the indiscriminate use of the whole dopamine vial, four (4%) were related to wrong pump infusion rates, and 18 (19%) were preparations having required strong support from the nurse investigator.

Among the nine errors committed when using the app, three (33%) were overdoses ranging from 10% to 99.96% (median 90% [69–97%; appendix). 68 (71%) of the errors were due to an inappropriate amount of drug drawn from the vial whereas three (3%) were due to inappropriate dilution with sodium chloride. Three (3%) of the remaining errors were due to the indiscriminate use of the whole dopamine vial, four (4%) were related to wrong pump infusion rates, and 18 (19%) were preparations having required strong support from the nurse investigator. 72 (75%) of 96 incorrect infusions were cumulative errors. Among these, nurses unable to prepare the drugs without strong support accounted for 58 (81%) of the errors and 16 (22%) contained wrong infusion rates. We observed no errors in the communication of the drug doses (ie, no errors in the prescription given by the doctor nor these prescriptions being misheard by the participants).

Among the nine errors committed when using the app, three (33%) were overdoses ranging from 10% to 99.96% (median 90% [69–97%; appendix). 68 (71%) of the errors were due to an inappropriate amount of drug drawn from the vial whereas three (3%) were due to inappropriate dilution with sodium chloride. One participant correctly chose dopamine but followed the wrong drug preparation instruction on the app (dobutamine instead of dopamine). Another participant used norepinephrine at 0·1 mg/mL instruction on the app (dobutamine instead of dopamine). A benefit from using the mobile app was observed in all centres but with some
heterogeneity in the effect size (appendix). The hospital size (tertiary vs regional) and nurses’ years of experience did not modify the intervention effect (appendix).

Mean time to drug preparation and time to drug delivery was shorter with the app than with the infusion-rates table in both study periods (table 2; appendix). Overall, time to drug preparation decreased by 45% with the app and time to drug delivery decreased by 40% (tables 2, 3). The shorter time to drug delivery with the mobile app was similar in both study arms, regardless of whether participants started the scenario with the app or the table (table 3). Shorter time to drug preparation and time to drug delivery were observed overall when using the app but with some heterogeneity (appendix) and without any difference between tertiary and regional hospitals (appendix). Nurses’ years of experience did not modify the intervention effect (appendix). Finally, the variability of individual recorded times was lower with the app than with the table (table 2; figure 2).

The questionnaire was completed by all participants. Participants rated the overall perceived stress before the scenario as 4·8 (SD 1·9) on the 10-point Likert scale. After scenario completion, they reported higher stress using the infusion-rates table than the app (8·6 [SD 1·6] vs 4·9 [2·0]; p<0·0001). The app obtained a mean overall satisfaction score of 9·4 (1·0) out of 10.

### Discussion

Early haemodynamic alterations after ROSC or shock states can require vasoactive support as continuous drug infusions. However, despite conversion methods intended to simplify their preparation in children, such infusions remain difficult to use and prone to medication errors. In this multicentre, randomised, controlled, crossover trial, we report lower medication error rates with the mobile app PedAMINES than with an internationally used infusion-rates table for the preparation of continuous drug infusions among paediatric emergency department nurses with little experience in preparing vasoactive infusions. This result was observed in both tertiary and regional hospitals, irrespective of nurses’ years of experience. Inter-individual variance was also reduced with the app, suggesting a worthwhile benefit of its use by nurses with different experience levels.
Paediatric emergency departments present a unique clinical practice environment that is especially at risk for the occurrence of medication errors, particularly when procedures such as continuous infusions preparation in critical situations are complex and uncommon. To date, there is a paucity of studies providing insight into the magnitude of errors made in continuous infusions during emergency medical situations in critical care settings, and especially in paediatric emergency departments. This lack of data might not reflect the true reality of this phenomenon. The 75% occurrence of medication errors we observed using the infusion-rates table is consistent with these studies. 70–73% errors have been observed in handwritten continuous infusions in neonatal and paediatric intensive care units where nurses are more exposed to critically ill children requiring vasoactive continuous infusions.1 Numerous interventions involving information technologies have been developed to improve the security of the medication process.20 However, apart from computerised physician order entry systems and so-called smart intravenous pumps, few robust data are available to measure their real impact on patient safety.21 Smart pumps are available and extensively used in the USA, but they are expensive and do not have features that help to prepare a syringe for a specific weight-based infusion rate for a drug supplied in a specific concentration. Furthermore, no conclusive evidence shows that smart pumps prevent medication errors and adverse drug events.22,23 Some authors have advocated to replace tasks inducing cognitive load during paediatric resuscitation as much as possible by automated actions to optimise patient care and diminish medication errors.24 Multiple dose calculators are available on the web or as smartphone apps but most are not evidence based. These programmes most often calculate the infusion rate for a specific drug concentration, whereas the PedAMINES app handles the conversion of specific drug concentrations in mL/h into an infusion rate in μg/kg per min. They also do not provide information on how to prepare the drug solution, whereas PedAMINES does. This is an important consideration for paediatric patients. Although emergency medication given to adults being resuscitated are often in prefilled ready-to-inject syringes containing a single dose adapted for most patients, drugs given to paediatric patients are typically provided in vials not adapted for this population. The correct dose must first be calculated before being drawn. Errors with infusions frequently result from mistakes during preparation due to wrong drug-volume calculations, imprecision of volume measurements, or incorrect mixing during dilution.25 At this stage, even small errors can have a large detrimental impact on the amount of drug delivered.26,27 We suggest that a mobile device app designed to help paediatric drug preparation might circumvent some of these flaws. Although the components contributing to survival from resuscitation are complex and numerous, survival relies in part on time to drug preparation.27 In a study28 with adults in cardiac arrest, the chance of ROSC was decreased by 4% for every 1-min delay in delivery of a vasopressor bolus dose. The immediate post-arrest period is also a time in which patients are at substantial risk of re-arrest. Although optimal management of post-ROSC has not been established, maintaining sufficient cardiac perfusion by vasopressor support during and immediately after cardiac arrest is recommended as part of a bundle of care to improve haemodynamic status, avoid or minimise any hypotension-related ischaemia to vital organs, and attempt to improve long-term survival and neurological outcomes.29-31 In our study, use of the app reduced time to drug preparation and time to drug delivery by 45% and 40%, respectively. The ability to reduce the delay from the moment the drug is prescribed might contribute to improving haemodynamic support.

Our study has some limitations. First, it was done during a resuscitation simulation-based scenario rather than tested in real-life situations. However, high-fidelity simulation is an essential method to teach resuscitation skills and technologies that cannot be practised during real cardiopulmonary resuscitation because the diversity among patients and their diseases makes such studies hard to standardise in critical situations.31 Moreover, standardising the scenario and the environment helped to avoid effect modifiers by limiting the influence of undesired variables on the outcomes. Second, the 5-min app training was dispensed just before the scenario. In real-life, the interval between training and actual use would probably be months. However, training with the app months before the study would have unblinded participants to its purpose and could have created a preparation bias. Finally, our findings might not be generalisable to providers with extensive experience in preparing vasoactive infusions, such as neonatal and paediatric intensive care unit nurses.

In conclusion, this randomised trial showed fewer medication errors and shorter times to drug preparation and delivery for continuous drug infusions when using a mobile app designed to help paediatric drug preparation compared with an infusion-rates table. A next step would be to determine in real-life studies whether the reduced occurrence of medication errors and time saved owing to the use of this app translates into similar results in clinical practice. The results generated from this simulation-based study might be of great importance and might be sufficient to change and improve future paediatric emergency and critical care practice.

Contributors

NS was the chief investigator, conceived the study, led the design, collected the data, contributed to the statistical outcome analyses, prepared the figures and tables, and drafted the paper. FE conceived the study, contributed to the design, and was responsible for the development of the project software PedAMINES. CC contributed to the design and carried out the trial analyses. CL contributed to the development of the project software. KH was a study investigator, coordinated the study in each trial centre, and was in charge of the
equipment and manikin. FH was a study investigator and collected the data. PL collected the data. LL contributed to the critical review of the paper. AG contributed to the design and contributed to the critical review of the paper. SM was the trial coordinator, was a study investigator, conceived the study, contributed to the design, operated the manikin, and oversaw the drafting of the paper. The authors vouch for the accuracy and completeness of the data and for the fidelity of the trial to the protocol. All authors commented upon and approved the final manuscript.

Declaration of interests

The Geneva University Hospitals are the owners of the app PedAMINES, which is available on the Google Play Store and the Apple App Store. All authors declare no competing interests.

Data sharing

Requests for data sharing should be submitted to the corresponding author for consideration. Access to de-identified data might be granted following review upon request to the Department of Paediatric Emergency Medicine of the Geneva Children’s Hospital.

Acknowledgments

This study was funded by the Swiss National Science Foundation. We thank the nurses for their contributions to the trial and Rosemary Sudan for providing editorial assistance.

References