Impact of Personal Protective Equipment on the Performance of Emergency Pediatric Procedures by Prehospital Providers

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ABSTRACT

**Background:** Personal protective equipment (PPE) is worn by prehospital providers (PHPs) for protection from hazardous exposures. Evidence regarding the ability of PHPs to perform resuscitation procedures has been described in adult but not pediatric models. This study examined the effects of PPE on the ability of PHPs to perform resuscitation procedures on pediatric patients.

**Methods:** This prospective study was conducted at a US simulation center. Paramedics wore normal attire at the baseline session and donned full Level B PPE for the second session. During each session, they performed timed sets of psychomotor tasks simulating clinical care of a critically ill pediatric patient. The difference in time to completion between baseline and PPE sessions per task was examined using Wilcoxon signed-rank tests.

**Results:** A total of 50 paramedics completed both sessions. Median times for task completion at the PPE sessions increased significantly from baseline for several procedures: tracheal intubation (+4.5 s; \( P = 0.01 \)), automated external defibrillator (AED) placement (+9.5 s; \( P = 0.01 \)), intraosseous line insertion (+7 s; \( P < 0.0001 \)), tourniquet (+8.5 s; \( P < 0.0001 \)), intramuscular injection (+21-23 s, \( P < 0.0001 \)), and pulse oximetry (+4 s; \( P < 0.0001 \)). There was no significant increase in completion time for bag-mask ventilation or autoinjector use.

**Conclusions:** PPE did not have a significant impact on PHPs performing critical tasks while caring for a pediatric patient with a highly infectious or chemical exposure. This information may guide PHPs faced with the situation of resuscitating children while wearing Level B PPE.

**Key Words:** pediatric patients, personal protective equipment, prehospital providers

The 2015 Ebola virus epidemic in West Africa renewed worldwide attention on the ability of health-care entities to handle highly infectious patients. Development and implementation of hospital protocols to prepare for Ebola and other exposures requiring high-level personal protective equipment (PPE) occurred throughout the United States and worldwide.\(^1\),\(^2\) These efforts included training health-care providers (HCPs) to care for infected HCPs being evacuated for treatment in their native countries. Following chemical attacks in the United Kingdom and Syria, attention has also increased on the protection of HCPs in scenarios involving mass poisonings. In 2020, these preparations now include protecting HCPs from the novel coronavirus.

PPE refers to barrier clothing, gloves, and/or headgear designed to protect an individual from exposure to biologic, chemical, radiological, or environmental hazards in the prehospital or hospital environment. The key focus of PPE in health care is to provide a barrier for mucocutaneous and/or respiratory protection, with a wide spectrum of equipment. The selection of the appropriate level of PPE for HCPs is based on the specific biohazard and environment in which the patient is treated, and for which an appropriate combination of respiratory and/or chemical protection should be anticipated, classified as Levels A through D from most to least protective.\(^3\)

As front-line medical professionals, prehospital providers (PHPs) have an important role to safely care for patients with potentially transmissible illness. When the identity of a prehospital hazardous material is unknown, fully encapsulating nonpermeable, gastight protective gear (Level A) is recommended.\(^4\) Level B PPE is visually similar to Level A gear, but used for the highest degrees of respiratory protection when a...
substance is known. Both of these forms of PPE are bulky, limit
tactile and auditory feedback, and restrict mobility.\textsuperscript{5} A few
publications on adult resuscitation describe the decreased
ability of providers to perform resuscitation procedures in
PPE,\textsuperscript{6,7} but research on the impact of PPE on procedural per-
formance by PHPs caring for children remains limited. It is
also unknown as to whether or how specific elements of care
delivery should be modified when providers must wear PPE
to resuscitate pediatric patients.

Prehospital emergency care of children is already challenging.
Life-threatening pediatric emergencies are low-frequency but
high-stakes events. In the United States, paramedics and emer-
gency medical technicians serve as PHPs, but only paramedics
perform advanced life support procedures for both adults and
children. Preparations for pediatric emergencies require even
more attention to ensure the correct selection of appropriately
sized equipment and calculation of weight-based medication
dosages. Because of the relative infrequency of life-threatening
emergencies in children, simulation is a useful adjunct for
PHPs to practice these skills. This study sought to examine
the impact of Level B PPE on the ability of paramedics to per-
form simulated clinical tasks when compared with regular
attire.

**METHODS**

**Participants**

This prospective single-arm study of PHPs was performed at
the simulation center of a suburban Level I trauma and tertiary
care center; the local institutional review board deemed the
study exempt. A total of 59 eligible paramedics from a subur-
ban fire and rescue department were enrolled in the study.
Paramedics were eligible for inclusion if they (a) had experi-
ence performing the procedures to be studied, (b) had experi-
ence wearing PPE, (c) had been in their current role for at least
1 year, and (d) had no contraindication to wearing PPE.
Prescreening included a wellness and blood pressure screening
on the morning of each session. Paramedics who failed to meet
criteria were excluded from participation. Exclusion criteria
also included status as an emergency medical services student,
reported problems with claustrophobia, or a history of diffi-
culty wearing PPE.

**Study Design**

In this single-group, prospective crossover, pre/post study, par-
ticipants served as their own controls to limit the impact of
interpersonal differences in procedural skill. Demographic
information collected on each participant included age, gen-
der, height, training/discipline, area of clinical practice, years
of clinical experience, prior experience with PPE, and use of
corrective lenses.

Each participant attended 2 sessions separated by a minimum
of 2 wk. During each session, each participant performed the
same sets of procedures in the same sequence. During session 1,
participants wore normal paramedic attire, and all procedures
were performed using standard nonsterile nitrile gloves. For
session 2, participants donned a Level B PPE ensemble typi-
cally worn by the county fire department; this ensemble
included a (1) Kipler Encapsulated Level B suit (nongas
tight-hooded garment), (2) a Scott AP50 4.5.2002 NFPA
(National Fire Protection Association)-compliant self-
contained breathing apparatus (SCBA) with facemask and
60-min duration cylinder, (3) Honeywell North Chemical
Resistant 4T453 B131/10 (13 mil) gloves, and (4) Tingley
TG-82330 hazardous materials boots. The participants per-
formed selected procedures in Guardian Neoprene smooth
Level A chemical hazard class gloves (35 mil). The order of
gloves used first was randomized. To avoid learning effects
in the PPE session when the 2 types of gloves were worn, data
only included the gloves that were worn first in the analysis.
The study flow is detailed in Figure 1.

**Procedures**

The specific procedures were selected based on their impor-
tance and ability to be assessed in a simulated setting. In all
cases, timing was measured by a member of the study team
who was physically present to observe and used a stopwatch.
The success of each task was based on the following predeter-
mined criteria:

A. **Airway skills**

1. Bag-mask ventilation (BMV): Participants were asked to per-
form BMV using a 0.5-L self-inflating bag and a preselected
appropriately sized mask on the Laerdal SimJunior manikin
(Laerdal, Wappingers Falls, NY). Participants were asked to
assemble the bag-mask and perform 3 ventilations that were
verified by chest rise. Successful completion was defined as
when the chest returned to baseline. Participants were given
5 min to complete the task. Direct observation was used as it
was found to be more accurate than software feedback during
the pilot testing.

2. Endotracheal intubation: Participants were asked to perform
direct laryngoscopy using a 4.5 ET tube and a Miller 1 blade,
placed in the same positions at the head of the bed for each
trial. Participants were instructed to stand at the head of the
bed before starting. Successful completion required manikin
intubation and ventilation within 3 min. Time to comple-
tion was defined as when the chest returned to baseline.

3. Pulse oximetry probe placement: Participants were asked to
place a pulse oximeter probe onto the toe of the manikin.
Successful completion required placement in less than a
minute. Time to completion was measured from the instruc-
tion to begin until probe was secured.

B. **Circulation Skills**

1. Intravenous (IO) needle placement: Participants were asked
to place an Arrow EZ-I0 using an Arrow EZ-I0 Power Driver
(Teleflex, Morrisville, NC) into the tibia of a Pediatric HAL

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**Figure 1.**

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FIGURE 1

Study Design.

Minimum of 2 weeks between sessions

<table>
<thead>
<tr>
<th>Session 1: Normal Attire</th>
<th>Session 2: Level B PPE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Station 1: Airway and breathing skills</strong>&lt;br&gt;Bag-mask-ventilation, tracheal intubation, pulse oximetry probe placement</td>
<td><strong>Station 1: Airway and breathing skills</strong>&lt;br&gt;Bag-mask-ventilation*, tracheal intubation, pulse oximetry probe placement</td>
</tr>
<tr>
<td><strong>Station 2: Circulation skills</strong>&lt;br&gt;Intraosseous needle, intramuscular (IM) injection (syringe), IM autoinjector, tourniquet placement</td>
<td><strong>Station 2: Circulation skills</strong>&lt;br&gt;Intraosseous needle*, IM injection (syringe)<em>, IM autoinjector, tourniquet placement</em></td>
</tr>
<tr>
<td><strong>Station 3: Automated external defibrillator (AED) skills</strong>&lt;br&gt;Pediatric AED skills, pad placement and defibrillation</td>
<td><strong>Station 3: AED skills</strong>&lt;br&gt;Pediatric AED skills, pad placement and defibrillation</td>
</tr>
</tbody>
</table>

Station 1

Station 2

Station 3
Effect of PPE on Paramedics’ Performance

C. Automated External Defibrillator Pad Application and Defibrillation

Participants were asked to place a pediatric adaptor into the automated external defibrillator (AED) device (Philips MRx; Koninklijke Philips N.V., Amsterdam, Netherlands) and then place the adhesive defibrillator pads onto the chest wall of the 5-year-old child manikin (MegaCode Kid, Laerdal, Wappingers Falls, NY). Successful completion was the delivery of a shock. Time was measured from the instruction to begin until the AED noted that a shock was delivered.

The interrater reliability for the primary time to completion measures was assessed through timing of the participant with 2 raters in a sample of visits. Twenty percent of participants were selected, and only PPE visits were used for this assessment.

A multiple-choice attitude survey was administered to participants at both sessions to assess whether (1) nonsterile gloves would interfere with procedures, (2) full-body PPE suits interfered with procedures, (3) PPE made it difficult for the provider to focus upon the procedure, (4) PPE was claustrophobic, (5) the provider would be slower performing procedures in the full-body PPE, and (6) the provider felt prepared to appropriately don Level B PPE. A 5-point anchored rating scale was used.

Analysis

Demographic and baseline characteristics were summarized with frequencies and percentages for categorical data and means and standard deviations (SDs) for continuous data. The primary outcome was time to procedure completion, comparing regular paramedic attire (baseline) and PPE; the secondary outcome was the attitude toward using PPE as assessed in the attitude survey.

For the primary outcome, the median difference in time of performing selected tasks with and without PPE was examined using Wilcoxon signed-rank tests. Differences between the completion times when paramedics wore Level A or B gloves were also compared using 2-sample median tests. The survey data were analyzed using paired t-tests comparing baseline and post-PPE session responses. Interrater reliability was assessed for agreement of raters at the PPE session using the intraclass correlation (ICC). All tests were 2-sided; \( P < 0.05 \) was considered to be statistically significant. Statistical analyses were performed using SAS software version 9.4 (SAS Institute Inc., Cary, NC).

Sample Size Calculation

The study was powered to detect a 10-s difference in time to completion between baseline and PPE sessions. The SDs of the differences between baseline and PPE sessions was assumed to be as high as 20.6 s with an assumed intra-individual correlation of 0.6 between assessments. Using these assumptions, a sample size of 48 participants was sufficient to provide >90% power for time to completion endpoints when assessing difference using 95% 2-sided confidence intervals. Significance levels used in the power analysis for this exploratory analysis were not adjusted for multiplicity.

RESULTS

A total of 59 eligible paramedics from a suburban fire and rescue department were enrolled; 50 participants attended the 2 study sessions and were included in the analysis. Participant demographics and reported levels of experience are provided in Table 1. The mean paramedic age was 39.6 (SD 6.9) years, and 94.0% were male. Almost 90% had previously worn PPE more than 10 times.

Table 2 summarizes the time to completion of each procedure in the order in which the tasks were performed, as well as the procedures that were performed in the 2 types of gloves. For procedures performed in full Level B PPE with Level B gloves only, the median time to completion increased by 4.5 s (\( P = 0.01 \)) for tracheal intubation, 4 s for pulse oximetry probe placement (\( P < 0.0001 \)), and

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9.5 s for AED pad placement and defibrillation \( (P = 0.01) \) compared with the baseline session with regular nitrile gloves.

In the comparison of Level A and Level B gloves, analysis was performed only on the type of glove worn first to minimize the effect of learning. The completion time between baseline and PPE visits for BMV was 1 s in both Level A \( (P = 0.5) \) and Level B \( (P = 0.01) \) glove types. The median time for IM injection preparation (drawing up the medication and removing the needle from the vial) increased 21 s in Level B gloves but 28.5 s in Level A gloves from baseline \( (P < 0.0001) \). The median time to deliver the IM medication (from drawing up the medication, injecting, and then removing the needle from the thigh) increased 23 s in Level B gloves \( (P = 0.0001) \) and 35 s in Level A gloves \( (P = 0.0001) \) from baseline. The median time to deliver an injection by means of autoinjector increased by 2 s in Level B gloves \( (P = 0.46) \), and 4 s in Level A gloves \( (P = 0.03) \) from the baseline. The median times for tourniquet placement increased by 8.5 s in Level B gloves, and 12.5 s in Level A gloves \( (P < 0.0001) \) from baseline (Table 2).

Of note, all participants were able to meet the defined standard for procedures except tracheal intubation and IM autoinjector, for which 3 and 1 participants, respectively, did not meet the standard in PPE. Interrater reliability was high (ICC > 0.86) for all tasks except AED implementation (ICC = -0.14).

**Attitudes Survey Results**

On the presession survey (Table 3), the participants agreed or strongly agreed that PPE would interfere with procedures and slow procedures down, and were neutral on the impact of PPE on their focus. They did not believe that the PPE was claustrophobic and believed they were prepared to don PPE. After the PPE session, the participants were significantly more likely to agree that nonsterile gloves affected procedures (mean change score = 1.7; \( P < 0.001 \)). They also felt more positively after the PPE session than at baseline regarding their preparedness to don PPE (mean change score = 0.5; \( P = 0.01 \)). The change in responses from baseline to post-PPE altered the overall group category from disagree to neutral for the impact of nonsterile gloves on procedural performance.

### TABLE 1

<table>
<thead>
<tr>
<th>Demographic and Baseline Characteristics</th>
<th>Participants ( (N = 50) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age (SD), years</td>
<td>39.6 (6.9)</td>
</tr>
<tr>
<td>Male sex</td>
<td>47 (94.0%)</td>
</tr>
<tr>
<td>Number of times PPE worn previously</td>
<td></td>
</tr>
<tr>
<td>1-2</td>
<td>1 (2.0%)</td>
</tr>
<tr>
<td>3-5</td>
<td>2 (4.0%)</td>
</tr>
<tr>
<td>6-10</td>
<td>3 (6.0%)</td>
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<tr>
<td>&gt;10</td>
<td>44 (88.0%)</td>
</tr>
</tbody>
</table>

Abbreviation: PPE, personal protective equipment; SD, standard deviation.

### TABLE 2

<table>
<thead>
<tr>
<th>Completion Time of Study Procedures</th>
<th>Level B</th>
<th>Level B</th>
<th>Level B</th>
<th>Level B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Task</td>
<td>Median (IQR)</td>
<td>Median (IQR)</td>
<td>Median (IQR)</td>
<td>Median (IQR)</td>
</tr>
<tr>
<td>-------------------------------</td>
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</tr>
<tr>
<td>Bag-mask ventilation</td>
<td>14 (12.2, 17)</td>
<td>151 (14.1, 19)</td>
<td>26 (15.1, 11.15)</td>
<td>61.5 (50.7, 77.1)</td>
</tr>
<tr>
<td>Endotracheal intubation</td>
<td>37 (32.4, 44)</td>
<td>43 (33.2, 62)</td>
<td>50 (33.2, 25)</td>
<td>50 (10.7, 51)</td>
</tr>
<tr>
<td>Pulse oximetry probe</td>
<td>49 (32.9, 41)</td>
<td>42 (35, 51)</td>
<td>50 (43.5, 51)</td>
<td>50 (43, 51)</td>
</tr>
<tr>
<td>Intravenous injection (syringe)</td>
<td>62 (46.6, 93)</td>
<td>62.5 (46.6, 93)</td>
<td>50 (43, 51)</td>
<td>50 (43, 51)</td>
</tr>
<tr>
<td>Automated external defibrillator</td>
<td>885 (56, 114)</td>
<td>97.5 (147, 114)</td>
<td>26 (16, 49)</td>
<td>26 (16, 49)</td>
</tr>
</tbody>
</table>

Abbreviations: IQR, interquartile range; PPE, personal protective equipment.

Wilcoxon signed-rank test.
DISCUSSION

The results revealed that, while Level B PPE did increase the time to completion of some procedures, it had little to no impact on the ability of paramedics to successfully perform all categories of resuscitation procedures in the pediatric model. Our study is the first to address this with PHPs.

Glove Type

Although glove type was not observed to impact paramedics' performance of BMV or IM autoinjection, the measured completion times of all tasks requiring fine motor skills were longer when performed in Level A gloves, and from baseline (Table 2). This might suggest the thickness of chemical protective gloves (CPG) as a factor in causing delays. After the PPE session, paramedics were more likely to believe that CPG interfered with procedures than with regular nitrile gloves. Both glove types were observed to interfere with tasks involving handling adhesive material such as tape and AED pads, but actions such as the tearing open of equipment packaging and the twisting of IV connector equipment were observed to be more difficult in CPG. However, differences of 4 s for tourniquet placement and 12 s for IM injection when performing these procedures in Level A versus Level B gloves are unlikely to be of clinical significance. This suggests that CPG thickness may not necessarily be the factor that inhibits these skills, but just the CPG themselves.

When comparing times of IM injection, paramedics wearing CPG took up to 35 s (Level A) and 23 s (Level B) longer than baseline, which was also statistically significant. While this is also unlikely to be clinically significant, CPG only caused an increase of 4 s from baseline in IM autoinjection in Level A gloves. A retrospective study investigating survival after epinephrine administration for nonshockable out-of-hospital arrest in 595 pediatric patients found that the odds of survival were 9% lower for every minute of delay in administration of epinephrine, with 71% of children receiving epinephrine by means of intraosseous. The odds ratio of survival when epinephrine was given late versus early was 0.43. The results of our study support that IM autoinjection is preferable to an IM injection when in CPG. Although not studied here, the insertion of a simple supraglottic device would also probably not be hindered by CPG as tracheal intubation was minimally impacted by CPG versus baseline.

To explore other reasons for PPE's impact on procedures, we observed the ability of the paramedics to see or hear instructions in the encapsulating hood and SCBA equipment. Level B PPE was chosen as it closely resembles Level A PPE with minor differences: Level A PPE has built-in gloves and is airtight once the seal is fastened (there are no vents). Level B PPE was chosen as it closely resembles Level A PPE. To explore other reasons for PPE's impact on procedures, we observed the ability of the encapsulating hood and SCBA equipment. To explore other reasons for PPE's impact on procedures, we observed the ability of the paramedics to see or hear instructions in the encapsulating hood and SCBA equipment. Level B PPE was chosen as it closely resembles Level A PPE with minor differences: Level A PPE has built-in gloves and is airtight once the seal is fastened (there are no vents). Level B PPE was chosen as it closely resembles Level A PPE.

LIMITATIONS

Variability in the level of provider experience would be a possible source of bias. As the intent of the study was to investigate the impact of the PPE on procedural performance, the study...
The decreased ability of the maybelle.kou@inova.org intubation VOL. 16/NO. 1 Hospital, 3300 Gallows Road, Falls Church, VA 22042 Performance Disaster Medicine and Public Health Preparedness overall completion of positions in PPE could complicate procedural performance (eg, medics would not have a world encounters and unpredictable environments, the para- with donning and doffing, and cleaning off their visor. In real- sions, each procedural task was simplified and performed with brand new equipment. Manikins were placed on stretchers at waist height. To remove the effect of retrieving equipment from a bag, it was laid out on countertops for easy access. The rooms were air-conditioned with ample space to move about. For safety reasons, a volunteer assisted each participant with donning and doffing, and cleaning off their visor. In real- world encounters and unpredictable environments, the paramedics would not have a “spotter” or be able to clean their own visor. It is possible that prolonged crouched or awkward positions in PPE could complicate procedural performance (eg, “on the floor” intubation11) or increase metabolic demand.12 The latter could not be studied due to the oxygen tank supply and our participants were not in PPE for greater than 1 h. Given current conditions where providers are in PPE for pro- longed periods of time, this would be useful to investigate. Last, the integrity or durability of the suit could be theoretically affected by kneeling on rough surfaces, but this was not identi- fied in the literature.

It could be argued that these procedures would not be under- taken in a prehospital response to a large-scale disaster involving a biohazard. The JumpSTART pediatric triage algorithm14 and operations plans for Hot Zone responses15 recommend the prioritization of resuscitation efforts when faced with multiple critical patients. This study supports that in the case of a single pediatric patient suffering from exposure to a biohazard, airway rescue, or autoinjection of antidote/pressor by an experienced provider would not be impacted by wearing Level B PPE.

This simulation study of a pediatric model adds to the literature to address the assumption that Level B PPE would impact the performance of resuscitative procedures by PHPs. However, it is still imperative that PHPs have access to pediatric procedural practice sessions while in PPE. At the very least, procedural training in CPG could be useful where PPE is a commodity or in scarce supply.

CONCLUSIONS
The use of Level B PPE made no statistically significant impact on experienced paramedics’ measured performance of BMV and IM autoinjection in a pediatric model and suggests auto- injection is preferable for resuscitation medication administra- tion when using CPG. While there was no clinically significant impact on the paramedics’ overall completion of the remainder of advanced resuscitative procedures, this study suggests that pediatric preparedness and training drills in Level B PPE would especially be of benefit for PHPs with less expe- rience. Preparedness to work in Level B PPE should continue to be emphasized, especially when the use of CPG is required, and is not a deterrent to resuscitation attempts on a pediatric patient in the right clinical scenario. This information pro- vides guidance to those preparing for events requiring Level B PPE in the care of acutely ill children.

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M.K.: study design, data collection, analysis and interpretation, drafting and revisions of the manuscript for important intellectual content and statistical expertise; A.J.D. and M.D.A.: study design, analysis and interpretation, drafting and manuscript revisions; S.K. and D.S.: study design, interpretation, acquisition of funding; M.N.: study design, data collection and interpretation of data; H.S., A.K., and M.S.: data collection, revisions; L.K. and J.D.: study design, manuscript revisions; J.C. and G.S.: statistical expertise.

Conflicts of Interest
None of the authors have any relevant conflicts of interest.

REFERENCES