The effect of simulation-based training on initial performance of ultrasound-guided axillary brachial plexus blockade in a clinical setting – a pilot study

Owen O’Sullivan1,2,3*, Gabriella Iohom1,2,3†, Brian D O’Donnell1,2,3† and George D Shorten1,2,3†

Abstract

Background: In preparing novice anesthesiologists to perform their first ultrasound-guided axillary brachial plexus blockade, we hypothesized that virtual reality simulation-based training offers an additional learning benefit over standard training. We carried out pilot testing of this hypothesis using a prospective, single blind, randomized controlled trial.

Methods: We planned to recruit 20 anesthesiologists who had no experience of performing ultrasound-guided regional anesthesia. Initial standardized training, reflecting current best available practice was provided to all participating trainees. Trainees were randomized into one of two groups: (i) to undertake additional simulation-based training or (ii) no further training. On completion of their assigned training, trainees attempted their first ultrasound-guided axillary brachial plexus blockade. Two experts, blinded to the trainees’ group allocation, assessed the performance of trainees using validated tools.

Results: This study was discontinued following a planned interim analysis, having recruited 10 trainees. This occurred because it became clear that the functionality of the available simulator was insufficient to meet our training requirements. There were no statistically significant difference in clinical performance, as assessed using the sum of a Global Rating Score and a checklist score, between simulation-based training [mean 32.9 (standard deviation 11.1)] and control trainees [31.5 (4.2)] (p = 0.885).

Conclusions: We have described a methodology for assessing the effectiveness of a simulator, during its development, by means of a randomized controlled trial. We believe that the learning acquired will be useful if performing future trials on learning efficacy associated with simulation based training in procedural skills.

Trial registration: ClinicalTrials.gov identifier: NCT01965314. Registered October 17th 2013.

Keywords: Ultrasound-guided regional anesthesia, Simulation, Validation, Virtual reality, Procedural training, Technology-enhanced learning

Background

The learning environment in which resident anesthesiologists acquire procedural skills has fundamentally changed. Training programmes are shorter and afford fewer training opportunities. Patient, institutional and regulatory expectations limit acceptance of trainees acquiring skills by “practicing” on patients. In the context of ultrasound-guided axillary brachial plexus blockade (USgABPB), we have demonstrated that anesthesiologists in Ireland perceive a lack of learning opportunity as being the most important impediment to procedural skill development [1].

It is indisputable that simulation will play an increasingly important part in the training and assessment of procedural skills [2]. Simulation offers trainees an opportunity to hone skills in a risk-free environment. Training bodies are attempting to move from traditional time-based training programmes to competency-based
training [3]. Since January 2010, the American Board of
Anesthesiology (ABA) has included simulation-based
training as a mandatory component of Maintenance of
Certification in Anesthesiology (MOCA) [4]. A recent
meta-analysis demonstrated that technology-enhanced
simulation-based training is associated with large posi-
tive effects on knowledge, skills, and behaviors, and
moderate effects on patient based outcomes [5].

To date, simulation in ultrasound-guided regional
anesthesia (UGRA) has largely been limited to tissue
(e.g. turkey breasts or cadavers) and non-tissue (e.g. gelatin
or tofu) phantoms [6,7]. Computer-based VR simulation
has been utilized effectively for training in a number of pro-
cedural domains, e.g. laparoscopic surgery [8] and colonos-
copy [9]. Grottke et al. [10] have previously described the
development of a virtual reality (VR) simulator for regional
anesthesia guided by peripheral nerve stimulation. Previous
work at our institution described the development of a
similar device simulating spinal anesthesia [11]. VR simula-
tion offers a number of advantages over the alternatives; (i)
variety of predefined standardized scenarios, (ii) multiple
anatomical variations, (iii) models do not degrade with
repeated needle insertion, (iv) realistic representations of
anatomy acquired via MRI, CT or ultrasound derived data,
(v) normal variation of a single anatomical site can be rep-
resented, and (vi) multiple anatomical sites (thus different
types of blocks) can be represented in a single simulator
[12]. We have participated in developing a VR visuo-haptic
simulator to train USgABPB, as part of a collaborative
project with the National Digital Research Centre
(www.ndrc.ie). The simulator is intended to render the
haptic (related to tactile and proprioceptive) sensa-
tions normally felt during manipulation of both needle
and ultrasound probe. We set out to assess the effect of
training USgABPB utilizing a novel prototype simu-
lator on skill transfer, during its development.

We hypothesized that VR-based training offers an
additional learning benefit over standard training (using
cadaveric dissection and human volunteers) in preparing
novice anesthesiologists to perform their first USgABPB
in the clinical setting. We carried out pilot testing of this
hypothesis using a prospective, single blind, randomized
control trial.

Methods
This prospective, randomized controlled trial was con-
ducted at Cork University Hospital and St Mary's Ortho-
paedic Hospital (Cork, Ireland). The Clinical Research
Ethics Committee of the Cork Teaching Hospitals approved
the study and the study was registered with ClinicalTrials.
gov (NCT01965314). All subjects, patients and anesthesi-
ologists, provided written informed consent. We planned to
recruit 20 residents (all College of Anaesthetists of Ireland
affiliated trainees) with no experience of performing UGRA.
The sample size was arbitrarily based on previous studies
indicating the effectiveness of VR simulation-based teach-
ing procedural skills to novices [8]. Subjects provided base-
line personal data, experience in practice of anesthesia
(years in training) and handedness. Each subject was asked
to categorize his/her (i) previous experience of peripheral
nerve blockade with peripheral nerve stimulation [0 = 0
blocks, 1 = 1–5 blocks, 2 = 5–10 blocks, 3 = 10–50 blocks,
4 = 50–100 blocks, 5 = 100 blocks] (ii) previous experience
of ultrasound-guided vascular access [0 = 0 procedures, 1 =
1–5 procedures, 2 = 5–10 procedures, 3 = 10–50 proce-
dures, 4 = 50–100 procedures, 5 = 100 procedures] (iii)
previous attendance at a peripheral nerve blockade course
(incorporating ultrasound-guided techniques) [0 = never,
1 = < 50 procedures, 2 = 50–100 procedures, 3 = 100–500
procedures, 4 = 500–1000 procedures, 5 = > 1000 procedures]
(iv) previous attendance at a peripheral nerve blockade course
(vi) multiple courses. Baseline visuo-spatial ability was
assessed using the card rotation, shape memory, and snowy
picture tests (Educational Testing Service) [13]. Psychomotor
ability was assessed using a grooved pegboard
(Lafayette Instruments, Lafayette, IN). Subjects were ran-
domly allocated (non-stratified) into 1 of 2 groups, (i) the
teacher group (CG) or (ii) the simulator trained group (SG)
using random number tables.

Common training
All participating anesthesiologists received standardized
training. These educational sessions took place in the
Department of Anatomy, University College Cork. The
educational sessions were attended by 4–6 trainees. A
single anesthesiologist (BOD) with expertise in both
teaching and performing the procedure delivered all ses-
sions and supervised the trainees during the hands-on
sessions. Each session comprised a number of compon-
ents, namely; (i) a didactic session, (ii) an hands on ses-
don with appropriately prepared cadaveric specimens,
(iii) ultrasound scanning of a volunteer, and (iv) a need-
ling skills session with tissue phantoms. Subjects were
taught to perform USGABPB using a technique as de-
scribed in Appendix IV V of ‘The American Society
of Regional Anesthesia and Pain Medicine and the
European Society of Regional Anaesthesia and Pain
Therapy Joint Committee Recommendations for Educa-
ion and Training in Ultrasound-Guided Regional
Anesthesia’ [14]. All ultrasound examinations performed
on volunteers or on patients entailed the use of a Sonoi-
site M Turbo (Sonosite, Bothell, WA, USA) (or similar
device) with a 7–12 MHz 38 mm linear probe. Following
the educational intervention, all subjects were asked to
give written feedback, by means of a standard form, on
the content and delivery of the session. On completion of
the common training those in the CG received no
further training and those in the SG went on to
complete a proficiency-based training period using a
prototype simulator.
Simulator training

The simulator was comprised of two PHANTOM Desktop devices (www.sensable.com, Wilmington, MA, USA), a desktop computer (Hewlett-Packard, www.hp.com), a liquid crystal display (LCD) monitor (Samsung Sync master 2233) capable of rendering 120 frames per second synchronized with a pair of 3D stereoscopic glasses (www.nvidia.co.uk), and the H3D API (www.sensegraphics.se). The SG subjects were asked to scan and perform procedure specific tasks on a virtual arm. The model of the arm was informed using 1.5 Tesla MRI DICOM datasets which generated skin and bone surfaces. A number of computer generated structures were added to this model based on typical anatomical positioning (The Science Picture Company, www.sciencepicturecompany.com, Dublin, Ireland). These were the axillary artery and three nerves (representing median, ulnar and radial nerves). The resultant image was thus a computer generated “animation”.

Before subjects began simulation-based training, 3 experts (each of whom had undertaken structured higher subspecialty training in regional anesthesia and maintained proficiency by performing at least 100 UGRA procedures during the previous year) performed each task under similar conditions on three consecutive occasions. The mean values of their performances set a proficiency level against which subsequent trainee performance was benchmarked.

Following initial familiarization with the simulator, lasting 50 – 60 minutes (duration partially due to the prototypal nature of the device), SG subjects were asked to complete 4 procedure specific tasks to a predefined proficiency level, 2 relating to ultrasound scanning (utilizing a single haptic device) and 2 relating to needle advancement under ultrasound guidance (concurrently controlling two haptic devices – see Figure 1). Computer generated feedback was given to the subject after each attempted performance of each task. Participants were required to meet proficiency levels on two consecutive attempts before passing each task. In order to complete simulation training the SG participants had to pass all 4 tasks. The tasks were specifically chosen to cover the pre-procedural scout scan and the needling component of USGABPB, while also permitting capture of behaviors likely to lead to significant clinical errors [15]. Table 1 outlines each task, the feedback given and the proficiency level which had to be met. There was no specified time limit to meet these requirements. Subjects were free to control the frequency and duration of use of the simulator. Following initial orientation, training on the simulator in this study was largely unsupervised. An investigator was immediately available to address any technical issues which may have arisen. A flow diagram of the study design is provided (Figure 2). Figures 3, 4, and 5 provide sample images, representative of (i) instructional material, (ii) automated feedback, and (iii) the automated login process.

Assessment

We aimed to assess the performance of the subjects first USGABPB within two weeks of completing the educational interventions. Patients recruited required anesthesia for forearm/wrist/hand surgery where USGABPB would ordinarily be offered as standard care. Informed patient consent was obtained. Study participant performance of USGABPB was closely supervised by a trained regional anesthesiologist. Intravenous sedation was administered as clinically indicated, and subsequent care of the patient may have included general anesthesia. Subjects were asked to perform USGABPB using an in-plane approach and short-axis view. The procedure was video recorded using a handheld video recording device (Flip Ultra, www.theflip.com) in a manner aimed to conceal both the identity of the patient and the identity of the anesthesiologists performing the block. Two experts in UGRA later evaluated the video data.

Patient inclusion and exclusion criteria were

Inclusion criteria: ASA grades I and II, age 18–80 years, capacity to consent, already consented for USGABPB, Body Mass index 20 – 26 kg/m²

Exclusion criteria: Parameters outside inclusion criteria, contraindication to regional anesthesia, language barrier, psychiatric history, pregnancy.

Outcome measures

The subject’s performances were assessed retrospectively based on a task specific, dichotomous, checklist and a behaviorally anchored 5-point global rating scale previously validated for this procedure (see Additional file 1) [16]. Two experts, experienced with this form of evaluation, carried out these assessments. The experts were
Table 1 Task, the feedback given and the proficiency level to be met

| Task                                           | Feedback                                                                 | Proficiency Level             |
|------------------------------------------------|--------------------------------------------------------------------------|-------------------------------|
| 1 Identify the 4 relevant structures represented at a point in the axilla | Number of structures correctly identified | All four structures identified |
| 2 Follow the course of two of these structures (median and ulnar nerves) from axilla towards the elbow, while keeping the structures in the centre of the virtual ultrasound screen | The amount (%) of the structure represented in the middle of the virtual ultrasound as a proportion of the total length of the structure (from axilla to elbow) (out of 100%) | Mean expert performance |
| 3 Advance a virtual needle towards a specified target (median nerve) keeping the needle in plane during advancement | The proportion (%) of needle advancement which occurred “in plane” as a proportion of the total distance the needle tip advanced in the virtual arm | Mean expert performance |
| 4 Trigger a virtual injectate at an appropriate distance from the target. | The distance from the needle tip to the target structure when injection triggered | Injection at a distance not less than the mean expert minimum distance and not more than the mean expert maximum distance. Needle tip must also be visualized at the time of triggering. |

Results

Having originally planned to recruit 20 trainees, this study was discontinued following a planned interim analysis. Ten trainee anesthesiologists were recruited from a university affiliated teaching hospital (Cork University Hospital) in July 2010, 4 to the Simulation group and 6 to the Control group. Recruitment was discontinued because, it became clear that the functionality of the available simulator was insufficient to meet our training requirements. Baseline participant data are summarized in Table 2. The results of visuo-spatial testing using Snowy Picture, Shape Memory and Card Rotation Tests and psychomotor assessment using the Perdue Pegboard are summarized in Table 3. Trainees in the SG did score significantly better in the Shape Memory Test than those in the CG, a measure of visual memory (23.3 (4.6) vs. 12.3 (4.6), p = 0.010). The differences in other visuo-spatial and psychomotor tests were not statistically significant.

Video data corruption occurred during the recording of two participants’ ultrasound-guided axillary brachial plexus blockade, rendering assessment impossible (both in CG). As a result, a comparison of primary and secondary outcome measures involved 8 participants, with 4 in each group (Table 4). There was no statistically significant difference in clinical performance between the groups, as assessed using the sum of the GRS and CHECKLIST scores. There was also no difference in the secondary outcomes measured. No participant completed the performance of the block independently. Data relating to procedural times, number of needle passes and block success/failure were therefore not available. All candidates in both groups were adjudged by expert consensus to have “failed” in their performance of the block.

Participant assessment of content and delivery of the traditional training portion is shown in Table 5. Trainees in the SG rated elements of traditional training higher than CG participants. However, the magnitude of the differences tended to be low.

There was a trend towards a greater interval from commencement of training (traditional training session) to block performance in the Simulation group compared to that in the Control group; however this was not statistically significant [SG 24.5 (16.1) [mean (std dev)], CG 6.5 (6.0) respectively, p = 0.054].
The inter-rater reliability of the assessment of trainee performance by review of video was 89.3% (Range 83.7-93.9%) for checklist scores and 27.8% (Range 0–66.7%) for GRS scores. The Kappa for checklist scores was 0.749 (p < 0.01) indicating a good level of agreement [17], while the Kappa for GRS scores was not statistically significant (Kappa = 0.037, p = 0.628), indicating poor inter-rater reliability [17]. Table 4 compares i) sum of global rating scale plus checklist scores, ii) global rating scale scores, and iii) checklist scores between the two groups. Participant confidence did not differ statistically between the group 2 (2.45), and 2.83 (2.64) [mean (std dev)] in the Simulation and Control Groups respectively (p = 0.587).

Discussion

We have described a methodology for assessing the effectiveness of simulator based training in improving novice trainee’s performance in a clinical setting, by means of a randomized control trial. Our pilot study found no difference in trainee performance between those who underwent standard training and those who received
supplemental simulation-based training. This may have been due to a Type 2 error or to limitations of the prototype simulator (which led to early discontinuation of the trial). We recruited half our a priori defined group of 20 trainees, at one time point (the recruitment of trainee anesthetists in our region occurring biannually). Having trained this cohort, a planned natural point of interim analysis arose. Simulated sono-anatomy was subject to a number of limitations (e.g. clinically relevant muscles/tendons/fat were not modeled), resulting in relevant structures being presented against a relevantly homogenous background. There are two reasons for this; 1. The technical requirements to generate simulated structures, such as biceps or coracobrachialis muscles/tendons, would be significant and were beyond the resources of our team, and 2. The computational requirements to render these secondary structures accurately in real-time, as the user scanned the virtual arm, would be beyond the capacity of the available computer processing units. As a result, it is likely that the simulator allowed for identification of structures in an unrealistic fashion (i.e. lacked fidelity). Indeed, one participant in the SG commented that she would have preferred to attempt to perform the block at an interval closer to the traditional training session, where she had
practiced scanning a real human volunteer. It is likely the simulator had a negative impact in teaching trainees sonoanatomy relevant to USgABPB. It is possible that this diminished any potential improvement in ultrasound-guided needle advancement. Following our interim analysis, we elected to suspend further recruitment as further use of our prototype simulator could have resulted in negative learning which could have negatively impacted on the care of real patients.

The recent Association for Medical Education in Europe (AMEE) Best Evidence in Medical Education (BEME) guide [18], highlighted outcome measures of education as one of the key areas requiring further research. This is the first study to look at the transfer of skills from VR simulation-based training to clinical practice, for an UGRA procedure. In their analysis of VR-based training for laparoscopic surgery, Sinitsky et al. [19] acknowledged that the science of setting proficiency levels is still ill-defined, describing it as “the most pressing issue”. We chose to set proficiency levels based on a limited number of attempts by our group of experts (mean of first three attempts following initial familiarization). Sinitsky et al. [19] also recommended that laparoscopic procedural skills are best learnt through distributed not massed practice. A one day intensive hands-on course on UGRA is an example of massed practice, whereas distributed practice is spread over a greater period of time (shorter practice sessions with long intervals between sessions).

In more general studies of the effectiveness of technology-enhanced learning on medical education, Cook [5,20] also suggests distributed practice is more effective than massed practice. The same authors also found an association between individualized learning and better non-time based skills outcomes [20]. Following the initial familiarization session, trainee’s use of the simulator in this study was self-regulated. As a result, participants could train at a rate which best suited them and was distributed across a number of sessions over a number of days.
Our study is subject to a number of limitations. Firstly, the prototype simulator used was inadequate, in terms of functionality, to meet the training requirements for teaching anesthesiologists, novice in UGRA. Our study sample was small and technical issues with video-recording decreased the size of the dataset acquired further. Inter-rater reliability between experts was poor for GRS scores. This is likely due to the relatively subjective nature of GRS assessment. This may have been improved by enhanced training on using the assessment tools. While inter-rater reliability was good for checklist scores, such tools are subject to a number of limitations. It is possible that an assessment tool which specifically captures clinically relevant errors would be more useful in assessing procedural skills. Such a tool would be particularly useful in providing formative feedback. In the absence of such a validated tool, we chose our primary outcome measure as a combination of GRS and checklist scores. The poor inter-rater reliability of the GRS component raises questions over the validity of our results. There was a difference in training time between the two groups. This difference related to the additional time it took participants in the simulation group to complete simulation training to the predefined proficiency level. It is possible that an improvement in performance in the SG could have been partially attributed to the increased training time, had this occurred. It is also possible that, in this novice population, elements of the traditional training were more important than those enhanced by the simulator training. In particular, when compared to the simulator-generated images, novices appeared overwhelmed by the amount of information they had to interpret in reality. The trend towards an increased interval from the traditional training to block performance in the SG may have had a negative impact on their performance. A number of elements of the traditional training session were rated lower by CG participants than by SG participants. This study does not look at cost of training as the simulator was under development and has no defined monetary value [21]. The simulator described utilizes haptic devices which are costly. Comparisons of haptic and non-haptic based in VR simulation has questioned the need for such devices when training laparoscopic surgical skills [22]. Future studies will need to address this question in training UGRA. Such analysis would have to include both the costs of the devices themselves and also the potential savings (e.g. those associated with faster training, more self-directed learning with lesser requirement for 1:1 teaching with trainers, lesser need for dedicated UGRA courses). The results of this study have informed the iterative development of the simulator. Ultrasound imagery in future prototypes will likely be based on real acquired ultrasound data [23] from which the simulator will be capable of rendering a real-time image. Rosenberg et al. have described mannequin-based simulators for ultrasound-guided regional anesthesia [24]. Based on our experience in this study, we have cautioned that the ultrasound imagery generated not be clinically relevant and could result in negative learning [25].
For the purpose of this study simulation-based training and traditional training were separated into two discrete components with no overlap (to control for possible confounding effects). In reality a more blended approach would be potentially more beneficial. This would likely lessen the negative impact of deficiencies of, for example, an inadequate training tool/simulator.

With increasing computational capacity and reduced cost, it is likely that simulation will move to a more personal environment where supervision is no longer a necessary component to the experience [26]. This may facilitate an individual gaining expertise through self-regulated deliberate practice. However establishing validity of such devices would be essential. The potential for a trainee to learn incorrect or dangerous techniques in an unsupervised simulated environment could have catastrophic results if transferred into the clinical domain [26]. To date, publications of simulation-based training in UGRA have largely been limited to descriptive pieces with few addressing transfer of skills into a clinical setting. In this study we attempted to address this deficit.

Conclusions

We believe that the information acquired during this pilot study will be useful in performing future trials on learning efficacy associated with simulation-based training in procedural skills. In particular, confirmation of a degree of fidelity in the challenges rendered by a simulator is a pre-requisite to carrying out such a study. We believe that failure to do so, could result in spurious results due to factors other than the training or educational value of the simulation-based programme.

### Table 3 Visuo-spatial and psychomotor testing

|                          | Simulation group (n = 4) | Control group (n = 6) | Mann-Whitney’s U-tests |
|--------------------------|-------------------------|-----------------------|------------------------|
| Snowy pictures (mean[std dev]) | 13.3 (5.6) | 10 (4.8) | p = 0.285 |
| Shape memory test       | 23.3 (4.6) | 12.3 (4.6) | p = 0.010* |
| Card rotation test      | 21 (15.3) | 6.67 (10.7) | p = 0.165 |
| Pegboard - Sum Averages Right + Left + Both Hands | 45.1 (8.0) | 43.1 (5.3) | p = 0.522 |
| Pegboard – Assembly     | 35.6 (7.8) | 32.3 (5.7) | p = 0.240 |

Legend: Visuo-spatial testing using Snowy Picture, Shape Memory and Card Rotation Tests (Educational Testing Service) and psychomotor assessment using the Grooved Pegboard (Lafayette Instruments).

* p<0.05.
Additional file

**Additional file 1:** Task Specific Checklist and Global Rating Scale for assessment Ultrasound Guided Axillary Brachial Plexus Block performance.

**Abbreviations**

ABA: American board of anesthesiology; AMEE: Association for medical education in Europe; ASA: American society of anesthesiologists; BEME: Best evidence in medical education; CG: Control group; GRS: Global rating scale; LCD: Liquid crystal display; MOCA: Maintenance of certification in anesthesiology; SG: Simulator group; UGRA: Ultrasound-guided regional anesthesia; USgAPB: Ultrasound-guided axillary brachial plexus block; VR: Virtual reality.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

OOS, GI, ROOD and GDS contributed to the conception and design of the study. OOS and GDS collected and analyzed the data. OOS and GDS drafted the manuscript. GI and ROOD made critical revisions of the manuscript. All authors read and approved the final manuscript.

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**Author details**

1. Department of Anaesthesia and Intensive Care Medicine, Cork University Hospital, Wilton, Cork, Ireland. 2. Department of Anaesthesia and Intensive Care Medicine, University College Cork, Cork, Ireland. 3. ASSERT for Health Centre, University College Cork, Cork, Ireland.

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