A randomised clinical trial of take-home laparoscopic training

Ebbe Thinggaard¹,², Flemming Bjerrum¹,², Jeanett Strandbygaard², Lars Konge¹ & Ismail Gögenur²

ABSTRACT

INTRODUCTION: Simulation-based training in surgery helps trainees master laparoscopic skills, and training at home on mobile box trainers may allow trainees to reach proficiency faster. The aim of this study was to examine the added effects of training at home.

METHODS: Participants were trainees from departments of surgery, gynaecology and urology who were recruited while taking part in a laparoscopic training course. The intervention consisted of added access to a mobile box trainer allowing participants to train at home.

RESULTS: During a one-year study period, 36 participants completed the trial. There was no statistically significant difference in the number of days it took to complete the course (86 days versus 89 days, p = 0.89) or in the final test scores of the two groups (493 versus 460, p = 0.07). A significant difference in the number of training sessions attended was found (5.8 versus 2.3, p < 0.001). Participants were able to reliably rate their own performance; the intraclass correlation coefficient was 0.86, p < 0.001.

CONCLUSIONS: Trainees who had access to training at home did not pass a test earlier or achieve a higher score at the end of a course than trainees who had no such access. Improved access to training at home allowed for shorter and more frequent sessions; however, testing and mandatory training requirements apparently determine training patterns. Trainees were able to reliably rate their own performance.

FUNDING: Equipment for the study was provided by the Copenhagen Academy for Medical Education and Simulation, Capital Region of Denmark.

TRIAL REGISTRATION: The study was exempt from ethical approval according to Danish legislation (H-3-2014-FSP31). The trial protocol was registered with www.clinicaltrials.gov prior to commencing the trial (NCT02243215).

Training on simulators has become part of how we train surgeons. Simulation training has been shown to improve patient outcomes and is a valuable addition to the traditional method of training surgeons at the operating table [1]. Although many surgical trainees and their patients have benefitted from these developments, barriers to simulation training remain. Studies have identified barriers such as access to simulators, time for training and financial constraints [2]. To overcome these barriers, simple mobile box trainers (BT) have been developed, which allow training at home at a time that suits the trainee [3]. Nonetheless, training at home without supervision poses new challenges [4]. Home training of laparoscopic skills has been shown to be feasible [5] However, providing trainees with the freedom to organise their training could change training patterns, allowing for more distributed training where trainees practice more frequently at shorter intervals. A distributed approach to training is beneficial for technical skills acquisition [6], and is also in line with educational principles of deliberate practice [7] and directed self-regulated learning (DSRL) [8]. The purpose of the present study was to examine the added effects of training at home. We looked at the number of training sessions at-home did not pass a test earlier or achieve a higher score at the end of a course than trainees who had no such access. Improved access to training at home allowed for shorter and more frequent sessions; however, testing and mandatory training requirements apparently determine training patterns. Trainees were able to reliably rate their own performance; the intraclass correlation coefficient was 0.86, p < 0.001.

METHODS

Setting

At the Copenhagen Academy for Medical Education and Simulation [9], doctors in speciality training participate in a basic laparoscopic skills training programme during the first year of their training. The course is a cross-speciality training programme for doctors from departments of gynaecology, urology and surgery [10]. The aim of the course is to prepare the course participants for their first supervised laparoscopic surgical procedure. The course consists of two formalised one-day courses separated by a period of self-regulated training on virtual reality simulators (VRS) and BT. The first part of the programme is an introductory course, which includes theoretical teaching imparted as traditional classroom training mixed with practical sessions to prepare the trainees for training on VRS and BT. After the introduction course, the participants go through a period of self-regulated training during which they book training sessions at the simulation centre and practice on both VRS and BT. At the simulation centre, they are assisted by a simulator technician who is able to give technical assistance and provide
Participants

The course participants consisted of doctors in the first year of their specialty training. Participants who had performed more than fifty laparoscopic procedures were excluded.

Intervention

The intervention consisted of the addition of home training on a mobile BT. The intervention group trained at the simulation centre and were also given a portable BT [12] allowing them to practice at home. The control group trained at the simulation centre only. Both groups had access to training on VRS at the simulation centre.

Randomisation

The primary investigator (ET) was responsible for inclusion of participants. After enrolment, participants were randomly allocated using a computer-generated allocation sequence (randomiser). The administrator at the simulation centre retrieved the allocation sequence and kept the sequence concealed until the allocation had been finalised.

Outcomes

All participants were given a training log to record their training. Based on information from the logbooks, we looked at the number of days from enrolment to passing the TABLT test, the time spent training and the number of training sessions attended. We also explored differences in the performance levels that participants reached on their final TABLT test and recorded the participants’ ability to rate themselves.

Statistical analysis

The sample size for the trial was calculated based on the assumption that the control group would pass the TABLT test after six weeks of practice (42 days), standard deviation (SD) ± 3 weeks (± 21 days). The intervention group was expected to pass after four weeks of practice (28 days), SD ± 3 weeks (± 21 days). Setting alpha at 0.05 and beta to 0.10, a total of 24 participants were required in each group. The trial was planned with a one-year inclusion period. Accounting for inaccuracies, we expected to include a total of 50 participants in the trial during the one-year study period during which six courses were planned with up to 72 course places. We used student’s t-test to analyse whether there was a significant level of difference in the above-mentioned measurements. A p-value below 0.05 was considered statistically significant for the primary outcome. To determine the reliability of the self-rated test, we compared participants’ ratings of their pre-test and the rating of a trained blinded rater. The intraclass correlation coefficient (ICC) was used to ex-
amine the reliability of the participants’ self-rating. A statistical software package was used (SPSS vs. 20.0, Chicago, IL).

**Trial registration:** The trial was submitted for evaluation to the Regional Ethics Committee, which determined that no approval was needed for the trial (H-3-2014-FSP31). The trial was also registered with clinicaltrials.gov prior to its commencement (NCT02243215), and it was conducted according to the CONSORT statement [13].

**RESULTS**

We included participants during a one-year period in which 50 doctors participated in the training course. Out of the 50 participants who took part in the course, 46 were enrolled in the study and 36 completed the course within the one-year study period. Four participants dropped out of the training course, and six participants were excluded from the study as they did not complete the training course during the one-year study period. Out of the 36 who completed the course, 18 were from the control group, and 18 were from the intervention group, see **Figure 1**. For the participants’ baseline characteristics, see **Table 1**. At the end of the one-year study period, we performed a new sample size calculation based on data available from the 36 participants, corresponding to 75% of the anticipated sample size. We found that 11,422 participants would be needed in each group which was not feasible, and therefore we decided to stop recruiting participants. We found no difference in the number of days from enrolment to the passing of the TABLT test (86 days versus 89 days, p = 0.89), time spent training on box trainers (302 minutes versus 218 minutes, p = 0.26) or between the test score (493 versus 460, p = 0.07) (**Table 2**). However, we did find a significant difference in the number of training sessions (5.8 versus 2.3, p < 0.001), see **Table 2**. There was a good reliability when comparing participants’ ratings of their pre-test and that of a blinded rater, ICC 0.86, p < 0.001.

**DISCUSSION**

In this study, we explored the added effect of training laparoscopic skills at home and found no difference in the number of days or in the time spent training to pass the TABLT test. However, we did find a significant difference in the number of training sessions (5.8 versus 2.3, p < 0.001), see **Table 2**. There was a good reliability when comparing participants’ ratings of their pre-test and that of a blinded rater, ICC 0.86, p < 0.001.

**TABLE 1**

**Participants’ baseline characteristics.**

<table>
<thead>
<tr>
<th></th>
<th>Intervention group</th>
<th>Control group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants, n</td>
<td>18</td>
<td>18</td>
<td>36</td>
</tr>
<tr>
<td>Age, yrs, median (range)</td>
<td>30 (25-36)</td>
<td>30 (25-46)</td>
<td>30 (25-46)</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Men/women</td>
<td>5/13</td>
<td>5/13</td>
<td>10/26</td>
</tr>
<tr>
<td>Speciality</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Surgery</td>
<td>6</td>
<td>5</td>
<td>11</td>
</tr>
<tr>
<td>Urology</td>
<td>3</td>
<td>3</td>
<td>6</td>
</tr>
<tr>
<td>Gynaecology</td>
<td>9</td>
<td>10</td>
<td>19</td>
</tr>
<tr>
<td>Dominant hand, n</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right/ left</td>
<td>16/2</td>
<td>16/2</td>
<td>32/4</td>
</tr>
</tbody>
</table>

**TABLE 2**

**Training on box trainers.**

<table>
<thead>
<tr>
<th></th>
<th>Group, mean (95%CI)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Intervention</td>
<td>control</td>
</tr>
<tr>
<td>Time to complete the course, days</td>
<td>86 (52-120)</td>
<td>89 (52-127)</td>
</tr>
<tr>
<td>Time spent training, min.</td>
<td>302 (189-414)</td>
<td>218 (112-223)</td>
</tr>
<tr>
<td>Training sessions, n</td>
<td>5.8 (4-7.5)</td>
<td>2.3 (1.5-3.1)</td>
</tr>
<tr>
<td>Final TABLT test score</td>
<td>493 (465-522)</td>
<td>460 (434-485)</td>
</tr>
</tbody>
</table>

CI = confidence interval; TABLT = Training and Assessment of Basic Laparoscopic Techniques.
Laparoscopic virtual reality simulator training [15], and learning curves, in particular, have been shown to improve by using distributed training compared with massed training [16]. Even though the ideal training interval for laparoscopic simulation training has not been established, it has been shown that short training intervals are superior to long training intervals [17]. The fact that participants with access to training at home did not reach a higher level on the test might be explained by the fact that they were instructed on how to rate their own performance during training. Therefore, they knew that it made sense for them to stop training when they reached a sufficient performance level. However, this was a deliberate choice of training strategy. Being able to rate your own performance allows for a more independent approach to training and has emerged from the instructional method called DSRL [8, 18], which is recommended for simulation training [19]. Principles of DSRL have shown to be useful in VRS mastoidectomy training [20]. This approach may also be of great value for training of laparoscopic skills at home. When considering unsupervised laparoscopic skills training at home, using DSRL as a strategy would allow for a structured training programme where trainees are in control of their training. In the present study, we showed that participants could reliably rate their own performance on the TABLT test. Being able to reliably rate your own test allows trainees to monitor their own training and provides them with a tool to apply self-regulatory skills.

**Limitations**

In this study, we chose to investigate the added effect of training at home on a simple mobile BT while also training in a simulation centre. As we did not wish to limit the participants’ access to training, it was not possible to compare the effect of only using home-based training with that of training only at a simulation centre. Having chosen a different design could have given us insight into the effects of training at home versus training at a simulation centre. However, this was beyond the scope of our study. In our sample size calculation, we used a beta of 0.10. Having chosen a beta of 0.20 might have allowed for our inclusion of participants to match that of our sample size calculation. In our training programme, we use both VRS and BT; mixing two training methods could cloud findings. A trial focusing on BT exclusively might have more clearly demonstrated potential benefits of training at home using a BT. However, examining the use of training at home as a supplement was a deliberate choice of study design. We chose to do the study under realistic circumstances as part of an existing laparoscopic training programme. The results of our study could help guide others that may consider incorporating take-home training in their laparoscopic training course. In the basic laparoscopy course, we also use a cross-specialty approach to laparoscopic training where doctors from different specialities practice together. Having participants from different specialities and with different levels of experience may have had an impact on the results. Using participants from different specialities increases the external validity as findings can be generalised across training programmes for different specialities. The participants in our study had different levels of experience prior to commencing the training programme. This makes the results of the trial applicable to trainees with different degrees of experience.

**CONCLUSIONS**

Take-home training of basic laparoscopic skills on a mobile box trainer allowed trainees to practice at their own convenience. The increased access to training did not result in trainees passing a test earlier or getting a higher score, but they did engage in shorter and more frequent training sessions. Testing and mandatory training requirements apparently determine training patterns. Trainees could reliably rate their own performance.

**CORRESPONDENCE:** Ebbe Thinggaard.
E-mail: ebbe.thinggaard@regionh.dk

**ACCEPTED:** 12 October 2018

**CONFLICTS OF INTEREST:** Disclosure forms provided by the authors are available with the full text of this article at Ugeskriftet.dk/dmj

**LITERATURE**


