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A randomised controlled trial


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Virtual reality exposure before elective day care surgery to reduce anxiety and pain in children

A randomised controlled trial

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BACKGROUND Pre-operative anxiety in children is very common and is associated with adverse outcomes.

OBJECTIVE The aim of this study was to investigate if virtual reality exposure (VRE) as a preparation tool for elective day care surgery in children is associated with lower levels of anxiety, pain and emergence delirium compared with a control group receiving care as usual (CAU).

DESIGN A randomised controlled single-blind trial.

SETTING A single university children’s hospital in the Netherlands from March 2017 to October 2018.

PATIENTS Two-hundred children, 4 to 12 years old, undergoing elective day care surgery under general anaesthesia.

INTERVENTION On the day of surgery, children receiving VRE were exposed to a realistic child-friendly immersive virtual version of the operating theatre, so that they could get accustomed to the environment and general anaesthesia procedures.

MAIN OUTCOME MEASURES The primary outcome was anxiety during induction of anaesthesia (modified Yale Pre-operative Anxiety Scale, mYPAS). Secondary outcomes were self-reported anxiety, self-reported and observed pain, emergence delirium, need for rescue analgesia (morphine) and parental anxiety.

RESULTS A total of 191 children were included in the analysis. During induction of anaesthesia, mYPAS levels (median [IQR] were similar in VRE, 40.0 [28.3 to 58.3] and CAU, 38.3 [28.3 to 53.3]; P = 0.862). No differences between groups were found in self-reported anxiety, pain, emergence delirium or parental anxiety. However, after adenoidectomy/tonsillectomy, children in the VRE condition needed rescue analgesia significantly less often (55.0%) than in the CAU condition (95.7%) (P = 0.002).

CONCLUSION In children undergoing elective day care surgery, VRE did not have a beneficial effect on anxiety, pain, emergence delirium or parental anxiety. However, after more painful surgery, children in the VRE group needed rescue analgesia significantly less often, a clinically important finding because of the side effects associated with analgesic drugs. Options for future research are to include children with higher levels of anxiety and pain and to examine the timing and duration of VRE.

TRIAL REGISTRATION Netherlands Trial Registry: NTR6116.

Published online 25 July 2019
Introduction

Pre-operative anxiety is very common in children. On the day of surgery, 50 to 70% of children experience anxiety that usually peaks during induction of anaesthesia. Pre-operative anxiety is associated with problematic induction of anaesthesia, risk of emergence delirium, increased pain and poorer recovery. Anxious children undergoing surgery, and their parents, are also at risk of posttraumatic stress symptoms. These adverse outcomes underscore the urgent need for effective interventions to reduce pre-operative anxiety.

A promising innovative intervention is virtual reality. Virtual reality is especially engaging for children, as they often become truly captivated by imaginative play. In our recent meta-analysis on virtual reality interventions in children undergoing medical procedures, we found that virtual reality is effective in reducing anxiety and pain. In most studies, virtual reality was used as a distraction tool during medical procedures. However, research has demonstrated that exposure is more effective than distraction in reducing anxiety. Virtual reality exposure (VRE) has already been proven effective in treating anxiety disorders, such as specific phobias (fear of spiders), but very limited research has been conducted on the effect of VRE as preparation for medical procedures.

VRE offers the chance to reduce pre-operative anxiety by exposing children to a realistic version of the operating theatre, in which they can get accustomed to the environment and procedures associated with anaesthesia. Until now, only two studies have applied VRE prior to surgery. In these studies, both including 69 children, the intervention took place on the day of surgery and consisted of either a 360° virtual reality tour of the operating theatre or a virtual reality game in which patients experienced the pre-operative process. Children in the control group received conventional education about the pre-operative process. Both studies were limited to pre-operative outcomes and found that children were significantly less anxious and more compliant in the VRE than in the control group. However, as pre-operative anxiety is associated with negative postoperative outcomes, such as increased pain and emergence delirium, it is also important to investigate postoperative effects of VRE.

This is the first randomised controlled trial (RCT) to study the effects of VRE on pre-operative anxiety, in addition to postoperative outcomes. The objective of the current study was to compare levels of anxiety during induction of anaesthesia (primary outcome), postoperative anxiety, pain, emergence delirium, rescue analgesia and parental anxiety (secondary outcomes) in children receiving VRE, with controls, and to identify predictors of VRE efficacy in children 4 to 12 years old undergoing elective maxillofacial, dental or ear-nose-throat (ENT) day surgery.

Materials and methods

The PREoperative Virtual reality Intervention to Enhance Wellbeing (PREVIEW) study was approved by the Medical Ethics Committee of the Erasmus Medical Centre (MEC-2016-626) on 30 November 2016 and registered at the Netherlands Trial Registry (NTR6116). This single-centre, single-blinded RCT was conducted in accordance with the CONSORT guidelines at the Erasmus MC-Sophia Children’s Hospital in the Netherlands, by the Departments of Child and Adolescent Psychiatry/Psychology, Paediatric Anaesthesiology, Maxillofacial, Dental and ENT Surgery. Written informed consent was obtained from all parents and from all children aged 12 years. Children aged 11 years and under gave permission orally.

Participants

Eligible participants were consecutive children aged 4 to 12 years undergoing elective maxillofacial, dental or ENT day care surgery between March 2017 and October 2018. Exclusion criteria were mental retardation, inability of parents to read or write Dutch, epilepsy, visual impairment, an American Society of Anaesthesiologists (ASA) physical status at least III and need for pre-operative anxiolytic medication.

Procedure

Eligible children and their parents were informed about the study by paediatric anaesthesiologists during pre-operative screening. Those interested received a patient information folder via e-mail. During this screening, the anaesthesiologists recommended that all children and parents should watch an informative online film at home about general anaesthesia according to the standard hospital protocol.

On the day of surgery, after informed consent was obtained, personal and medical data were collected by the research assistant and baseline anxiety and problem behaviour were assessed (T1) (Fig. 1). Next, the research assistant randomly allocated children to the VRE intervention, which the children received together with usual care, or to the control group, in which children only received care as usual (CAU). Block randomisation was performed, stratified by type of surgery: adenoidecomy and/or tonsillectomy, insertion of tympanostomy tubes, maxillofacial and dental procedures or other ENT procedures. After randomisation, the VRE intervention took place in a separate room, under the guidance of the research assistant. Afterwards, children were admitted to the day care unit. Children in the CAU group were admitted to the day care unit directly after randomisation. Assessments after randomisation were performed by the blinded researcher (RE) in the holding area (T2) and during induction of anaesthesia (T3), and by a blinded recovery nurse, postoperatively, in the recovery room (T4). One assessment per time point was
performed. In the recovery room, assessments took place during later phases of awakening. Parents were present from early phases of awakening (one parent per child). Anaesthesiologists were blinded to group allocation. Postoperatively parents were encouraged and reminded by phone and e-mail to complete online questionnaires at home, via a secure website, on the third day after surgery, but were allowed to do so until 2 weeks after surgery (T5). An overview of the process can be found in Fig. 1.

**Virtual reality intervention**

The VRE tool encompasses a highly realistic virtual environment that is modelled according to the real operating theatre and medical staff. The virtual environment is computer-generated, interactive and child-friendly. It was presented to the child for approximately 15 min via an HTC Vive (HTC Corporation, Xindian, New Taipei, Taiwan) head-mounted display and was also displayed on a personal computer monitor, so the accompanying parent could see what the child was viewing. We developed two versions, for children aged 4 to 7 and 8 to 12 years, in order to attune explanations to a child’s developmental level. The storyline begins in the holding area (Fig. 2a). A receptionist welcomes the child and shows a video on a virtual tablet that explains that one of the child’s parents will stay with him/her until the child is anaesthetised and shows the hospital gowns they will be wearing to the operating room. Next, the child is transported, in a hospital bed, into the corridor of the operating room, by an anaesthesiologist and a nurse anaesthetist (Fig. 2b). After arrival in the operating room, the child can point at different instruments with a motion tracked controller so that the nurse anaesthetist can explain what these are used for (Fig. 2c). Then, the child moves onto the operating table and preparation for anaesthesia takes place. The programme is able to show both intravenous (i.v.) and inhalational induction. After induction, the operating room fades out and the recovery room fades in (Fig. 2d). Here, the nurse anaesthetist shows another video that
explains what kind of feelings the child might experience after surgery, for example nausea. For a more detailed overview of the storyline, as well as the technical hardware and software specifications, we refer to the trial article.¹⁶

**Anaesthesia protocol**

None of the children received pre-operative anxiolytic premedication. EMLA cream (lidocaine/prilocaine) or Rapydan (lidocaine/tetracaine) plasters were applied on the back of the hands, 30 to 60 min before going to the operating room. Induction of anaesthesia took place in the operating room in the presence of a parent or a guardian. Children were lying down or sitting on the operating table but were also permitted to sit on the parent’s lap. After placement of the electrocardiography electrodes, pulse oximeter and blood pressure cuff, anaesthesia was induced, i.v. or by inhalation if i.v. cannulation was declined or i.v. access was unsuccessful. For i.v. induction, a peripheral i.v. catheter was placed in the back of the hand, and i.v. propofol 2 to 4 mg kg⁻¹ and fentanyl 1 to 2 μg kg⁻¹ were administered. For inhalation induction, sevoflurane in a mixture of oxygen and air was administered by mask. In these cases, i.v. cannulation took place after induction, after which i.v. fentanyl 1 to 2 μg kg⁻¹ was administered. Depending on the surgical procedure, a laryngeal mask airway (LMA) or an endotracheal tube (ETT) was inserted. Before intubation, the child received a muscle relaxant. Anaesthesia was maintained with sevoflurane 0.7 to 1.0 minimal alveolar concentration (MAC) in air and oxygen. During surgery, i.v. fentanyl was administered at the discretion of the anaesthesiologist. At the end of the procedure, first doses of i.v. paracetamol 20 mg kg⁻¹ and diclofenac 1 mg kg⁻¹ were administered. If needed, i.v. morphine 0.1 mg kg⁻¹ was also administered. After extubation, children were brought to the recovery area. Rescue analgesia, extra morphine, could be administered by the recovery nurse according to perceived clinical need. Standard postoperative analgesics were prescribed: paracetamol 90 mg kg⁻¹ per day orally or rectally and diclofenac 3 mg kg⁻¹ per day orally or rectally.

**Assessment instruments**

An overview of the well validated assessment instruments at each time point is provided in Fig. 1.

**Child anxiety**

The primary outcome was child anxiety during induction of anaesthesia (T3) assessed with the modified Yale Preoperative Anxiety Scale (mYPAS).¹⁸ The mYPAS is considered the gold standard in observational instruments to assess pre-operative anxiety in children¹⁸ and was completed at three timepoints (T1, T2 and T3). The mYPAS consists of 27 items divided into five domains: activity, emotional expressivity, state of arousal, vocalisation and use of parents. Scores range from 23.33 to 100, with higher scores indicating higher levels of anxiety. The domains have good to excellent interobserver and intra-observer reliability.¹⁸ The research assistant (T1) and blinded researcher (T2 & T3) were trained in administering the mYPAS with standardised instructions. Children indicated their own anxiety level on a visual

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**Fig. 2**

Virtual reality environment. (a) The receptionist welcomes the child to the holding area. (b) The operating room, where the child receives information about different instruments (pulse oximeter, blood pressure cuff and anaesthesia mask). (c) The child wakes up in the recovery room.
analogue scale (VAS)\textsuperscript{19} prior to anaesthesia and after surgery (T1, T2, T4 and T5).

**Child pain and emergence delirium**

Postoperative pain was reported by three informants. Children reported their pain intensity (T4 and T5) with the six-faces revised Faces Pain Scale (FPS-r): range 0 to 10.\textsuperscript{20} A blinded recovery nurse assessed pain intensity (T4) with the Face, Legs, Activity, Cry and Consolability (FLACC) scale: range 0 to 10.\textsuperscript{21} Parents assessed their child’s pain (T5) by completing the Parents’ Postoperative Pain Measure (PPPM): range 0 to 15.\textsuperscript{22} Emergence delirium was assessed (T4) with the Paediatric Anaesthesia Emergency Delirium (PAED) scale by a blinded recovery nurse: range 0 to 20.\textsuperscript{23}

**Child behaviour problems**

At T1, parents completed the Child Behaviour Checklist (CBCL) to assess pre-operative emotional and behavioural problems during the past 6 months.\textsuperscript{24,25} Either the 1.5 to 5 years of age version with 100 items (for 4 to 5-year-old participants) or the 6 to 18 years of age version with 113 items (for 6 to 12-year-old participants) was used. T-scores for total scores were computed.

**Parental anxiety**

The State–Trait Anxiety Inventory (STAI) is a self-reporting instrument that contains two separate scales for trait and state anxiety.\textsuperscript{26} Scores on both Likert-type scales range from 20 to 80. Parents completed the state form directly after induction of anaesthesia (T3).

**Statistical analyses**

Intention-to-treat (ITT) analyses were performed for all randomised participants. We used two-way imputation to adjust for missing item data. The Shapiro–Wilk test was used to test the assumption of normal distribution. Non-normally distributed continuous variables were compared between conditions using the Mann–Whitney U test. Categorical variables were analysed with the Eijlers et al.\textsuperscript{732} scale: range 0 to 10.21 Parents assessed their child with the Face, Legs, Activity, Cry and Consolability (FLACC) scale: range 0 to 10.\textsuperscript{21} Parents assessed their child’s pain (T5) by completing the Parents’ Postoperative Pain Measure (PPPM): range 0 to 15.\textsuperscript{22} Emergence delirium was assessed (T4) with the Paediatric Anaesthesia Emergency Delirium (PAED) scale by a blinded recovery nurse: range 0 to 20.\textsuperscript{23}

**Sample size calculation**

A sample size of 100 patients per group was sufficient to compare the primary outcome, anxiety during induction of anaesthesia (mYPAS) between the intervention and control groups, with a Cohen’s d of 0.4 (small to medium effect size), an alpha of 0.05 (two-tailed) and a power of 0.85. A sample size of 100 patients in the intervention group was sufficient to perform regression analyses with six predictor variables, a small to medium effect size and a power of 0.85.

**Results**

No significant differences were found between groups in patient and surgical characteristics (all P > 0.05).

Between March 2017 and October 2018, 393 children were assessed for eligibility, of whom 193 children did not participate. Reasons for non participation were did not meet the inclusion criteria (n = 35), did not want to participate (n = 109) or for other reasons such as inability to contact, postponement or cancellation of surgery before data collection (n = 49) (Fig. 3). Two hundred children were enrolled in the study (VRE: n = 100, CAU: n = 100). Nine children were excluded because of accidental unblinding (n = 5), noncompliance with the anaesthetic protocol (n = 2), no data collection at T2 and T3, due to logistical reasons (n = 1), or cancelled surgery (n = 1). Therefore, 191 participants were included in the data analyses (VRE: n = 94, CAU: n = 97). Baseline characteristics of all participants are given in Table 1. Twenty-one children in the VRE condition discontinued the intervention by taking off the virtual reality headset. Ad-hoc analyses showed that this group consisted of an equal number of boys (n = 11, 52.4%) and girls (n = 10, 47.6%), with a median age of 5.0 years [4.5 to 6.3]. More specifically, 71.4% of these children were 4 or 5 years old.

For the data collection at T5, most parents (56.0%) had completed the online questionnaires on the fourth day after surgery. On the eighth day after surgery, almost all parents (82.7%) had completed the questionnaires. The final percentage of completed questionnaires after 14 days was 91.6%. No significant correlations (Spearman’s ρ) were found between day of completion and postoperative outcomes (P = 0.228, P = 0.577, and P = 0.721, for VAS, FPS-r and PPPM, respectively). Therefore, T5 data from all postoperative days (3 to 14) were combined and included in the analysis.

**Child anxiety**

At baseline (T1), in the holding area (T2) and during induction of anaesthesia (T3), mYPAS scores were similar in CAU and VRE (P = 0.697, P = 0.765 and P = 0.862, respectively). Self-reported VAS scores were also comparable between conditions at different time points (P = 0.407 at T1, P = 0.753 at T2, P = 0.735 at T4 and P = 0.727 at T5) (Table 2).
VR exposure before paediatric day care surgery

**Fig. 3**

**Excluded** (n = 193)
- Not meeting inclusion criteria (n = 35)
- Declined to participate (n = 109)
- Other reasons, e.g. unable to contact or surgery postponed or cancelled before data collection (n = 49)

**Allocated to VRE intervention** (n = 100)
- Received allocated intervention (n = 100)
- Discontinued intervention (child wanted to take off VR headset) (n = 23)

**Allocated to CAU** (n = 100)

**Analysed** (n = 97)
- Excluded from analysis, due to accidental unblinding (n = 1); non-compliance with anaesthetic protocol (n = 1); no data collection at T2 and T3 (n = 1)

**Table 1** Patient and surgical characteristics

<table>
<thead>
<tr>
<th></th>
<th>VRE (n = 94)</th>
<th>CAU (n = 97)</th>
<th>P (^a)</th>
<th>VRE (n = 73)</th>
<th>P (^f)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>8.3 [5.7 to 10.2]</td>
<td>7.5 [5.6 to 10.7]</td>
<td>0.938 (^a)</td>
<td>9.0 [6.4 to 10.7]</td>
<td>0.064 (^a)</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>45 (47.9)</td>
<td>56 (57.7)</td>
<td>0.172 (^b)</td>
<td>34 (46.6)</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>49 (52.1)</td>
<td>41 (42.3)</td>
<td>0.141 (^b)</td>
<td>39 (53.4)</td>
<td>0.073 (^b)</td>
</tr>
<tr>
<td>ASA physical status</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>72 (76.6)</td>
<td>65 (67.0)</td>
<td>0.915 (^b)</td>
<td>58 (79.5)</td>
<td>0.477 (^b)</td>
</tr>
<tr>
<td>II</td>
<td>22 (23.4)</td>
<td>32 (33.0)</td>
<td></td>
<td>15 (20.5)</td>
<td></td>
</tr>
<tr>
<td>Type of surgery</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adenoidectomy and/or tonsillectomy</td>
<td>20 (21.3)</td>
<td>23 (23.7)</td>
<td>0.499 (^b)</td>
<td>12 (16.4)</td>
<td>0.674 (^b)</td>
</tr>
<tr>
<td>Tympanostomy tubes</td>
<td>23 (24.5)</td>
<td>23 (23.7)</td>
<td></td>
<td>17 (23.3)</td>
<td></td>
</tr>
<tr>
<td>Maxillofacial and dental procedures</td>
<td>23 (24.5)</td>
<td>26 (26.8)</td>
<td></td>
<td>18 (24.7)</td>
<td></td>
</tr>
<tr>
<td>Other ENT procedures</td>
<td>28 (29.8)</td>
<td>25 (25.8)</td>
<td></td>
<td>26 (35.6)</td>
<td></td>
</tr>
<tr>
<td>Induction method</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inhalation</td>
<td>54 (57.4)</td>
<td>51 (52.6)</td>
<td></td>
<td>36 (49.3)</td>
<td></td>
</tr>
<tr>
<td>Intravenously</td>
<td>40 (42.6)</td>
<td>46 (47.4)</td>
<td></td>
<td>37 (50.7)</td>
<td></td>
</tr>
<tr>
<td>Total problem behaviour (CBCL)</td>
<td>47.0 [41.0 to 56.0]</td>
<td>46.0 [39.0 to 53.0]</td>
<td>0.251 (^a)</td>
<td>47.0 [41.0 to 55.0]</td>
<td>0.268 (^a)</td>
</tr>
<tr>
<td>Parental education level</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>5 (5.3)</td>
<td>2 (2.1)</td>
<td></td>
<td>3 (4.1)</td>
<td></td>
</tr>
<tr>
<td>Medium</td>
<td>30 (31.9)</td>
<td>35 (36.1)</td>
<td></td>
<td>25 (34.2)</td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>59 (62.8)</td>
<td>60 (61.9)</td>
<td></td>
<td>45 (61.6)</td>
<td></td>
</tr>
</tbody>
</table>

Values are median [interquartile range] or frequency (percentage); ASA, American Society of Anaesthesiologists; CAU, care as usual; CBCL, Child Behaviour Checklist; VRE, virtual reality exposure. "Mann–Whitney U-test. "χ² test. "Children who were allocated to the VRE condition (n = 94). "Intention-to-treat analyses. "Children who completed the VRE intervention. "Per-protocol analyses (ad-hoc).
The only significant predictor of anxiety during induction of anaesthesia was pre-operative parental state anxiety $[F(1,85) = 5.05, \ P = 0.027]$. Higher parent anxiety levels prior to surgery were related to higher child anxiety levels during induction, in the VRE group. The linear regression model accounted for 11.3% of the variance in anxiety during induction of anaesthesia $[F(6,85) = 1.80, \ P = 0.109]$.

Child pain and emergence delirium

No differences in pain levels were found between VRE and CAU, neither when self-reported with FPS-r ($P = 0.699$ at T4, $P = 0.454$ at T5), nurse-observed with FLACC ($P = 0.669$) nor parent-observed with PPPM ($P = 0.410$). Further investigation of pain levels in the recovery room (T4) indicated that there were no differences between VRE and CAU in the proportion of children experiencing considerable levels of pain (FPS-r > 3 or FLACC > 3$^{27,28}$) when self-reported with the FPS-r (VRE: $n = 24$, CAU: $n = 23$, $P = 0.211$), nor when nurse-observed with the FLACC (VRE: $n = 4$, CAU: $n = 5$, $P = 0.549$). No differences were found in emergence delirium symptoms between conditions ($P = 0.266$), nor in proportion of children experiencing considerable levels of emergence delirium symptoms (PAED > 10)$^{23}$ (VRE: $n = 3$, CAU: $n = 1$, $P = 0.505$).

No significant predictors of postoperative pain were found in the model. The linear regression model accounted for 7.4% of the variance in self-reported pain in the recovery room, $F(6,81) = 1.07, \ P = 0.386$.

Rescue analgesia

Overall, there was no difference in need for rescue analgesia between VRE and CAU ($P = 0.131$). When analysing rescue analgesia for each type of surgery separately, 11 out of 20 (55.0%) children in the VRE group who underwent adenoidectomy and tonsillectomy needed significantly less frequent rescue analgesia than the 22 out of 23 (95.7%) children in the CAU group ($\chi^2 = 9.91$, $P = 0.002$). No differences in rescue analgesia were found for the other three types of surgery.

Parental anxiety

No differences in parental anxiety during induction of anaesthesia were found between groups, either when self-reported (STAI-state) ($P = 0.753$), or when observed (VAS) ($P = 0.418$).

Ad-hoc analyses

We did not replace the 21 children who discontinued the VRE intervention, in line with intention-to-treat principles. However, because this concerns a substantial

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**Table 2** Anxiety, pain, rescue analgesia and emergence delirium levels, and parental anxiety levels in both groups

<table>
<thead>
<tr>
<th>Anxiety</th>
<th>VRE (n = 94)</th>
<th>CAU (n = 97)</th>
<th>P $^a$</th>
<th>VRE (n = 73)</th>
<th>P $^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>mYPAS (observed)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T1</td>
<td>28.3 [23.3 to 31.7]</td>
<td>26.7 [23.3 to 32.5]</td>
<td>0.697$^{a}$</td>
<td>28.3 [23.3 to 30.0]</td>
<td>0.636$^{a}$</td>
</tr>
<tr>
<td>T2</td>
<td>28.3 [23.3 to 36.7]</td>
<td>28.3 [23.3 to 41.7]</td>
<td>0.765$^{a}$</td>
<td>26.7 [23.3 to 36.7]</td>
<td>0.129$^{a}$</td>
</tr>
<tr>
<td>T3</td>
<td>40.0 [28.3 to 58.3]</td>
<td>38.3 [28.3 to 53.3]</td>
<td>0.862$^{a}$</td>
<td>36.7 [27.5 to 48.3]</td>
<td>0.266$^{a}$</td>
</tr>
<tr>
<td>VAS (self-reported)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T1</td>
<td>3.0 [0.1 to 5.0]</td>
<td>1.5 [0.0 to 5.0]</td>
<td>0.407$^{a}$</td>
<td>3.0 [0.5 to 5.0]</td>
<td>0.209$^{a}$</td>
</tr>
<tr>
<td>T2</td>
<td>3.0 [1.0 to 5.8]</td>
<td>3.5 [0.0 to 6.0]</td>
<td>0.753$^{a}$</td>
<td>3.5 [1.0 to 6.0]</td>
<td>0.999$^{a}$</td>
</tr>
<tr>
<td>T4</td>
<td>0.0 [0.0 to 2.0]</td>
<td>0.0 [0.0 to 2.0]</td>
<td>0.735$^{a}$</td>
<td>0.3 [0.0 to 2.0]</td>
<td>0.466$^{a}$</td>
</tr>
<tr>
<td>T5</td>
<td>0.5 [0.0 to 1.0]</td>
<td>0.3 [0.0 to 2.0]</td>
<td>0.727$^{a}$</td>
<td>0.5 [0.0 to 1.0]</td>
<td>0.620$^{a}$</td>
</tr>
<tr>
<td>Pain</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FPS-r (self-reported)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T4</td>
<td>2.0 [0.0 to 4.0]</td>
<td>2.0 [0.0 to 2.5]</td>
<td>0.699$^{a}$</td>
<td>2.0 [0.0 to 4.0]</td>
<td>0.769$^{a}$</td>
</tr>
<tr>
<td>T5</td>
<td>0.0 [0.0 to 2.0]</td>
<td>0.0 [0.0 to 2.0]</td>
<td>0.484$^{a}$</td>
<td>0.0 [0.0 to 2.0]</td>
<td>0.551$^{a}$</td>
</tr>
<tr>
<td>FLACC (observed, T4)</td>
<td>0.0 [0.0 to 0.0]</td>
<td>0.0 [0.0 to 0.0]</td>
<td>0.669$^{a}$</td>
<td>0.0 [0.0 to 0.0]</td>
<td>0.739$^{a}$</td>
</tr>
<tr>
<td>PPPM (observed, T5)</td>
<td>3.0 [0.0 to 5.0]</td>
<td>3.0 [1.0 to 8.0]</td>
<td>0.410$^{a}$</td>
<td>3.0 [0.0 to 5.8]</td>
<td>0.502$^{a}$</td>
</tr>
<tr>
<td>Rescue analgesia$^b$ (T4)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overall</td>
<td>28.9 (28.8)</td>
<td>39 (40.2)</td>
<td>0.131$^{b}$</td>
<td>22 (30.1)</td>
<td>0.175$^{b}$</td>
</tr>
<tr>
<td>Adenoidectomy and tonsillectomy</td>
<td>11 (55.0)</td>
<td>22 (95.7)</td>
<td>0.002$^{b}$</td>
<td>6 (50.0)</td>
<td>0.001$^{b}$</td>
</tr>
<tr>
<td>Tympanostomy tubes</td>
<td>0 (0.0)</td>
<td>1 (4.3)</td>
<td>0.31$^{b}$</td>
<td>0 (0.0)</td>
<td>0.384$^{b}$</td>
</tr>
<tr>
<td>Maxillofacial and dental procedures</td>
<td>4 (17.4)</td>
<td>6 (23.1)</td>
<td>0.622$^{b}$</td>
<td>4 (22.2)</td>
<td>0.947$^{b}$</td>
</tr>
<tr>
<td>Other ENT procedures</td>
<td>13 (46.4)</td>
<td>10 (40.0)</td>
<td>0.637$^{b}$</td>
<td>12 (22.2)</td>
<td>0.657$^{b}$</td>
</tr>
<tr>
<td>Emergence delirium</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PAED (observed, T4)</td>
<td>7.0 [5.0 to 9.0]</td>
<td>6.0 [5.0 to 9.0]</td>
<td>0.266$^{b}$</td>
<td>7.5 [5.0 to 9.0]</td>
<td>0.223$^{b}$</td>
</tr>
<tr>
<td>Parental anxiety</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>STAI-state (self-reported, T3)</td>
<td>41.0 [34.5 to 48.5]</td>
<td>40.5 [33.0 to 50.0]</td>
<td>0.753$^{b}$</td>
<td>38.5 [34.0 to 45.75]</td>
<td>0.579$^{b}$</td>
</tr>
<tr>
<td>VAS (observed, T3)</td>
<td>3.0 [2.0 to 5.0]</td>
<td>3.5 [2.0 to 5.0]</td>
<td>0.418$^{b}$</td>
<td>3.0 [2.0 to 4.0]</td>
<td>0.171$^{b}$</td>
</tr>
</tbody>
</table>

Values are median [interquartile range] or frequency (percentage). FLACC, Face, Legs, Activity, Cry and Consolability; FPS-r, Faces Pain Scale revised; mYPAS, modified Yale Preoperative Anxiety Scale; PAED, Paediatric Anaesthesia Emergency Delirium; PPPM, Parents’ Postoperative Pain Measure; STAI, State-Trait Anxiety Inventory; T1, hospital admission (baseline); T2, holding area; T3, induction of anaesthesia; T4, recovery room, when children were fully awake, in the presence of a parent; T5, at home, within 2 weeks after surgery; VAS, Visual Analogue Scale. $^a$ Mann–Whitney $U$-test. $^b$ Per-protocol analyses (ad-hoc). $^c$ Need for rescue analgesia (yes or no) was also analysed for each type of surgery, separately.
number of children, we repeated the analyses per-protocol, in which we compared the children in the VRE group who completed the intervention ($n = 73$) with children in the CAU group ($n = 97$). These analyses did not produce significantly different results when compared with the intention-to-treat analyses (Tables 1 and 2; the two columns on the right).

**Discussion**

This single-blinded RCT, with a sample of 191 children, was designed to investigate the effect of fully immersive VRE in children undergoing elective day care surgery. No significant differences were found between VRE and CAU in child anxiety, pain, emergence delirium or parental anxiety. However, after VRE, children undergoing the most painful surgical procedure needed significantly less rescue analgesia compared with CAU. Lastly, levels of parental anxiety did not differ between VRE and CAU.

Virtual reality has previously been investigated as a means of improving health outcomes and previous studies have found that virtual reality reduced pain and anxiety in children undergoing different medical procedures. Most of these studies showed virtual reality being successfully used as a method of distraction. Because these studies were small, often not blinded and lacked standardised assessments, chance findings and a degree of bias could not be ruled out. Previously studied medical procedures that included oncological and burn wound care were more complex and painful compared with the procedures in our study. This is reflected by the relatively small proportion of our patients who experienced substantial levels of pain. It may be possible that VRE is more effective prior to more problematic surgery, with higher levels of anticipated anxiety and pain, compared with elective day care surgery. This is supported by our finding that only children who underwent the most painful type of surgery needed less rescue analgesia after VRE. This finding is of great clinical importance, because rescue analgesia such as morphine has several side effects, including nausea, vomiting and dizziness. Therefore, administering rescue analgesia may be associated with slower postoperative recovery. In our study, pain levels at T4 were similar in VRE and CAU groups. However, by that time, rescue analgesia would have already been administered if needed. It is possible that, despite substantial pain levels, no treatment effect was found because of adequate pain management. A final explanation for the absence of effects on anxiety and pain is that more time was needed between VRE and surgery for children to process the information. Children require up to 1 week for the processing of information about peri-operative processes. Therefore, VRE may be more effective if taken up no earlier than a week prior to surgery, perhaps even in multiple sessions, or via a mobile application for smartphones. This dispenses with an extra hospital visit and requires no hospital staff for the intervention, resulting in no extra healthcare costs. Considering the intervention only takes 15 min, it is achievable to implement VRE even in a busy clinical setting. However, it might be preferable to limit exposure to children who are most at risk for high levels of anxiety and pain, because these are the children who might benefit the most from VRE.

Two studies by Ryu et al. who used VRE prior to elective day care surgery, found positive effects. These studies were methodologically sound and included an acceptable number of participants. We offer several reasons for the discrepancy in results compared with our study. First, during induction of anaesthesia, Ryu et al. considered compliance, whereas our study considered anxiety. Compliance and anxiety are known to be different concepts, and even though patients were more compliant, they might still have been anxious during induction of anaesthesia. This is in line with their finding that distress levels in the operating room were not affected by playing a VR game pre-operatively. Second, the VRE group actually consisted of VRE and CAU because all the children in our study, including those in the VRE group, received routine care in a hospital setting that places great emphasis on patient comfort, in line with patient-centred and family-centred care. More specifically, all children and their parents received a pre-operative visit from a paediatric anaesthesiologist, during which elaborate education was provided along with a suggestion that the child and parents should watch an informative online movie about general anaesthesia. In addition, according to routine practice, children were not separated from their parents during anaesthetic induction and all parents were with their children throughout the recovery room stay. Successful routine care, in both the VRE and CAU groups, might have resulted in relatively low anxiety levels. A cut-off score of mYPAS at least 30 indicates high anxiety. In the current study, median anxiety levels in the CAU condition were 26.7 at baseline and 28.3 in the holding area. In comparison, Ryu et al. found substantially higher median levels of anxiety, also measured with the mYPAS, during baseline (CAU: 51.7 and 46.7). Unfortunately, it is not possible to make the same comparison for anxiety during induction of anaesthesia, as Ryu et al. did not use the mYPAS during induction. However, the comparisons between studies at admission (baseline) and in the holding area indicate that, overall, anxiety scores in our study were low, making it potentially more difficult to detect treatment effects. Finally, the lack of strong game design elements in our VRE intervention may explain the absence of results. Patients in the most recent study by Ryu et al. faced different challenges and received rewards whilst playing the VR game. These game elements are associated with

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greater engagement and education, so possibly also with a greater anxiolytic effect.

Strengths and limitations
The strengths of this study include the large sample size, limited missing data, use of internationally established standardised assessment tools, blinding of the medical and research staff and the narrow range of surgical procedures (elective day care surgery).

This study also has some limitations. First, in the recovery room, only one assessment took place. Multiple assessments, on entering the recovery room, and again after 5, 10 and 15 min would have provided a more comprehensive insight into the postoperative effects of VRE. Second, we did not include a survey on the subjective experience of the VRE, such as satisfaction, in children or their parents. Third, 21 children discontinued the intervention by taking off the headset. We found that the majority (71.4%) of these children were 4 or 5 years old. Wearing the rather large and heavy headset may have been uncomfortable, or the intervention may have been too lengthy for younger children, who overall have a limited attention span. Finally, by excluding patients who received anxiolytic premedication, we excluded the most anxious children in our study. Hospital policy dictates that anxiolytic premedication is not given unless, for example, it is after a previous traumatic experience with anaesthetic induction. Therefore, these cases can be considered exceptions and excluding them probably did not influence our results.

Conclusion
No significant differences were found between VRE and CAU in child anxiety, pain, or emergence delirium, or parental anxiety. However, after VRE, less rescue analgesia was needed after painful surgery. Considering the side effects of rescue analgesia, this means that VRE could be associated with increased patient comfort and a decreased need for postoperative care. It is possible that we did not find an effect of VRE on the other outcomes because we only investigated relatively mild procedures, the VRE intervention and surgery were too close to each other in time, and anxiety levels prior to induction of anaesthesia were relatively low. This is in line with the fact that more compelling results have been found in previous studies that either applied virtual reality to more complex procedures or to patient groups with higher levels of anxiety prior to induction.

Future research
On the basis of our results and conclusions, an option for future research is to investigate VRE in children with higher levels of pre-operative anxiety or to investigate VRE prior to more complex procedures with higher levels of expected postoperative pain. Second, when investigating postoperative effects of VRE, it would be valuable to make multiple assessments in the recovery room, as well as collecting information on nausea, vomiting and length of stay. Finally, more research is needed on the inclusion of game elements and the timing of VRE in relation to the day of surgery.

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Presentation: none.

References

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