Special issue: Research report

Out-of-body experience in vestibular disorders — A prospective study of 210 patients with dizziness

Christophe Lopez a,⁎ and Maya Elzière b

a Aix Marseille Univ, CNRS, LNIA, FR3C, Marseille, France
b Centre des Vertiges, Hôpital Européen, Marseille, France

Article history:
Received 31 December 2016
Reviewed 19 February 2017
Revised 11 May 2017
Accepted 30 May 2017
Published online xxx

Keywords:
Vestibular system
Out-of-body experience
Sensory conflict
Bodily self-consciousness
Self-location

Abstract

Out-of-body experiences (OBEs) are states during which people experience their centre of awareness as located outside of their physical body, along with the sensation of seeing the environment from an elevated viewpoint. OBE is encountered in epilepsy, migraine and depersonalization, and it is not an uncommon experience in the general population. Current neuroscientific models of bodily self-consciousness consider that OBE are related to a failure to integrate visual, somatosensory and vestibular signals. These models have highlighted the importance of visual-vestibular mismatch in OBE. Case reports from older clinical literature suggest that vestibular disorders may precipitate OBE, but we were lacking population-based evidence that OBE is related to vestibular disorders. The present observational, prospective study describes otoneurological, neuropsychological and phenomenological correlates of OBE in the largest sample of patients with dizziness to date (n = 210) compared to a group of age- and gender-matched controls with no history of dizziness (n = 210). We show a significantly higher occurrence of OBE in patients with dizziness (14%) than in healthy participants (5%). Most of the patients experienced OBE only after they started having dizziness for the first time. OBE in patients with dizziness were mainly related to peripheral vestibular disorders. We also identify depersonalization-derealization, depression and anxiety as the main predictors of OBE in patients with dizziness, as well as a contribution of migraine. Depersonalization-derealization was the only significant predictor of OBE in healthy controls. Altogether, our data indicate that OBE in patients with dizziness may arise from a combination of perceptual incoherence evoked by the vestibular dysfunction with psychological factors (depersonalization-derealization, depression and anxiety) and neurological factors (migraine).

© 2017 Elsevier Ltd. All rights reserved.
1. Introduction

Out-of-body experiences (OBEs) are states during which people experience their “self,” “mind,” or centre of awareness, as located outside of their physical body. During an OBE, people seem to be fully awake and often report a sensation of floating along with the impression of seeing the environment from an elevated position (Blackmore, 1982; Blanke & Dieguez, 2009; Brugger, 1997). OBE is not an uncommon experience in the general population and it is also encountered in conditions such as epilepsy, migraine and depersonalization (Blanke & Dieguez, 2009; Blanke & Mohr, 2005). This has resulted in a surge of interest from neurologists and neuroscientists over the past two decades to understand OBEs and provide a better comprehension of the sensorimotor and neurophysiological foundations of self-consciousness (Blanke, 2012; Brugger, 1997; Ionta et al., 2011; Kessler & Braithwaite, 2016; Metzinger, 2009).

Current neuroscientific models of bodily self-consciousness propose that accurate integration of visual, tactile, proprioceptive, interoceptive, motor and vestibular signals supports the experience of an embodied self (Blanke, 2012). OBEs are thus seen as a failure to coherently integrate these signals. A large body of evidence from neurology and research in healthy participants support this proposition. First, OBEs have been evoked in patients during electrical stimulations of the temporo-parietal junction, where they very likely interfere with multisensory processing (Blanke, Ortigue, Landis, & Seck, 2002; Bos, Spoor, Smits, Schouten, & Vincent, 2016). Second, OBEs due to epilepsy or stroke are often associated with complex multisensory illusions, such as visual sensations (including autocopy; the sensation of looking at one’s own body), somesthetic sensations (the perceived shape and size of the body is distorted), and vestibular sensations (Blanke & Mohr, 2005; Blanke, Landis, Spinelli, & Seck, 2004; Devinsky, Feldmann, Burrowes, & Bromfield, 1989; Lopez, Halje, & Blanke, 2008; Lopez, Heydrich, Seck, & Blanke, 2010). Third, recent studies in neurologically normal participants show that the perceived location of the “self” is altered when conflicts are created between visual and tactile signals (Ehrsson, 2007; Ionta et al., 2011; Lenggenhager, Mouthon, & Blanke, 2009; Lenggenhager, Tadi, Metzinger, & Blanke, 2007). These paradigms can evoke the feeling that the participants’ viewpoint is disembodied and that they self-identify with a distant avatar.

The vestibular contributions to the sense of self and embodiment have been poorly described when compared to the role of vision and touch, despite the crucial role of the vestibular system in the perception of self-motion and orientation (Blanke, 2012; Ferré & Haggard, 2016; Lenggenhager & Lopez, 2015; Lopez et al., 2008). In a recent study, healthy participants who received low-intensity galvanic stimulation of the vestibular nerves were more likely to adopt an embodied perspective to perceive letters traced on their forehead (Ferré, Lopez, & Haggard, 2014). The authors proposed that low-intensity vestibular stimulation increases the natural tendency of the vestibular system to anchor the self to the body, suggesting a vestibular contribution to embodied self-location. Another way to examine the vestibular contribution to embodiment would be to investigate patients with vestibular disorders. If vestibular signals are central for anchoring the self to the body (Ferré et al., 2014), patients suffering from vestibular disorders may be more prone to OBE.

An older clinical literature review only found some cases of patients with dizziness reporting abnormal sense of embodiment, and extremely rare cases of full-blown OBE in vestibular disorders (reviewed in Lopez, 2013). In the 19th Century, Krishaber (1873) was probably the first to report abnormal self and bodily perceptions in patients with dizziness, followed by Bonnier (1905), who described apparent dissociations between the self and the body. One of Bonnier’s patients reported “he was divided into two persons, one who had not changed posture, and another new person on his right, looking somewhat outwardly. Then the two somatic individuals approached each other, merged, and the vertigo disappeared.” Three decades later, Skworzoff (1931) established a link between vestibular disorders and illusory perceptions of doubles (autoscopic phenomena), as one of his patients suffering from dizziness reported seeing and feeling every day his own double. Schilder (1935) proposed that normal vestibular functions are required for a normal body schema and described several patients with dizziness who experienced abnormal perceptions of their body shape and size. Yet, these patients did not receive systematic otoneurological examinations. The sensations often reported by Krishaber (1873), Bonnier (1905) and Schilder (1935) that the self feels strange, unreal and disconnected from the body in patients with dizziness are reminiscent of symptoms of depersonalization. There is evidence that Menière’s disease can evoke symptoms resembling depersonalization (e.g., “I feel like I’m outside of myself. I feel like I’m not in myself”; Grigsby & Johnston, 1989) and that depersonalization is more frequent and severe in patients with dizziness (Jauregui-Renaud, Sang, Gresty, Green, & Bronstein, 2008; Tschan, Wiltink, Adler, Beutel, & Michal, 2013; Sang, Jauregui-Renaud, Green, Bronstein, & Gresty, 2006; reviewed in Jauregui-Renaud, 2015). Depersonalization is also often associated with anxiety and depression during dizziness (Tschan et al., 2013). Because of the lack of detailed phenomenology of disembodiment and the absence of systematic OBE questionnaires in older case reports (Bonnier, 1905; Grigsby & Johnston, 1989; Krishaber, 1873; Schilder, 1935; Skworzoff, 1931), the relation between vestibular disorders, OBE and depersonalization remains unclear. Finally, we note the recent description of one patient with a unilateral vestibular dysfunction who experienced OBEs (Kaliuzhna, Vibert, Grivaz, & Blanke, 2015). When tested in visuo-tactile conflicts known to evoke disembodied self-location (Lenggenhager et al., 2007), the patient reported a stronger feeling of elevated, disembodied self-location than control participants, suggesting a role of vestibular signals in OBE. In conclusion, despite anecdotal cases collected over more than a century, we lack convincing evidence of full-blown disembodiment related to vestibular disorders, as there has been to date no systematic neuropsychological and otoneurological investigations of OBE in a large population of patients with dizziness.

The present study describes otoneurological, neuropsychological and phenomenological correlates of OBE in the largest sample of patients with dizziness to date. We first
aimed at measuring the occurrence and describing the phenomenology of OBE in patients with dizziness, as done for epilepsy (Devinsky et al., 1989), stroke (Ionta et al., 2011), migraine (Podoll & Robinson, 1999), near-death experience (van Lommel, Wees, Meyers, & Elfferich, 2001), and sleep paralysis (Cheyne & Girard, 2009). Second, the study aimed at clarifying the relations between depersonalization and OBE in patients with dizziness and healthy participants.

## 2. Methods

### 2.1. Participants

This prospective, observational study was conducted in patients referred to our otoneurological centre for dizziness and vertigo, defined as having the feeling of spinning, swaying or tilting, that they were off balance, or that the room around them was spinning (Brandt, Dieterich, & Strupp, 2013). We included 210 patients with dizziness (143 females and 67 males; mean age ± SD: 51 ± 15.6 years) who matched the inclusion criteria (age over 18, ability to read and understand the questionnaires, no severe neurological or psychiatric disorders) and gave their consent to participate in the study. Depending on the patient’s symptoms, otoneurological examination included audiological assessment, videonystagmographic examination of spontaneous nystagmus, positional nystagmus, head shaking test (HST), pendular rotatory test and caloric test, video head impulse test (VHIT), cervical vestibular-evoked myogenic potentials (cVEMPs), and MRI with contrast injection. Forms of dizziness and the most common aetiologies are summarized in Table 1. 111 patients had peripheral vestibular disorders including benign paroxysmal positional vertigo (BPPV), Menière’s disease, perilymphatic fistula and inner ear malformations, or other acute unilateral vestibular disorders, as the most common aetiology. 24 (11%) patients had central forms of dizziness, 14 (6%) had somatoform (psychogenic) dizziness, and 4 (2%) had dizziness from another origin. The exact origin of dizziness was unknown in 57 (27%) patients, some of which had somatoform (psychogenic) dizziness, and 4 (2%) had unknown origin.

### 2.2. Methods

#### 2.2.1. Otoneurological examinations

Otoneurological examinations included the detection of spontaneous nystagmus (SN), positional nystagmus (PN) and nystagmus revealed by the HST using videonystagmography (numbers indicate the occurrence of positive findings at the time of the examination by the otoneurologist, not during the history of disease). Functioning of the peripheral vestibular apparatus was measured using the VHIT, caloric vestibular stimulation (CVS) and measurements of cVEMP over the sternocleidomastoid muscles. Numbers in the last three columns indicate the number of patients (out of those in which the test have been conducted) with abnormal vestibular responses, irrespective of aetiology. Forms of dizziness and main otoneurological findings in the sample of patients are presented in Table 1. Patients with abnormal vestibular responses during the VHIT for the lateral semicircular canals (gain < .8), decreased caloric vestibular responses (deficit > 25%), and absent or asymmetrical cVEMP amplitudes were considered as having vestibular dysfunction. The exact origin of dizziness was unknown in 57 (27%) patients, some of which presented with nystagmus and objective signs of vestibular dysfunction (Table 1). Otoneurological examinations showed that 147 patients presented with a spontaneous, positional or HST nystagmus at the time of the examination, and 81 patients had objective signs of vestibular dysfunctions, as evidenced by decreased gain of the horizontal vestibulo-ocular reflex during the VHIT (gain < .8), decreased caloric vestibular response (deficit > 25%), and absent or asymmetrical cVEMP amplitudes. Patients were compared to 210 age-, gender-, and smoking-matched (146 females and 64 males) healthy control participants with no history of otoneurological and psychiatric disorders. Control participants were recruited at the same otoneurological centre, and included partners, family members and acquaintances that accompanied the patients (n = 109), as well as, hospital staff (n = 101). Table 2 summarizes the characteristics of the two populations in terms of age, gender, smoking and drinking habits, education level, marital status, employment status and history of migraine. Patients with dizziness and healthy controls were matched for age, gender, education level and smoking habits.
Table 2 – Demographic and clinical characteristics of patients with dizziness and healthy controls. Mean ± SD is reported. Education level according to the French education system; Level 1: before high school; Level 2: accomplished high school; Level 3: two years after high school; Level 4: Bachelor’s degree, Level 5: Master’s degree, Engineering degree, PhD, MD.

<table>
<thead>
<tr>
<th>Patients with dizziness</th>
<th>Healthy controls</th>
<th>Patients vs. controls</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All (n = 210)</td>
<td>Without OBE (n = 181)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>51.0 ± 15.6</td>
<td>51.9 ± 15.8</td>
</tr>
<tr>
<td>Females/Males</td>
<td>143/67</td>
<td>119/62</td>
</tr>
<tr>
<td>Sex ratio</td>
<td>2.13</td>
<td>1.92</td>
</tr>
<tr>
<td>Highest education level</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Level 1</td>
<td>34%</td>
<td>34%</td>
</tr>
<tr>
<td>Level 2</td>
<td>18%</td>
<td>19%</td>
</tr>
<tr>
<td>Level 3</td>
<td>22%</td>
<td>20.5%</td>
</tr>
<tr>
<td>Level 4</td>
<td>13%</td>
<td>14%</td>
</tr>
<tr>
<td>Level 5</td>
<td>13%</td>
<td>12.5%</td>
</tr>
<tr>
<td>Employment status (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Employed</td>
<td>61%</td>
<td>59.5%</td>
</tr>
<tr>
<td>Student</td>
<td>1.5%</td>
<td>1%</td>
</tr>
<tr>
<td>Retired</td>
<td>22%</td>
<td>24%</td>
</tr>
<tr>
<td>Unemployed</td>
<td>15.5%</td>
<td>15.5%</td>
</tr>
<tr>
<td>Marital status (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Single</td>
<td>21%</td>
<td>19%</td>
</tr>
<tr>
<td>Married/couple</td>
<td>56.5%</td>
<td>57%</td>
</tr>
<tr>
<td>Divorced/uuowed</td>
<td>22.5%</td>
<td>24%</td>
</tr>
<tr>
<td>Smokers (%)</td>
<td>21.9%</td>
<td>20.2%</td>
</tr>
<tr>
<td>Alcohol consumption (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No alcohol</td>
<td>63.5%</td>
<td>62%</td>
</tr>
<tr>
<td>1 to 5 units/week</td>
<td>31.5%</td>
<td>33.5%</td>
</tr>
<tr>
<td>6 to 10 units/week</td>
<td>3.5%</td>
<td>3.5%</td>
</tr>
<tr>
<td>&gt;10 units/week</td>
<td>1.5%</td>
<td>1%</td>
</tr>
<tr>
<td>History of dizziness (months)</td>
<td>68.5 ± 103.2</td>
<td>70.9 ± 108.5</td>
</tr>
<tr>
<td>Migraine (%)</td>
<td>35.1%</td>
<td>31.9%</td>
</tr>
</tbody>
</table>

whereas they differed with respect to employment status, marital status and alcohol consumption (i.e., alcohol consumption was lower in patients with dizziness than in controls). Our institutional Ethics Committee (Hôpital Européen, CEDP-HE16.03) approved all procedures for this observational study.

2.2. Data recording

The same otoneurologist recruited patients and controls, and procedures were explained to both groups similarly. Participants filled out an OBE questionnaire whose purpose was explicitly written: This questionnaire is about out-of-body sensations that you may have experienced while you were awake, that is outside the periods of sleep or dream, and outside any consumption of alcohol or drug. OBE was evaluated using Palmer’s questionnaire (Palmer, 1979), later used by Terhune (2009): Have you ever had an experience in which you felt that “you” were “outside of” or “away from” your physical body; that is, the feeling that your consciousness, mind or centre of awareness was at a different place than your physical body? (If in doubt, please answer “no”). Participants ticked a “no” or “yes” box. Those who answered “yes” were invited to answer 5 more questions. Healthy controls answered these 5 questions once. Patients were asked whether they experienced OBE before they started having dizziness for the first time, and/or after they started having dizziness. Each patient answered these 5 questions for both periods:

1. During the OBE, have you had visual experiences? Possible answers were: “no”, “I am unsure”, “yes”, and “yes, and during this experience I saw my own body from the outside”.
2. During the OBE, have you had the feeling that the shape or size of your own body, or body parts, had changed? Possible answers were: “no”, “I am unsure”, and “yes” (participants could indicate which body part felt distorted).
3. During the OBE, have you had the feeling that you moved (sensation of motion, lightness, elevation, etc.)? Possible answers were: “no”, “I am unsure”, and “yes” (participants could indicate the nature of the sensation).
4. How long did the sensations of being out of body last? Possible answers were: “a few seconds”, “a few minutes”, “a few hours”, “about a day”, “more than a day”, and “more than a week”.
5. How many times did you have OBEs?

The introductory sentence ensured that we did not collect instances of OBEs that occurred during dreams or lucid dreams, neither during alcohol or drug consumption. The phrase “If in doubt, please answer ‘no’” reduced the risk of false
positives, but may have underestimated the occurrence of OBEs.

Participants also filled out the Hospital Anxiety and Depression Scale (Zigmond & Snaith, 1983), which includes seven items tapping into anxiety and seven items tapping into depression. Anxiety and depression scores each range from 0 to 21. Finally, patients filled out the Cambridge Depersonalization Scale (Sierra & Berrios, 2000) which includes 29 items about depersonalization and derealization symptoms. Patients were asked whether they have had these 29 symptoms since the start experiencing dizziness for the first time (as in Sang et al., 2006), whereas control participants were asked whether they have had these experiences during the last 6 months (as in Sierra & Berrios, 2000). The questionnaire was introduced as follows: This questionnaire describes strange and ‘funny’ experiences that normal people may have in their daily life. We are interested in their: (a) frequency, i.e., how often have you had these experiences [since you had dizziness for the first time (patients)/over the last 6 months (controls)]; and (b) their approximate duration (Sierra & Berrios, 2000). For each item, participants indicated the frequency of the experience on a scale ranging from 0 (“never”) to 4 (“all the time”), and the duration of the experience on a scale from 1 (“few seconds”) to 6 (“more than a week”). The introductory paragraph indicated: If you are not sure, give your best guess. For each item, the global score (sum of the frequency and duration) ranged from 0 to 10. The total score for the Cambridge Depersonalization Scale ranges from 0 to 290.

2.3. Data analysis

After stratification of patients and controls according to their answer to the Palmer's questionnaire (with vs without OBE), we compared their respective socio-economical and clinical data using two-tailed t-tests for independent samples for quantitative variables and χ² tests for qualitative variables.

A multivariate analysis of covariance (MANCOVA) was calculated to clarify the relations between depersonalization-derealization, depression and anxiety in patients and healthy controls, with the occurrence of OBE (IBM SPSS Chicago, IL, USA). Depersonalization-derealization, depression and anxiety scores were the dependent variables, whereas group (patients, controls), OBE (with OBE, without OBE), gender, and migraine were fixed factors. Age, marital status, smoking habits, alcohol consumption, education, and employment status were the covariates. When MANCOVA revealed an overall effect of a variable (i.e., significant Pillai's trace) on the dependent variables, univariate ANOVAs were conducted to determine which of the significant MANCOVA variables generated the significant multivariate effects.

In addition, we carried out receiver operating characteristic (ROC) curve analyses and binary logistic regression analyses (IBM SPSS Chicago, IL, USA) to determine how good the predictors (depersonalization-derealization score, anxiety score, depression score and each socio-economical and health variable) are to classify patients and controls as reporting OBE or not (for similar procedures in neuropsychology, see Gaser, Franke, Klöppel, Koutsouleris, Sauer, & Alzheimer's Disease Neuroimaging Initiative, 2013; Moura et al., 2017; Whelan-Goodinson, Ponsford, & Schönberger, 2009). For the ROC curve analysis, we evaluated the area under the curve (AUC) as a measure of the accuracy of the classification. An AUC of .5 reflects results from a random classifier whereas an AUC of 1 reflects perfect sensitivity and specificity of the classifier. Discrimination is considered to have failed for an AUC between .5 and .6, to be poor for an AUC between .6 and .7, fair for an AUC between .7 and .8, good for an AUC between .8 and .9, and excellent for an AUC between .9 and 1. Subsequently, we carried out binary logistic regression analyses, recommended for the analysis of independent variables that can take two possible values (i.e., with OBE, without OBE). Analyses used the Enter method, with OBE as the binary dependent variable, and with the following characteristics as covariates: depersonalization-derealization score, anxiety score, depression score, age, gender, migraine, marital status, smoking habits, alcohol consumption, education, employment status. Patients had additional covariates regarding the nature of their disease, including history of the disease, form of dizziness, and presence of a nystagmus. For all analyses, differences with a p < .05 were considered significant.

3. Results

3.1. Occurrence of OBE

We observed that 29 out of 210 patients with dizziness (14%), and 11 out of 210 healthy controls (5%), reported having had an OBE. Importantly, patients were significantly more likely to report an OBE than healthy controls (χ² = 7.41, p < .01; Fig. 1A). The statistical power was 89% according to SAS V9.4 Power Procedure (SAS Institute Inc. Cary, NC). Another important finding was that the occurrence of OBE in patients was significantly modulated by the onset of dizziness (χ² = 9.92, p < .01): most patients with OBE (n = 14, 48%) had OBE only after they experienced dizziness for the first time. Seven patients with OBE (24%) reported having had an OBE before and after they experienced dizziness, while 8 (28%) had OBE only before they experienced dizziness (Fig. 1B).

3.2. Phenomenology of OBE

Fig. 2 summarizes the phenomenological content of OBE. During their OBE, most of the patients experienced vestibular sensations (59%); e.g., sensations of elevation and lightness, “sensation of being attracted by a spiral, like in a tunnel”, “sensation of entering my body, like in an envelope, from the top”), while some experienced visual sensations (43%). Only few patients reported a change in the perceived shape and size of their body, which affected their entire body, arms or head (15%; e.g., “It feels as if my whole body was very small”, “My cheek was bigger”), and autoscopy (8%; e.g., “I saw myself, smaller, from the top”). Regarding the duration of the OBE, patients report it lasting for a few seconds (42.5%) to a few minutes (46%) or had these sensations several times for longer than a week (11.5%). For patients who remembered the occurrence of OBEs, 87% reported multiple OBEs. The pattern of sensations was similar in healthy control participants (Fig. 2). The frequency of vestibular and visual sensations, as well as, body schema distortion and autoscopy in healthy controls did not differ from that reported in patients (all χ² < 1.07 and p > .3).
3.3. Characteristics of individuals reporting OBE

Otoneurological findings in the 29 patients with OBE are summarized in Table 3. We found that OBE was mainly due to peripheral vestibular disorders \((n = 23)\), including Menière’s disease, vestibular neuritis, perilymphatic fistula and BPPV, with objective signs of an organic dysfunction. OBE associated to unilateral peripheral vestibular disorders was related to the left ear in eight patients and to the right ear in four patients (these proportions did not differ).

After stratification of the patients and controls according to their answer to the Palmer’s questionnaire (Table 1), we found that patients with OBE differed from patients who never had OBE in that they were significantly younger (mean \(\pm\) SD: 44.9 \(\pm\) 12.8 vs 51.9 \(\pm\) 15.8 years; \(t = 2.35, p < .05\)) and more prone to migraine (53.6\% vs 31.9\%; \(\chi^2 = 4.92, p < .05\)). There was a higher prevalence of OBEs in female \((n = 24)\) than male \((n = 5)\) patients, but the proportion did not differ from that in patients without OBE (statistical trend: \(\chi^2 = 3.33, p = .068\)). The prevalence of OBEs was not related to education level, employment status, marital status, smoking and drinking habits, or the history of the disease.

We compared vestibular responses in patients with and without OBE using the gain of the vestibulo-ocular reflex during the VHIT for the lateral semicircular canals (done in 14/29 patients with OBE and 79/181 patients without OBE) and the percentage of vestibular loss at the caloric test (done in 11/29 patients with OBE and 67/181 patients without OBE). Patients with and without OBE had similar gain at the VHIT (right ear: \(t = 1.14, p = .26\); left ear: \(t = .27, p = .79\)) and similar vestibular loss at the caloric test (\(t = .02, p = .98\)). Additional analyses showed that 20/29 (69\%) patients with OBE and 127/181 (70.2\%) patients without OBE presented with a spontaneous, positional, or a nystagmus at the HST (the proportions did not differ: \(\chi^2 = .017, p = .90\)). Thus, clinical data available indicate similar vestibular function in patients with and without OBE.

In healthy controls, there was no difference between participants with OBE and participants without OBE regarding their age, gender, education level, employment status, marital status, smoking and drinking habits, or history of migraine.

Fig. 1 — Occurrence of OBE. (A) Histogram shows higher occurrence of OBE in patients \((n = 29)\) than control participants \((n = 11)\). Asterisk indicates a significant difference between the two groups \((\chi^2\) test). (B) Histogram shows the proportion of patients who reported OBE before, after, or before and after they experienced dizziness for the first time. Asterisk indicates that the distribution of OBE depends significantly on the occurrence of dizziness \((\chi^2\) goodness-of-fit test), with most patients reporting OBE only after they started experiencing dizziness.

3.3. Characteristics of individuals reporting OBE

Otoneurological findings in the 29 patients with OBE are summarized in Table 3. We found that OBE was mainly due to peripheral vestibular disorders \((n = 23)\), including Menière’s disease, vestibular neuritis, perilymphatic fistula and BPPV, with objective signs of an organic dysfunction. OBE associated to unilateral peripheral vestibular disorders was related to the left ear in eight patients and to the right ear in four patients (these proportions did not differ).

After stratification of the patients and controls according to their answer to the Palmer’s questionnaire (Table 1), we found that patients with OBE differed from patients who never had OBE in that they were significantly younger (mean \(\pm\) SD: 44.9 \(\pm\) 12.8 vs 51.9 \(\pm\) 15.8 years; \(t = 2.35, p < .05\)) and more prone to migraine (53.6\% vs 31.9\%; \(\chi^2 = 4.92, p < .05\)). There was a higher prevalence of OBEs in female \((n = 24)\) than male \((n = 5)\) patients, but the proportion did not differ from that in patients without OBE (statistical trend: \(\chi^2 = 3.33, p = .068\)). The prevalence of OBEs was not related to education level, employment status, marital status, smoking and drinking habits, or the history of the disease.

We compared vestibular responses in patients with and without OBE using the gain of the vestibulo-ocular reflex during the VHIT for the lateral semicircular canals (done in 14/29 patients with OBE and 79/181 patients without OBE) and the percentage of vestibular loss at the caloric test (done in 11/29 patients with OBE and 67/181 patients without OBE). Patients with and without OBE had similar gain at the VHIT (right ear: \(t = 1.14, p = .26\); left ear: \(t = .27, p = .79\)) and similar vestibular loss at the caloric test (\(t = .02, p = .98\)). Additional analyses showed that 20/29 (69\%) patients with OBE and 127/181 (70.2\%) patients without OBE presented with a spontaneous, positional, or a nystagmus at the HST (the proportions did not differ: \(\chi^2 = .017, p = .90\)). Thus, clinical data available indicate similar vestibular function in patients with and without OBE.

In healthy controls, there was no difference between participants with OBE and participants without OBE regarding their age, gender, education level, employment status, marital status, smoking and drinking habits, or history of migraine.

Fig. 2 — Phenomenology of OBE. The occurrence of the associated sensations did not differ significantly between patients with dizziness and healthy controls \((\chi^2\) tests). “Body schema” refers to the perception of a distorted shape and size of the body.
Table 3 – Demographic and clinical characteristics of patients reporting OBEs. BPPV: benign paroxysmal positional vertigo; PPPD: persistent postural-perceptual dizziness; SN: spontaneous nystagmus; PN: positional nystagmus; HST: nystagmus evoked by the head shaking test.

<table>
<thead>
<tr>
<th>Occurrence of OBE with respect to the onset of the disease</th>
<th>Gender</th>
<th>Age (years)</th>
<th>Otoneurological findings and diagnosis</th>
<th>History of the disease (months)</th>
<th>Nystagmus at the time of examination</th>
<th>Migraine</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1 Only after</td>
<td>Female</td>
<td>30</td>
<td>PPPD, experience of room tilt illusions</td>
<td>204</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>P2 Only before</td>
<td>Female</td>
<td>46</td>
<td>Left vestibular neuritis, herpes infection</td>
<td>9</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>P3 Only after</td>
<td>Female</td>
<td>32</td>
<td>Right Menière’s disease</td>
<td>&gt;180</td>
<td>HST</td>
<td>No</td>
</tr>
<tr>
<td>P4 Only after</td>
<td>Male</td>
<td>42</td>
<td>Left perilymphatic fistula</td>
<td>72</td>
<td>PN</td>
<td>No</td>
</tr>
<tr>
<td>P5 Only after</td>
<td>Female</td>
<td>28</td>
<td>Chronic vestibulopathy of unknown origin, herpes infection</td>
<td>12</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>P6 Only after</td>
<td>Male</td>
<td>25</td>
<td>PPPD</td>
<td>4</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>P7 Only after</td>
<td>Male</td>
<td>44</td>
<td>Cerebrospinal fluid hypotension</td>
<td>4</td>
<td>HST</td>
<td>–</td>
</tr>
<tr>
<td>P8 Only after</td>
<td>Female</td>
<td>49</td>
<td>Unknown</td>
<td>53</td>
<td>PN</td>
<td>Yes</td>
</tr>
<tr>
<td>P9 Only before</td>
<td>Female</td>
<td>67</td>
<td>Left acute unilateral vestibular disorder</td>
<td>11</td>
<td>PN</td>
<td>Yes</td>
</tr>
<tr>
<td>P10 Only after</td>
<td>Male</td>
<td>33</td>
<td>Vestibulopathy of unknown origin</td>
<td>10</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>P11 Only before</td>
<td>Female</td>
<td>19</td>
<td>Right vestibulopathy</td>
<td>3</td>
<td>SN</td>
<td>No</td>
</tr>
<tr>
<td>P12 Only before</td>
<td>Female</td>
<td>55</td>
<td>Right lateral semicircular canal BPPV, arachnoid cyst, experience of room tilt illusions</td>
<td>24</td>
<td>HST</td>
<td>No</td>
</tr>
<tr>
<td>P13 Before + after</td>
<td>Female</td>
<td>36</td>
<td>Chronic vestibulopathy of unknown origin</td>
<td>8</td>
<td>HST</td>
<td>Yes</td>
</tr>
<tr>
<td>P14 Only after</td>
<td>Male</td>
<td>45</td>
<td>History of BPPV, migraines with aura</td>
<td>3</td>
<td>PN</td>
<td>Yes</td>
</tr>
<tr>
<td>P15 Only after</td>
<td>Female</td>
<td>53</td>
<td>Left Menière’s disease</td>
<td>36</td>
<td>HST</td>
<td>Yes</td>
</tr>
<tr>
<td>P16 Before + after</td>
<td>Female</td>
<td>64</td>
<td>Chronic vestibulopathy with positional vertigo, herpes infection</td>
<td>2</td>
<td>PN</td>
<td>No</td>
</tr>
<tr>
<td>P17 Only after</td>
<td>Female</td>
<td>65</td>
<td>Left acute unilateral vestibulopathy, left perilymphatic fistula operated</td>
<td>72</td>
<td>SN, PN, HST</td>
<td>Yes</td>
</tr>
<tr>
<td>P18 Only before</td>
<td>Female</td>
<td>51</td>
<td>Right posterior canal BPPV</td>
<td>1</td>
<td>PN</td>
<td>No</td>
</tr>
<tr>
<td>P19 Only before</td>
<td>Female</td>
<td>59</td>
<td>Chronic vestibulopathy of unknown origin</td>
<td>36</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>P20 Only after</td>
<td>Female</td>
<td>46</td>
<td>Unknown</td>
<td>20</td>
<td>PN</td>
<td>No</td>
</tr>
<tr>
<td>P21 Before + after</td>
<td>Female</td>
<td>43</td>
<td>Destruction of the left lateral semicircular canal</td>
<td>108</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>P22 Only after</td>
<td>Female</td>
<td>56</td>
<td>Left semicircular canal dehiscence, positional vertigo</td>
<td>96</td>
<td>SN</td>
<td>Yes</td>
</tr>
<tr>
<td>P23 Only after</td>
<td>Female</td>
<td>44</td>
<td>PPPD</td>
<td>27</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>P24 Before + after</td>
<td>Female</td>
<td>36</td>
<td>Vestibulopathy of unknown origin</td>
<td>4</td>
<td>PN</td>
<td>Yes</td>
</tr>
<tr>
<td>P25 Only before</td>
<td>Female</td>
<td>26</td>
<td>Vestibulopathy of unknown origin, tinnitus</td>
<td>180</td>
<td>SN, HST</td>
<td>No</td>
</tr>
<tr>
<td>P26 Before + after</td>
<td>Female</td>
<td>60</td>
<td>Unilateral vestibulopathy of unknown origin</td>
<td>144</td>
<td>SN</td>
<td>Yes</td>
</tr>
<tr>
<td>P27 Before + after</td>
<td>Female</td>
<td>57</td>
<td>Vestibulopathy of unknown origin, tinnitus</td>
<td>168</td>
<td>PN</td>
<td>Yes</td>
</tr>
<tr>
<td>P28 Before + after</td>
<td>Female</td>
<td>44</td>
<td>Recurrent positional vertigo, tinnitus, herpes infection</td>
<td>24</td>
<td>PN</td>
<td>No</td>
</tr>
<tr>
<td>P29 Only before</td>
<td>Female</td>
<td>46</td>
<td>Left lateral semicircular canal BPPV</td>
<td>48</td>
<td>PN</td>
<td>Yes</td>
</tr>
</tbody>
</table>
3.4. Relation of OBE to depersonalization, depression and anxiety

Using Pillai’s trace in the MANCOVA, we found a significant effect of OBE ($V = .77$, $F_{3,320} = 8.937$, $p < .0001$) and group ($V = .46$, $F_{3,320} = 5.087$, $p < .005$) on depersonalization-derealization, depression and anxiety scores. No other significant main effects or interactions were observed. Thus, OBE and group were used as factors for separate univariate ANOVAs on the outcome variables (Fig. 3A).

A first univariate ANOVA on the depersonalization-derealization score revealed a significant main effect of group, with patients having significantly higher depersonalization-derealization scores than controls ($F_{1,416} = 44.997$, $p < .00001$). Importantly, there was a main effect of OBE, with significantly higher depersonalization-derealization scores in participants who reported OBE ($F_{1,416} = 75.912$, $p < .00001$). There was also a significant group × OBE interaction ($F_{1,416} = 19.600$, $p < .0001$), which is shown in Fig. 3A. Post-hoc analyses revealed that depersonalization-derealization scores were higher in patients with OBE than in patients without OBE (88.6 ± 65.3 vs 25.5 ± 24.8; planned comparison: $p < .0001$). Similarly, depersonalization-derealization scores were higher in controls with OBE than without OBE (35.1 ± 29.6 vs 14.4 ± 15; $t = 4.17$, $p < .0001$).

A second univariate ANOVA on the depression score showed a significant main effect of group, with patients having significantly higher depression scores than controls ($F_{1,412} = 29.903$, $p < .00001$). There was also a main effect of

Fig. 3 – Relation between OBE and depersonalization-derealization, depression and anxiety. (A) Histograms illustrate the average depersonalization-derealization (DD) scores from the Cambridge Depersonalization Scale (Sierra & Berrios, 2000) and the average depression and anxiety scores (mean ± standard error of the mean) from the Hospital Anxiety and Depression scale (Zigmond & Snaith, 1983). Asterisks indicate a significant difference between participants who experienced OBE (coloured bars) and participants who did not report OBE (hatched bars), and sharp signs indicate a significant difference between patients with dizziness and healthy controls (post-hoc planned comparisons). (B) ROC curves for detecting the occurrence of OBE in patients with dizziness and healthy controls.

Please cite this article in press as: Lopez, C., & Elzière, M., Out-of-body experience in vestibular disorders – A prospective study of 210 patients with dizziness, Cortex (2017), http://dx.doi.org/10.1016/j.cortex.2017.05.026
OBE, whereby participants with OBE had significantly higher depression scores than participants without OBE \((F_{4,412} = 4.898, \ p < .05)\). In addition, there was a statistical trend for the group \(\times\) OBE interaction \((F_{4,412} = 2.922, \ p = .088)\). When exploring this statistical trend using post-hoc tests (Fig. 3A), we found that patients with OBE differed significantly from patients who never had OBE in that they had significantly higher depression scores \((8.8 \pm 3.8\ vs\ 6.0 \pm 4.3; \ p < .005)\), whereas healthy controls with and without OBE had similar depression scores \((3.8 \pm 3.1\ vs\ 3.5 \pm 3.0; \ p = .72)\).

Finally, a univariate ANOVA on the anxiety score showed a significant main effect of group, with significantly higher anxiety scores in patients with respect to controls \((F_{4,411} = 19.454, \ p < .00005)\). In addition, we found a statistical trend for the main effect of OBE, with a trend for higher anxiety scores in participants with OBE than without OBE \((F_{4,411} = 3.361, \ p = .067)\). There was also a statistical trend for the group \(\times\) OBE interaction \((F_{4,411} = 3.781, \ p = .053)\). Post-hoc analysis of this nearly significant interaction (Fig. 3A) showed that patients with OBE differed from patients who never had OBE in that they had significantly higher anxiety scores \((11.9 \pm 4.8\ vs\ 9.3 \pm 4.6; \ p < .005)\). By contrast, healthy controls with and without OBE had similar anxiety scores \((7.3 \pm 4.2\ vs\ 7.4 \pm 3.5; \ p = .94)\).

3.5. Precipitating factors of OBE

Although the analyses reported above indicate that patients with and without OBE differ for some demographic and clinical data, we wanted to establish which variables successfully discriminate between participants with and without OBE, using ROC curve analysis and binary logistic regression.

The ROC curve analysis showed that three measures were relevant in discriminating between patients with and without OBE (Fig. 3B). Depersonalization-derealization score revealed excellent discrimination with AUC = 0.859 \((p < .00001; \ 95\%\ confidence\ interval\ (CI): \ .760–.959)\) and a standard error of .051. Depression was a fair predictor of OBE with AUC = 0.737 \((p = .001; \ 95\%\ CI: \ .628–.846)\) and a standard error of .056. Anxiety was a poor predictor of OBE with AUC = 0.688 \((p = .007; \ 95\%\ CI: \ .575–.801)\) and a standard error of .058. There was a poor contribution of the form of dizziness (peripheral, central, somatoform, unknown) with AUC = 0.605 \((p = .135; \ 95\%\ CI: \ .465–.745)\) and a standard error of .071, as well as, of the gender with AUC = 0.602 \((p = .144; \ 95\%\ CI: \ .477–.728)\) and a standard error of .064. There was no reliable discrimination based on age \((AUC = .406; \ p = .180)\), migraine \((AUC = .588; \ p = .210)\), marital status \((AUC = .405; \ p = .177)\), smoking habits \((AUC = .556; \ p = .472)\), alcohol consumption \((AUC = .449; \ p = .467)\), education \((AUC = .501; \ p = .888)\), employment status \((AUC = .466; \ p = .626)\), history of the disease \((AUC = .490; \ p = .888)\), and presence of a nystagmus \((AUC = .533; \ p = .637)\).

In healthy controls (Fig. 3B), the ROC curve analysis showed that only the depersonalization-derealization score revealed fair classification with AUC = .719 \((p = .015; \ 95\%\ CI: \ .550–.888)\) and a standard error of .086. In contrast with patients, there was no reliable prediction from depression \((AUC = .541; \ p = .648)\) and anxiety \((AUC = .466; \ p = .706)\). These data however, should be taken with caution given the small number of healthy controls reporting OBE.

Binary logistic regression analysis in patients revealed that the depersonalization-derealization score was the only significant predictor of OBE \((\beta = -.060, \ Wald's \chi^2 = 9.865, \ p < .005, \ odds\ ratio: .942)\) while migraine was second, although not significant \((\beta = -1.461, \ Wald's \chi^2 = 1.873, \ p = .171, \ odds\ ratio: .223)\). This binary logistic regression model correctly classified 90.3% of patients regarding their report of an OBE, with 66.7% of true-positive, 95.6% of true-negative, 4.4% of false-positive and 33% of false-negative (Hosmer and Lemeshow goodness-of-fit test: \(\chi^2(8) = 8.943, \ p = .347\); Cow and Snell \(R^2 = .397\); Nagelkerke \(R^2 = .643\)).

Binary logistic regression analysis in healthy controls also revealed that the depersonalization-derealization score was the only significant predictor of OBE \((\beta = -.063, \ Wald's \chi^2 = 6.111, \ p < .05, \ odds\ ratio: .939)\). The second (although not significant) predictor was anxiety \((\beta = .202, \ Wald's \chi^2 = 2.101, \ p = .147, \ odds\ ratio: 1.224)\). This binary logistic regression model correctly classified 94.5% of healthy controls. Yet, there was a low number of true-positive (18.2%) and a high number of false-negative (81.8%), with a low number of true-negative (99.4%) and false-positive (6%) (Hosmer and Lemeshow goodness-of-fit test: \(\chi^2(8) = 10.668, \ p = .347\); Cow and Snell \(R^2 = .397\); Nagelkerke \(R^2 = .643\)). Thus, classification in the controls does not seem as reliable as in patients and the small number of healthy participants reporting OBE imposes caution with the interpretation of this result.

4. Discussion

This prospective study conducted in a large sample of participants shows that there is a significantly higher occurrence of OBE in patients with dizziness than in gender- and age-matched healthy controls. We discuss the relation between vestibular disorders, disembodied experience, depersonalization-derealization, anxiety and depression in the framework of current multisensory models of self-consciousness.

4.1. Dizziness increases the occurrence of OBE

Our data provide support to intuitions from older literature that proposed relations between dizziness and disembodied experience (Bonnier, 1905; Krishaber, 1873; Menninger-Lerchenthal, 1935; Schilder, 1935; Skworzoff, 1931). While these studies did not quantify the occurrence of OBE in patients with dizziness, we showed that 14% of the patients have had at least one OBE, which is a significantly higher occurrence than in healthy controls (5%).

Notably, OBE was not frequent in either group, indicating the reliability of the sensorimotor and cognitive mechanisms of embodiment. Earlier studies estimated that OBEs occur in about 10% of the general population (Blackmore, 1982; Blanke & Dieguez, 2009). The occurrence of OBE ranged from 8 to 50%, depending on the tested population, but most studies were conducted in student populations (see Blackmore (1982) for a detailed review). For example, Palmer (1979) found OBE in 25% of the students versus in 14% of the other inhabitants of Charlottesville, Virginia, whereas Green (1968) found OBE in 19% of the Southampton University students. Another survey
found OBE in 8% of the Icelandic population (Haraldsson, Gudmundsdottir, Ragnarsson, Loftsson, & Jonsson, 1977). The occurrence of OBE in our sample of healthy participants is lower than in previous studies. Importantly, our participants were selected on the basis that they had no history of dizziness, otoneurological or psychiatric disorders, which was not the case in previous investigations. Given that dizziness is a common condition affecting 11–30% of the population (Bigelow, Semenov, du Lac, Hoffman, & Agrawal, 2016; Neuhauser, 2007), previous studies have inevitably measured OBE in participants with dizziness. Our study allows measuring the occurrence of OBE separately for participants with and without dizziness, showing that dizziness precipitates OBE.

4.2. How vestibular disorders precipitate OBE

We observed that the majority of the patients experienced OBE only after they experienced dizziness for the first time. This indicates that dizziness is a triggering factor of OBE. Multiple diseases of various aetiologies can evoke dizziness (Brandt et al., 2013). Here, we established that peripheral vestibular disorders were the most common cause of OBE in 29 patients, including vestibular neuritis, Menière’s disease, perilymphatic fistula and BPPV (Table 3). Importantly, we found objective signs of vestibular disorders in these patients (spontaneous, positional or HST nystagmus and decreased vestibulo-ocular reflex), indicating that OBE involves organic dysfunctions. Yet, vestibular functions did not differ between patients with and without OBE, indicating that additional factors were involved, such as depersonalization-derealization, depression and anxiety (see below). In three patients, OBE was related to PPPD, which is close to phobic postural vertigo and chronic subjective dizziness (Dieterich & Staab, 2017). While vestibular pathologies are associated with deficits in postural control and gaze stabilization, spatial navigation and cognition, we provide the first population-based evidence that vestibular disorders can also evoke abnormal forms of embodiment.

The phenomenology of OBE indicates that most of the patients with dizziness and healthy participants experienced concomitant vestibular sensations. Our findings confirm and extend previously reported association between OBE and vestibular sensations (Brugger, 1997; Grüsser & Landis, 1991; Menninger-Lerchenthal, 1935). This association is particularly evident from self-reports in neurological patients (Blanke et al., 2004; Devinsky et al., 1989; Heydrich, Lopez, Seeck, & Blanke, 2011; Hécaen & de Ajuriaguerra, 1952; Lopez et al., 2010), healthy participants (Blackmore, 1982; Green, 1968), and during sleep paralysis (Cheyne & Girard, 2009). Other evidence comes from presurgical evaluation of epilepsy, showing that electrical stimulation of the temporoparietal cortex evokes both vestibular sensations and OBE (Blanke et al., 2002; Penfield & Jasper, 1954). Finally, experimental inductions of “OBE-like” experiences in healthy participants were characterized by self-location distant from the physical body, together with sensations of floating (Jonta et al., 2011; Lenggenhager et al., 2009). Altogether, this indicates that vestibular signals crucially contribute to the experience of embodiment.

What is the pathophysiology of OBE in vestibular disorders? According to Blanke and colleagues (Blanke & Mohr, 2005; Blanke et al., 2004; Lopez et al., 2008), OBEs result from a multisensory disintegration in personal space (i.e., a failure to integrate visual body-related information and somatosensory information) and a disintegration between personal and extrapersonal space (i.e., a failure to integrate vestibular and visual information). This model emphasizes the conflict between vestibular and visual information and in this respect OBE has been referred to as a “visuo-vestibular splitting of the somatosensory body image” (Grüsser & Landis, 1991). Regarding the neuroanatomical bases of OBE, Grüsser and Landis (1991) proposed that OBE “is a combination of visual and vestibular hallucinations involving the posterior insula and retrolenticular cortex”, as the parieto-insular cortex is a visuo-vestibular-somatosensory area and the core of the vestibular cortex (Guldin & Grüsser, 1998; Lopez, Blanke, & Mast, 2012). The parieto-insular cortex is also the most frequently damaged area in neurological patients reporting OBE (Jonta et al., 2011). We propose that during acute vestibular disorders (e.g., Menière’s disease, vestibular neuritis, BPPV), the central nervous system receives erroneous signals from the inner ear regarding body motion and orientation (e.g., sensation of body inclination to the right) that are conflicting with visual and somatosensory signals indicating another body position and motion (e.g., the body and the environment are upright). This sensory conflict would lead to perceptual incoherence and ultimately distort bodily self-consciousness (Lopez, 2013). The model holds true for patients with PPPD, also characterized by abnormal multisensory processing (Holte et al., 2015). For some patients with dizziness, perceptual incoherence may evoke full-blown OBE, probably because of additional precipitating factors (i.e., depersonalization-derealization; see below), whereas for other patients perceptual incoherence may evoke simpler forms of depersonalization without full-blown OBE (Tschan et al., 2013).

4.3. Relations between depersonalization-derealization and OBE

Multivariate and univariate analyses established relations between depersonalization-derealization scores and the occurrence of OBE in both patients and controls. We found that dizziness was associated with significantly higher depersonalization-derealization scores than in healthy controls, confirming previous investigations but in a larger sample (Jauregui-Renaud, 2015; Jauregui-Renaud et al., 2008; Kolev, Georgieva-Zhostova, & Berthoz, 2014; Sang et al., 2006). In addition, depersonalization-derealization scores were significantly higher in both patients and controls with OBE compared to those without OBE. Importantly, ROC curve analysis and binary logistic regression identified depersonalization-derealization as the main predictor of OBE in patients and healthy controls.

Patients with OBE had particularly high depersonalization-derealization scores, whose average was above the cut-off value (70) for clinical depersonalization-derealization, but lower than the average score (119) of psychiatric patients diagnosed with depersonalization-derealization disorders.
A previous investigation in patients with vestibular disorders showed higher scores of depersonalization-derealization in anxious patients than in patients with low anxiety, establishing relations between mood disorders and self-perception (Kolev et al., 2014). As anxiety was also a significant predictor of OBE (see below), we suggest that anxiety and depersonalization-derealization may combine to precipitate OBE in patients with dizziness.

Depersonalization-derealization was the only significant predictor of OBE in healthy controls. However, since only 11 controls reported an OBE, this result should be taken with caution. In line with our findings, a study by Murray and Fox (2005) showed that healthy participants with OBE report stronger somatoform dissociation, self-consciousness and body dissatisfaction when compared to participants without OBE. Another investigation in university students showed that individuals with OBE reported more perceptual anomalies associated with body-distortion processing (Braithwaite, Samson, Apperly, Broglia, & Hulleman, 2011).

### 4.4. Influence of anxiety and depression

Overall patients with dizziness were more anxious and depressed than healthy controls. The reciprocal relations between dizziness and anxiety/depression are well established in the clinical literature: there is a high comorbidity of vestibular disorders with anxiety/depression, and anxious patients are more likely to experience vertigo (Bigelow et al., 2016; Eckhardt-Henn et al., 2008; Tschan et al., 2013; Yardley, Masson, Verschuer, Haacke, & Luxon, 1992).

The ROC curve analysis indicated that depression was a predictor of OBE in patients with dizziness. Depression was also significantly higher in patients with OBE when compared to patients without OBE. Depression was similar in controls with and without OBE. We found no strong evidence in the literature to link OBE and depression. Bunning and Blanke (2005) reviewed cases of OBE in psychiatric patients and found that OBE was very rarely associated with depression. Depression has been related to autoscopic hallucination, but the origin of the viewpoint was not disembodied (Dening & Berrios, 1994). A retrospective study in 550 patients with depression (n = 100), mania and schizophrenia, showed that depression was never associated with OBE (McGilchrist & Cutting, 1995). This study found OBE only in acute schizophrenia. Our study provides the first evidence that depression, combined with other factors, precipitates OBE in patients with dizziness.

The ROC curve analysis also indicated that anxiety was a predictor of OBE in patients. We found that patients with OBE had significantly higher anxiety scores than patients without OBE, whereas anxiety scores were identical in healthy controls with and without OBE. This is consistent with previous comparisons of personality traits in healthy participants with and without OBE showing no differences with regards to their general anxiety (reviewed in Blanke & Dieguez, 2009) or social anxiety (Murray & Fox, 2005). The impact of anxiety on embodiment may differ in patients with dizziness, given their high level of anxiety. Mohr and Blanke (2005, p. 192) hypothesized that “anxiety is most closely linked to the form of [autoscopic phenomena] that is characterized by complete disembodiment (as in OBE), also attributable to the vestibular involvement in anxiety and OBEs”. Patients with high anxiety may be more prone to OBE, similarly to healthy individuals who experience OBE during stressful or life threatening situations (Blanke & Dieguez, 2009; Brugger, Regard, Landis, & Oelz, 1999).

### 4.5. Other factors

Migraine was more frequent in patients who reported OBE and was a predictor contributing only marginally to OBE. Migraine has been related to OBE in older clinical literature (Lippman, 1953; Podoll & Robinson, 1999) and is often associated with dizziness (Lempert & Neuhauser, 2009). When compared to patients with other vestibular disorders, patients with vestibular migraine have the highest risk of psychiatric comorbidities, including high anxiety and depression (Eckhardt-Henn et al., 2008; Lempert & Neuhauser, 2009). Patients with migraine seem to be more sensitive to motion, as they have lower thresholds for vestibular perception (Lewis, Priesol, Nicoucar, Lim, & Merfeld, 2011). More work is needed to understand how patients with migraine interpret self-location and self-motion during situations of perceptual incoherence. We note that migraine is more frequent in females, which could explain why there tend to be more females in our sample of patients reporting OBE.

Finally, patients with OBE were younger than patients without OBE. This was not the case in healthy controls. Previous research has consistently found that there is no age or gender difference in the occurrence of OBE (Blackmore, 1982). Given that depersonalization-derealization scores were higher in patients with OBE, and because depersonalization generally occurs in younger individuals (Simeon et al., 1997), this may explain partly why the group of patients with OBE was younger.

### 4.6. Limitations of the study

A limitation of the present study is that it relies on self-report assessment. Patients experienced difficulty reporting the phenomenology of their OBE, as they did not fill out the questionnaire immediately after an OBE. Patients reported complex experiences that may have occurred years ago, with inevitable imprecision of memories (see Blackmore (1982) for a detailed account). A strength of the study was that all patients with dizziness had a clinical diagnosis by an otoneurologist. Yet, we lack systematic recordings of all vestibular parameters in all patients, allowing detailed comparison of saccular, utricular and semicircular canals functions in patients with and without OBE. This is incompatible with otoneurological routine. For example, there is no systematic indication for BPPV to record vestibular-evoked myogenic potentials. In addition, we note several limitations regarding our sample of healthy controls. First, there was no otoneurological investigation in the healthy controls, which is practically undoable in this large sample. Second, only 11 healthy participants reported OBEs and therefore average demographic data, depersonalization-derealization, anxiety and depression scores calculated in this subpopulation should be taken with caution. Future studies should be conducted in larger samples.
of healthy participants to identify more precisely the predictors of their OBEs. Finally, a strength of our study was the large sample size (n = 420) and the inclusion of participants with a large range of age and socio-economical origins, different from the students populations usually tested in OBE studies.

5. Conclusions

Our study measured the occurrence of OBE in patients with dizziness and healthy participants. The vast majority of participants never experienced an OBE, indicating that the OBE-prone subpopulations in patients with dizziness and healthy controls, which cumulate several precipitating factors of otoneurological and psychological origin (Mohr & Blanke, 2005). Patients who have had an OBE also exhibit comorbid depersonalization-derealization, anxiety and depression. Altogether, our data indicate that OBE in patients with dizziness may arise from a combination of perceptual incoherence (due to conflicting vestibular signals with other sensory signals about body orientation and motion) mainly with depersonalization-derealization, depression and anxiety, as well as migraine.

Acknowledgements

The research leading to these results has received funding from the People Programme (Marie Curie Actions) of the European Union’s Seventh Framework Programme (FP7/2007–2013) under REA grant agreement number 333607 (“BODILYSELF, vestibular and multisensory investigations of bodily self-consciousness”). C. Lopez’s research is supported by a grant from the VolkswagenStiftung (Grant no. 89434: “Finding Perspective: Determining the embodiment of perspectival experience”). The study has received support from the Hôpital Européen. We thank Dr A. Pavlidou and Dr. R. Ackerley for helpful comments on the manuscript and statistical analyses.

References


