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The effect of normalization of sodium on bone turnover markers in patients with epilepsy. A randomized single-blinded placebo-controlled trial

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ABSTRACT

Hyponatremia [p[Na] <136 mmol/L] is an independent risk factor for decreased bone mineral density (BMD). However, whether hyponatremia represents a surrogate marker, or a direct causal relationship to bone loss remains unknown. The aim of the study was to investigate the effect of salt replacement therapy on bone turnover markers (BTM) and BMD in patients with epilepsy and chronic hyponatremia. This prospective single-blinded randomized trial investigated serum BTM and BMD, evaluated by Dual Energy X-ray Absorptiometry (DXA), in 21 patients at baseline and following three months of salt replacement therapy. Patients with two consecutive measurements of hyponatremia prior to baseline and no known osteoporosis were included from the epilepsy out-patient clinic at Rigshospitalet, Denmark. Seven patients were randomized to placebo and 14 to salt intervention. The baseline p[Na] was 134 (130.5–140) mmol/L (median (IQR)). All patients had BTM within age-specific reference ranges at baseline. Following 3 months of intervention with 3–9 g of salt daily there was no difference in levels of procollagen type 1 N-terminal propeptide (PINP) or C-terminal cross-linking telopeptide of type 1 collagen (CTX) between placebo and intervention. Nor was there any difference in BMD evaluated at the lumbar spine (L2-L4) or at the femoral neck or total hip. In our study, salt replacement did not affect BTM nor BMD. However, due to the small size of the study, more studies are needed to further investigate this.

1. Introduction

Patients with epilepsy have a greatly increased risk of developing osteoporosis [1]. As many as 80% of patients with epilepsy have decreased bone mineral density (BMD) and more than 30% have actual osteoporosis [2]. The high occurrence appears to be caused by multiple pathophysiological mechanisms where also hyponatremia seems to play a role [3]. Through the past decade accumulating evidence have shown a strong association between hyponatremia and the occurrence of osteoporosis and fractures [4–6]. Several antiepileptic drugs (AEDs), especially enzyme-inducing AEDs (EIAEDs) such as carbamazepine and oxcarbazepine, are known to cause hyponatremia in up to 46% of treated individuals [7–10]. Despite the high occurrence of hyponatremia, the association with osteoporosis in patients with epilepsy has, to our knowledge, not previously been investigated in a prospective clinical trial. Furthermore, the effect of correcting hyponatremia with salt supplements on bone turnover and BMD is unknown. Therefore, the aim of this study was to evaluate the effect on bone turnover of salt replacement therapy in patients with epilepsy and hyponatremia, evaluated by serum bone turnover markers (BTM) cross-linked N-telopeptide of type 1 collagen (CTX) and procollagen type 1 amino-terminal propeptide (PINP-intact), and bone mineral density (BMD), evaluated by Dual Energy X-ray Absorptiometry (DXA) scan.

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2. Materials and methods

2.1. Inclusion and exclusion criteria

The inclusion criteria were AED treated epilepsy through a minimum of two years, known chronic hyponatremia (p[Na]<136 mmol/L) in two consecutive measurements, age between 18 and 80 years, fluent in Danish and having signed a form of written consent. Patients were excluded if they had known osteoporosis, defined by the WHO definition of site-specific T-score ≤−2.5 standard deviation (SD) in any of the following sites: lumbar spine L1-L4, femoral neck and of the total femur. Patients were likewise excluded if they were undergoing treatment for osteoporosis, were known with the diagnosis SIADH, were pregnant or breastfeeding, had severe concomitant disease like cancer or ischemic heart disease, had a known alcohol, drug or medicine abuse or were known with any disease affecting their calcium metabolism (including primary hyperparathyroidism, thyrotoxicosis, myxedema, severe vitamin D deficiency or severe decreased kidney function).

2.2. Study design

Patients came fasting (except water and usual medication) to the clinic on the day of randomization. Here a blood sample was taken between 7.30 and 10am to avoid circadian influence on BTM. All samples were measured on Vitros® 5,1 FS/5600 Ortho Clinical Diagnostics, Albertslund, Denmark. The patients were asked question on past medical history, both epilepsy- and osteoporosis-related. At the end of the interview BMD was assessed by DXA scan. Randomization was done by the patient choosing or intervention group. Randomization was done by the patient choosing

2.3. Endpoints and statistics

The primary endpoint of the study was the change in bone resorption marker cross-linked N-telopeptide of type 1- collagen (CTX) from baseline visit to visit 6. Secondary endpoints were the bone formation marker procollagen type 1 amino-terminal propeptide (PINP-intact), and BMD evaluated by Dual Energy X-ray Absorptiometry (DXA). Due to the exploratory nature of the trial 21 patients were recruited. Patients were randomized 2:1 in favor of the intervention group. A 2:1 inclusion ratio (active vs. control treatment) was chosen in order to increase acceptability for patients to enroll in the study. Since the primary endpoint was relative changes in BTM from baseline to visit 6 men and pre- and postmenopausal women were analyzed together.

To characterize the population all descriptive variables and biochemical parameters were tested for normal distribution and expressed as either mean and standard deviation (SD) or median and interquartile range (IQR) depending on the distribution of data or as N-value and percentage (N (%)). Statistical differences between the placebo and intervention group were for parametric continuous data tested using an independent samples t-test and for non-parametric continuous data a Mann-Whitney U test was used. For categorical data Pearson chi-square test or Fisher’s exact test were used.

To test for any differences in BTM and T-scores between the groups an independent samples t-test was done for T-scores and for the BTM a Mann-Whitney U test was done. Furthermore, to test for changes from baseline to visit 6 within the two groups, for T-scores a Paired-Samples t-test was done. Furthermore, to test for the effects of the intervention on BTM and BMD that could not be contributed to the effects of time a factorial repeated

![Diagram of experiment outline](image-url)
measures ANOVA was done. Data was analyzed using the SPSS software package (version 22.0) (IBM, Armonk, New York). A p-value < 0.05 (two-sided) was considered statistically significant.

2.4. Ethics

The study was approved by the Danish Committee on Health Research Ethics (57353) and by the Danish Data Protection Agency (2012-58-0004). Informed consent was obtained from all individual participants included in the study. No experiments and procedures were done that conflict with the Helsinki Declaration of 1975 (revised 2000). The study was registered at clinicaltrials.gov (NCT03371199).

3. Results

3.1. Baseline characteristics, biochemical parameters, BTM and DXA scans

A total of 21 patients were included in the study. Their osteoporosis-related baseline characteristics and biochemical parameters are shown in Table 1. There was no difference between the groups in age, ethnicity, weight, height, osteoporosis-related characteristics or use of EIAEDs. Nor was there any difference in epilepsy-related characteristics or other biochemical parameters (data not shown). For all patients, the two consecutive measurements of p[Na] < 136 mmol/L prior to inclusion were for the first measurement, 132 ± 3 mmol/L (mean ± SD) and for

### Table 1

| Patient characteristics | All patients (N = 21) | Placebo group (N = 7) | Intervention group (N = 14) | Statistical significance Placebo vs Intervention |
|-------------------------|-----------------------|-----------------------|-----------------------------|-------------------------------------------------|
| Female sex (N (%))      | 16 (76.2)             | 6 (85.7)              | 10 (71.4)                   | 0.624                                           |
| Age at baseline (years) | 48 (42.5–65.5)        | 48 (35–63)            | 48.5 (42.75–68.5)           | 0.799                                           |
| Ethnicity (N (%))       |                       |                       |                             |                                                 |
| Caucasian               | 20 (90.9)             | 6 (85.7)              | 14 (100)                    |                                                 |
| Asian                   | 1 (4.8)               | 1 (14.3)              | –                           | 0.333                                           |
| Weight (kg) (mean (SD)) | 78.3 (15.0)           | 79.2 (17.8)           | 77.8 (14.1)                 | 0.839                                           |
| Height (cm) (mean (SD)) | 170 (8.4)             | 166 (5.4)             | 172 (9.0)                   | 0.129                                           |
| BMI (kg/m²) (mean (SD)) | 27.0 (4.7)            | 28.8 (6.9)            | 26.1 (3.0)                  | 0.352                                           |
| Family disposition to osteoporosis (N (%)) | 17 (81) | 7 (100) | 10 (71.4) | 0.255 |
| Other disease relevant for osteoporosis (N (%)) | 4 (19) | 0 (0) | 4 (28.6) | 0.255 |
| Has the patient ever received treatment with steroids (N (%)) | 21 (100) | 7 (100) | 14 (100) | 0.05 |
| History of low energy fractures (N (%)) | 21 (100) | 7 (100) | 14 (100) | 0.05 |
| History of high energy fractures (N (%)) | 6 (0) | 0 (0) | 0 (0) | 0.05 |
| Intake of calcium (mg/day) (median (IQR)) | 29 (5-54) | 14 (0-60) | 33.5 (14-25-41.00) | 0.799 |
| Intake of vitamin D (μg/day) (median (IQR)) | 1169 (804-1521.5) | 965 (900-1540) | 1210 (676-1543) | 0.743 |
| Previous or current smoking (N (%)) | 10 (47.6) | 4 (57.1) | 6 (42.9) | 0.659 |
| Current alcohol consumption (units/week) (median (IQR)) | 11 (52.4) | 3 (42.9) | 8 (57.1) | 0.659 |
| Use of EIAEDs (N (%)) | 2 (9.5)               | 1 (14.3)              | 1 (7.1)                     | 0.572                                           |
| Bone-related biochemical parameters at baseline |                        |                       |                             |                                                 |
| Cal (mmol/L) (mean (SD)) | 1.18–1.32             | 1.18 (0.04)           | 1.19 (0.06)                 | 1.18 (0.03)                                     | 0.531 |
| Vitamin D (D₃) (mmol/L) (median (IQR)) | >50 | 71.0 (49–80) | 46.5 (30.75–77.25) | 71.0 (56.0-81.5) | 0.106 |
| TSH (μIU/L) (median (IQR)) | 0.400–4.80 | 1.92 | 2.32 (0.75–2.43) | 1.38 (1.22–2.07) | 0.322 |
| PTH (pmol/L) (median (IQR)) | 1.48–7.63 | 4.30 | 4.20 (3.28–5.57) | 4.34 (3.21–5.35) | 0.856 |
| Phosphate (mmol/L) (median (IQR)) | 0.76–1.41 | 1.11 | 1.14 (1.09–1.26) | 1.09 (0.94–1.21) | 0.360 |
| Mg (mmol/L) (mean (SD)) | 0.71–0.94 | 0.81 (0.07) | 0.81 (0.07) | 0.80 (0.08) | 0.837 |
| ALP (U/L) (mean (SD)) | 35–105 | 60.4 (23.5) | 69.1 (27.2) | 56.0 (21.2) | 0.237 |
| BALP (μg/L) (mean (SD)) | 8.3–29.4 (F) | 15.2 (5.0) | 16.9 (6.7) | 14.3 (4.0) | 0.363 |

Table 1. Osteoporosis-related descriptive data and blood samples for all patients, the placebo group and the intervention group. All data is number and percentage (N (%)), mean and standard deviation (SD) or median and interquartile range (IQR), when appropriate. Statistical differences between the placebo group and intervention group at baseline were for normally distributed data evaluated by Independent Samples t-test, for continuous but not normally distributed variables the Mann-Whitney U test was applied. For categorical data the Pearson chi-square test of Fisher’s exact test was used, when appropriate. Other diseases relevant for osteoporosis was defined as rheumatoid arthritis, type 1 diabetes, hyperthyroidism, hypogonadism, gastro-intestinal disease, chronic liver disease, pulmonary disease, and organ transplantation. Enzyme-inducing antiepileptic drugs (EIAEDs), Thyroid-stimulating hormone (TSH), parathyroid-stimulating hormone (PTH), ionized calcium (Ca), magnesium (Mg), alkaline phosphatase (ALP), bone specific alkaline phosphatase (BALP).
the second $133 \pm 3$ mmol/L (mean ± SD). Plasma [Na] at baseline was
below reference interval for both the placebo p[Na] 133 (130–141)
mmol/L (median (IQR)) and intervention group p[Na] 136.5
(130.75–139.5) mmol/L (median (IQR)), with no difference between the
groups ($p = 0.458$). Both the placebo group and the intervention groups
had decreased levels of plasma aldosterone with a median of 109 pmol/L
and 50.5 pmol/L [ref. interval: 130–859], respectively. Plasma osmo-
lality was decreased for both the placebo, $263.3 \pm 30.7$ mmol/kg (mean ± SD)), and the intervention group, $277.5 \pm 18.1$ mmol/kg (mean ± SD)) [ref. interval: 280–300].

For levels of ionized calcium (CaI) both groups were in the lower end
of the reference range (1.18–1.32 mmol/L), placebo, $1.19 \pm 0.06$ mmol/

![A) Procollagen type 1 N-terminal propeptide (P1NP) and C-terminal cross-linking telopeptide of type 1 collagen (CTX) levels in the patients with age-adjusted reference ranges [11].](image1)

![B) Lumbar L1-L4 T-scores at Baseline](image2)

![Femoral Neck and Femur Total T-scores at Baseline](image3)

**Fig. 2.** Comparison of bone turnover markers and bone mineral density (BMD) in the study population with the background population. a: Procollagen type 1 N-terminal propeptide (P1NP) and C-terminal cross-linking telopeptide of type 1 collagen (CTX) levels in the patients with age-adjusted reference ranges [11]. b: Bone mineral density for the lumbar spine L1-L4, and femoral neck and total femur with cut-off limit for osteoporosis.
L (mean ± SD), and intervention, 1.18 ± 0.03 nmol/L (mean ± SD), with no difference between the groups (p = 0.531). Vitamin D levels (cut-off for sufficient plasma 25-hydroxy vitamin D > 50 nmol/L) in the placebo group, 46.5 (30.75–77.25) nmol/L (median (IQR)), showed hypovitaminosis D as opposed to in the intervention group, 71.0 (56.0–81.5) nmol/L (median (IQR)).

All men, across the two groups, except one (20%) with decreased levels of CTX, had baseline plasma concentration of CTX and P1NP within age-specific reference intervals [11]. Two women (13%) had decreased levels of CTX, and one (6%) had decreased levels of P1NP [11] (Fig. 2a).

Due to the inclusion criteria all patients had site-specific T-score > −2.5 SD, with mean T-scores for the lumbar L1-L4 region, the femoral neck and the total femur of 0.6 ± 0.7 SD (mean ± SD), −0.4±0.2 SD (mean ± SD) and 0.1 ± 0.2 SD (mean ± SD), respectively (Fig. 2b).

3.2. The effects of three months of sodium replacement therapy

Seven patients were randomized to the placebo group and 14 to the intervention group. Of the 14 patients in the intervention group 7 patients were treated with 3g of salt daily, 3 patients with 9g of salt daily and 4 patients with 9g of salt and 20 mg of furosemide daily. Through the experiment there was only a significant difference in p[Na] at visit 3 between the placebo group 133.7 ± 4.9 mmol/L (mean ± SD) and intervention group 138.8 ± 3.8 (mean ± SD) (p = 0.016) (Fig. 3). Following 3 months of intervention the p[Na] for the placebo group was 137.1 ± 3.7 mmol/L (mean ± SD) and for the intervention group 137.7 ± 2.9 mmol/L (mean ± SD), with no statistical difference (p = 0.706). There was a borderline significant difference in p[Na] following the intervention from baseline to visit 6 for the intervention group (p = 0.050). The compliance in the placebo and intervention group was ≥77% and ≥85%, respectively.

The results of the BTM and DXA scans are summarized in Table 2. There was no difference between the groups in any of the parameters. Furthermore, there was no difference from baseline to visit 6 in the placebo or the intervention groups in any of the parameters (Fig. 4). When looking specific at visit 3, where a significant difference in p[Na] was present, there is no difference between the placebo and intervention groups nor any changes from baseline to visit 6 (Fig. 4 a). In the placebo groups there was an increase in CTX (p = 0.028) but not in P1NP (p = 0.310) from baseline to visit 3 (Fig. 4a).

The factorial repeated measures ANOVA shows that, when testing the effect of the intervention on all measures of the same variable over time, there was no effect of the intervention that could not be explained by time, except for femoral neck BMD, where there was an independent effect of the intervention over time (p = 0.040) (data not shown).

For the bone-related biochemical parameters the placebo group had a significantly higher plasma phosphate level, 1.24 ± 0.20 mmol/L (mean ± SD), than the intervention group, 1.09 ± 0.12 mmol/L (mean ± SD) (p = 0.044), at visit 6. Levels of TSH increased from baseline to visit 6 in the placebo group, 2.32 (0.75–2.43) 10^{-3} IU/L (median (IQR)), to, 3.50 (0.89–4.37) 10^{-3} IU/L (median (IQR)) (p = 0.028), while levels of PTH decreased from 4.20 (3.28–5.57) pmol/L (median (IQR)) to 3.60 (2.40–4.25) pmol/L (median (IQR)), (p = 0.018). For the intervention group there was a significant increase in plasma PTH levels from 4.34 (3.21–5.35) pmol/L (median (IQR)) to 5.60 (3.54–6.22) pmol/L (median (IQR)) (p = 0.041) and in plasma osmolality from 278 ± 18 mmol/kg (mean ± SD)) to 291 ± 17 mmol/kg (mean ± SD)) (p = 0.027). There was no difference in CaI between the groups at visit 6 nor when comparing within the groups from baseline to visit 6 (data not shown).

Comparing the 24-h urine sample collected at baseline to the one collected at visit 6, the intervention group had a significant increase in U [Na] from 80.9 ± 3.51 mmol/day (mean ± SD) at baseline to 123.9 ± 40.55 mmol/day (mean ± SD)) at visit 6 (p = 0.001), whereas the placebo groups did not change from baseline 83.1 ± 26.9 mmol/day (mean ± SD) to visit 6 66.7 ± 26.6 mmol/day (mean ± SD) (p = 0.159) (data not shown).

4. Discussion

To our knowledge this is the first study investigating how hyponatremia affects BTM in patients with epilepsy and whether salt supplements have any effect on BTM in patients with epilepsy and hyponatremia. We did not find elevated levels of the BTM, P1NP and CTX, in patients with epilepsy and chronic hyponatremia and no osteoporosis (Fig. 2a) [11]. Elevated BTM in patients with epilepsy have been found previously however, not in relation to hyponatremia but possibly do to treatment with AEDs [12,13]. There was no change following 3 months of salt replacement in P1NP or CTX levels between the groups nor any changes from baseline to visit 6 (Fig. 4a). We did find a significant difference between baseline and visit 3 in P1NP and CTX levels for the intervention group but only in P1NP levels in the placebo group. There was no difference in T-scores in the lumbar L1-L4 region,

![Fig. 3. Plasma [Na] for the placebo (dotted line) and intervention group (full line) from pre-inclusion values (Pre-Na 1 and Pre-Na 2) to visit 6. Circles represents mean value and error bars standard error of the mean (SEM). The dashed line represents the definition of hyponatremia (p[Na] 136 mmol/L). The asterix represents a significant difference between intervention and placebo group.](image-url)
Hyponatremic-induced osteoporosis in patients with epilepsy

Studies have shown that hyponatremia is associated with fractures and decreases BMD in the general population [4-6,14,15]. Metabolic bone disease has long been known as a grave and frequent co-morbidity to epilepsy and so has chronic hyponatremia, which is associated with certain AEDs [7,16,17]. Not until recently have the two however been combined and the occurrence of moderate and severe hyponatremia [p-Na<129 mmol/L] demonstrated as a risk factor for osteoporosis in patients with epilepsy [5]. Our finding showed no difference in BTM at baseline compared to age-specific reference intervals [11]. However, it should be noted that the patients included in the study did not have osteoporosis, and it is possible that we have selected patients with excellent bone health or a group of “non-progressors” that are less susceptible to adverse effects on bone. The high occurrence of metabolic bone disease in patients with epilepsy had been associated with the use of AEDs and especially those that are EIAEDs [18-21]. These drugs, such as carbamazepine, oxcarbazepine and eslicarbazepine acetate are hypothesized to induce isoenzymes of the livers cytochrome P450 enzyme-system and thereby accelerating hydroxylation of vitamin D and often resulting in hypovitaminosis D in the treated patients [22,23]. This is supported by our finding that the placebo patients had low levels of vitamin D and that the mean level of ionized calcium for all patients was equivalent to the lower end of the reference interval. An interesting overlap exists between the EIAEDs, primarily associated with metabolic bone disease and the AEDs that frequently cause chronic hyponatremia. This complicates the interpretation as it could both mean that hyponatremia plays an independent role in the development of metabolic bone disease or simply serves as a surrogate marker for patients at high risk of developing osteoporosis due to other mechanisms.

A valid question is whether the study population truly represents patients with chronic hyponatremia. The patients were included based on two hyponatremic measurements (Fig. 3), however both the intervention group and the placebo group increased in p[Na] up to baseline indicating that two consecutive measurements of hyponatremia are inadequate to detect patient with chronic hyponatremia and our patients may therefore not be truly chronic hyponatremic. Our patients p[Na] may merely fluctuate around the lower end of the reference interval which is an important point when considering that hyponatremia-induced osteoporosis in patients with epilepsy was demonstrated in patients with moderate and severe hyponatremia (p ≤ 129 mmol/L). The same is found in preclinical studies where the effects in rats is found at p[Na] around 110 mmol and in osteoclasts at media [Na] at 131 mmol/L and more so at media [Na] at 129 mmol/L and lower [5,24]. Furthermore, it is possible that hyponatremia in general does not pose such a great problem in patients with epilepsy as anticipated, which is further illustrated by the fact that we were hardly able to include any patients with moderate or severe hyponatremia.

The question of timely relation is equally important when dealing with hyponatremia-induced osteoporosis. The chronic hyponatremia and the development of osteoporosis are two interlinked chronic conditions evolving over long time periods [6]. It is therefore possible that including patients based on two prior measurements of hyponatremia does not accurately represent the time aspect required for hyponatremia-induced osteoporosis to evolve. This might be a reason why no effects on BTM is detectable.

4.2. The effects of salt supplements on BTM and BMD in patients with epilepsy and hyponatremia

The occurrence of hyponatremia-induced osteoporosis has mainly been shown as associations in population-based studies limiting the possibility for drawing conclusions about causality and whether the changes are reversible. There are to our knowledge no studies investigating the reversibility of hyponatremia-induced osteoporosis. In the current study we found no effect on BTM or BMD evaluated by DXA scan following three months of salt supplements. However, when looking at the p[Na] for the two groups through the experiment (Fig. 3) there is a clear increase in the intervention group as well as the placebo group, which spontaneously increase in p[Na] over time. However, the timely difference between the two should be noted, the intervention group increases in p[Na] immediately following the intervention and are normonatremic already at visit 2, whereas the placebo group’s increase to normonatremia does not occur until visit 5. This could suggest that the patients included were not truly chronic hyponatremic, however there is no strict definition of chronic hyponatremia in this context. An important point, however, is the increase in urine sodium output in the femoral neck or total femur between the groups following the intervention or when comparing within the groups from baseline to visit 6 (Fig. 4b). In the factorial repeated measures ANOVA there was a significant effect of the intervention over time for BMD at the femoral neck.

4.1. Hyponatremia-induced osteoporosis in patients with epilepsy

Table 2. Levels of procollagen type 1 N-terminal propeptide (P1NP) and C-terminal cross-linking telopeptide of type 1 collagen (CTX) expressed as median and interquartile range (IQR). Lumbar L1-L4, femoral neck T-score, femoral neck T-score and total femur T-score expressed as mean and standard deviation (SD). Statistical differences between the placebo group and intervention group were for P1NP and CTX tested with a Mann-Whitney test and for Lumbar L4, femoral neck and total femur an Independent Samples t-test was done.

| Visit | Placebo group | Intervention group | Statistical significance | Placebo vs Intervention |
|-------|---------------|--------------------|--------------------------|-------------------------|
| 1     | P1NP (ng/L)   | CTX (ng/L)         |                          |                         |
|       | (median (IQR))| (median (IQR))     | 28.3 (23.9-36.2)         | 0.079                   |
|       |               |                    | 37.1 (31.8-43.3)         |                         |
| 2     | P1NP (ng/L)   | CTX (ng/L)         |                          |                         |
|       | (median (IQR))| (median (IQR))     | 131 (80-273)             | 0.172                   |
|       |               |                    | 212 (144.9-315.3)        |                         |
| 3     | P1NP (ng/L)   | CTX (ng/L)         |                          |                         |
|       | (median (IQR))| (median (IQR))     | 0.0 (1.0)                | 0.169                   |
|       |               |                    | 0.9 (1.4)                |                         |
| 4     | Femoral Neck (SD) | Femoral Neck (SD) | -0.7 (1.1)              | 0.342                   |
|       | (mean (SD))   | (mean (SD))        | -0.3 (0.9)               |                         |
| 5     | Total Femur (SD) | Total Femur (SD)  | 0.0 (1.1)                | 0.676                   |
|       | (mean (SD))   | (mean (SD))        | 0.2 (1.0)                |                         |
intervention group from baseline to visit 6, which is not found in the placebo group. This demonstrates that the intervention group did have an increase in salt intake and the placebo group did not.

The unexpected increase in p[Na] in the placebo group limits the possibilities for any general conclusions on the effects of salt supplements in patients with epilepsy and chronic hyponatremia. Furthermore, it should be noted that a time span of 3 months, with great likelihood, is far too short a period to see any changes in BMD evaluated by DXA scan. Evaluating changes in BMD by DXA usually requires time periods of two years or more. Our finding of a difference in BTM between baseline and visit 3 within the intervention and placebo group suggest that BTM might be affected by changes in p[Na] levels, however no generalizable conclusion should be based on this finding.

Our finding of a significant interaction over time on femoral neck BMD in the factorial repeated measures ANOVA is most likely a random finding. The difference is derived mainly by a decrease in the placebo groups BMD and not by an increase in BMD in the intervention group. However, the decrease in BMD cannot be attributed to hyponatremia hence the previously mentioned increase in p[Na] in the placebo groups. Furthermore, it is unlikely that an effect should be seen in femoral neck and not also in total femur just as any changes in BMD in three months is unlikely to be found.

A valid point to consider is the choice of examining effects of hyponatremia by using BTM. To our knowledge no literature exists on how BTM are affected by hyponatremia or how fast any changes would be detectable thus further studies are warranted.

4.3. Strengths and limitation

The study’s main strength is the thorough characterization of the intervention group from baseline to visit 6, which is not found in the placebo group. This demonstrates that the intervention group did have an increase in salt intake and the placebo group did not.

The unexpected increase in p[Na] in the placebo group limits the possibilities for any general conclusions on the effects of salt supplements in patients with epilepsy and chronic hyponatremia. Furthermore, it should be noted that a time span of 3 months, with great likelihood, is far too short a period to see any changes in BMD evaluated by DXA scan. Evaluating changes in BMD by DXA usually requires time periods of two years or more. Our finding of a difference in BTM between baseline and visit 3 within the intervention and placebo group suggest that BTM might be affected by changes in p[Na] levels, however no generalizable conclusion should be based on this finding.

Our finding of a significant interaction over time on femoral neck BMD in the factorial repeated measures ANOVA is most likely a random finding. The difference is derived mainly by a decrease in the placebo groups BMD and not by an increase in BMD in the intervention group. However, the decrease in BMD cannot be attributed to hyponatremia hence the previously mentioned increase in p[Na] in the placebo groups. Furthermore, it is unlikely that an effect should be seen in femoral neck and not also in total femur just as any changes in BMD in three months is unlikely to be found.

A valid point to consider is the choice of examining effects of hyponatremia by using BTM. To our knowledge no literature exists on how BTM are affected by hyponatremia or how fast any changes would be detectable thus further studies are warranted.

5. Conclusion

In conclusion, in patients with epilepsy and hyponatremia and no osteoporosis, 3 months of treatment with sodium tablets did not result in significantly higher sodium concentrations compared to the placebo group but instead higher concentration of sodium in the urine. There were also no beneficial effects on BTM or BMD. This suggests, that sodium replacement therapy is insufficient when trying to restore the sodium plasma balance in patients with epilepsy. Future studies are
warranted to further understand the meaning of hyponatremia in patients with epilepsy and its consequences for bone loss.

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**Declaration of competing interest**

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