Antidepressants and the risk of hyponatremia: a Danish register-based population study

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ABSTRACT

Objective: To examine the association between classes of antidepressants and hyponatraemia, and between specific antidepressants and hyponatraemia.


Setting: The North Denmark Region.

Participants: In total, 638 352 individuals were included.

Primary and secondary outcome measures: Plasma sodium was obtained from the LABKA database. The primary outcome was hyponatraemia defined as plasma sodium (p-sodium) below 135 mmol/L and secondary outcome was severe hyponatraemia defined as plasma sodium below 130 mmol/L. The association between use of specific antidepressants and hyponatraemia was analysed using multivariable Poisson regression models.

Results: An event of hyponatraemia occurred in 72 509 individuals and 11.36% (n=6476) of these events happened during treatment with antidepressants. Incidence rate ratios and CIs for the association with hyponatraemia in the first p-sodium measured after initiation of treatment were for citalopram 7.8 (CI 7.42 to 8.20); clomipramine 4.93 (CI 2.72 to 8.94); duloxetine 2.05 (CI 1.44 to 2.92); venlafaxine 2.90 (CI 2.43 to 3.46); mirtazapine 2.95 (CI 2.71 to 3.21); and mianserin 0.90 (CI 0.71 to 1.14).

Conclusions: All antidepressants except mianserin are associated with hyponatraemia. The association is strongest with citalopram and lowest with duloxetine, mirtazapine and mianserin.

BACKGROUND

The consumption of antidepressants has increased significantly in most Organisation for Economic Co-operation and Development (OECD) countries since 2000. Denmark is among the OECD countries with the highest consumption.1

Hyponatraemia is one of the many well-known side effects of antidepressants.2 5 Even mild hyponatraemia is associated with instability and falls, reduced cognitive function, osteoporosis and increased morbidity and mortality.4–13

The kidneys control the renal excretion of water and electrolytes and keep serum osmolality and sodium within a narrow range. Several hormones influence the renal handling of water and electrolytes, the most important being arginine vasopressin, also known as antidiuretic hormone (ADH).

ADH binds to V2 receptors in the kidney, and thereby leads to an increase in reabsorption of water in the collecting ducts. The main stimuli for ADH is plasma osmolality and low plasma volume, but a number of non-specific stimuli like stress and nausea can also result in elevated ADH concentrations.14 15

The presence of hypotonic hyponatraemia always implies excess of water, and hyponatraemia essentially is a water balance disorder.16

In most cases of hypotonic hyponatraemia, ADH secretion is increased due to one of the following reasons: (1) appropriate response to decreased effective plasma volume as in hypovolaemia, cirrhosis and heart failure; (2) ectopic production as seen in neoplastic diseases resulting in syndrome of inappropriate ADH (SIADH); or (3) syndrome of inappropriate antidiuresis (SIAD) either due to a drug induced increase in ADH production (SIADH) or an enhanced renal response to ADH, the nephrogenic syndrome of antidiuresis.16
In a cross-sectional study of elderly users of antidepressants, Mannesse et al studied the mechanisms of hyponatremia in 34 patients with hyponatremia. Data on serum and urinary sodium were available in 30 patients, and of these 23 were classified as having either SIADH or probable SIADH. However, only in a minority of these patients was ADH not suppressed, and the authors concluded that the mechanism of hyponatremia produced by antidepressants might be either an elevated ADH or an increased renal response to ADH. Some previous studies have examined the relationship between hyponatremia and antidepressants. However, most studies are small and observational and only few have had the power to examine whether specific antidepressants carry a higher or lower risk of hyponatremia. The number of case reports and small observational studies with hyponatremia concerning selective serotonin reuptake inhibitors (SSRIs) is substantially higher than the number of case reports and observational studies with other antidepressants, but it is not clear whether this is due to a true difference in incidence of hyponatremia.

A recent review concluded that current evidence suggests a relatively higher risk of hyponatremia with SSRIs and venlafaxine compared to tricyclic antidepressants (TCA) and mirtazapine, but for several antidepressants, data were insufficient to determine the risk of hyponatremia.

The aim of this study was to examine the association between specific classes of antidepressant medication and hyponatremia and between specific drugs within these classes and hyponatremia. We examined the most commonly used drugs among TCA, SSRI, serotonin–norepinephrine reuptake inhibitors (SNRIs) and noradrenergic and specific serotonergic antidepressants (NaSSAs).

METHODS
By the time of birth or immigration, every Danish citizen gets a permanent personal identification number through the Danish Civil Registration System. This makes it possible to link information from multiple registers at the individual level. The current study is a register-based population study where we linked exposure to antidepressant therapy at the individual level. The Danish National Patient Register, the Danish register of Medicinal Products Statistics and laboratory data. Since 1968, the Danish Civil Registration has been complete regarding information about dates of birth and death for all Danish citizens. The Danish National Patient Register contains records of all hospitalisations since 1977. The Danish Register of Medicinal Products Statistics consists of all prescription claims from Danish pharmacies since 1995, including medicine strength, quantity and date. The prescriptions are classified according to the Anatomical Therapeutic Chemical system (ATC).

Laboratory data
Information concerning p-sodium was obtained from the LABKA database containing results of all blood samples from hospitalised patients and outpatients submitted for analyses to hospital laboratories in the Northern and Central Denmark. From the database, information on the date and result of the analysis was retrieved.

Population
All individuals born before 1998 and living in The North Denmark Region in the period from 1 January 1998 to 31 December 2012 were included. Individuals born after 1 January 1998 were excluded.

We continuously included individuals who immigrated to The North Denmark Region in the study period. Individuals were followed until end of study, emigration, death or an event of hyponatremia, whichever occurred first.

Outcome assessment
Primary outcome was an event of hyponatremia defined as plasma sodium (p-sodium) below 135 mmol/L. Secondary outcome was severe hyponatremia defined as p-sodium below 130 mmol/L. Since the prescription register does not include medicine administered during hospital stay, hyponatremia during hospitalisation was excluded; however, the first blood test after admission was included if it was drawn on the day of admission.

Antidepressant therapy
We decided to include only the most commonly used antidepressants in Denmark. Information regarding antidepressant therapy was retrieved from the Danish National Prescription Register and included the following drugs (ATC codes in brackets): Amitriptyline (N06AA09), Clomipramine (N06AA04) and Nortriptyline (N06AA10), Citalopram (N06AB04), Escitalopram (N06AB10), Fluoxetine (N06AB03), Paroxetine (N06AB05) and Sertraline (N06AB06), Duloxetine (N06AX21), Venlafaxine (N06AX16) and Mirtazapine (N06AX11), Mianserin (N06AX03). Treatment was defined in a time-dependent manner; thus, we only considered individuals as exposed if they were covered by a prescription claim. Every prescription in the prescription register contains data on tablet strength and number of tablets. To determine exposure status, we defined minimum, maximum and typical daily dosage for each drug and tablet strength. On the basis of this information, we calculated length of treatment by creating an algorithm that used up to five consecutive prescriptions. These calculations were exclusively based on previous prescription claims. When it was a patient’s first prescription claim, the typical daily dosage was used and the amount of claimed medications was divided by the daily dosage to determine length of treatment. When a patient who was covered by a previous prescription claimed a new one, we calculated a new daily dosage by dividing the amount of claimed medications in the previous period by the time between prescription claims. If the calculated new daily dosage was higher than the stated maximum dosage, this dosage was used instead. It was presumed that the remaining tablets were kept and...
consumed immediately after the last prescription coverage ended.

**Comorbidities and concurrent pharmacotherapy**

The following comorbidities were identified as possible confounders of the association (The International Statistical Classification of Diseases and Related Health Problems 10th Revision, (ICD10) codes in brackets): chronic lung (J00–J18, J40–J47, J60–J70, J80, J82–J84, J95–J99), liver (K70–K77), kidney (N18, N19), and intracranial (I60–I69) diseases, heart failure (I50) and cancer (C). These comorbidities were accessed from the Danish National Patient Register. All comorbidities were identified from 1993 and throughout the study period. Furthermore, we identified treatment with antiepileptic drugs or diuretics since these treatments could potentially confound the association. As for the antidepressants drugs, we defined treatment in a time-dependent manner, and treatment length was calculated in the same way.

**Statistics**

The incidence rate ratio (IRR) of hyponatremia associated with exposure to antidepressant therapy was analysed in a time-dependent multivariable Poisson regression model. Before the analyses, we divided observational time into 1 year timebands. Age was updated at the beginning of each timeband and afterwards categorised into 10-year intervals. At the time of a new diagnosis or change in medication, the observational time was divided into a ‘pre’ and a ‘post’ period, ensuring a timely correct classification of comorbidities and exposure to pharmacotherapy. Hereafter, binary variables (current use or no use) were created for each antidepressant. Furthermore, we divided time after first prescription into five timebands: 0–14, 15–30, 31–90, 91–180 and >180 days.

We performed five different multivariable Poisson analyses: (1) estimating IRR for having p-sodium <135 with current use of antidepressants; (2) estimating IRR for having p-sodium <130; (3) estimating IRR for having p-sodium measured after initiating treatment with the different antidepressants; (4) estimating IRR for hyponatremia after initiating treatment (in each of the previously mentioned timebands) with one of the included antidepressants. Owing to few events in some of the strata in this analysis, we explored the association between hyponatremia and four classes of antidepressants (TCAs, SSRIs, SNRIs and NaSSAs); (5) estimating IRR for hyponatremia in the first p-sodium measurement after initiating treatment. For the unexposed individuals we used their first p-sodium measurement after inclusion. Lastly, we made a subgroup analysis, only including individuals who had p-sodium measured. To avoid conditioning on the future, this was done in a logistic regression analysis instead of a time-dependent regression. In this analysis, we estimated the ORs for hyponatremia with current use of antidepressants, as for the Poisson regression analyses an individual was only considered exposed if covered by a prescription claim. All analyses were adjusted for age (in 10-year intervals), sex, comorbidities and concurrent pharmacotherapy with diuretics, antiepileptic drugs and other antidepressants; furthermore, all Poisson regression analyses were adjusted for time.

All analyses were performed using SAS V.9.2 (SAS Institute).

**RESULTS**

In total, 638 352 individuals were included in the study, and they were on average followed for 10.98 years. Of these, 15.50% (n=98 976) initiated treatment with an antidepressant drug. An event of hyponatremia occurred in 72 509 (11.36%) individuals, and 8.90% (n=6476) of these events happened during treatment with antidepressants. Population characteristics for individuals exposed to the different antidepressants during follow-up and for unexposed individuals are shown in table 1.

During follow-up, 60.28% (n=103 109) of the study population had their p-sodium measured. 75% of individuals who started treatment with antidepressants had p-sodium measured at some point during follow up, but only 31% had p-sodium measured within the first 6 months after initiation of treatment. The percentage of individuals who had p-sodium measured within 6 months after initiating treatment varied between drugs; the data are available in online supplementary appendix A.

**Figure 1** shows the association between use of specific antidepressants and an event of hyponatremia and the chance of having serum sodium measured while being in treatment with an antidepressant. Drugs associated with a high chance of having p-sodium measured were also strongly associated with the finding of hyponatremia. We therefore performed a logistic regression analysis in the subpopulation of individuals who had p-sodium measured. This subpopulation included 403 109 individuals; 20.25% of these (n=81 618) were at some point in treatment with an antidepressant. This subpopulation had their p-sodium measured. Figure 2 shows the association between use of antidepressants and hyponatremia in this subpopulation. The pattern of associations between use of specific antidepressants and hyponatremia in the subpopulation is very similar to the results in the total population. Clomipramine (OR 4.15, CI 2.56 to 6.74) and nortriptyline (OR 2.44, CI 1.99 to 3.00) are strongly associated with hyponatremia, but there are few events leading to uncertainty in data and wide CIs. Mianserin (OR 1.25, CI 0.96 to 1.62), mirtazapine (OR 1.53, CI 1.39 to 1.69) and escitalopram (OR 1.12, CI 0.98 to 1.29) have a weaker association with hyponatremia than with citalopram (OR 2.34, CI 2.20 to 2.48). In order to avoid possible bias due to some individuals having p-sodium measured several times and thereby possibly increasing the chance of catching an event of hyponatremia, we performed a Poisson analysis to estimate IRR of hyponatremia in the first measurement of p-sodium after having initiated treatment with an antidepressant. This...


**DISCUSSION**

In this large register-based study including ~10% of the total Danish population living in a geographically well-defined area (The North Denmark Region), we investigated the association between different antidepressants and incidence of hyponatremia. We observed that all the included antidepressants except mianserin were significantly associated with hyponatremia.

For all drugs, we found the highest IRR for hyponatremia during the first 14 days of treatment. This is consistent with previous studies and, with the assumed SIAD mechanism since the distribution and effect of ADH, occurs within a few hours. This has been demonstrated in studies concerning thiazide where changes in plasma osmolality and p-sodium were observed within 6 h after a single dose of thiazide.

**Definition of hyponatremia**

In this study, we defined hyponatremia as p-sodium <130 mmol/L. We have chosen this definition due to the fact that even mild hyponatremia during recent years has been shown to be of clinical importance.

Recently, an expert panel judged even mild hyponatremia to be of clinical importance due to association with adverse outcomes including mortality. A German study found mean p-sodium 131 mmol/L in psychiatric inpatients who were deemed to have symptomatic hyponatremia. Symptoms were primarily neuropsychiatric such as confusion, dizziness, gait impairments and falls. Hyponatremia due to any cause demonstrates an imbalance in the homeostatic system that normally controls p-sodium, and therefore even mild hyponatremia warrants caution when considering medical treatment and medical advice concerning liquid and nutrition. One study even showed the level of p-sodium when patients were admitted to hospital to be associated with in-hospital mortality with a cut-off at p-sodium of <138 mmol/L.

**Comparison of drug classes**

We have performed several analyses in order to minimise the bias that is inherent in an observational study. Analysis showed a slightly different pattern with highest IRR for citalopram (7.80, CI 7.42 to 8.20); the results are shown in figure 3. We also investigated the occurrence of hyponatremia relative to the time of initiating treatment with an antidepressant. For all antidepressants, the association between treatment and occurrence of hyponatremia was strongest in the first 2 weeks after initiation and thereafter gradually declined; the data are shown in figure 4.

Finally, we looked at the occurrence of severe hyponatremia, defined as p-sodium <130 mmol/L. The pattern of association was not different from mild hyponatremia, but due to fewer events the results are less clear with an overlap of CIs for most drugs; the data are available in online supplementary appendix B.
Most of the analyses presented show similar results, but there are slight differences between the analyses presented. When comparing the specific drugs, we consider the analysis of association between drug treatment and hyponatremia in the first p-sodium measured to give the most valid and less biased results (Figure 3). In this analysis, we have minimised the bias that may result from some individuals having p-sodium measured many times and some having p-sodium measured just once. Furthermore, on the basis of knowledge concerning pathophysiology of hyponatremia, we would expect hyponatremia due to an antidepressant to be present in the first p-sodium measured. TCAs and SSRIs in this analysis show IRRs for hyponatremia at the same level. The initial reports of antidepressants associated with hyponatremia concerned TCA, but only a few later studies have compared the incidence of hyponatremia in TCAs with other antidepressants. In a large register-based study, Coupland et al found a weaker association between hyponatremia and TCAs compared to SSRIs. However, Coupland’s study included only persons above the age of 65. We cannot therefore directly compare the results of Coupland’s study with those of our study.

The SNRIs including duloxetine and venlafaxine have lower IRRs. This is contradictory to another study that examined venlafaxine. However, this study had a very different population (patients in a psychiatric department) and only a few venlafaxine cases. To the best of

![Figure 1](http://bmjopen.bmj.com/) Incidence rate ratios of having an episode of hyponatremia while being in treatment with an antidepressant and incidence rate ratios of having p-sodium measured while being in treatment with an antidepressant.

![Figure 2](http://bmjopen.bmj.com/) ORs for hyponatremia for specific antidepressants in the subpopulation of individuals (n=403 109) who had p-sodium measured.
our knowledge, duloxetine has previously been studied only in a case report including five cases.35

The NaSSAs include mianserin and mirtazapine. The IRR for mirtazapine was similar to that for duloxetine and venlafaxine. In contrast, mianserin was not associated with hyponatremia. On the basis of case reports and results from case series, it is generally assumed that mirtazapine has a lower risk of hyponatremia than SSRIs.20 24 Our results support this assumption since the IRR for mirtazapine is significantly lower compared to SSRIs, but not compared to SNRIs. Mianserin and mirtazapine both block α2-adrenergic receptors, thereby increasing the level of norepinephrine. Interestingly, the α2-adrenergic system has been shown to be involved in polydipsia in patients with schizophrenia in a small pilot study where mianserin was used as treatment for polydipsia.36 The fact that mirtazapine also blocks postsynaptic serotonergic receptors might explain the difference between mianserin and mirtazapine.

Mechanism of antidepressant-induced hyponatremia

It is assumed that the mechanism of antidepressant hyponatremia is SIAD with or without elevated ADH.17 However, no studies have explored the influence of antidepressant-induced hyponatremia on thirst. One study has demonstrated that there is a downward setting of the osmotic threshold for thirst in patients with SIADH.37 Thiazide induced hyponatremia, which is also believed to be induced by an SIADH mechanism, has been associated with increased thirst.30 Combined with the assumption that the α2-adrenergic system is involved in polydipsia, this may explain why mianserin is not

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<table>
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<th>Antidepressants</th>
<th>No. of events</th>
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<td>2.90 [2.43-3.46]</td>
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<td>Mianserin</td>
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<td>0.90 [0.71-1.14]</td>
</tr>
<tr>
<td>Mirtazapine</td>
<td>652</td>
<td>2.95 [2.71-3.21]</td>
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*Adjusted for age, time, sex, comorbidities and concurrent pharmacotherapy with diuretics, antiepileptic drugs and other antidepressants.

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**Figure 3** Incidence rate ratios for hyponatremia in the first measurement of p-sodium level after initiating treatment with a specific antidepressant.

**Figure 4** Incidence rate ratios for hyponatremia in timebands after initiating antidepressant therapy with drugs belonging to tricyclic antidepressants (TCA), selective serotonin reuptake inhibitors (SSRI), serotonin–norepinephrine reuptake inhibitors (SNRIs), and noradrenergic and specific serotonergic antidepressants (NaSSAs).
associated with hyponatremia and mirtazapine to a lesser degree than other antidepressants.

**Clinical relevance and implications**

Incidence of hyponatremia increases with age, which is not surprising seen in the light of age-associated changes in water homeostasis.\(^3\)\(^6\)\(^8\)\(^9\) Even mild hyponatremia influences performance on cognitive as well as balance and mobility testing in elderly individuals.\(^1\)\(^5\) Chronic hyponatremia is associated with osteoporosis and high fracture risk.\(^40\)\(^41\) Furthermore, mild hyponatremia is an indicator of a water balance disorder, which under different circumstances may develop into severe and life-threatening hyponatremia. This is underlined by the fact that patients, who for any reason have had hyponatremia once, are likely to have recurrent hyponatremia as a response to different diseases and stimuli, and lower baseline p-sodium is a risk factor for developing hyponatremia.\(^42\)\(^43\) Even mild hyponatremia therefore warrants cautious evaluation and, if possible, should be abolished. The results from our study can inform the choice of antidepressants, especially when treating patients who are known to have a high risk of hyponatremia like, for instance, elderly people and persons who have previously had hyponatremia.

**Strengths and limitations**

The main strengths of this study are the large and well-defined population and high-quality registers, which made it possible to examine specific antidepressants. The limitations are mainly those that are inherent in an observational study, that is, residual confounding due to differences between patients and controls not accounted for because of lack of information on other variables that could possibly influence the risk of hyponatremia like, for instance, eating and drinking habits. Although we have adjusted for comorbidities and drugs known to be able to induce hyponatremia, we cannot fully exclude residual confounding due to other reasons for hyponatremia. Not all persons who were treated with antidepressants had p-sodium measured and measurement may have been performed due to symptoms of hyponatremia. On the other hand, the symptoms of mild hyponatremia are often not recognised, and in elderly persons they may even be mistaken for symptoms of normal ageing (forgetfulness, dizziness, drowsiness). If the patients who had p-sodium measured represent a selection of more symptomatic patients, we would overestimate the rates of hyponatremia. However, an analysis including only those individuals in the population who had p-sodium measured showed similar results, which means that comparison between drugs should be safe. Thus, we consider the comparison of drugs valid, but we cannot be certain about the absolute level of IRRs for the specific drugs. The study population is Caucasian, and therefore generalisation of the results to different populations should be done with caution.

Comedication can increase or decrease the functional activity of hepatic cytochrome enzymes (CYP) and, as such, inhibit or induce the metabolic pathway of antidepressants.\(^44\)\(^45\) Since the CYP activity determines the plasma concentration of antidepressants and/or their metabolites that will be achieved as a function of the dose administered, drug–drug interactions may change the risk of dose-dependent adverse drug events. In this study, potential drug–drug interactions were not taken into account.

**Conclusion**

There is an increased risk of hyponatremia for all included antidepressants, except for mianserin. The strongest association between hyponatremia and antidepressants is found in SSRLs. The association with SNRLs is slightly lower and the weakest association is with NaSSAs. The risk of hyponatremia is highest within the first 2 weeks of treatment. Considering the possible serious adverse effects of chronic hyponatremia, we recommend routine measurement of p-sodium shortly after initiating treatment with any antidepressant.

**Contributors** KBL-M, EAH, CT-P and GG designed the research. CT-P and GG were involved in the acquisition of data. KBL-M performed the research. All authors analysed the data. EAH, AHH and KBL-M wrote the manuscript. All authors contributed to a critical revision of the manuscript.

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**Competing interests** CT-P declares grants and personal fees from Cardiome, Merck, Sanofi and Daiichi, as well as grants from BMS, all outside the submitted work. GG is supported by an unrestricted clinical research scholarship from The Novo Nordisk Foundation. All other authors declare that they have no relationships with companies that might have an interest in the submitted work in the previous 3 years and none of the authors have non-financial interests that may be relevant to the submitted work.

**Ethics approval** This study was approved by the Danish Data Protection Agency (J.nr. 2007-58-015/I. Suite nr: Region H 00916 GEH-2010-001). In Denmark, no ethics approval is required for retrospective register-based studies.

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Data sharing statement** Data stem from Danish National registers and cannot be shared.

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