Clinical experience with Tolvaptan outpatient use. Cost and effectiveness in 9 cases

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Abstract

Introduction: Tolvaptan introduction has constituted the main therapeutic novelty in the management of hyponatremia in recent years. Objective: To describe the experience with this drug at Complejo Asistencial Universitario de León, Spain. Method: Retrospective, observational study of tolvaptan outpatient use in a tertiary care hospital from March 2014 to August 2017. Results: A total of 9 patients were treated with tolvaptan in the outpatient setting. Eunatremia was reached in 24 hours by 23.1 %. After tolvaptan administration, a reduction in days of hospitalization was recorded (361 vs. 70; p = 0.007), especially in those days of hospitalization that were attributable to hyponatremia (306 versus 49; p = 0.009). Conclusions: Long-term use of tolvaptan appears to be safe and is associated with a decrease in days of hospitalization.

KEY WORDS: Tolvaptan. Hyponatremia. Sodium. Syndrome of inappropriate antidiuretic hormone secretion.

Experiencia clínica con el uso ambulatorio de tolvaptan. Costes y efectividad en nueve casos

Resumen

Introducción: La introducción de tolvaptan ha supuesto la principal novedad en el tratamiento de la hiponatremia en los últimos años. Objetivo: Describir la experiencia con tolvaptan en el Complejo Asistencial Universitario de León, España. Método: Estudio observacional retrospectivo de utilización ambulatoria de tolvaptan en un hospital de tercer nivel, de marzo de 2014 a agosto de 2017. Resultados: Fueron tratados con tolvaptan de forma ambulatoria nueve pacientes, 23.1 % alcanzó eunatremia en 24 horas. Posterior a la administración de tolvaptan se registró reducción en días de hospitalización (361 versus 70, p = 0.007), especialmente por hiponatremia (306 versus 49, p = 0.009). Conclusiones: El uso a largo plazo de tolvaptan parece ser seguro y se relaciona con descenso en los días de hospitalización.


Introduction

Hyponatremia, which is a decrease in serum sodium concentration below 135 mmol/L, is the most prevalent electrolyte disorder in the outpatient and hospital setting.1 Its main cause is inadequate secretion of antidiuretic hormone, pathophysiologically caused by inability to suppress vasopressin secretion.2 Mild or moderate hyponatremia (serum sodium [SNa] = 120-135 mmol/L), which in many cases is chronically maintained, is associated with an increase in hospital3 and outpatient mortality,4 as well as with...
an increase in instability, falls and fractures. The need to address its treatment has been emphasized in various consensus documents and clinical practice guidelines, which propose water restriction or increasing aquaresis with loop diuretics, urea or vasopressin V2 receptor antagonists, which constitute the main therapeutic novelty in its management; in Europe, tolvaptan (TV) is available.

**Contribution to the scientific literature**

The purpose of this study was to assess costs and effectiveness of tolvaptan outpatient use at Complejo Asistencial Universitario de León, Spain, between March 2014 and August 2017, with the purpose to contribute some knowledge on the use of TV.

The hypothesis was proposed that hospitalization episodes and emergency department visits decrease with the use of TV with regard to the period without TV treatment. The elevated cost of this drug and the fact that it constitutes the main novelty in the treatment of hyponatremia in recent years motivated this report.

**Method**

Retrospective, observational study on the use of TV on an outpatient basis in a tertiary care hospital. All patients older than 18 years that were treated with tolvaptan between March 2014 and August 2017 were included. Data were collected from patient medical records. The information was anonymously handled.

The study variables were age, gender, reason for admission or visit to the emergency department, hospitalization and emergency department visit episodes before and after outpatient treatment, cause of hyponatremia, initial and maintenance dose, baseline and last available SNa measurement, treatment duration and treatment-related direct costs, calculated based on the manufacturer retail price, dispensed doses and number of days of treatment.

Statistical analysis was performed with Stata, version 14. The variables without normal distribution were described as medians and interquartile ranges and were compared using Mann-Whitney’s U-test or Kruskal-Wallis test. Categorical variables were summarized as percentages and compared using the chi-square test. For all tests, statistical significance was considered with a p-value < 0.05.

This study was carried out in accordance with the guidelines established in the Fortaleza Declaration and all procedures performed on human subjects were approved by the Ethics and Clinical Research Committee of the hospital. Since this was a retrospective study, and given the difficulty to obtain informed consent of each patient, this was not applied.

**Results**

In total, nine patients received TV on an outpatient basis; their characteristics are summarized in Table 1.

The maintenance dose was lower than indicated in the summary of product characteristics (15 mg/day) in seven patients, with eunatremia being achieved with
2.1 mg/day (15 mg/seven days) in 11.1 %, with 3.75 mg/day (7.5 mg/48 hours) in 22.2 % and with 7.5 mg/day (7.5 mg/24 hours) in 55.6 %.

There were no episodes of hypernatremia (Na > 145 mmol/L) or acute renal failure during the follow-up.

When the periods of outpatient treatment with TV were compared in each patient with regard to equivalent periods of time prior to the start of treatment, we found a reduction in total days of hospitalization (361 versus 70, p = 0.007) at the expense of hospitalizations due to hyponatremia (11 vs 306, p = 0.009), with a trend towards a reduction in the number of total hospitalization episodes (19 versus 7, p = 0.072) and especially in those attributable to hyponatremia (14 versus 4, p = 0.051), as well as in episodes of aid at the emergency department for all causes (18 versus 11, p = 0.302) and in those directly attributable to hyponatremia (15 versus 5, p = 0.076).

Outpatient treatment total cost during the analyzed period was € 170,875.7 and mean cost per patient was € 17,930.6 (3,733.7-30,766.1).

Table 2 shows the itemized expenditure per diagnosis and lists the cost and the recorded difference in assessed events: admission episodes, days of hospitalization and episodes requiring visits to the emergency department.

Discussion

In the analyzed patients, TV outpatient use was shown to be safe and efficacious in the reduction of clinical events, even with doses lower than those indicated in the summary of product characteristics. Owing to the SALTWATER study, we have data available on the safety and efficacy of TV chronic use with regard to the evolution of natremia; however, experience is limited in terms of clinical benefits associated with the correction of natremia, which is restricted to the publication of clinical cases. In our patients, a follow-up of 3,993 days was accumulated, with a median of 277 days and a maximum individual follow-up of 1,089 days (2.98 years), which constitutes one of the strengths of the study. The use of TV appears to be associated with a significant decrease in days of hospitalization associated with hyponatremia and, overall, with any pathology; in addition, we identified a decrease in emergency department visits and admission episodes in patients treated with tolvaptan.

Of note, the majority of patients maintain eunatremia with the use of doses lower than those indicated in the summary of product characteristics. Along the same lines, the San Carlos Clinical Hospital group reported their follow-up results in 16 patients on outpatient treatment with TV for at least three years; they found similar outcomes in patients treated with tolvaptan.

It should be taken into account that the results vary significantly depending on the etiology of hyponatremia; for example, cancer patients attend the emergency department more often once they are treated, probably due to progression of the underlying disease. On the other hand, this is a non-randomized retrospective study and the number of patients is small,
which constitutes the most important limitation of our study.

In the cost analysis, the expenses generated by ambulatory follow-up of the treated patients (medical appointments, laboratory tests, transportation, etc.) have not been taken into account, and neither have those related to admission or care at the emergency department; therefore, studies specifically designed to analyze the cost-effectiveness of this treatment would be necessary.

Conclusions

Long-term use of tolvaptan appears to be safe and could be related to a decrease in days of hospitalization due to hyponatremia.

Funding

This investigation did not receive public or commercial sector subsidy.

Conflicts of interest

David E. Barajas Galindo and Emilia Gómez Hoyos declare having received honoraria from Otsuka Pharmaceutical Co. Ltd. for other works unrelated to this report.

References