

In-centre nocturnal haemodialysis: The experience of a centre

Hemodiálise noturna: Experiência de um centro

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■ ABSTRACT

Background and Objectives: As a result of improved clinical and quality-of-life outcomes, in-centre nocturnal haemodialysis emerges as one of the alternatives to conventional haemodialysis. Even today, little is known about the flow of patients through in-centre nocturnal haemodialysis programmes or about patient's survival or technique failure. This study sought to address this gap in knowledge. **Materials and Methods:** Retrospective, descriptive study that included all patients in in-centre nocturnal haemodialysis programme, between January 1995 and December 2011, which involved 51 patients. Descriptive data at baseline were presented as means or percentages of the total. We used Kaplan–Meier survival curves for patient survival. **Results:** Patients underwent for more than 5.5 hours thrice weekly sessions and remained in the technique for a mean period of 59.5 ± 73 months. The annual input and output of patients in the programme was on average of to 2.1 and 2.5 patients, respectively. Forty-three patients discontinued the technique: kidney transplantation in 24, conventional haemodialysis in 11 and peritoneal dialysis in three patients, with mean times in in-centre nocturnal haemodialysis of 37.4 ± 46.4 , 85.3 ± 93 and 59.7 ± 79.7 months, respectively. Throughout the period under review, five patients died (9.8%). Unadjusted survival at 20 years was 65% in patients who remained in in-centre nocturnal haemodialysis. **Conclusion:** In-centre nocturnal haemodialysis is associated with a higher survival rate, notwithstanding the possibility of selection bias resulting from patient's profile that opts for this modality. The relative contribution of patient selection versus effect of therapy on outcomes requires evaluation in future prospective clinical trials.

Key-words: Chronic renal disease; flow patient technique; in-centre nocturnal haemodialysis; patient survival.

■ RESUMO

Introdução e Objectivos: Como consequência dos resultados da sobrevida e qualidade de vida, o programa de hemodiálise noturna em centro surge como uma alternativa à hemodiálise convencional. Ainda hoje pouco se sabe sobre o fluxo de doentes, sobrevida e abandono da técnica. Este estudo pretendeu abordar essa lacuna no conhecimento. **Métodos:** Estudo descritivo de desenho retrospectivo que incluiu todos os doentes em programa de hemodiálise noturna da clínica, entre Janeiro de 1995 e Dezembro de 2011. Foram avaliados

51 doentes. Os dados descritivos foram apresentados como médias ou percentagens do total. Foram utilizadas as curvas de Kaplan-Meier para a sobrevida dos doentes. **Resultados:** Os doentes realizaram sessões com duração maior ou igual a 5,5 horas 3 vezes por semana e permaneceram na técnica por um período médio de $59,5 \pm 73$ meses. A entrada e saída anual de doentes do programa foi em média de 2,1 e 2,5, respectivamente. Quarenta e três doentes mudaram de técnica: 24 por transplante renal, 11 para hemodiálise diurna e três para diálise peritoneal, com tempos médios de permanência de $37,4 \pm 46,4$, $85,3 \pm 93$ e $59,7 \pm 79,7$ meses, respectivamente. Durante todo o período em análise faleceram cinco doentes (9,8%). A sobrevida não ajustada aos 20 anos foi de 65% nos doentes que permaneceram em hemodiálise nocturna. **Conclusão:** A hemodiálise nocturna está associada a uma aumento da sobrevida, pese embora existir um viés de selecção resultante do perfil dos doentes que optam por esta modalidade. A contribuição relativa de selecção de doentes *versus* o efeito da técnica sobre os resultados necessita de avaliação em ensaios clínicos prospectivos.

Palavras Chave: Doença renal crónica; fluxo de doentes; hemodiálise nocturna em centro; sobrevida dos doentes.

INTRODUCTION

The significant morbidity and mortality among long-term haemodialysis (HD) patients^{1,2} has increased the interest in more intensive dialysis regimens³. The Hemodialysis (HEMO) Study failed to show a significant benefit of high dialysis doses three times per week⁴. These results renewed interest in the contribution of dialysis time (i.e., treatment duration) to outcomes in haemodialysis^{5,6}. Home nocturnal haemodialysis (HNHD) and short-daily haemodialysis (SDHD) are the most extensively studied forms of intensified dialysis, although the actual cardiovascular and quality of life benefits of these modalities remain controversial⁷⁻¹⁰.

In-centre nocturnal haemodialysis (INHD) has emerged as a potential “in-between SDHD and HNHD” therapy by providing more flexibility for patients during the day and offering some of the shortcomings attributed to both SDHD and HNHD¹⁰. The INHD, which is typically administered thrice weekly for more than 5.5 hours (generally about 8 hours) per session in the dialysis unit during the overnight hours^{5,6}, has emerged as a viable mechanism for the administration of intensified dialysis. It offers the theoretical advantages of prolonged dialysis with the benefits of nursing supervision and the non-disruption of productive daytime hours¹⁰.

In the last few years, better laboratory and patient outcomes have been observed in INHD compared with three times per week conventional haemodialysis (CHD)^{3,6,11-17}.

In-centre nocturnal haemodialysis was instituted in our clinic in 1983 as a pioneer of this technique in Portugal. Even today, little is known about the flow and survival of patients in INHD programmes. This study sought to address this gap in knowledge.

MATERIALS AND METHODS

This is a retrospective study of all patients who participated in the INHD programme in our clinic, from January 1995 until December 2011, which included 51 patients.

The INHD programme performed HD using the Gambro AK 100 HD machine and, subsequently, AK 200. Treatment sessions were administered for more than 5.5 hours thrice weekly, with a dialysate flow rate (Qd) of 600 ml/min and a prescribed blood flow rate (Qb) of 300 ml/min.

Descriptive data at baseline were presented as means or percentages of the total. We used Kaplan-Meier survival curves for patient survival.

RESULTS

The study cohort included 51 patients who were treated by INHD in our facilities during the study

period. The patients characteristics at admission in INHD are shown in Table I.

Table I

Baseline patient characteristics

Characteristics at Baseline	
Patients (n)	51
Age (yr; mean $\pm$ SD)	36.88 $\pm$ 10.68
Male gender (%)	78
Ethnic group (%)	
Black	17.65
White	76.47
other	5.88
Education level (n)	
unknown	7
fourth grade	7
sixth grade	4
ninth grade	7
high school	14
college	12
Work status (n)	
unknown	4
student	2
employed	39
unemployed	6
Cause of CKD (%)	
diabetes	11.76
hypertension	7.84
glomerulonephritis	47.06
hereditary/cystic	9.8
other (includes unknown)	23.53
Vascular access (%)	
fistula	92.16
graft	7.84
catheter	0

During the study period, the movement of patients coming in and out of the technique over the years was constant, with an average of incoming and outgoing of 2.1 and 2.5 patients, respectively (Fig. 1). A mean of 9.6 patients per year remained in the programme.

Forty-three of the total 51 patients discontinued the technique: 24 patients underwent kidney transplantation, 11 switched for conventional HD and three changed for peritoneal dialysis (PD). From patients transferred to conventional haemodialysis, one chose home haemodialysis and two needed hospital haemodialysis. Throughout the period under review five patients died. Figure 2 shows motives for dropping out during the study years. The causes to move from

the programme to peritoneal dialysis were: vascular access failure in two patients and the wish for greater autonomy in the other patient. About the two patients who went to hospital haemodialysis, one returned to the hospital unit (where he had previously been on haemodialysis) by non-adaptation to INHD because discomfort and consequent insomnia. The other patient required hospital haemodialysis for clinical deterioration. Desire of a more flexible schedule was the main reason for the patient who chose home haemodialysis. In the case of the eight patients who switch from INHD to conventional HD, the motives were: insomnia in two, indiscipline and disturbance of other patients in one and psychosocial causes in five (patient's choice, family convenience, geographic relocation to a region where INHD was not offered, nocturnal work/change for the night shift, retirement).

The global mean time on INHD was 59.5 $\pm$ 73 months (median 32 months) during the follow-up period. Table II illustrates the mean time to discontinuing INHD in accordance to patient's destination.

Table II

Mean and median times on INHD, according to the destination

Dropout motive	INHD duration (months)	
	Mean	Median
Renal transplantation n = 24 (47%)	37.4 $\pm$ 46.4	23
Conventional haemodialysis n = 11 (21.56%)	85.3 $\pm$ 93	46
Peritoneal dialysis n = 3 (5.9%)	59.7 $\pm$ 79.7	24
Deaths n = 5 (9.8%)	101.4 $\pm$ 100	34
In-centre nocturnal haemodialysis n = 8 (15.7%)	39.9 $\pm$ 52.6	23

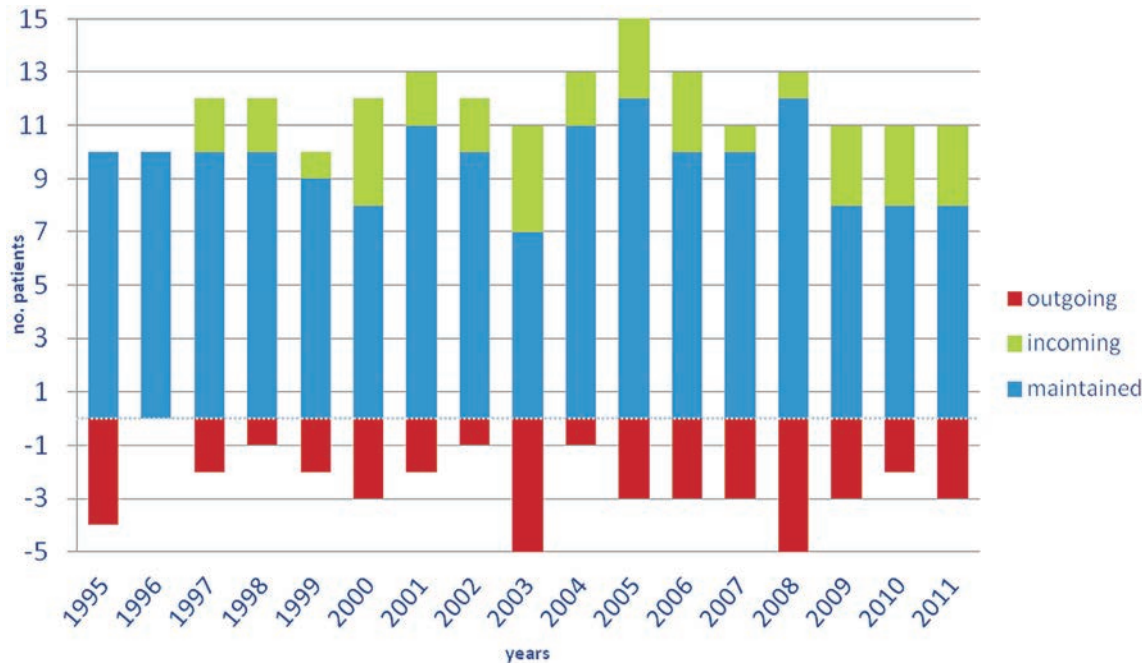
Regarding the survival of patients who remained in INHD, Kaplan–Meier unadjusted survival curve (Fig. 3) indicates 20 years patient survival rate of 65%. Patients were censored upon changing modality.

DISCUSSION

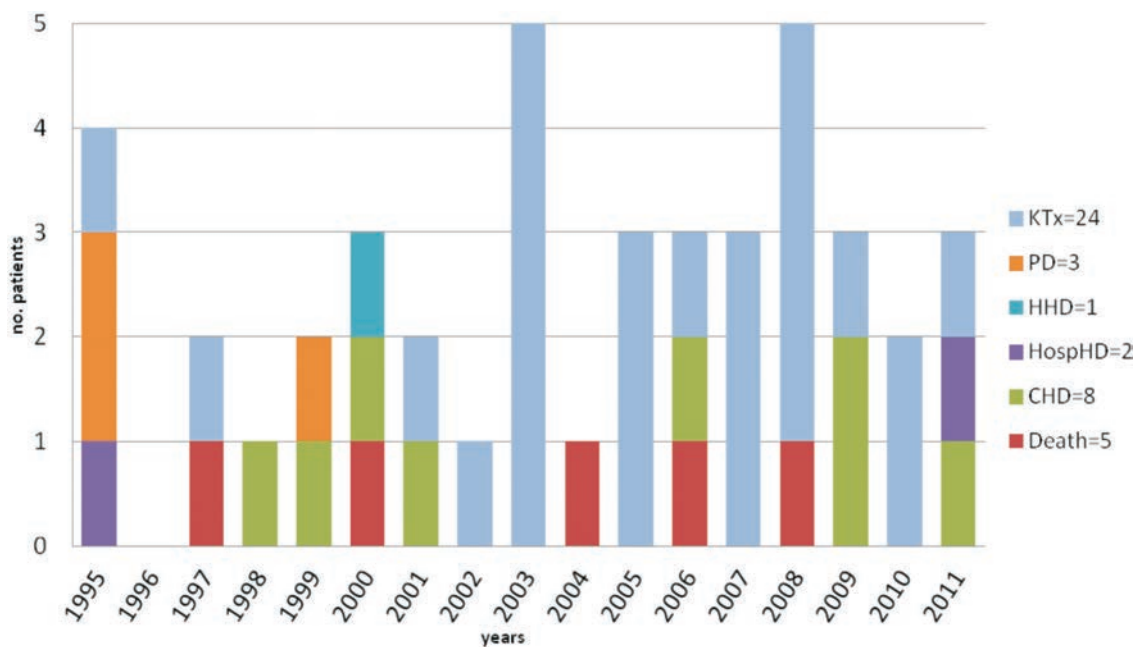
The characteristics of patients treated by INHD at baseline revealed that they were young (36.9 years),

Figure 1

Patients movement over the years.

**Figure 2**

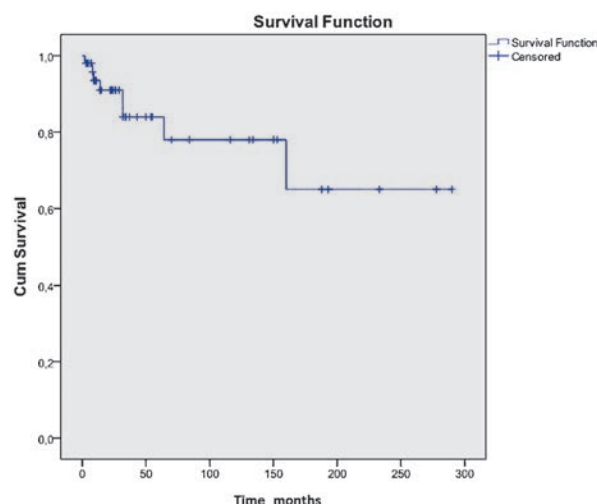
Dropout reasons over the years.



KTx – kidney transplant; PD – peritoneal dialysis; HHD – home haemodialysis; HospHD – hospital haemodialysis; CHD – conventional haemodialysis.

Figure 3

INHD patients unadjusted survival.



fistula was the preferential vascular access (92.2%) and only 12% were diabetic, aspects quite different when compared with the national registration of patients on HD, with a mean age of dialysis patients of 66.9 years, 70.7% of fistulae and 28.2% of diabetics¹⁸.

Patients movement over the study years was constant and there was no significant increase in the INHD programme. The main reason for abandoning the technique was kidney transplantation. The switch of dialysis technique was recorded in 15 patients (11 to CHD and three to PD) after a long period of stay in INHD. Vascular access failure; desire for greater self-sufficiency and flexible schedule; discomfort and insomnia, and psychosocial factors were the most common reasons for patient dropout. Our data are according to the literature, in which trouble sleeping either due to the uncomfortable chair or from insomnia and missing being at home with their relatives are the most frequent motives for abandoning INHD¹¹.

Twenty years patient unadjusted survival rate of 65% obtained in our study was an excellent outcome, especially when compared with the national¹⁸ and international^{19,20} registration of patients on haemodialysis, the former with an annual global mortality of 12%.

Several studies have analysed patient survival on INHD. In Tassin, French patients have been prescribed

a regimen of 8-hour sessions, thrice weekly. The results showed improved survival, with an unadjusted survival for all patients at 10 years of 54% on INHD compared to 31% on CHD (21-23). These exceptional outcomes have been the drive for prolongation of dialysis above conventional session durations¹⁰. Ok *et al.*¹⁷ performed a prospective multicentre cohort study that matched 247 INHD patients with 247 period-prevalent CHD patients, which found a 72% relative risk reduction for mortality in the INHD cohort. The U.S. Fresenius Medical Care dialysis network developed the largest study of INHD patients reported to date. An initial study compared 655 prevalent INHD recipients with 15,334 patients on CHD⁵. One year unadjusted mortality was lower in INHD patients, although after adjustment for case mix and vascular access type this association was no longer obvious. The significant dilution of the association between INHD and survival in the adjusted analysis underlined the presence of considerable confounding factors, like medical opinion and patient self-selection⁵. To overcome this limitation, the authors matched 746 patients who converted to INHD to 2,062 CHD recipients using a propensity score-based matching. The results showed that INHD was associated with a significant reduction in all-cause mortality⁶.

In a simplistic way, extending the mean treatment time differentiates CHD (≥ 12 hours per week) from INHD (≥ 16.5 hours per week). Longer treatment time has been associated with improved outcomes, with a better survival in short daily haemodialysis with length of time of ~ 15 hours versus approximately 10.5 hours^{9,24,25}. The first direct effect was a marked increase in urea clearance, represented by eKt/V , that overcomes the dialyzers with less surface area, lower blood and dialysate flow rates^{5,6}.

Our study design is observational and with no control population, which is a strong limiting factor that does not allow proving any causation. Moreover, there are several characteristics that may predispose patients to self-select nocturnal therapy, such as being educated, well-informed and working. The majority of our patients (64.7%) had an education level equal to or greater than ninth grade and 76.5% were working active patients. These features may predispose better outcomes *per se*^{5,6,10} and, in association with a patient's younger age, could also explain the elevated transplantation rate observed in our INHD patients. Nevertheless, it is also possible

that extended survival may have contributed for the larger number of renal transplants^{5,6}.

To date, observational studies demonstrated an association between INHD and improved survival. However these results should be interpreted with caution because patients who are on INHD are a selected (or self-selected) group¹⁰.

The published data provide an overview of INHD therapy in comparison with CHD, that established associations but are not conclusive^{5,6}. Additional studies are needed, preferably prospective and randomized, to evaluate this therapeutic option. Such a trial is unlikely to happen, as has already been shown by recruitment sample size failure of the randomized Frequent Hemodialysis Network (FHN) Nocturnal Trial⁸. It is understandable that when faced with hypotheses that implicate two diametrically opposed dialysis schedules, both with large implications on lifestyle, most patients are reluctant to leave this decision to the trial allocation¹⁰.

In summary, patient willingness and logistical issues will still continue to guide patient's selection for INHD. The characteristics of patients who opt for INHD are not representative of the general CHD population, such that the relative contributions of patient selection versus effect of therapy on outcomes remain to be explained.

Conflict of Interest Statement: None declared.

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