A cost of illness study of hypoglycaemic events in insulin-treated diabetes in the Netherlands

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ABSTRACT

Objectives Patients with diabetes mellitus are at a risk for hypoglycaemia. Besides the burden of hypoglycaemia for patients, hypoglycaemia poses an economic burden to society. The aim of this study was to calculate the per patient societal costs of hypoglycaemia among patients with type 1 diabetes (T1DM) and type 2 diabetes (T2DM) on insulin therapy in the Netherlands.

Methods To calculate the costs of hypoglycaemia, data from the Global Hypoglycaemia Assessment Tool (HAT) study were used. Dutch patients were selected from the HAT study database and data regarding healthcare resource use, informal care use and productivity losses were combined with Dutch unit costs to calculate the per patient 4-week costs of patients experiencing hypoglycaemia. Besides these 4-week costs, costs per hypoglycaemic event were calculated by dividing the study population total 4-week costs by the total number of events in this period.

Results Mean 4-week total costs of hypoglycaemia amounted to €163 (SD, €870) in T1DM and €134 (SD, €364) in T2DM. While productivity costs were the most important cost driver of hypoglycaemia in patients with T1DM (accounting for 72% of the total costs), costs of hypoglycaemia in patients with T2DM were almost entirely driven by costs within the healthcare sector (accounting for 98% of the total costs). Mean costs of a severe hypoglycaemic event were €823 and €508 in T1DM and T2DM, respectively, whereas mean costs of a non-severe event were almost zero.

Conclusions This study showed that the economic burden of severe hypoglycaemia is substantial. The prevention of hypoglycaemia could therefore not only reduce the burden for patients, but also the economic burden to society.

INTRODUCTION

Worldwide 382 million individuals suffer from diabetes mellitus.1 The estimated prevalence of diabetes in the Netherlands was 1.1 million in 2015.2 Between 1991 and 2014, the prevalence of diabetes in men has more than doubled, and the prevalence in women has increased by 50%.3 The prevalence is expected to increase further because of the ageing population and due to an increase in the number of people being overweight or having other risk factors for diabetes.3

Patients with diabetes mellitus are at risk for hypoglycaemia, which is caused by medications used to treat diabetes mellitus, such as insulin or oral hypoglycaemic medications, combined with reduced carbohydrate intake or increased activity. Khunti et al have shown that the rate of any hypoglycaemic event was 73.3 events per patient-year for type 1 diabetes (T1DM) and 19.3 events per patient-year for type 2 diabetes (T2DM) treated with insulin.4 Hypoglycaemia is mostly associated with mild symptoms, such as sweating, dizziness or headache, but hypoglycaemia can also lead to unconsciousness or coma. Moreover, hypoglycaemia is associated with more future events.5

Besides the burden for patients, hypoglycaemia poses an economic burden to society. Hypoglycaemic events can lead to healthcare use, such as ambulance transport, hospital admissions and clinical appointments. Healthcare costs of hypoglycaemia have been estimated in different countries,6-10 but the costs of hypoglycaemia in the Netherlands are unknown. In addition to the healthcare costs,
hypoglycaemia is expected to lead to societal costs due to lost productivity and time costs for informal caregivers, but these estimates are also lacking. The aim of this study was therefore to calculate the per patient societal costs of hypoglycaemia among insulin-treated patients with T1DM and T2DM in the Netherlands.

**METHODS**

To calculate the costs of hypoglycaemia, data from the Global Hypoglycaemia Assessment Tool (HAT) study were used. The primary objective of the HAT study was to determine the percentage of patients experiencing at least one hypoglycaemic event during the 4-week follow-up period among insulin-treated patients with T1DM and T2DM. The HAT study was a non-interventional, multicentre, 6-month retrospective and 4-week prospective study. Patients were invited to participate by their healthcare provider during routine scheduled clinical appointments if they fulfilled the following inclusion criteria: (1) having T1DM or T2DM and treated with insulin for >12 months; (2) being ≥18 years at the time of the survey and (3) giving informed consent to participate in the study. Non-ambulatory patients were excluded. The HAT study was conducted in 24 countries. To calculate the costs of hypoglycaemia in the Netherlands, Dutch patients were selected from the HAT study database. In the Netherlands, 150 general practitioners and 18 hospitals (including 2 academic hospitals) participated in the study. Patients originated from urban and rural areas from all over the Netherlands. Patients were invited to participate during routine clinical visits and were not actively approached to participate otherwise.

In the HAT study, a Self-Assessment Questionnaire (SAQ) was used, consisting of two parts. In the first part, data about baseline demographics and treatment were collected. Additionally, questions regarding knowledge, awareness and perceptions of hypoglycaemia were asked. Lastly, information about the history of severe hypoglycaemia over the last 6 months and non-severe hypoglycaemia over the last 4 weeks was collected. A severe event was defined as hypoglycaemia which requires assistance from another person to administer carbohydrate and/or glucagon. Hypoglycaemia resulting in hospital admission and hypoglycaemia requiring assistance from medical personnel but not requiring hospital admission were also regarded as severe in this study. Non-severe hypoglycaemia was defined as hypoglycaemia managed by the patient alone. Part 1 of the questionnaire was provided to patients during a routine clinical appointment with a healthcare professional, and patients were asked to complete and return the questionnaire during the visit. In the next 4 weeks, patients were asked to fill in a diary to capture hypoglycaemic events. After 4 weeks, patients were asked to complete part 2 of the questionnaire, which assessed the history of both severe and non-severe hypoglycaemia in the previous 4 weeks. Both part 1 and part 2 of the SAQ also included questions regarding the impact of hypoglycaemia on healthcare and informal care utilisation and productivity. The patient questionnaire is provided in the online supplementary appendix 1. Figure 1 shows the design of the study.

Costs of hypoglycaemic events within the healthcare sector were calculated by combining healthcare utilisation as derived from the SAQ with Dutch unit costs. Unit costs were derived from the Dutch costing manual, except for the costs of extra blood glucose tests (these were derived from a report by the Dutch National Healthcare Institute about the reimbursement and quality of blood glucose test material) and the costs of glucose administered by medical professionals during a hypoglycaemic event (these were derived from medicijnkosten.nl). Besides

![Figure 1](https://example.com/screenshot.png)
the costs within the healthcare sector, informal care costs were calculated by combining informal care use with time costs of informal caregivers which were also derived from the Dutch costing manual.\textsuperscript{12} In order to calculate the costs of hypoglycaemic events within the healthcare sector and the costs of informal care, the following assumptions were made: (1) patients admitted to a hospital as a result of a hypoglycaemic event arrived at the hospital with an ambulance; (2) 75% of patients requiring assistance from medical personnel but not requiring hospital admission was treated at home by ambulance personnel and 25% was treated by a general practitioner (only applicable to patients with severe hypoglycaemia); (3) patients requiring assistance from medical personnel received an intravenous infusion (or injection) with glucose; (4) patients requiring assistance from another person (non-medical) to administer carbohydrate and/or glucagon were given assistance by an informal caregiver and this assistance was assumed to take 1 hour for patients with T1DM and 3 hours for patients with T2DM; (5) patients with T1DM who had an additional clinical appointment with a doctor or nurse visited the hospital, whereas patients with T2DM who had an additional clinical appointment with a doctor or nurse visited the general practitioner.

Productivity costs were calculated by combining productivity losses as derived from the SAQ with hourly productivity costs from the Dutch costing manual.\textsuperscript{12} Hourly productivity costs were €32 for female patients and €38 for male patients. To calculate the productivity costs of hypoglycaemic events, two additional assumptions were made: (6) a working day was assumed to be 8 hours (for patients with a full-time and part-time job); (7) patients arriving at work late or leaving work early were absent from work for half a day. Productivity costs were zero for students and patients who were unemployed or patients who were retired from work.

If a question regarding resource use remained unanswered, it was assumed that the patient did not use this specific type of resource use. Missing data regarding the frequency of healthcare use (if a patient indicated that he had used a specific type of resource use) were handled by imputing the mean of the available data. In this way, all patients were included in the calculation of total costs including patients with missing values on some, but not all variables.

Base case analyses present the per patient 4-week costs as derived from part 2 of the SAQ (ie, the 4-week prospective study). Results were reported separately for patients with T1DM and T2DM. Within these two categories, a further distinction was made between patients with hypoglycaemia including at least one severe event and patients with non-severe events only. Results of part 1 of the SAQ (ie, 6-month retrospective study) were provided in the online supplementary materials.

The impact of assumptions 2, 4 and 7 on the total 4-week costs of patients experiencing hypoglycaemia were tested in univariate sensitivity analyses. First, the proportion of patients treated at home by a general practitioner or ambulance personnel was varied, that is, 50% or 0% treated at home by a general practitioner (instead of 25%) and 50% or 100% treated by ambulance personnel (instead of 75%). Second, the duration of assistance by an informal caregiver was doubled, that is, 2 hours for T1DM (instead of 1 hour) and 6 hours for T2DM (instead of 3 hours). Third, the hours that patients were absent from work if they indicated that they arrived at work late or left work early were halved, that is, 2 hours instead of 4 hours. In addition to these assumptions, the impact of higher unit costs (+20%) of an ambulance was studied.

Besides the per patient total 4-week costs of patients experiencing hypoglycaemia, costs were calculated per hypoglycaemic event. To calculate these costs, the study population total 4-week costs were divided by the total number of events in this period. The total number of events was derived from the SAQ and patient diary. Since number of events according to the SAQ and patient diary differed, two point estimates of the costs of an event were provided.

Additionally, the total costs of hypoglycaemia among insulin-treated patients with T1DM and T2DM in the Netherlands were estimated using information on the number of insulin-users in the Netherlands. According to the Dutch Foundation for Pharmaceutical Statistics (SFK), there were 310 000 insulin-users in the Netherlands in 2014.\textsuperscript{15} Assuming that all patients with T1DM use insulin and that the estimated prevalence of diabetes in the Netherlands was 1.1 million in 2015, and 9% (ie, 99 000 patients) had T1DM,\textsuperscript{2} the estimated number of insulin-users is 99 000 in T1DM and 211 000 in T2DM (ie, 310 000 minus 99 000). These numbers were then multiplied by the proportion of patients experiencing hypoglycaemia, and the per patient 4-week costs (extrapolated to yearly costs) of these latter patients.

All costs were reported in Euro 2016. Wherever necessary, costs were adjusted to 2016 values using the consumer price index derived from Statistics Netherlands.\textsuperscript{16}

RESULTS

Patient and disease characteristics

Six hundred and thirty-three patients returned part 2 of the SAQ, that is, 142 patients with T1DM and 491 patients with T2DM. Table 1 shows the patient and disease characteristics. The median age was 46 years for patients with T1DM and 68 years for patients with T2DM. The duration of diabetes was 18.6 years (SD, 12.1) and 14.2 years (SD, 8.0) for patients with T1DM and T2DM, respectively. Patients with T2DM used insulin, on average, for 7.7 years (SD, 6.1). Levels of glycaemic control (ie, HbA1c) were similar across patients with T1DM and T2DM (59.3 mmol/mol (7.3%) vs 57.7 mmol/mol (7.6%)).

In the 4 weeks after baseline, 92% of the patients with T1DM and 43% of the patients with T2DM experienced one or more hypoglycaemic events. Of these patients, 10% and 15% experienced at least one severe event, and almost all these patients (ie, 98% and 96%) experienced...
at least one non-severe event. Fifty-one per cent and 32% of the patients with T1DM and T2DM who experienced hypoglycaemia, respectively, experienced (a) nocturnal hypoglycaemic event(s).

Healthcare utilisation and informal care utilisation

Table 2 shows the healthcare utilisation and informal care utilisation of those patients who experienced hypoglycaemia in the 4 weeks after baseline (patients without hypoglycaemia were excluded from the analyses). Only 43 patients (13%) of the population had at least one event requiring (1) hospital admission (T1DM, 2%; T2DM, 10%); (2) assistance from medical personnel but not requiring hospital admission (T1DM, 1%; T2DM, 2%); and/or (3) assistance from another person to administer carbohydrate and/or glucagon (T1DM, 9%; T2DM, 6%). Most patients (T1DM, 98%; T2DM, 96%) also had one or more events that they could manage themselves.

Actions undertaken by patients as a result of hypoglycaemic events are also presented in table 2. Fifteen per cent (T1DM, 21%; T2DM 11%) of the population addressed a hypoglycaemic event at the next scheduled clinic visit, 2% (T1DM, 2%; T2DM 1%) attended an additional clinical appointment with a doctor or nurse, 8% (T1DM, 7%; T2DM 8%) made additional telephone contacts with a doctor or nurse, 1% (T1DM, 1%; T2DM 1%) consulted another healthcare professional and 68% (T1DM, 70%; T2DM 67%) did not consult a doctor, nurse or healthcare professional if they experienced a hypoglycaemic event (note that this is the average of patients with hypoglycaemia including at least one severe event or non-severe events only).

Also, 26% (T1DM, 27%; T2DM 25%) of the population increased the quantity of carbohydrates or number of snacks in diet, 7% (T1DM, 8%; T2DM 6%) reduced the amount of sport or physical exercise, 29% (T1DM, 39%; T2DM 22%) decreased the insulin dose, 15% (T1DM, 16%; T2DM 14%) skipped the insulin dose, 53% (T1DM, 58%; T2DM 50%) increased the number of blood glucose checks per day and 11% (T1DM, 16%; T2DM 8%) made any other change to diabetes treatment.

Per patient 4-week healthcare utilisation and informal care costs

Table 3 presents the 4-week healthcare utilisation costs and informal care costs of patients who experienced hypoglycaemia.
### Table 2  Healthcare utilisation, informal care utilisation and additional actions of patients experiencing hypoglycaemia in 4 weeks after baseline

| Patients indicated the following hypoglycaemic events (in the last 4 weeks)† | All (n=332) | T1DM (n=130) | T2DM (n=202) |
|---|---|---|---|
| Event resulting in hospital admission, n (%) | 23 (7) | 3 (2) | 3 (23) |
| Times admitted, mean (SD) | 1.0 (0.0) | 1.0 (0.0) | 1.0 (0.0) |
| Admission length (days), mean (SD) | 1.1 (0.4) | 1.7 (1.2) | 1.7 (1.2) |
| Event requiring assistance from medical personnel but not requiring hospital admission, n (%) | 6 (2) | 1 (1) | 1 (8) |
| Number of episodes, mean (SD) | 1.0 (0.0) | 1.0 (0.0) | 1.0 (0.0) |
| Event requiring assistance from another person to administer carbohydrate and/or glucagon (informal care), n (%) | 24 (7) | 12 (9) | 12 (92) |
| Number of episodes, mean (SD) | 1.8 (1.3) | 1.5 (0.8) | 1.5 (0.8) |
| Event managed by the patient, n (%) | 322 (97) | 128 (98) | 12 (92) |
| Number of episodes, mean (SD) | 5.1 (5.2) | 8.6 (6.3) | 10.5 (8.1) |
| Event occurred at night, n (%) | 130 (39) | 66 (51) | 9 (69) |
| Number of episodes, mean (SD) | 2.1 (1.8) | 2.5 (2.2) | 3.3 (3.1) |

**Patients indicated the following actions (if they experienced a hypoglycaemic event in the last 4 weeks)**

| Addressed the hypoglycaemic event at next scheduled clinic visit, n (%) | 50 (15) | 27 (21) | 6 (46) |
| Attended additional clinical appointments with doctor/nurse, n (%) | 5 (2) | 2 (2) | 1 (8) |
| Number of appointments, mean (SD) | 1.7 (0.8) | 2.3 (0.9) | 3.0 (0.0) |
| Rescheduled clinic appointment for an earlier time, n (%) | 1 (0) | 1 (1) | 1 (8) |
| Made additional telephone contacts with doctor/nurse, n (%) | 26 (8) | 9 (7) | 5 (38) |
| Number of contacts, mean (SD) | 2.0 (1.2) | 2.2 (1.3) | 2.6 (1.5) |
| Consulted another healthcare professional, n (%) | 2 (1) | 1 (1) | 0 (0) |
| Did not consult a doctor/nurse/healthcare professional, n (%) | 227 (68) | 91 (70) | 5 (38) |
| Increased the quantity of carbohydrates or number of snacks in diet, n (%) | 86 (26) | 35 (27) | 3 (23) |
| Reduced the amount of sport or exercise, n (%) | 24 (7) | 11 (8) | 3 (23) |
| Decreased insulin dose, n (%) | 96 (29) | 51 (39) | 7 (54) |
| Skipped insulin dose, n (%) | 49 (15) | 21 (16) | 2 (15) |
| Increased the number of blood glucose checks per day, n (%) | 176 (53) | 76 (58) | 8 (62) |
| Number of checks, mean (SD) | 2.0 (0.7) | 1.9 (0.8) | 2.1 (0.8) |
| Number of days, mean (SD) | 4.3 (5.1) | 5.5 (4.1) | 5.5 (3.9) |
| Made any other changes to diabetes treatment, n (%) | 38 (11) | 21 (16) | 3 (23) |

*Note that these categories include patients with (a) non-severe event(s) only (and two patients (1 T1DM patient and 1 T2DM patient) with a nocturnal event only).
†Note that the percentages do not count to 100%, because patients can have multiple hypoglycaemic events.

NSH, non-severe hypoglycaemia; SH, severe hypoglycaemia; T1DM, type 1 diabetes; T2DM, type 2 diabetes.
hypoglycaemia in the 4 weeks after baseline. Mean total 4-week healthcare and informal care costs were €98 (SD, €325). Mean costs were higher for patients with T2DM than for patients with T1DM; €131 (SD, €363) compared with €46 (SD, €248). Mean costs of patients with hypoglycaemia including at least one severe event were €426 (SD, €696) and €863 (SD, €512) in T1DM and T2DM, respectively, compared with €4 (SD, €16) and €4 (SD, €12) in T1DM and T2DM, for patients with non-severe events only.

Productivity losses

Productivity losses are presented in table 4. Of the patients with a full-time or part-time job, 4% reported sick leave (T1DM, 5%; T2DM, 2%) with a mean of 6 days (SD, 8). Additionally, 8% (T1DM, 12%; T2DM, 2%) and 6% (T1DM, 9%; T2DM, 0%) of the patients with a full-time or part-time job arrived at work late or left work early for ≥1 days, respectively.

Per patient total 4-week costs

Table 5 shows the per patient total 4-week costs of patients who experienced hypoglycaemia. Mean total 4-week costs were €145 (SD, €613). Mean costs were €163 (SD, €870) for patients with T1DM and €134 (SD, €364) for patients with T2DM. While productivity costs were the most important cost driver of hypoglycaemia in patients with T1DM (accounting for 72% of the total costs), costs of hypoglycaemia in patients with T2DM were almost entirely driven by costs within the healthcare sector (accounting for 98% of the total costs).

Mean costs of patients with hypoglycaemia including at least one severe event were €1401 (SD, €2,497) and €863 (SD, €512) in T1DM and T2DM, respectively, compared with €26 (SD, €81) and €7 (SD, €41) in T1DM and T2DM for patients with non-severe events only. The mean total costs of patients with T1DM experiencing at least one severe event were largely driven by two patients, one with sick leave from work for 21 days and one with sick leave from work for 10 days.

Sensitivity analyses

Results of the sensitivity analyses suggest that the impact of the assumptions on the mean total 4-week costs of patients experiencing hypoglycaemia was limited. If the proportion of patients treated at home by ambulance personnel was decreased to 50%, total 4-week costs would decrease from €414 (SD, €98) to €205 (SD, €65) in T1DM and T2DM, respectively, compared with €26 (SD, €81) and €7 (SD, €41) in T1DM and T2DM for patients with non-severe events only. The mean total costs of patients with T1DM experiencing at least one severe event were largely driven by two patients, one with sick leave from work for 21 days and one with sick leave from work for 10 days.
Cost per hypoglycaemic event
Mean costs per hypoglycaemic event are presented in table 5 and amounted to €19 for patients with T1DM and €47 for patients with T2DM (€16 and €46 if the number of events as registered in the patient diary was used (note that the number of events according to the patient diary were consistently higher than in the SAQ)). Mean costs of a severe event were €828 and €552 in T1DM and T2DM, respectively (€552 and €1036 if the number of events as registered in the patient diary was used), whereas the costs of a non-severe event were almost zero.

Total cost of hypoglycaemia in the Netherlands
Given that 92% of the T1DM population experience at least one hypoglycaemic event in a 4-week period, and that the mean total 4-week costs are €163 (95% CI €12 to €314), total 4-week costs in the entire Dutch T1DM population were estimated to amount to €14.8 million. Total 4-week costs in the entire Dutch T2DM population were estimated to amount to €12.2 million given that 43% of the insulin-treated T2DM population experience at least one hypoglycaemic event and that the mean total 4-week costs are €134 (95% CI €83 to €184). Total yearly costs of hypoglycaemia were estimated to be €352.3 million for the entire patient T1DM and T2DM population in the Netherlands (healthcare costs, €201.4 million; informal care costs, €8.3 million; productivity costs, €142.6 million). Using the boundaries of the 95% CI of the per patient total 4-week costs, total yearly costs of hypoglycaemia in the Netherlands were estimated to range from €112.5 to €590.8 million.

Per patient total costs based on part 1 of the SAQ
Mean total 6-month costs of patients experiencing hypoglycaemia including at least one severe event were €1132 (SD, 2933) for patients with T1DM and €586 (SD, 1439) for patients with T2DM using part 1 of the questionnaire, corresponding to €189 and €98 per month. The costs within the healthcare sector are the most important cost driver of hypoglycaemia in both patients with T1DM and T2DM (see the online supplementary tables 1–4).

DISCUSSION
This is the first study providing information on the costs of hypoglycaemia among insulin-treated patients with T1DM and T2DM in the Netherlands. The study shows that the economic burden is substantial. Mean 4-week costs of patients who experienced hypoglycaemia were €145 (SD, €613). In the Netherlands in 2011, the costs of diabetes care amounted to 1.7 billion Euros.17 Healthcare costs of hypoglycaemia in the Netherlands were estimated
to be €201.4 million per year, corresponding to 12% of the total healthcare costs of diabetes in the Netherlands. The total yearly costs of hypoglycaemia in the Netherlands, including informal care costs and productivity costs, were estimated to be €352.3 million.

The healthcare costs of hypoglycaemia in the Netherlands might seem relatively high with respect to the total costs of diabetes, but the total costs of diabetes might miss important diabetes-related costs, such as the costs of complications (eg, heart attack, stroke, eye problems and kidney disease). Total yearly costs of hypoglycaemia have been estimated in different countries, and all of these estimates are considerably lower than the costs we estimated. A comparison of healthcare costs between countries is complicated by practice variation, differences in the incentives to physicians and institutions and differences in relative and absolute prices. These differences should be taken into account when making international comparisons of healthcare costs. Nevertheless, possible explanations for the observed differences might be: first, one Swedish study was limited to the total yearly costs of T2DM, whereas we estimated total yearly costs of both T1DM and T2DM. Furthermore, this study only assessed costs of hypoglycaemic events that were classified as severe events in our study. Second, some studies were limited to healthcare costs, while we also took into account informal care costs and productivity costs. Third, different assumptions were made regarding the incidence of hypoglycaemia: while one study assumed an event rate of 0.09 per patient with T2DM per year (ie, 0.24 in insulin users and 0.04 in patients using oral anti-diabetic agents), the incidence among insulin-treated patients with T2DM in our study varied from 571 to 584 events in the 4-week follow-up period (ie, 15.8–16.2 per patient per year). Fourth, the number of patients in other countries was considerably lower than in the Netherlands.

Interestingly, the costs per hypoglycaemic event as calculated in our study were considerably lower than the costs as estimated in other studies. Again it should be noted that comparing healthcare costs between countries is complicated. Nevertheless, one of the explanations for the difference is the definition of a severe event which was limited to events resulting in hospitalisation in one study. An Italian study presented resource utilisation and productivity losses per person-years, which complicates comparisons of their estimates of cost per episode.

### Table 5 Per patient total 4-week costs of patients experiencing hypoglycaemia and costs per event

|                              | All (n=332) | T1DM (n=130) | Patients with NSH (n=13) | Patients with T2DM (n=202) | Patients with NSH (n=172) |
|------------------------------|-------------|--------------|--------------------------|----------------------------|---------------------------|
| **Per patient total 4-week costs** |             |              |                          |                            |                           |
| Costs within the healthcare sector (including informal care), mean costs (SD) (min–max) | €98 (€325) (€0–€2101) | €46 (€248) (€0–€2101) | €426 (€696) (€14–€2101) | €4 (€16) (€0–€153) | €131 (€363) (€0–€1654) |
| Productivity costs, mean costs (SD) (min–max) | €47 (€412) (€0–€6577) | €117 (€652) (€0–€6577) | €975 (€905) (€0–€6677) | €22 (€77) (€0–€638) | €3 (€36) (€0–€610) |
| **Per patient total 4-week costs of patients experiencing hypoglycaemia, mean costs (SD) (min–max)** | €145 (€613) (€0–€8678) | €163 (€870) (€0–€8678) | €1401 (€2497) (€0–€8678) | €26 (€81) (€0–€638) | €134 (€364) (€0–€1654) |
| **Costs per event** |             |              |                          |                            |                           |
| Study population total 4-week costs of patients experiencing hypoglycaemia | €48 140 | €21 190 | €18 213 | €30 42 | €27 068 |
| Number of events based on SAQ* | 1700 | 1129 | 22 | 981 | 571 | 51 | 473 |
| Total costs per event (based on SAQ) | €28 | €19 | €828 | €3 | €47 | €508 | €3 |
| Number of events based on patient diary§ | 1923 | 1339 | 33 | 1139 | 584 | 25 | 483 |
| Total costs per event (based on patient diary¶) | €25 | €16 | €552 | €3 | €46 | €1036 | €2 |

*The number of events excludes nocturnal events in order to prevent double counting. The number of events in patients with at least one severe event excludes non-severe events.

†Two patients experienced three events requiring assistance from another person to administer carbohydrate and/or glucagon, and two patients had two of these events, which increases the total number of events in the population.

‡Six patients experienced more than one event requiring assistance from another person to administer carbohydrate and/or glucagon, with one patients having six of these events.

§Number of events in patients with SH and NSH are limited to the number of severe and non-severe events, respectively, as reported within these groups.

¶Please note that the SD could not be calculated because costs per event were calculated by dividing total population costs by the total number of events in the population, but is expected to be substantial.

NSH, non-severe hypoglycaemia; SH, severe hypoglycaemia; T1DM, type 1 diabetes; T2DM, type 2 diabetes.
to the estimates in the current study. The cost estimates in that study seem to be higher because of a longer stay in hospital and larger productivity losses for patients and caregivers. In contrast to the Italian study, work days lost by family members are not taken into account in our study, to avoid possible double counting with informal care time. A Danish study found lower costs per hypoglycaemic event than the costs we calculated, but in this study, costs were limited to healthcare costs.

Some limitations to the data and methods deserve mentioning. First, there is large variation in the per patient costs. This can be explained by the skewed distribution of the data, that is, a small number of patients has very high costs (e.g., while the mean total 4-week costs are €145 (SD, 613), 25 out of 322 patients (8%) have costs >€1000). This is frequently observed in costing studies. Further research, preferably with larger sample sizes, should confirm the results of our study.

Second, there is a large discrepancy between the per patient costs of patients experiencing hypoglycaemia including at least one severe event between part 1 and part 2 of the SAQ. The monthly costs based on part 1 of the questionnaire are much lower than the costs based on part 2 (i.e., €188 and €100 vs €1401 and €863 in T1DM and T2DM, respectively). There might be several explanations: first, the results from part 1 might suffer from recall bias, since patients were asked about healthcare use in the previous 6 months and productivity losses in the previous year and as a consequence they might not remember using healthcare or being absent from work. As a result, the costs based on part 1 of the SAQ might be underestimated. Second, the results from part 2, especially those related to T1DM, are largely driven by two patients (i.e., one with sick leave from work for 21 days and one with sick leave from work for 10 days).

Third, not all patients (88%) who participated in the study and completed part 1 of the SAQ filled in part 2 of the SAQ and the diary. A comparison of the baseline characteristics (i.e., age, gender, years with diabetes and years on insulin) of the patients who did and did not complete part 2 did not show significant differences, suggesting that the patients completing part 2 are representative of all patients participating in the HAT study (although non-significant findings might be related to small sample size). The only significant difference we found was a difference in age, that is, the patients who did not complete part 2 were younger than the patients who did complete part 2. This is explained partly by the type of diabetes among the two groups, that is, 15% of the patients with T1DM did not complete part 2 compared with 8% of the patients with T2DM. However, since all results are reported separately for patients with T1DM and T2DM, the impact of the potential under-representation of patients with T1DM is limited. It is further assumed that the samples (T1DM and T2DM) reflect the total insulin-treated T1DM and T2DM population (treated with insulin for at least 12 months) in the Netherlands, because all consecutive patients fulfilling the inclusion criteria were enrolled during routine scheduled clinical appointments with their healthcare provider. Demographic properties (age and gender) were similar in our study compared with the overall diabetes population in the Netherlands. Studies in other Western European countries show similar results with respect to the incidence and prevalence of hypoglycaemic events. The HAT study only included patients using insulin for at least 12 months. As patients who started using insulin more recently receive extra training about the risks of hypoglycaemia, they might be more focused on hypoglycaemic events and might be more in control when these occur. Consequently, the number of hypoglycaemic events might be lower compared with patients who are treated with insulin for >12 months. Since the number of newly treated patients (<12 months insulin use) is likely to be low, the impact on the results would be limited. Nonetheless, these patients were excluded from the analyses, so that the sample consisted of patients in which changes in insulin treatment were not expected.

Fourth, the incidence of hypoglycaemic events according to the patient diary was higher than reported in the SAQ. As mentioned by Khunti et al., results from the SAQ might suffer from recall bias and the number of events might therefore be underestimated. On the contrary, double counting might occur in the SAQ in case a patient had an event resulting in hospital admission or an event requiring assistance from medical personnel but not requiring hospital admission and required assistance from another person for the same event. As a consequence, results from the SAQ might be overestimated. Although we present the costs per event based on both incidences, we believe that the cost per event based on the incidence as estimated from the SAQ is more reliable, as data on healthcare utilisation and frequency of hypoglycaemic events were derived from the same source. Nevertheless, the number of severe events seems relatively high, that is, 22 events in 13 patients with T1DM and 51 events in 30 patients with T2DM in a 4-week period. Two patients with T1DM experienced three events requiring assistance from another person to administer carbohydrate and/or glucagon and two patients had two of these events, which increases the total number of events in the population. Similarly, there were six patients with T2DM with more than one event requiring assistance from another person to administer carbohydrate and/or glucagon with one patients having six of these events.

Fifth, productivity losses due to a reduced ability to work efficiently (i.e., presenteeism) were not taken into account, because no data about presenteeism were available. Since it seems plausible that patients who experienced a hypoglycaemic event but are not absent from work will work less efficiently, total productivity costs may be underestimated. Additionally productivity costs may be underestimated, because productivity losses of unpaid work were neglected (e.g., household activities). In contrast, productivity losses could be overestimated because patients might have attributed productivity losses to hypoglycaemia that were actually related to other
causes (eg, comorbidities). However, as the questionnaire specifically asks for productivity losses that were related to hypoglycaemia, such a potential overestimation is likely to be limited. Future studies could use a control group consisting of patients with diabetes but without hypoglycaemia to address this issue.

In conclusion, this study showed that the economic burden of hypoglycaemia is substantial, especially the burden of severe events. Important differences between the costs of hypoglycaemia in T1DM and T2DM were found. While productivity costs are the most important cost driver of hypoglycaemia in patients with T1DM, costs within the healthcare sector are the most important cost driver of hypoglycaemia in patients with T2DM. As the costs of hypoglycaemia are substantial, the prevention of hypoglycaemia could not only reduce the burden for patients, but also the economic burden to society, if the costs of prevention are smaller than the costs of hypoglycaemia. Increasing awareness among physicians about the frequent occurrence of hypoglycaemia is essential to prevent hypoglycaemia; as patients might not always report these for various reasons, including fear or shame, or because they do not see the relevance, physicians might be unaware of (the frequency of) hypoglycaemic events. Furthermore, it is important to incorporate the costs of hypoglycaemia in cost-effectiveness analysis of treatments for diabetes, for example, to study potential cost-savings if treatment reduces the number of (severe) hypoglycaemic events. However, given the large variation in costs per event, the results from this study should be used carefully and tested using sensitivity analyses to acknowledge the uncertainty around our estimates.

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Patient consent Obtained.

Ethics approval The Global HAT study was conducted in accordance with the Declaration of Helsinki; Ethical Principles for Medical Research Involving Human Patients. The protocol was subject to review/approval by appropriate country-specific regulatory agencies/ethics committees.

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Data sharing statement The data generated during and/or analysed during the current trial are available from the corresponding author on reasonable request.

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