Cost-effectiveness of insulin degludec versus insulin glargine U100 in adults with type 1 and type 2 diabetes mellitus in Bulgaria

CURRENT STATUS: ACCEPTED

Monika Russel-Szymczyk
Novo Nordisk Pharma Sp. z o.o.
zmns@novonordisk.com Corresponding Author
ORCID: https://orcid.org/0000-0003-3958-5280

Vasil Valov
Novo Nordisk Pharma EAD

Alexandra Savova
Faculty of Pharmacy, Medical University Sofia, Bulgaria

Manoela Manova
Faculty of Pharmacy, Medical University Sofia, Bulgaria

DOI: 10.21203/rs.2.14752/v2

SUBJECT AREAS
Endocrinology & Metabolism

KEYWORDS
Cost-effectiveness, Diabetes, Hypoglycaemia, ICER, Insulin degludec, QALY
Abstract

Background This study investigates the cost-effectiveness of insulin degludec (degludec) versus biosimilar insulin glargine U100 (glargine U100) in patients with type 1 (T1DM) and type 2 diabetes mellitus (T2DM) in Bulgaria.

Methods A simple, short-term model was used to compare the treatment costs and outcomes associated with hypoglycaemic events with degludec versus glargine U100 in patients with T1DM and T2DM from the perspective of the Bulgarian National Health Insurance Fund. Cost-effectiveness was analysed over a 1-year time horizon using data from clinical trials. The outcome measure was the incremental cost-effectiveness ratio (ICER).

Results Degludec was highly cost-effective versus glargine U100 in people with T1DM and T2DM in Bulgaria. The ICERs were estimated to be 4,493.68 BGN/quality-adjusted life year (QALY) in T1DM, 399.11 BGN/QALY in T2DM on basal oral therapy (T2DM BOT ) and 7,365.22 BGN/QALY in T2DM on basal bolus therapy (T2DM B/B ), which all fall below the cost-effectiveness threshold of 39,619 BGN in Bulgaria. Insulin costs were higher with degludec in all three patient groups, however these were partially offset by savings from a reduction in hypoglycaemic events with degludec versus glargine U100. Sensitivity analysis demonstrated that the results were robust and largely insensitive to variations in input parameters. At a willingness-to-pay threshold of 39,619 BGN/QALY, the probability of degludec being cost-effective versus glargine U100 was 60.0% in T1DM, 99.4% in T2DM BOT and 91.3% in T2DM B/B .

Conclusion Degludec is a cost-effective alternative to biosimilar glargine U100 for patients with T1DM and T2DM in Bulgaria. Degludec could be of particular benefit to those patients suffering recurrent hypoglycaemia and those who require additional flexibility in the dosing of insulin.

Background

The treatment of diabetes and its complications represents a major economic burden to healthcare systems worldwide. An estimated 425 people aged 20–79 years worldwide had diabetes in 2017 with an associated healthcare expenditure of approximately $727 billion [1]. This is projected to reach $776 billion by 2045. The mean diabetes-related expenditure in Bulgaria per person with diabetes is
$798 per year [1]. The treatment of diabetes-related complications, such as cardiovascular disease, retinopathy and neuropathy, makes up the largest part of the direct medical costs associated with diabetes care, with less than 10% spent on insulin and anti-diabetic drugs [2, 3]. Indirect costs associated with diabetes, such as absenteeism from work and reduced productivity while at work are also considerable [4].

The goal of diabetes treatment is to achieve good glycaemic control to prevent or delay macro- and microvascular complications and reduce cardiovascular and all-cause mortality [5, 6]. All people with type 1 diabetes (T1DM) require insulin. The Bulgarian Society of Endocrinology recommends the use of intensive insulin therapy with a basal-bolus regimen for people with T1DM [7]. Type 2 diabetes (T2DM) is a progressive disease and although glycaemic control can initially be achieved with the use of other anti-hyperglycaemic agents, a large proportion of people will eventually require insulin to achieve glycaemic targets [8]. The Bulgarian Society of Endocrinology guidelines recommend a patient-centred approach, and indicate that the treatment should be individually adjusted to patient age, disease progression, existing co-morbidities and preference [7]. Insulin is the most effective therapy for reducing blood sugar [8, 9], however, despite clear guidelines [10, 11], glycaemic control remains sub-optimal (HbA$_{1c}$ >7%) in a substantial number of patients [12–15]. In Bulgaria, approximately 50% of patients with T2DM in specialist care and 57% of patients in primary care have HbA$_{1c}$ $\geq$6.5%, and for many patients regular testing and subsequent treatment adjustment is uncommon [16]. Insulin is often underutilised, as it is not initiated in a timely manner, titrated properly, or intensified appropriately [17, 18]. Key barriers to the use of insulin include hypoglycaemia and fear of hypoglycaemia, weight gain and restrictive treatment regimens [19].

Insulin degludec (degludec) is a basal insulin with a duration of action of more than 42 h, and a distinct, slow absorption mechanism, which results in a flat and stable action profile [20, 21]. Degludec has a four times lower day-to-day variability in glucose-lowering effect compared with insulin glargine (glargine) U100 and U300 [22–24]. Due to its stable and long action profile, degludec allows for flexibility in the timing of insulin administration and allows people to advance or delay
administration with no impact on short-term glycaemic control and minimal risk of hypoglycaemia [25].

The efficacy and safety of degludec have been evaluated in a large scale clinical trial programme (BEGIN) which included more than 9,000 people with T1DM and T2DM spanning the entire treatment spectrum of insulin treatment for T1DM and T2DM [26]. Meta-analyses of the phase 3a clinical trials demonstrated that degludec mediated equivalent reductions in HbA$_1c$ with a lower risk of hypoglycaemia compared with glargine U100 at a significantly lower total daily insulin dose in T1DM and T2DM with basal-only insulin [27, 28]. These results have been confirmed in real-world studies demonstrating better glycaemic control and fewer episodes of hypoglycaemia in patients switching from glargine U100 or other basal insulins to degludec [29, 30].

With the ever-increasing constraints on the healthcare budget, new interventions should represent good value for money. Understanding both the economic and clinical impact of an intervention helps decision makers determine resource use and optimal care for patients. Cost-effectiveness analyses assess the value of interventions by comparing the relative cost and outcomes. The relative differences are typically presented as an incremental cost-effectiveness ratio (ICER), which is the difference of costs of two interventions divided by the difference in health effects. A commonly accepted measure of effectiveness in cost-effectiveness analyses is the quality-adjusted life year (QALY) which takes into account both life expectancy and health-related quality of life [31]. The incremental cost per additional QALY gained allows decision makers to broadly compare across disease areas and allocate healthcare resources for maximal economic and clinical benefits. A financial threshold is often set where cost-effectiveness is accepted. In Bulgaria, 3x the gross domestic product (GDP) per capita, as suggested by the World Health Organization [32], is the commonly accepted threshold (39,619 BGN/QALY).

Recent cost-effectiveness analyses have demonstrated that degludec is cost-effective versus glargine U100 (Lantus®) in the United Kingdom and Serbia [33, 34]. Biosimilar insulin glargine U100 (Abasaglar®) has recently entered the basal insulin analogue market in Bulgaria, expanding therapeutic options for patients. There are currently no head-to-head data of degludec versus
Abasaglar®, however, cost-effectiveness analyses can be conducted using available data and plausible assumptions.

The objective of this study was to assess the cost-effectiveness of degludec versus biosimilar glargine U100 in the treatment of adults with T1DM and T2DM in Bulgaria from the perspective of the Bulgarian National Health Insurance Fund (NHIF).

**Methods**

**Model Overview**

This cost-effectiveness model compared degludec with glargine U100 in three separate patient groups: T1DM using basal-bolus therapy (T1DM<sub>B/B</sub>); T2DM using basal-oral therapy (T2DM<sub>BOT</sub>); T2DM using basal-bolus therapy (T2DM<sub>B/B</sub>). Biosimilar glargine U100 (Abasaglar®) was chosen as the most appropriate comparator in the economic analyses as glargine is the most widely used basal insulin analogue in Bulgaria and biosimilar glargine U100 serves as reference price per daily defined dose (DDD).

The analysis assumes that Abasaglar® has the same efficacy and safety as originator glargine U100 (Lantus®).

For the current cost-effectiveness analysis, a simple and transparent short-term (1-year time horizon) model was developed in Microsoft® Excel 2010 (Microsoft Corp., Redmond, WA, US) to capture the direct medical costs associated with insulin treatment and hypoglycaemia (Figure 1). The effectiveness (QALY) was calculated by multiplying the disutility per hypoglycaemic event by the number of events in each treatment group.

The short-term model is appropriate as the treat-to-target trial design enforces a similar level of glycaemic control across comparators and no difference in terms of glycaemic control (difference in HbA<sub>1c</sub>) is expected. Therefore, there is no rationale for long-term modelling based on reductions in HbA<sub>1c</sub>. Although the cost-effectiveness of degludec was analysed in a short-term setting and based on data from 1-year clinical studies, results are not only applicable for the cost-effectiveness of degludec in the first year of treatment. As the model can be replicated for subsequent years, the outcomes represent the average annual cost-effectiveness in a steady state. As the time horizon was 1 year, no
discounting was applied.

Figure 1. Overview of cost-effectiveness model

Abbreviations: Δ, change in; HC, healthcare, HRQoL, health-related quality of life; ICER, incremental cost-effectiveness ratio; QALY, quality-adjusted life year; SMBG, self-monitored blood glucose

Clinical Data

Insulin Dose

Units of insulin used per day for glargine U100 were taken from daily insulin doses used in clinical practice in Bulgaria [35]. The degludec/glargineU100 dose ratio was derived from a meta-analysis of insulin dose [27]. The degludec dose was calculated using the dose ratio to allow for adjustment of covariate factors such as trial, treatment, anti-diabetic therapy at screening, age, sex, region, and baseline dose (Table 2). The insulin dose and dose ratio was solely used to calculate costs and was not included as a clinical outcome.

Hypoglycaemia Event Rates

Real-world non-severe hypoglycaemic event rates were derived from a large-scale multi-national survey of patients and physicians, and severe event rates were derived from a published observational study from the UK Hypoglycaemia Study Group (UKHSG) [36, 37]. These rates, derived from observational studies, provide a better estimation of real life event rates than clinical trial data which can be biased in the selection of patients and the treatment setting. The real-world hypoglycaemic event rates were taken as the base-case event rates in the glargine U100 group. Event rates for the degludec group were calculated using the degludec/glargine U100 hypoglycaemia rate ratios derived from a meta-analysis of hypoglycaemia which are adjusted for trial, type of diabetes, treatment, anti-diabetic therapy at screening, sex, region and age [27]. Only significant results were used for the calculation; in case of non-significant results, a rate ratio of one was used in the calculation (Table 3).

Perspective

All costs and resources used were estimated from a healthcare payer perspective of the Bulgarian NHIF.

Data Used in the Model
Direct Treatment Costs

The cost of insulin was calculated based on the pharmacy selling price (pharmacy purchase price, including VAT) in 2018, in BGN. The cost of needles and self-monitoring of blood glucose (SMBG) tests are not reimbursed by NHIF and are paid out of pocket by patients.

Cost of Hypoglycaemia

The direct costs of treating severe and non-severe hypoglycaemic events were estimated based on the resource use data collected in the clinical trial programme. For non-severe events, data were derived from completed hypoglycaemia safety questionnaires. For severe events, data were obtained from the serious adverse events case reports [38].

For patients with severe hypoglycaemia events, it was assumed that all patients who were hospitalised used the ambulance service to get to hospital, and all patients used glucagon to recover. It was assumed that the number of patients contacting a hospital or healthcare professional following a hypoglycaemic event, and the number of SMBG test strips used after an event, were similar between degludec and glargine U100 and that the differences between treatment-related costs of hypoglycaemia events stemmed only from the differences in hypoglycaemia rates between the two interventions.

The average costs for hypoglycaemic events are summarised in Table 1. It was assumed that the cost of treating a severe or non-severe hypoglycaemic event (diurnal and nocturnal) was similar regardless of whether the event was experienced by patients with T1DM, T2DM_{BOT} or T2DM_{B/B}.

Utility Data

QALYs were calculated by applying a disutility [39] per hypoglycaemic event. Disutilities incurred per hypoglycaemic event were taken from a large-scale time trade off (TTO) study, which reported a disutility of 0.0565 for a severe event (with no significant difference between daytime and nocturnal severe events) and a disutility of 0.0041 and 0.0067 for non-severe daytime and nocturnal events, respectively (a significant difference in utility was demonstrated for nocturnal vs daytime non-severe events) [39]. The disutility per hypoglycaemic event was multiplied by the number of events observed in each treatment group.
The analysis included an additional QALY benefit for flexible dosing associated with degludec. According to a study by Boye et al. [40] flexible dosing is associated with a 0.006 utility gain, which was applied to the QALY benefit of degludec.

**Sensitivity Analyses**

One-way and probabilistic sensitivity analyses were conducted to assess the impact of varying key assumptions and outcomes used in the base case analysis. (Table 4).

**Results**

**Costs**

The total costs for the three treatment groups are presented in Table 5.

In T1DM, the total costs were estimated to be 3,143.28 BGN per patient per year in the degludec group and 3,073.92 BGN per patient per year for glargine U100, with approximately 40% of costs attributable to insulin and the remainder primarily due to severe hypoglycaemia. Total costs were 69.37 BGN higher (2.3% higher) in the degludec group than the glargine U100 group, which is due to the increased insulin costs with degludec, partially offset (47.19 BGN) by lower costs of hypoglycaemia due to a significantly lower rate of non-severe nocturnal hypoglycaemia versus glargine U100.

In T2DM\textsubscript{BOT}, the total costs per patient per year were 1.6% higher in the degludec group (1,101.32 BGN) versus the glargine U100 group (1,083.97 BGN). The incremental costs were mainly driven by the increased cost of insulin, partially offset by lower costs of hypoglycaemia in the degludec group. Lower costs of hypoglycaemia were driven by significant reductions in the number of non-severe nocturnal and severe hypoglycaemic events in the degludec group versus glargine U100.

In the T2DM\textsubscript{B/B} group, the total costs per person per year were 304.21 BGN (16.4%) higher in the degludec group (2,156.67 BGN) versus the glargine U100 group (1,852.47 BGN). The incremental costs with degludec were caused by increased cost of insulin which were partially offset by lower costs of non-severe hypoglycaemia. In the clinical trials, the dose of basal insulin in the degludec treatment arm was higher than in the glargine U100 arm [27] which drives the higher incremental costs in this group.

**Incremental cost-effectiveness**
The incremental costs per QALY gained with degludec versus glargine U100 was estimated at 4,499 BGN, 399 BGN and 7,365 BGN in T1DM, T2DM\_BOT and T2DM\_B/B, respectively. In all three settings, degludec was highly cost-effective versus glargine U100 with the ICER values falling considerably below the cost-effectiveness threshold assumed for Bulgaria (39,619 BGN) (Table 5).

**Sensitivity analysis**

One-way sensitivity analyses demonstrate that the results are robust and largely insensitive to variations in input parameters (Supplementary Table S 1–Table S 3). The parameter with most influence on the ICER was the rate of hypoglycaemia. In T1DM, degludec remained highly cost-effective versus glargine U100 in all scenarios tested. The ICERs ranged between 45 and 19,781 BGN/QALY gained, with hypoglycaemic event rates having most influence on the ICER. When the number of non-severe and severe hypoglycaemic events was increased to those reported by Ericsson et al. [41], the ICER decreased to 45 BGN/QALY gained. Conversely, when the lower hypoglycaemic event rates reported in clinical trials [27, 28] were used, the ICER increased to 5,472 BGN/QALY gained. Assuming no difference in the number of hypoglycaemic events between degludec and glargine U100 had the greatest effect on cost-effectiveness, increasing the ICER to 19,781 BGN/QALY gained. However, even in this scenario, degludec remained cost-effective vs glargine U100.

In T2DM\_BOT, the favourable cost-effectiveness results were invariant to changes in most of the input parameters tested. ICERs ranged between 32 BGN/QALY and 36,739 BGN/QALY gained. Using a lower number of hypoglycaemic events as reported in the clinical trials [27, 28], resulted in an increase in the ICER to 24,817 BGN/QALY gained but degludec remained cost-effective versus glargine U100 in this scenario. Conversely, using higher rates for severe hypoglycaemic events reported by Ericsson et al [41], degludec became dominant versus glargine U100. Assuming no difference in hypoglycaemic event rates between degludec and glargine U100 had the greatest impact on the ICER, which increased to 36,739 BGN/QALY gained.

In T2DM\_B/B, the ICER was stable to reasonable variations in input parameters and was below the commonly accepted threshold for cost-effectiveness in almost all scenarios tested. The ICER increased when lower non-severe hypoglycaemic event rates were assumed (ICER 8,141 BGN/QALY gained).
gained) and when assuming no difference in the non-severe hypoglycaemic event rates between degludec and glargine U100 (ICER 13,826–15,236 BGN/QALY gained). The ICER exceeded the cost-effectiveness threshold when no difference in the hypoglycaemia rate ratios was assumed (63,239 BGN/QALY).

**Probabilistic sensitivity analysis**

To account for the uncertainty in the results caused by variation in data inputs, probabilistic sensitivity analyses were conducted. The cost-effectiveness acceptability curves demonstrated that, at a willingness-to-pay threshold of 39,619 BGN/QALY, the probability of degludec being cost-effective versus glargine U100 was 60.0% in T1DM, 99.4% in T2DM<sub>BOT</sub> and 91.3% in T2DM<sub>B/B</sub> (Figure 2).

**Figure 2. Probabilistic sensitivity analyses acceptability curves**

**Discussion**

There are limited data on the cost-effectiveness of insulin treatments in Bulgaria. With the increasing number of insulin treatment options, decision making based on economic evidence is essential to maximise health outcomes while effectively managing restrained budgets.

In this simple, short-term cost-effectiveness analysis, degludec was demonstrated to be cost-effective versus biosimilar glargine U100 in people with T1DM (ICER 4,498.68 BGN), T2DM<sub>BOT</sub> (ICER 399.11 BGN) and T2DM<sub>B/B</sub> (ICER 7,365.22 BGN). In all three patient groups insulin costs with degludec were higher than with glargine U100. However, these costs were offset by a reduction in the number of hypoglycaemic events with degludec in all three patient groups.

Cost-effectiveness analyses usually model the long-term impact of diabetes interventions on disease-related complications as a function of the differences in glycaemic control. However the data used in this model were derived from treat-to-target trials and glycaemic control was similar across both arms, thus the use of a long-term model based on differences in HbA<sub>1c</sub> was not appropriate.

Therefore, this simple, transparent short-term model focuses on other important aspects associated with insulin therapy, including hypoglycaemia and insulin dosing. Although this model only reflects a 1-year time horizon, it not only represents the cost-effectiveness of degludec vs glargine U100 within
the first year of treatment, but it can also be replicated for subsequent years, representing annual cost-effectiveness. This is supported by the insensitivity of the ICER to changes in the time horizon in the sensitivity analyses. The model has previously been used to evaluate the cost-effectiveness of degludec versus glargine U100 in patients with T1DM and T2DM in different settings [33, 34, 41–44].

In the United Kingdom (UK) and Denmark, degludec was found dominant versus glargine U100 in T1DM and T2DM\textsubscript{BOT} and highly cost-effective in T2DM\textsubscript{B/B} [33, 42]. Similarly, in Serbia and Sweden, degludec was cost-effective versus glargine U100 in all three patient groups [34, 41]. These results are consistent with those observed in the current study.

Hypoglycaemia can have a major impact on patient’s quality of life and lead to significant psychological and physical morbidity and in severe cases death [45, 46]. Additionally, people with recurrent hypoglycaemic episodes are at risk of developing impaired awareness of hypoglycaemia which can cause patients to miss early symptoms and treat hypoglycaemia, increasing the risk of severe hypoglycaemic events [47]. A recent real world multi-national, non-interventional study assessed the prevalence of hypoglycaemia worldwide, including Bulgaria. In Eastern Europe the estimated overall annual rate of hypoglycaemic events was 66.9 per person per year in T1DM and 23.7 per person per year in T2DM [48]. The prevalence of nocturnal and severe hypoglycaemia were 9.8 and 4.5 per person per year in T1DM and 4.0 and 2.2 per person per year in T2DM. The nocturnal hypoglycaemic event rates in T1DM and severe events in T1DM and T2DM in Bulgaria are higher than those used in this model which, based on the impact of hypoglycaemic event rates demonstrated in the sensitivity analyses, suggests that degludec may be even more cost-effective in real world clinical practice in Bulgaria.

The real-world cost effectiveness of switching patients with T1DM from other basal insulins (glargine U100, insulin detemir and NPH insulin) to degludec has been investigated using the IQVIA CORE diabetes model from the perspective of the United Kingdom and Sweden [30, 49]. The CORE diabetes model is a lifetime Markov model predicting diabetes complications over time in patient populations representative of clinical practice and calculates the resulting economic impact. From both the UK and Swedish perspective, degludec was dominant versus other basal insulins, which was mainly
driven by the significant reduction in HbA$_{1c}$ and lower rates of hypoglycaemia with degludec [30, 49]. Hypoglycaemia and the fear of hypoglycaemia are key barriers to the initiation and intensification of insulin regimens [50]. As a result of fear of hypoglycaemia, approximately 52% of people with T1DM and 41% of people with T2DM reduce their insulin dose following a hypoglycaemic event [51]. This compromises glycaemic control and puts patients at risk of serious long-term complications, such as cardiovascular disease, renal disease, retinopathy, neuropathy and amputations [52]. The unique pharmacological profile of degludec with a flat and stable action profile is associated with lower rates of hypoglycaemia [20, 22]. In phase 3a clinical trials comparing degludec with glargine U100, degludec demonstrated equivalent reductions in HbA$_{1c}$ with significantly lower rates of hypoglycaemia. In T1DM, degludec was associated with a 17% lower rate of non-severe nocturnal events, while in T2DM$_{\text{BOT}}$ and T2DM$_{\text{B/B}}$, rates of non-severe nocturnal hypoglycaemia were decreased by 36% and 25%, respectively [27]. Additionally, the rate of non-severe daytime events was 17% lower with degludec in T2DM$_{\text{B/B}}$, and the rate of severe event hypoglycaemia was 86% lower in T2DM$_{\text{BOT}}$ [27, 28]. The SWITCH 1 and SWITCH 2 trials, two phase 3b studies, sought to confirm the efficacy and safety of degludec in people with T1DM and T2DM with an increased risk of hypoglycaemia [53, 54]. In these studies, degludec achieved equivalent reductions in HbA$_{1c}$ with a significantly lower rate of overall symptomatic and severe hypoglycaemia versus glargine U100. These trials are more representative of patients in regular clinical practice than the phase 3a clinical trials, which excluded patients with recurrent hypoglycaemia. Data from SWITCH trials have recently been used to demonstrate that degludec is a cost-effective alternative to glargine U100 in the UK [55]. Furthermore, the hypoglycaemia benefit of degludec has been confirmed in real-world studies which demonstrate that switching to degludec from other basal insulin regimens is associated with significantly improved glycaemic control and a reduction in the rate of non-severe and severe hypoglycaemic events in T1DM and T2DM [29, 30].

In Bulgaria, SMBG tests are not paid by the healthcare payer but are paid out-of-pocket by patients,
and were therefore not considered in this analysis. Due to the long duration of action with a flat and stable action profile and the lower day-to-day variability versus glargine U100, fewer SMBG tests are needed for titration and maintenance with degludec [20, 22, 56]. Fewer SMBG tests would be cost saving for the patient, although this was not assessed in the present analysis. The long and stable action profile of degludec also allows for flexibility of dosing time without compromising efficacy or risk of hypoglycaemia [25]. This flexible dosing option may provide an additional benefit especially for those people who have difficulties adhering to their treatment regimens (e.g. shift workers, frequent travellers, patients requiring help with insulin injections). Here, an estimate of the utility benefit of flexible dosing with degludec was included in the analysis and a utility gain of 0.006 derived from the study by Boye et al. [40] was applied for degludec. This can be considered a conservative estimate as a large time trade-off study identified a utility gain of 0.016 associated with flexible dosing of basal insulin and 0.013 vs fixed basal-bolus regimens [57].

It is important to acknowledge the limitations associated with the current analysis. With biosimilar glargine U100 being a relatively new-to-market insulin, no head-to-head trial data are available comparing degludec vs biosimilar glargine U100 and the analysis was based on currently available data and plausible assumptions, and the results should be interpreted accordingly. The analysis used hypoglycaemic event rates derived from published economic analyses using this model [33]. This could have resulted in an underestimation of the cost-effectiveness of degludec vs glargine U100 in Bulgaria, as a recent study suggests that rates of hypoglycaemia in real-world clinical practice in Bulgaria may be higher [48]. Additionally, the data used in this model to inform hypoglycaemia rate ratios originate from meta-analyses of phase 3a clinical trials [27, 28] to increase the statistical power of the analyses and increase the reliability of the data, and assumes replication of such rates in clinical practice. Except for hypoglycaemia, adverse events are not considered in our analysis, as head-to-head clinical trials have shown that degludec and glargine have similar safety profiles [26]. Furthermore, the cardiovascular outcomes trial DEVOTE showed that insulin degludec was non-inferior to insulin glargine in terms of major adverse cardiovascular events [58]. The clinical trials which informed this analysis used a treat-to-target approach in which patients were titrated until glycaemic
targets were reached. In clinical practice, glycaemic targets are often not met for a variety of reasons, including non-adherence, and missed follow-up appointments. However, sensitivity analyses demonstrate that the results are robust to a wide variation in parameters, supporting the validity of the results.

**Conclusions**

This short-term cost-effectiveness analysis demonstrates that degludec is a cost-effective alternative to biosimilar glargine U100 for patients with T1DM and T2DM in Bulgaria. Degludec could be of particular benefit to those patients suffering recurrent hypoglycaemia and those who require additional flexibility in the dosing of insulin.

**Abbreviations**

| Abbreviation | Definition |
|--------------|------------|
| BGN          | Bulgarian LEV |
| CI           | Confidence interval |
| DDD          | Daily defined dose |
| GDP          | Gross domestic product |
| HC           | Healthcare |
| HRQoL        | Health-related quality of life |
| ICER         | Incremental cost-effectiveness ratio |
| NHIF         | National Health Insurance Fund |
| QALY         | Quality-adjusted life year |
| SMBG         | Self-monitored blood glucose |
| NS           | Non-significant |
| T1DM         | Type 1 diabetes mellitus |
| T1DM_{B/B}   | T1DM using basal-bolus therapy |
| T2DM         | Type 2 diabetes mellitus |
| T2DM_{B/B}   | T2DM using basal-bolus therapy |
| T2DM_{BOT}   | T2DM using basal-oral therapy |
| TTO          | Time trade off |

**Declarations**

**Ethics approval and consent to participate**

Not applicable

**Funding**

The study itself and the article processing charges were funded by Novo Nordisk. All authors had full access to all data in this study and take complete responsibility for the integrity of the data and accuracy of the data analysis.

**Author contributions**

All named authors (MR, VV, AS, MM) meet the International Committee of Medical Journal Editors (ICMJE) criteria for authorship for this article, take responsibility for the integrity of the work as a
whole, and have given their approval for this version to be published.

Data availability
All data on which the conclusions of the manuscript are based are presented in the main paper or additional supporting files.

Acknowledgements
Medical Writing and Editorial Assistance
The authors acknowledge medical writing and editorial support by Katrin Nather and Carrie Fidler of DRG Abacus (sponsored by Novo Nordisk).

Competing interests
Alexandra Savova and Manoela Manova have nothing to declare. Monika Russel-Szymczyk and Vasil Valov are employees of Novo Nordisk.

Availability of data and materials
All data generated or analysed during this study are included in this published article/as supplementary information files.

Ethics approval and consent to participate
This article does not contain any studies with human participants or animals performed by any of the authors.

References
1. International Diabetes Federation (IDF). Diabetes atlas - 8th edition. Available at: http://www.diabetesatlas.org/.

2. American Diabetes Association. Economic costs of diabetes in the U.S. in 2012. Diabetes Care. 2013 Apr;36(4):1033–46.

3. Jonsson B. Revealing the cost of Type II diabetes in Europe. Diabetologia. 2002;45(7):S5–12.

4. American Diabetes Association. Economic costs of diabetes in the U.S. in 2017. Diabetes Care. 2018 May;41(5):917–28.

5. Stratton IM, Adler AI, Neil HA, Matthews DR, Manley SE, Cull CA, et al. Association of glycaemia with macrovascular and microvascular complications of type 2 diabetes (UKPDS 35): prospective observational study. BMJ. 2000 Aug 12;321(7258):405–12.

6. The Diabetes Control and Complications Trial Research Group. The effect of intensive treatment of
diabetes on the development and progression of long-term complications in insulin-dependent diabetes mellitus. The Diabetes Control and Complications Trial Research Group. N Engl J Med. 1993 Sep 30;329(14):977-86.

7. Bulgarian Society of Endocrinology. Good clinical practice guidelines on diabetes mellitus. 2016.

8. Davies MJ, D’Alessio DA, Fradkin J, Kernan WN, Mathieu C, Mingrone G, et al. Management of hyperglycemia in type 2 diabetes, 2018. A consensus report by the American Diabetes Association (ADA) and the European Association for the Study of Diabetes (EASD). Diabetes Care. 2018 Dec;41(12):2669–701.

9. Giugliano D, Maiorino M, Bellastella G, Chiodini P, Ceriello A, Esposito K. Efficacy of insulin analogs in achieving the hemoglobin A1c target of <7% in type 2 diabetes: meta-analysis of randomized controlled trials. Diabetes Care. 2011 Feb;34(2):510-7.

10. International Diabetes Federation (IDF) Clinical Guidelines Task Force. Global guideline for type 2 diabetes, 2012. Available at https://www.idf.org/e-library/guidelines/79-global-guideline-for-type-2-diabetes

11. American Diabetes Association. Standards of medical care in diabetes–2018. Diabetes Care. 2018 Jan;41(Suppl 1):S1-S153.

12. Blak BT, Smith HT, Hards M, Curtis BH, Ivanyi T. Optimization of insulin therapy in patients with type 2 diabetes mellitus: beyond basal insulin. Diabet Med. 2012 Jul;29(7):e13–20.

13. de Pablos-Velasco P, Parhofer KG, Bradley C, Eschwege E, Gonder-Frederick L, Maheux P, et al. Current level of glycaemic control and its associated factors in patients with type 2 diabetes across Europe: data from the PANORAMA study. Clin Endocrinol (Oxf). 2014 Jan;80(1):47-56.

14. Govan L, Wu O, Briggs A, Colhoun HM, Fischbacher CM, Leese GP, et al. Achieved levels of HbA1c and likelihood of hospital admission in people with type 1 diabetes in the Scottish population: a study from the Scottish Diabetes Research Network Epidemiology Group. Diabetes Care. 2011 Sep;34(9):1992–7.

15. NHS Health & Social Care Information Centre. National Diabetes Audit 2011–2012; Report 1: Care Processes and Treatment Targets, available at http://content.digital.nhs.uk/catalogue/PUB12421/nati-
16.Cokolic M, Lalic NM, Micic D, Mirosevic G, Klobucar Majanovic S, Lefterov IN, et al. Patterns of diabetes care in Slovenia, Croatia, Serbia, Bulgaria and Romania. Wiener klinische Wochenschrift. 2017 2017/03/01;129(5):192–200.

17.Calvert MJ, McManus RJ, Freemantle N. Management of type 2 diabetes with multiple oral hypoglycaemic agents or insulin in primary care: retrospective cohort study. Br J Gen Pract. 2007;57(539):455–60.

18.Rubino A, McQuay LJ, Gough SC, Kvasz M, Tennis P. Delayed initiation of subcutaneous insulin therapy after failure of oral glucose-lowering agents in patients with Type 2 diabetes: a population-based analysis in the UK. Diabet Med. 2007;24(12):1412–8.

19.Peyrot M, Rubin RR, Lauritzen T, Skovlund SE, Snoek Fj, Matthews DR, et al. Resistance to insulin therapy among patients and providers: results of the cross-national Diabetes Attitudes, Wishes, and Needs (DAWN) study. Diabetes Care. 2005;28(11):2673–9.

20.Heise T, Nosek L, Bottcher SG, Hastrup H, Haahr H. Ultra-long-acting insulin degludec has a flat and stable glucose-lowering effect in type 2 diabetes. Diabetes Obes Metab. 2012 Oct;14(10):944–50.

21.Haahr H, Heise T. A review of the pharmacological properties of insulin degludec and their clinical relevance. Clin Pharmacokinet. 2014 Sep;53(9):787–800.

22.Heise T, Hermanski L, Nosek L, Feldman A, Rasmussen S, Haahr H. Insulin degludec: four times lower pharmacodynamic variability than insulin glargine under steady-state conditions in type 1 diabetes. Diabetes Obes Metab. 2012 Sep;14(9):859–64.

23.Heise T, Norskov M, Nosek L, Kaplan K, Famulla S, Haahr HL. Insulin degludec: Lower day-to-day and within-day variability in pharmacodynamic response compared with insulin glargine 300 U/mL in type 1 diabetes. Diabetes Obes Metab. 2017 Jul;19(7):1032–9.

24.Heise T, Kaplan K, Haahr HL. Day-to-Day and Within-Day Variability in Glucose-Lowering Effect Between Insulin Degludec and Insulin Glargine (100 U/mL and 300 U/mL): A Comparison Across Studies. J Diabetes Sci Technol. 2018 Mar;12(2):356–63.

25.European Medicines Agency. Tresiba® (insulin degludec) Summary of Product Characteristics.
Available at: https://www.ema.europa.eu/documents/product-information/tresiba-epar-product-information_en.pdf.

26. Wangnoo SK, Chowdhury S, Rao PV. Treating to target in type 2 diabetes: the BEGIN trial programme. J Assoc Physicians India. 2014 Jan;62(1 Suppl):21–6.

27. Vora J, Christensen T, Rana A, Bain SC. Insulin degludec versus insulin glargine in type 1 and type 2 diabetes mellitus: a meta-analysis of endpoints in phase 3a trials. Diabetes therapy: research, treatment and education of diabetes and related disorders. 2014 Dec;5(2):435–46.

28. Ratner R, Gough SC, Mathieu C, Prato SD, Bode B, Mersebach H, et al. Hypoglycaemia Risk With Insulin Degludec Compared With Insulin Glargine in Type 2 and Type 1 Diabetes: A Pre-planned Meta-Analysis of Phase 3 Trials. Diabetes Obes Metab. 2013 Nov 6;15(2):175–84.

29. Siegmund T, Tentolouris N, Knudsen ST, Lapolla A, Prager R, Phan TM, et al. A European, multicentre, retrospective, non-interventional study (EU-TREAT) of the effectiveness of insulin degludec after switching basal insulin in a population with type 1 or type 2 diabetes. Diabetes Obes Metab. 2018 Mar;20(3):689–97.

30. Evans M, McEwan P, Foos V. Insulin degludec early clinical experience: does the promise from the clinical trials translate into clinical practice—a case-based evaluation. J Med Econ. 2015 Feb;18(2):96–105.

31. Sassi F. Calculating QALYs, comparing QALY and DALY calculations. Health Policy Plan. 2006 Sep;21(5):402–8.

32. Macroeconomics and health: investing in health for economic development. Report of the Commission on Macroeconomics and Health. Geneva: World Health Organization; 2001. Available from: http://apps.who.int/iris/bitstream/10665/42435/1/924154550X.pdf.

33. Evans M, Chubb B, Gundgaard J. Cost-effectiveness of insulin degludec versus insulin glargine in adults with type 1 and type 2 diabetes mellitus. Diabetes therapy: research, treatment and education of diabetes and related disorders. 2017 Apr;8(2):275–91.

34. Lalic N, Russel-Szymbczyk M, Culic M, Tikkanen CK, Chubb B. Cost-Effectiveness of Insulin Degludec Versus Insulin Glargine U100 in Patients with Type 1 and Type 2 Diabetes Mellitus in Serbia. Diabetes
therapy: research, treatment and education of diabetes and related disorders. 2018 Jun;9(3):1201-16.

35. Doneva M, Valov V, Borisova A-M, Tankova T, Savova A, Manova M, et al. Comparative Analysis of the Cost of Insulin Treated Patients in Bulgaria. Biotechnology & Biotechnological Equipment. 2013 2013/01/01;27(2):3748-52.

36. UK Hypoglycaemia Study Group. Risk of hypoglycaemia in types 1 and 2 diabetes: effects of treatment modalities and their duration. Diabetologia. 2007 Jun;50(6):1140-7.

37. Brod M, Rana A, Barnett AH. Impact of self-treated hypoglycaemia in type 2 diabetes: a multinational survey in patients and physicians. Curr Med Res Opin. 2012 Dec;28(12):1947-58.

38. Heller SR, Frier BM, Herslov ML, Gundgaard J, Gough SC. Severe hypoglycaemia in adults with insulin-treated diabetes: impact on healthcare resources. Diabet Med. 2016 Jul 16;33:471–7.

39. Evans M, Khunti K, Mamdani M, Galbo-Jorgensen CB, Gundgaard J, Bogelund M, et al. Health-related quality of life associated with daytime and nocturnal hypoglycaemic events: a time trade-off survey in five countries. Health Qual Life Outcomes. 2013 Jun 3;11(1):90.

40. Boye KS, Matza LS, Walter KN, Van Brunt K, Palsgrove AC, Tynan A. Utilities and disutilities for attributes of injectable treatments for type 2 diabetes. Eur J Health Econ. 2011 Jun;12(3):219-30.

41. Ericsson A, Pollock RF, Hunt B, Valentine WJ. Evaluation of the cost-utility of insulin degludec vs insulin glargine in Sweden. J Med Econ. 2013 Dec;16(12):1442–52.

42. Pollock RF, Tikkanen CK. A short-term cost-utility analysis of insulin degludec versus insulin glargine U100 in patients with type 1 or type 2 diabetes in Denmark. J Med Econ. 2017 Mar;20(3):213-20.

43. Evans M, Wolden M, Gundgaard J, Chubb B, Christensen T. Cost-effectiveness of insulin degludec compared with insulin glargine for patients with type 2 diabetes treated with basal insulin - from the UK health care cost perspective. Diabetes Obes Metab. 2014 Apr;16(4):366–75.

44. Evans M, Wolden M, Gundgaard J, Chubb B, Christensen T. Cost-effectiveness of insulin degludec compared with insulin glargine in a basal-bolus regimen in patients with type 1 diabetes mellitus in the UK. J Med Econ. 2015 Jan;18(1):56–68.

45. Frier BM. The economic costs of hypoglycaemia. Br J Diabetes Vasc Dis. 2011;11(Suppl 1):S10-S2.
46. Khunti K, Davies M, Majeed A, Thorsted BL, Wolden ML, Paul SK. Hypoglycemia and risk of cardiovascular disease and all-cause mortality in insulin-treated people with type 1 and type 2 diabetes: a cohort study. Diabetes Care. 2015 Feb;38(2):316–22.

47. Graveling AJ, Frier BM. Impaired awareness of hypoglycaemia: a review. Diabetes Metab. 2010 Oct;36 Suppl 3:S64–74.

48. Khunti K, Alsifri S, Aronson R, Cigrovski Berkovic M, Enters-Weijnen C, Forsen T, et al. Rates and predictors of hypoglycaemia in 27,585 people from 24 countries with insulin-treated type 1 and type 2 diabetes: the global HAT study. Diabetes Obes Metab. 2016 Sep;18(9):907–15.

49. Landstedt-Hallin L, Gundgaard J, Ericsson A, Ellfors-Zetterlund S. Cost-effectiveness of switching to insulin degludec from other basal insulins: evidence from Swedish real-world data. Curr Med Res Opin. 2017 Apr;33(4):647–55.

50. Peyrot M, Barnett AH, Meneghini LF, Schumm-Draeger PM. Insulin adherence behaviours and barriers in the multinational Global Attitudes of Patients and Physicians in Insulin Therapy study. Diabet Med. 2012 May;29(5):682–9.

51. Khunti K, Alsifri S, Aronson R, Cigrovski Berkovic M, Enters-Weijnen C, Forsen T, et al. Impact of hypoglycaemia on patient-reported outcomes from a global, 24-country study of 27,585 people with type 1 and insulin-treated type 2 diabetes. Diabetes Res Clin Pract. 2017 Aug;130:121–9.

52. Diabetes UK. Facts and stats 2016. Available at: https://www.diabetes.org.uk/Documents/Position%20statements/DiabetesUK_Facts_Stats_Oct16.pdf.

53. Lane W, Bailey TS, Gerety G, Gumprecht J, Philis-Tsimikas A, Hansen CT, et al. Effect of Insulin Degludec vs Insulin Glargine U100 on Hypoglycemia in Patients With Type 1 Diabetes: The SWITCH 1 Randomized Clinical Trial. JAMA. 2017 Jul 04;318(1):33–44.

54. Wysham C, Bhargava A, Chaykin L, de la Rosa R, Handelsman Y, Troelsen LN, et al. Effect of Insulin Degludec vs Insulin Glargine U100 on Hypoglycemia in Patients With Type 2 Diabetes: The SWITCH 2 Randomized Clinical Trial. JAMA. 2017 Jul 04;318(1):45–56.

55. Evans M, Mehta R, Gundgaard J, Chubb B. Cost-effectiveness of insulin degludec vs. insulin glargine U100 in type 1 and type 2 diabetes mellitus in a UK setting. Diabetes therapy: research,
treatment and education of diabetes and related disorders. 2018 Oct;9(5):1919–30.

56. Philis-Tsimikas A, Brod M, Niemeyer M, Ocampo Francisco AM, Rothman J. Insulin Degludec Once-Daily in Type 2 Diabetes: Simple or Step-Wise Titration (BEGIN: Once Simple Use). Adv Ther. 2013 Jun;30(6):607–22.

57. Evans M, Jensen HH, Bogelund M, Gundgaard J, Chubb B, Khunti K. Flexible insulin dosing improves health-related quality-of-life (HRQoL): a time trade-off survey. J Med Econ. 2013 Nov;16(11):1357–65.

58. Marso SP, McGuire DK, Zinman B, Poulter NR, Emerson SS, Pieber TR, et al. Efficacy and safety of degludec versus glargine in type 2 diabetes. N Engl J Med. 2017 Aug 24;377(8):723–32.

59. Ostenson CG, Geelhoed-Duijvestijn P, Lahtela J, Weitgasser R, Markert Jensen M, Pedersen-Bjergaard U. Self-reported non-severe hypoglycaemic events in Europe. Diabet Med. 2014 Jan;31(1):92–101.

Tables

Table 1. Total costs of an average severe/non-severe hypoglycaemia event

|                    | Cost per hypoglycaemic event in patients with T1DM, T2DM<sub>BOT</sub>, T2DM<sub>B/B</sub> |
|--------------------|------------------------------------------------------------------------------------------------|
|                    | Non-severe daytime       | Non-severe nocturnal | Severe                        |    |
| TOTAL (BGN)/event (2014) | 0.66                      | 33.60                | 514.36                        |    |
| TOTAL (BGN)/event (inflated to 2018) | 0.65                      | 33.19                | 508.10                        |    |

Abbreviations: B/B, basal-bolus; BGN, Bulgarian LEV; BOT, basal oral therapy; T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

Table 2. Basal and bolus insulin use
| Treatment group | Observed glargine U100 (units/day) | Dose ratio (degludec/glargine U100) | Calculated degludec (units/day) |
|-----------------|-----------------------------------|-------------------------------------|-------------------------------|
| **T1DM<sub>B/B</sub>, total dose** | | | |
| Basal insulin | 28.11 | 0.87* | 24.46 |
| Bolus insulin | 37.13 | 0.88* | 32.67 |
| **T2DM<sub>BOT</sub>, total dose** | | | |
| Basal insulin | 28.11 | 0.90* | 25.30 |
| Bolus insulin | Not relevant | Not relevant | Not relevant |
| **T2DM<sub>B/B</sub>, total dose** | | | |
| Basal insulin | 28.11 | 1.08* | 30.36 |
| Bolus insulin | 37.13 | Not significant | 37.13 |

Abbreviations: B/B, basal-bolus; BOT, basal oral therapy; T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

* p<0.05; NS, non-significant; in the case of non-significant results, a relative rate of one was used in the calculation

Table 3. Calculation of hypoglycaemia event rates
|                | Non-severe hypoglycaemia | Severe hypoglycaemia |
|----------------|--------------------------|----------------------|
| **T1DM**       |                          |                      |
|                | **Daytime**              | **Nocturnal**        |
| Total events/patient/year for glargine U100<sup>a</sup> | 30.42                    | 8.52                    | 3.20                    |
| Degludec/glargine U100 hypoglycaemic event rate ratio<sup>b</sup> | Not Significant | 0.83* | Not significant |
| Calculated degludec hypoglycaemic event rate | 30.42 | 7.07 | 3.20 |
| **T2DM<sub>BOT</sub>** |                          |                      |
|                | **Daytime**              | **Nocturnal**        |
| Total events/patient/year for glargine U100<sup>a</sup> | 23.12 | 13.38 | 0.10 |
| Degludec/glargine U100 hypoglycaemic event rate ratio<sup>b</sup> | Not significant | 0.64* | 0.14* |
| Calculated degludec hypoglycaemic event rate | 23.11 | 8.57 | 0.01 |
| **T2DM<sub>B/B</sub>** |                          |                      |
|                | **Daytime**              | **Nocturnal**        |
| Total events/patient/year for glargine U100<sup>a</sup> | 30.42 | 8.52 | 0.70 |
| Degludec/glargine U100 hypoglycaemic event rate ratio<sup>b</sup> | 0.83* | 0.75* | Not significant |
| Calculated degludec hypoglycaemic event rate | 25.25 | 6.39 | 0.70 |

Abbreviations: B/B, basal-bolus; BOT, basal oral therapy; CI, confidence interval; insulin glargine;
T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

* p<0.05; NS, non-significant; in the case of non-significant results, a relative rate of one was used in the calculation

\(^{a}\) Taken from Brod et al. [37] and UK Hypoglycaemia Study Group [36]

\(^{b}\) Taken from Vora et al. [27]

Table 4. Sensitivity analyses conducted

| Parameter                          | Base case                      | Sensitivity analysis                                                                 |
|------------------------------------|--------------------------------|--------------------------------------------------------------------------------------|
| Time horizon                       | 1 year                         | The time horizon was increased to 5 years.                                            |
| Hypoglycaemia rates                | Published literature:          | Additional published event rates [27, 28, 41, 59] were investigated, including those reported in the clinical trial programme. |
|                                    | Brod et al. [37]                |                                                                                      |
|                                    | UK Hypoglycaemia Study Group   |                                                                                      |
|                                    | [36]                           |                                                                                      |
|                                    | Vora et al. [27]                |                                                                                      |
| Mortality incidence after severe hypoglycaemia | 1.12% mortality risk | The mortality risk was decreased to zero.                                              |
| Cost of hypoglycaemia              | Derived from clinical trial programme [38] | Healthcare costs were increased and decreased by 10%.                                  |
| Insulin dose                       | Published literature:          | A sensitivity analysis was conducted assuming no difference in insulin dose between the treatments. |
|                                    | Vora et al. [27]                |                                                                                      |
|                                    | Doneva et al. [35]              |                                                                                      |
| Flexible dosing utility            | 0.006 (Boye et al. [40])        | Several sensitivity analyses were conducted to assess the impact of flexible dosing with degludec: one using an alternative utility value for flexible dosing (0.0130 [57]); one where only 50% of patients gained a utility benefit of flexible dosing; and one where there was no utility gain from dosing flexibility. |

Table 5. Total cost per patient per year and incremental cost-effectiveness
|                   | T1DM                      | T2DM\_BOT                  | T2DM\_B/B                     |
|-------------------|---------------------------|----------------------------|-------------------------------|
|                   | Degludec (BGN/year) Glargine U100 (BGN/year) Incremental cost (BGN/year) | Degludec (BGN/year) Glargine U100 (BGN/year) Incremental cost (BGN/year) | Degludec (BGN/year) Glargine U100 (BGN/year) Incremental cost (BGN/year) |
| Insulin           | 1,296.67  1,180.11  116.56 | 794.93  574.23  220.70 | 1,574.88  1,196.94  377.95 |
| Hypoglycaemia events | 1,846.61  1,893.80  -47.19 | 306.40  509.74  -203.34 | 581.79  655.53  -73.74 |
| Non-severe daytime events | 19.42  19.42  0.00 | 15.02  15.02  0.00 | 16.35  19.69  -3.35 |
| Non-severe nocturnal events | 230.41  277.60  -47.19 | 284.26  443.94  -159.68 | 211.17  281.56  -70.39 |
| Severe events     | 1,596.78  1,596.78  0.00 | 7.11  50.78  -43.67 | 354.28  354.28  0.00 |
| Total costs       | 3,143.28  3,073.92  69.37 | 1,101.32  1,083.97  17.35 | 2,156.67  1,852.47  304.21 |
| QALYs             | 0.5722  0.5568  0.0154 | 0.7490  0.7055  0.0435 | 0.6893  0.6480  0.0413 |
| ICER (cost/QALY)  | 4,498.68 | 399.11 | 7,365.22 |

Abbreviations: B/B, basal-bolus; BGN, Bulgarian Lev; BOT, basal oral therapy; ICER, incremental cost-effectiveness ratio; QALY, quality-adjusted life years; T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

Figures
Figure 1

Overview of cost-effectiveness model. Abbreviations: Δ, change in; HC, healthcare, HRQoL, health-related quality of life; ICER, incremental cost-effectiveness ratio; QALY, quality-adjusted life year; SMBG, self-monitored blood glucose
Figure 2

Probabilistic sensitivity analyses acceptability curves

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

Tresiba_CE_Bulgaria_Supplementary_information.docx