DIVE/DPV registries: benefits and risks of analog insulin use in individuals 75 years and older with type 2 diabetes mellitus

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ABSTRACT

Introduction The aims of this study were to characterize insulin-treated individuals aged ≥75 years with type 2 diabetes using basal insulin analogs (BIA) or regular insulins (human insulin (Hi)/neutral protamine Hagedorn (NPH)) and to compare the benefits and risks.

Research design and methods The analysis was based on data from the DPV (Diabetes-Patienten-Verlaufs-Dokumentation) and DIVE (Diabetes Versorgungs-Evaluation) registries. To balance for confounders, propensity score matching for age, sex, diabetes duration, body mass index and hemoglobin A1c (HbA1c) as covariates was performed.

Results Among 167,300 patients aged ≥75 years with type 2 diabetes (mean age, 80.3 years), 9601 subjects used insulin regimens with basal insulin (Hi/NPH or BIA). Of these 8022 propensity score-matched subjects were identified. The mean diabetes duration was ~12 years and half of the patients were male. At the time of switch, patients provided with BIA experienced more dyslipidemia (89.3% vs 85.9%; p=0.002) and took a greater number of medications (4.3 vs 3.7; p<0.001) and depression was more prevalent (8.4% vs 6.5%; p=0.01). Aggregated to the most actual treatment year, BIA was associated with a higher percentage of patients using basal-supported oral therapy (42.6% vs 14.4%) and intensified conventional insulin therapy (44.3% vs 29.4%) and lower total daily insulin doses (0.24 IU/kg/day vs 0.30 IU/kg/day; p<0.001). The study did not reveal significant differences in efficacy (HbA1c 7.4% vs 7.3%; p=0.06), hospitalizations (0.7 vs 0.8 per patient-year (PY); p=0.15), length of stay (16.3 vs 16.1 days per PY; p=0.53), or rates of severe hypoglycemia (4.07 vs 4.40 per 100 PY; p=0.88), hypoglycemia with coma (3.64 vs 3.26 per 100 PY; p=0.88) and diabetic ketoacidosis (0.01 vs 0.03 per 100 PY; p=0.36).

Conclusion BIA were used in more individually and patient-centered therapy regimens compared with Hi/NPH in patients with a mean age of 80 years. Both groups were slightly overtreated with mean HbA1c <7.5%. The risk of severe hypoglycemia was low and independent of insulin type. Further analyses of elderly patients with type 2 diabetes are needed to provide evidence for best practice approaches in this age group.

INTRODUCTION

We previously described that type 2 diabetes (T2DM) treatment in elderly patients was characterized by rather low hemoglobin A1c (HbA1c) values and an increased risk of hypoglycemia.1 2 Several studies show that elderly patients aged >75 years with T2DM are at increased risk of severe or fatal hypoglycemia and have higher hospitalization rates for hypoglycemic events.3 4

In contrast to neutral protamine Hagedorn (NPH) insulin, basal insulin analogs (BIA) showed a reduced risk of hypoglycemia in elderly and obese patients and in patients with type 2 diabetes.
Clinical care/Education/Nutrition

with renal impairment and cardiovascular disease (CVD). They have also been associated with reduced rates of nocturnal hypoglycemia, which is often unrecognized in the elderly. Further, long-acting insulin formulations can be used with a once daily dosing, providing sufficient glycemic control.

Contrary to what we would have expected on the background of elderly patients and treatment characteristics, the elderly were treated more frequently with regular human insulin (HI) (from 35.0% of those aged 70–79 years old to 37.9% of those aged ≥90 years old) than with BIA (from 29.7% of those aged 70–79 years old to 23.5% of those aged ≥90 years old) in our previous study. This is unlike a global trend in diabetes treatment where insulin analogs are increasingly being used in a general diabetes patient population.

To explore the determinants and potential benefits of use of basal insulin in these elderly patients with T2DM, we had a closer look at patients who were at least 75 years of age and were receiving either basal HI/NPH insulin and compared the treatment outcomes with patients receiving BIA. We used propensity score matching in order to balance the characteristics of the patients at the time of switch to insulin (baseline). Our study aimed to address the following specific questions: (1) Do baseline patient characteristics, risk factors and comorbidities differ by type of insulin treatment? (2) Do concomitant antidiabetic drugs differ in relation to the basal insulin used? (3) Do efficacy parameters (eg, HbA1c, fasting blood glucose (FBG) and insulin dose) differ by type of insulin treatment? (4) Do safety parameters (eg, number of hospitalizations, mean length of stay, risk of hypoglycemia and hypoglycemia with coma, diabetic ketoacidosis (DKA)) differ by type of insulin treatment?

RESEARCH DESIGN AND METHODS

Data collection

Prospective longitudinal, standardized routine data were obtained from the prospective, multicenter German patient databases on diabetes mellitus: DPV (Diabetes-Patienten-Verlaufsdocumentation) and DIVE (Diabetes Versorgungs-Evaluation).

DPV data on patients with diabetes mellitus are collected during routine examinations using DPV software and the anonymized data are sent to Ulm University for inclusion into the database every 6 months. Detailed information on the documentation systems has been published previously. This analysis includes data from 159 DPV centers.

The DIVE registry was established in Germany in 2011. Consecutive patient data are collected from German centers and these patients continue to be followed up. Data are entered into an online database using DPV software. All patients provided written informed consent. This study comprises data of 101 DIVE study sites.

Patient population for the analysis

Patients were sampled in September 2020 and were included in the current analysis if they had at least one clinic visit per year and their documented therapy was available. We selected patients of at least 75 years with T2DM with basal insulin (human or analog) or NPH insulin treatment. Patients aged <75 years, with other forms of diabetes, using an insulin pump or with prandial insulin only were excluded. The final study comprised 8022 propensity score-matched (PSM) subjects for baseline and 3836 PSM subjects for follow-up analysis. The mean follow-up period was 2.3±2.1 years for the HI/NPH group and 2.6±2.1 years for BIA.

Definitions

Hypertension was defined as ≥145 mm Hg for systolic blood pressure and/or a diastolic blood pressure of ≥85 mm Hg and/or antihypertensive treatment. Body mass index (BMI) was defined as bodyweight divided by the square body height in kg/m². Dyslipidemia was defined as a low-density lipoprotein cholesterol of ≥100 mg/dL (≥2.6 mmol/L) without further risk factors and ≥70 mg/dL (≥1.8 mmol/L) in patients with CVD or chronic kidney disease or those receiving lipid-lowering drug treatment. Comorbidities were grouped into (patient-reported or physician-reported) microvascular and macrovascular diseases. The former included any form of retinopathy, blindness, nephropathy, renal failure, dialysis or neuropathy. The latter included transient ischemic attack/prolonged reversible ischemic neurologic deficit, stroke, coronary artery disease, myocardial infarction and peripheral arterial disease. Polypharmacy was determined as a numerical-only definition of >10 medications that were used concomitantly. The DIVE/DPV databases were searched with focus on drug categories according to the PRISCUS listing. Medications comprised prescription medications and routine use of over-the-counter medications.

HbA1c values were standardized to the Diabetes Control and Complications Trial. For severe hypoglycemia, the definition of the American Diabetes Association Workgroup on Hypoglycemia (‘an event requiring assistance by another person to actively administer carbohydrates, glucagon or other resuscitative actions’) was applied. Hypoglycemia with coma was defined as loss of consciousness or occurrence of seizures. DKA was defined as pH <7.3 and/or bicarbonate <15 mmol/mol or hospitalization for DKA. Insulin regimen was categorized as follows: (1) basal-supported oral therapy (BOT) (basal insulin only); (2) supplementary insulin therapy (SIT) (prandial insulin only); (3) conventional insulin therapy (CT) (prandial and basal insulin combined, up to three injection time points); (4) intensified conventional insulin therapy (ICT) (prandial and basal insulin combined, ≥3 injection time points); and (5) continuous subcutaneous insulin infusion (CSII). To stratify patients using basal insulin, patients on SIT and CSII were...
excluded. Patients on BOT, CT or ICT were identified as these therapy regimens use a long-acting (basal) insulin or premixed/co-formulation insulin. Patients on basal insulin were divided by use of HI/NPH and BIA. Insulin treatment initiation had to be documented between 2000 and June 2020, with HbA1c and BMI data available ±3 months from the time of insulin initiation. Baseline data for patients were aggregated within ±3 months of insulin initiation. Follow-up data were aggregated to the most recent documented treatment year without therapy switch (excluding the baseline period). Treatment was considered continuous if there was a maximum gap of 120 days between two clinic visits, with the same treatment being documented before and after the gap.

Statistics

Data from the two registries were combined and analyzed as a single data set. Data were then aggregated as medians in the year of intensification with insulin (baseline). Categorical variables are presented as percentages and continuous variables are presented as mean±SD. Rates and event rates are presented per 100 patient-years (PY). Differences of non-PSM variables were analyzed using Wilcoxon test for continuous variables or χ² test for binary variables. P values were adjusted for multiple testing using the Bonferroni step-down method.

Propensity score matching was used to ensure that both the HI/NPH therapy regimen group and the BIA group had similar baseline characteristics. Differences between the treatment groups and propensity score for insulin analogs were estimated according to a former analyses of the DIVE/DPV registries. The multivariable logistic regression model includes age, sex, duration of diabetes, BMI and HbA1c level as covariates. For each patient, the probability (propensity score) for HI/NPH and BIA was estimated from the logistic model based on the patient’s specific covariate values. Matching was conducted with a one-to-one matching process (greedy-matching algorithm) and caliper width 0.2. To evaluate balancing of matched variables, standardized differences were assessed. A standardized difference of <10% indicates a negligible difference in the mean. In order to compare hospitalization times and lengths of stay between the matched groups, a standard Poisson regression model with the logarithm of the individual time under risk as offset was used. Event rates of severe hypoglycemia, hypoglycemic coma, DKA and severe DKA were calculated and compared using negative binomial regression with the logarithm of the individual time under risk as offset. For HbA1c a linear regression model was used. Further, incidence rate ratios and 95% CIs were calculated for DKA, hypoglycemia and hypoglycemia with coma.

A p value <0.05 was considered statistically significant. Statistical analysis was performed using SAS V.9.4.

RESULTS

Patient population

The DIVE/DPV registries included 167 300 patients with T2DM aged 275 years at the time of data export. Of these, 29 800 (17.8%) patients initiated insulin therapy between 2000 and June 2020, and 11 108 (6.6%) used therapy regimens with basal insulin. The final analysis comprised 9601 patients in the entire cohort (n=5583 on regular insulin and n=4018 on insulin analogs) and 8022 PSM patients (n=4011 in both treatment groups) (figure 1).

Patient baseline characteristics

For analysis of baseline characteristics 4011 matched pairs were available. The standardized differences for the matched variables were highest for HbA1c (4.5% for the entire cohort and 2.0% for the matched cohort) and age (~6.1% and 0.6%) but remained largely under 10%, demonstrating a negligible difference between both treatment groups (table 1). The mean patient age was 80.3 years in both matched treatment groups, half of the patients were male, the mean diabetes duration was approximately 12 years, and the mean baseline HbA1c was 8.6% for both insulins. There were only minor significant differences in the entire cohort as well as in the matched cohort for non-matched variables at time of insulin therapy initiation.

FBG did not differ between the groups (10.6 mmol/L vs 10.7 mmol/L, p=0.50 for the entire cohort; 10.6 mmol/L vs 10.8 mmol/L for the PMS cohort, p=0.42). The bodyweight of BIA patients in the entire cohort was slightly higher (79.4 kg vs 80.6 kg; p=0.003), but this difference was not seen in the PMS cohort (80.2% vs 80.6%; p=1.00). In the PMS cohort, the proportion of patients with dyslipidemia was higher in the BIA group (85.9% vs 89.3%; p=0.001). The proportions of other risk factors such as hypertension (71.0% vs 73.5%) and smoking (3.5% vs 3.4%) were largely similar. The only exception was the mean number of medications, which was significantly higher in the BIA group compared with the HI/NPH group in (both cohorts 3.7 vs 4.3; p<0.001), whereas the proportion of patients with polypharmacy was borderline significantly lower in the HI/NPH group of the entire cohort (0.5% vs 1.1%; p=0.05), but not in the matched cohort (0.6% vs 1.1%; p=0.26). Furthermore, in the PSM cohort depression was more prevalent with BIA treatment (5.5% vs 8.4%; p=0.01).

There were small differences in the prevalence of comorbidities such as dementia (HI/NPH 7.9% vs BIA 8.8%), cancer (HI/NPH 5.0% vs BIA 5.8%) and microvascular and macrovascular diseases (HI/NPH 85.3% vs BIA 83.6% and HI/NPH 44.6% vs BIA 44.4%), but these were not statistically significant.

The proportion of care-dependent patients (HI/NPH 5.2% vs BIA 7.1%) was low and there were no differences observed between both treatment groups. Marginally more than one-third of the patients participated in a diabetes management program (HI/NPH 37.1% vs BIA 37.9%).
Concomitant antidiabetic drugs at follow-up
Detailed information on concomitant antidiabetic treatment was available for 49.5% of the HI/NPH subjects and 46.1% of the BIA group for the most recent treatment year (follow-up) (table 2). Most of the patients on regular insulin were treated with CT (56.2% vs 13.2%; p<0.001), whereas BOT and ICT were more frequent with BIA (14.4% vs 42.6% and 29.4% vs 44.3%; both p<0.001).

Rapid-acting insulin was added to basal insulin concomitantly in 79.4% of HI/NPH and 53.5% of BIA patients. Rapid-acting HI was preferred in the HI/NPH group versus the BIA group (61.3% vs 19.3%). The reverse was true for rapid-acting analogs (18.1% vs 34.1%) (both p<0.001).

Oral antidiabetic drugs and glucagon-like peptide-1 (GLP-1) agonists were more frequently used in patients using insulin analogs. Usage proportions were highest for metformin (23.3% vs 28.7%), dipeptidyl peptidase-4 (DPP-4) inhibitors (10.0% vs 25.8%) and sulfonylureas (10.8% vs 11.2%). With the exception of the latter and glucosidase inhibitors, differences in non-insulin antidiabetic therapy were highly significant between both groups. However, percentages of patients taking glucosidase inhibitors (0.8% vs 0.9%), GLP agonists (0.8% vs 2.2%) and sodium/glucose cotransporter 2 (SGLT-2) inhibitors (0.6% vs 3.6%) were low.

Efficacy and safety parameters
A comparison of follow-up data in 1984 matched patients on HI/NPH to 1852 patients on insulin analog was used to determine differences in glycemic control, rates of severe hypoglycemia and DKA (table 3).

Glycemic control and hospitalizations
Patients in the BIA group needed less total daily insulin (0.30 IU/kg/day vs 0.22 IU/kg/day; p<0.001). The prandial to total insulin ratio was higher in the BIA group compared with the regular insulin group (0.83 vs 0.70; p<0.001), demonstrating a higher use of prandial insulin in the BIA group. The resulting median HbA1c was similar with HI/NPH treatment and BIA treatment (7.3% vs 7.4%; p=0.06). FBG did not differ between the groups (both 8.4 mmol/L; p=0.85).

The mean number of hospitalizations per PY was slightly lower in the BIA compared with the HI/NPH group, but did not reach statistical significance (0.7 vs 0.8; p=0.15). The mean length of stay per PY did not differ between the groups (HI/NPH 16.1 days vs BIA 16.3 days; p=0.53).

Hypoglycemia and DKA
Of the HI/NPH patients 48.6% had HbA1c values below the median HbA1c and no severe hypoglycemia or hypoglycemia with coma was observed. The same was true for 51.5% of the BIA group (p=0.09 HI/NPH vs BIA). Overall, 2.8% of subjects on regular insulin and 2.4% on insulin analogs suffered at least one severe hypoglycemia event. Event rates for severe hypoglycemia, however, tended to be lower in the BIA group but did not reach statistical significance (HI/NPH 4.4 and BIA 4.1 per 100 PY; p=0.88). Severe hypoglycemia with coma was
Table 1  Characteristics of patients aged >75 years with HI/NPH vs BIA at baseline

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>Standardized difference, %</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>Standardized difference, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>80.6±4.2</td>
<td>80.3±4.2</td>
<td>-6.1</td>
<td>80.2±4.1</td>
<td>80.3±4.2</td>
<td>0.6</td>
</tr>
<tr>
<td>Male, %</td>
<td>40.0</td>
<td>50.0</td>
<td>-5.7</td>
<td>47.0</td>
<td>50.0</td>
<td>0.0</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td>28.8±5.3</td>
<td>29.0±5.4</td>
<td>2.2</td>
<td>29.0±5.3</td>
<td>28.9±5.4</td>
<td>-1.8</td>
</tr>
<tr>
<td>Diabetes duration, years</td>
<td>12.0±9.9</td>
<td>11.8±10.1</td>
<td>-1.3</td>
<td>11.9±9.9</td>
<td>11.8±10.1</td>
<td>-1.0</td>
</tr>
<tr>
<td>HbA1c, %</td>
<td>8.5±2.1</td>
<td>8.6±2.1</td>
<td>4.5</td>
<td>8.6±2.2</td>
<td>8.6±2.1</td>
<td>2.0</td>
</tr>
<tr>
<td>HbA1c, mmol/mol</td>
<td>69.9±23.4</td>
<td>71.0±23.4</td>
<td></td>
<td>70.4±23.5</td>
<td>70.9±23.2</td>
<td></td>
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Other variables at baseline

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>P value</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bodyweight, kg</td>
<td>79.4±15.9</td>
<td>80.6±16.3</td>
<td>0.003</td>
<td>80.2±15.9</td>
<td>80.6±16.2</td>
<td>1.000</td>
</tr>
<tr>
<td>FBG, mmol</td>
<td>10.6±5.9</td>
<td>10.7±5.6</td>
<td>0.496</td>
<td>10.6±5.9</td>
<td>10.8±5.6</td>
<td>0.423</td>
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</table>

Risk factors

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>P value</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypertension, %</td>
<td>71.2</td>
<td>73.5</td>
<td>0.100</td>
<td>71.0</td>
<td>73.5</td>
<td>0.125</td>
</tr>
<tr>
<td>BMI &gt;30 kg/m², %</td>
<td>36.1</td>
<td>37.9</td>
<td>0.496</td>
<td>37.6</td>
<td>37.8</td>
<td>1.000</td>
</tr>
<tr>
<td>Dyslipidemia, %</td>
<td>86.0</td>
<td>89.3</td>
<td>&lt;0.001</td>
<td>85.9</td>
<td>89.3</td>
<td>0.001</td>
</tr>
<tr>
<td>Smokers, %</td>
<td>3.6</td>
<td>3.4</td>
<td>1.000</td>
<td>3.5</td>
<td>3.4</td>
<td>1.000</td>
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</table>

Functional status

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>P value</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Care dependency, %</td>
<td>5.5</td>
<td>7.2</td>
<td>0.090</td>
<td>5.2</td>
<td>7.1</td>
<td>0.055</td>
</tr>
<tr>
<td>Polypharmacy (&gt;10 medications), %</td>
<td>0.5</td>
<td>1.1</td>
<td>0.049</td>
<td>0.6</td>
<td>1.1</td>
<td>0.261</td>
</tr>
<tr>
<td>Diabetes management program, %</td>
<td>36.4</td>
<td>37.8</td>
<td>0.809</td>
<td>37.1</td>
<td>37.9</td>
<td>1.000</td>
</tr>
<tr>
<td>Mental status, %</td>
<td>Dementia</td>
<td>8.0</td>
<td>8.8</td>
<td>0.827</td>
<td>7.9</td>
<td>8.8</td>
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<tr>
<td></td>
<td>Depression</td>
<td>6.6</td>
<td>8.4</td>
<td>0.007</td>
<td>6.5</td>
<td>8.4</td>
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</table>

Comorbidities, %

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>P value</th>
<th>HI/NPH</th>
<th>BIA</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cancer</td>
<td>5.0</td>
<td>5.8</td>
<td>0.809</td>
<td>5.0</td>
<td>5.8</td>
<td>1.000</td>
</tr>
<tr>
<td>Microvascular disease*</td>
<td>85.2</td>
<td>83.6</td>
<td>0.225</td>
<td>85.3</td>
<td>83.6</td>
<td>0.352</td>
</tr>
<tr>
<td>Macrovascular disease†</td>
<td>44.5</td>
<td>44.3</td>
<td>1.000</td>
<td>44.6</td>
<td>44.4</td>
<td>1.000</td>
</tr>
</tbody>
</table>

Data are presented as mean±SD or %.
Bold numbers indicate p-values <0.05.
*Includes any form of retinopathy, blindness, nephropathy, renal failure, dialysis or neuropathy.
†Includes transient ischemic attack/prolonged reversible ischemic neurologic deficit, stroke, coronary heart disease, myocardial infarction and peripheral arterial disease.
BIA, basal insulin analog; BMI, body mass index; FBG, fasting blood glucose; HbA1c, hemoglobin A1c; HI, human insulin; NPH, neutral protamine Hagedorn; PS, propensity score.

documented in DPV only and observed in 2.8% of the HI/NPH group and 3.0% of the BIA group with event rates of 3.3 and 3.6 per 100 PY (p=0.88). The percentage of patients with DKA was low and the event rates did not differ (HI/NPH 0.3% vs 0.1% of patients and 0.28 vs 0.05 per 100 PY; p=0.36).
Compared with patients prescribed HI/NPH. Further-of medications and were at higher risk of depression BIA had a worse metabolic profile, took a greater number with T2DM with a mean age of 80 years. insulin and insulin analogs in a population of patients our knowledge, this is the first study comparing regular (severe hypoglycemia and DKA) of basal insulin therapy analysis focused on the benefits (eg, HbA1c) and risks safety parameters with different insulin therapies. This

6 BMJ Open Diab Res Care 2021;9:e002215. doi:10.1136/bmjdrc-2021-002215

more, prescription of BIA was associated with more

Patient population

Patients with BIA tended to have more risk factors at baseline. This became significant in the slightly higher number of medications taken (4.3 vs 3.7) and higher proportion of patients with dyslipidemia in the BIA group. Although lipid targets are more likely to be met in the elderly, it is assumed that diabetic dyslipidemia in the elderly is undertreated.23 24 T2DM and obesity are also linked with depression,25 which also was more prevalent in the BIA group. Interestingly, the proportion of patients with polypharmacy (defined in this study as >10 medications) was only 0.6% and 1.1% in the HI/NPH and BIA groups, respectively. Although some medications are documented only in the DPV registry and not in DIVE, data indicate that this risk factor plays a minor role in the patient population.

Among the insulin-treated patients in this study the proportion of patients with hypertension was high (71.0% for HI/NPH and 73.5% for BIA). The dysregulation of neurohumoral and neuroimmune pathways contributes to the pathophysiology of both T2DM and hypertension; thus, there is a bidirectional association between macrovascular and microvascular systems.26 In line with this, we observed in this insulin-treated population a higher proportion of microvascular disease (85.3% vs 83.6%) compared with our previous DIVE/DPV analysis (also including insulin-naïve subjects), which revealed proportions from 59.8% of patients aged 70–79 years old to 50.4% of those aged >90 years.2

In Germany care-dependent elderly live either at home (assisted by family, caregivers or home care providers) or in ‘shared housing arrangements’ and nursing homes.27 The prevalence of care dependency rises from 8% among those aged 75–79 years old to 76% among those aged >90 years old.28 Surprisingly, in our study only 5.2% and 5.5% of patients with T2DM at comparable age were care-dependent. A previous study reported that care-dependent patients with T2DM are more likely to be treated with insulin compared with independent patients.29 However, a large proportion of nursing home residents are treated by general practitioners and not diabetologists and are therefore probably not documented in the registries.29 Furthermore, the support of self-management and therapy adherence and the intensified lifestyle training (eg, 37% of patients included in disease management programs) may lead to a higher proportion of independent elderly patients compared with the above-mentioned German statistics.29

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Table 2  Antidiabetic treatment in patients aged >75 years with regular vs analog insulin (follow-up)

<table>
<thead>
<tr>
<th></th>
<th>HI/NPH n=1984</th>
<th>BIA n=1852</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Insulin treatment strategy, %</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BOT</td>
<td>14.4</td>
<td>42.6</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>CT</td>
<td>56.2</td>
<td>13.2</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>ICT</td>
<td>29.4</td>
<td>44.3</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Rapid-acting insulin, %</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Human</td>
<td>61.3</td>
<td>53.5</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Analogs</td>
<td>18.1</td>
<td>34.1</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Non-insulin antidiabetic treatment, %</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Metformin</td>
<td>23.3</td>
<td>28.7</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Sulfonylurea</td>
<td>10.8</td>
<td>11.2</td>
<td>1.000</td>
</tr>
<tr>
<td>Glucosidase inhibitors</td>
<td>0.8</td>
<td>0.9</td>
<td>1.000</td>
</tr>
<tr>
<td>Glinides</td>
<td>3.7</td>
<td>6.1</td>
<td>0.002</td>
</tr>
<tr>
<td>DPP-4 inhibitor</td>
<td>10.0</td>
<td>25.8</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>GLP-1 agonist</td>
<td>0.8</td>
<td>2.2</td>
<td>0.001</td>
</tr>
<tr>
<td>SGLT-2 inhibitor</td>
<td>0.6</td>
<td>3.6</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

CT: prandial and basal insulin combined (up to 3 injection time points); ICT: prandial and basal insulin combined (>3 injection time points).

Bold numbers indicate p-values <0.05.

BIA, basal insulin analog; BOT, basal insulin-supported oral therapy; CT, conventional insulin therapy; DPP-4, dipeptidyl peptidase-4; GLP-1, glucagon-like peptide-1; HI, human insulin; ICT, intensified conventional insulin therapy; NPH, neutral protamine Hagedorn; SGLT-2, sodium/glucose cotransporter 2.

Incidence rate ratios and 95% CIs show no differences in risk of severe hypoglycemia (0.95, 95% CI 0.34 to 2.61), hyperglycemic events with coma (1.16, 95% CI 0.25 to 5.27) and DKA (0.19, 95% CI 0.01 to 6.80) between regular insulins compared with insulin analogs (figure 2).

DISCUSSION

The aims of this analysis were to describe the characteristics of a large cohort of almost 10 000 patients aged ≥75 years with T2DM and to compare efficacy and safety parameters with different insulin therapies. This analysis focused on the benefits (eg, HbA1c) and risks (severe hypoglycemia and DKA) of basal insulin therapy with NPH and/or HI compared with insulin analogs. To our knowledge, this is the first study comparing regular insulin and insulin analogs in a population of patients with T2DM with a mean age of 80 years.

At the time of switching to insulin, patients prescribed BIA had a worse metabolic profile, took a greater number of medications and were at higher risk of depression compared with patients prescribed HI/NPH. Furthermore, prescription of BIA was associated with more individually and patient-centered therapy regimens like BOT and ICT and lower total daily insulin doses. The study described an improvement in HbA1c control with insulin therapy for both insulin treatments, but did not reveal any significant differences in efficacy or safety with regard to type of insulin.
Rather than focusing on glycemic control in elderly patients with T2DM, it is important to maintain quality of life and patients’ ability to self-manage their diabetes. Current guidelines recommend simplifying treatment (eg, by using basal insulin and avoiding the additional use of regular and rapid-acting insulin) and to use drug classes with low hypoglycemic effect.13 30–32 BOT was used three times more frequently in the BIA group compared with HI/NPH (42.6% vs 14.4%) and the BIA...
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Group needed lower total daily insulin levels. Further, in our study additional rapid-acting insulins were needed less often in the BIA group compared with the HI/NPH group. Thus, data indicate that current guidelines advising a simple patient-approached therapy regimen to maintain self-management abilities in the elderly were better reflected by using a BOT regimen with insulin analogs.13

One study indicates a longer persistence on BOT therapy in elderly patients with T2DM with BIA before intensifying to ICT compared with patients with NPH.33 Further, with BOT additional antidiabetic drugs are necessary,44 explaining the higher percentages of concomitant medication in the BIA group compared with HI/NPH.

In contrast to that, more than half of the HI/NPH group used a CT insulin regimen (56.2%). CT necessitates fixed meal times and is useful in less active or care-dependent patients.34 It is likely that patients who have used this treatment strategy for long periods of time were not switched as they aged. In contrast to CT, the more complex ICT is used in active patients with T2DM with good mental and functional status but poor glycemic control.35 The higher proportion of ICT in the BIA group indicates that a higher proportion of patients had difficulty stabilizing their diabetes and explains the higher prandial to total insulin ratio. Another reason for the higher ICT proportion with BIA might be the higher risk of hypoglycemia with ICT and physicians might have prescribed BIA due to its promoted antihypoglycemic properties. Furthermore, the BIA group needed lower total daily insulin levels, but a higher percentage of patients were treated with newer non-insulin antidiabetic agents. For example, the proportion of DPP-4 inhibitors was twice as high compared with the HI/NPH group (10.0% vs 25.8%). In studies, DPP-4 inhibition was twice as high compared with non-

Limitations

The strength of this analysis is the large number of elderly patients aged ≥75 years with a matched group comprising over 8000 patients in a real-world setting and a mean age of 80.3 years. However, in an observational study residual selection bias despite the effect of propensity score matching is possible, for example by the sole involvement of specialized diabetes centers in the DIVE/DPV registries. Newer concepts such as frailty and sarcopenia are rarely or not reported in the registries due to lack of consensus on diagnostic criteria; for example, International Classification of Diseases-10 for sarcopenia was only established in 2016.42 45 Furthermore, the analysis included follow-up data from 2000. This may lead to an under-representation of newer therapy options, such as non-insulin antidiabetic drugs (SGLT-2 inhibitors, GLP-1 agonists) and the use of insulin analogs. Finally, we did not distinguish between premixed and split usage of insulins, which might also have an implication on outcomes.

CONCLUSIONS

Compared with patients on regular insulins patients treated with insulin analogs used more flexible therapy regimens with overall lower doses of total daily insulin. The high percentage of BOT and the lower use of rapid-acting insulin in the BIA group reflect the current
guidelines that recommend a simplified therapy in the elderly to maintain self-management abilities.

With regard to glycemic control, there might be a slight overtreatment in this elderly population with high proportions of macrovascular and microvascular diseases. Therapy goals, therefore, should be adapted continuously and individually to patients’ needs. The risk of severe hypoglycemic events is low and comparable between the two insulin types in patients with T2DM aged ≥75 years. However, further analyses of elderly patients are urgently needed to provide evidence for best practice medical care in this age group.

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Contributors
RW, JG, SS, BR, PMJ and DE contributed to data collection. GvM, PB reports to handling and interpretation of data, writing of the manuscript or decision to submit the article for publication.

Competing interests
PB reports to having received consultancy honoraria from Sanofi and Abbott.

Patient consent for publication
Not required.

Ethics approval
The DPV initiative, which was established in 1995, was approved by the ethics committee of the University of Ulm (no. 202-09 on August 14, 2009), and data collection was approved by local review boards. The DIVE registry was established in Germany in 2011. The protocol was approved by the ethics committee of the Medical School of Hannover (no. 6003 on August 25, 2011).

Provenance and peer review
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Data availability statement
Data are available upon reasonable request. Data may be obtained from a third party and are not publicly available. The data sets generated and analyzed during the current study are not publicly available due to data privacy but are available from the corresponding author on reasonable request.

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