Examining correlates of treatment satisfaction for injectable insulin in type 2 diabetes: lessons learned from a clinical trial comparing biphasic and basal analogues

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Abstract

Background: Successfully managing diabetes is a complex process that includes addressing issues of drug efficacy, safety and treatment satisfaction. Additionally, the combined impact of patient/disease characteristics and treatment outcomes on treatment satisfaction is not well understood. The purpose of this study was to examine the impact of age, weight, gender, co-morbid conditions, diabetes history, treatment burden, efficacy (HbA1c) and side effects (weight gain, hypoglycemic events) on patients' appraisal of treatment satisfaction using linear regression models.

Methods: Data from a multi-center, randomized clinical trial comparing the efficacy/safety of biphasic insulin aspart 70/30 (BIAsp 70/30) vs. glargine (Glar) among insulin naïve type 2 patients were analyzed. Subjects were between ages 18–75, with baseline HbA1c > 8% and BMI ≤ 40 kg/m^2 (N = 233). Treatment satisfaction was assessed by the Insulin Treatment Satisfaction Questionnaire (ITSQ).

Results: When factors were examined independently, multiple significant relationships (age, co-morbidity, hypoglycemic events, and weight gain) with overall and/or domains of treatment satisfaction were found. However, when all significant relationships were examined together, only neuropathy, treatment efficacy, and number of hypoglycemic events maintained their previous significance.

Conclusion: By examining predictors independently, significant relationships were identified. However, not all findings remained significant when examined in combination with each other. Thus, to more accurately characterize the impact of factors on treatment satisfaction, a more comprehensive approach may be necessary. By improving patient treatment satisfaction, the efficacy of treatments, as well as critical treatment outcomes such as compliance and cost of care should be improved.
Characteristics such as age, gender, time with diabetes, and prevalence and incidence of co-morbid conditions may act as mediators that influence these appraisals.

Static factors such as patient and disease characteristics are generally constant or fixed at any given point in time. Characteristics such as age, gender, level of education, ethnicity and duration of diabetes have been shown to account for some of the differences in patients' treatment satisfaction with their diabetes care when provided by endocrinologists compared to generalists [15]. However, data on the relationship of these factors to treatment satisfaction is often times conflicting, and many significant relationships do not endure when examined in combination with other important factors. For example, studies have found that greater satisfaction is associated with decreasing age, while others have indicated that younger patients are less satisfied with their treatment [16-18]. The occurrence of diabetes-related complications has been found to be associated with lower treatment satisfaction with diabetes medication, however, this association was no longer found when age, therapy and HbA1c level were also taken into consideration [17]. Insulin-treated patients have been found to have significantly greater treatment satisfaction than non-insulin treated patients with diabetes, though this significant difference was also lost after adjustments were made for age, gender, body mass index and duration of diabetes [19].

When examining treatment outcome factors which influence the appraisal of treatment satisfaction in diabetes, the most universal finding is that treatment satisfaction has generally been shown to be greater for those treatments that are most efficacious [13,17,20]. Additionally, clinical wisdom assumes that patients prefer treatments with few injections and few side effects such as weight gain or hypoglycemic events, although the data supporting these assumptions is not extensive and is contradictory [1]. For treatments with equal efficacy and no significant differences in weight gain or occurrence of hypoglycemic events, significant differences in treatment satisfaction have not been found [21]. Despite significant improvement in treatment satisfaction after 7 months of efficacious treatment (improved HbA1c) with decreased hypoglycemic events and increased weight, a significant relationship between weight gain, HbA1c and treatment satisfaction has not been found [22]. Interestingly, fewer number of daily injections has been shown both to improve treatment satisfaction and to have no impact on treatment satisfaction [23,24]. Further, continuous subcutaneous insulin infusion has been shown to be both superior to multiple daily injections, as well as to have no impact in determining patient treatment satisfaction [25,26].

Thus, it remains unclear what part of our beliefs about the relationships between patient/disease characteristics, clinical wisdom and treatment satisfaction in diabetes is myth versus reality. The purpose of this study was to examine, in the context of a randomized clinical trial (RCT), the correlative relationship between treatment satisfaction, patient/disease characteristics, and key treatment outcomes that impact the appraisal of treatment satisfaction. By understanding the myths vs. the realities of these relationships, clinicians will be in a better position to identify patients at risk for decreased treatment satisfaction and tailor treatment plans to maximize treatment satisfaction. By improving patient treatment satisfaction, the efficacy of diabetes treatments as well as other critical treatment outcomes such as compliance and cost of care is likely to be improved.
Methods

Procedures
Baseline and end of study (28 weeks) data from the INITIATE trial, a multi-center (25 U.S. sites) open-label RCT comparing the efficacy and safety of twice-daily (BID) biphasic insulin aspart 70/30 (BIAsp 70/30) vs. once-daily, bedtime (QD) glargine (Glar) among insulin naive type 2 patients failing oral medication were analyzed. Eligible subjects had type 2 diabetes inadequately treated, controlled on oral antidiabetic agents (OADs), were insulin naive, and male or female between the ages of 18–75 with baseline HbA1c > 8% and BMI ≤ 40 kg/m². Patients with a history of recurrent, severe hypoglycemia, hepatic or renal insufficiency, cardiac disease, acute or chronic metabolic acidosis, active proliferative retinopathy or intolerance to metformin were excluded from the study. The first phase of the study was a 4 week run-in period with metformin with patients with a fasting blood glucose level of ≤ 140 mg/dL, or any single value ≤ 170 mg/dL considered a run-in failure. Patients who successfully completed the run-in period were then randomized to receive equal doses of BIAsp 70/30 within 15 minutes before breakfast and dinner in addition to their current metformin or Glar at bedtime in addition to their current metformin treatment. Subjects were told to perform daily blood glucose self-monitoring before breakfast and dinner. Insulin doses were titrated weekly for the first 12 weeks, then every 2 weeks thereafter according to a predefined titration algorithm based on blood glucose levels for the 3 days preceding a visit. Post-randomization, subjects were seen in the office weekly for the first month, and then again at weeks 8, 12, 16, 20, 24 and 28. Phone visits were conducted for weeks in which there was no office visit. Patients in the INITIATE trial signed informed consent forms and the study complied with FDA Good Clinical Practices. The study protocol had IRB approval.

Measures
The following data were derived from and/or generated during the trial and were used for the present analyses:

- Patient characteristics and diabetes history including age (as a continuous variable), sex, ethnic group, time with diabetes (at baseline as a continuous variable), body mass index (BMI) and co-morbid conditions (classified as diabetes-related condition and as total number of conditions). These data were collected at the baseline visit.

- The Insulin Treatment Satisfaction Questionnaire (ITSQ) [11]: a validated measure consisting of 22 items that comprehensively assess treatment satisfaction for persons with diabetes on insulin. The measure was developed as a patient-reported outcome based on interviews with patients and clinical experts regarding treatment satisfaction as well as information in the literature. Satisfactory factor structure and internal consistency as well as adequate test-retest reliability, construct and discriminant validity for the ITSQ have been demonstrated [11]. In addition to an Overall score, the items make up five domains of satisfaction: Inconvenience of Regimen (IR-5 items), Lifestyle Flexibility (LF-3 items), Glycemic Control (GC-3 items), Hypoglycemic Control (HC-5 items) and Insulin Delivery Device (DD-6 items). All items are scored on a seven-point Likert-like response scale ranging from "not at all" to "extremely" (as worded appropriately for item). Items in the Glycemic and Hypoglycemic Control subscales are worded so that the respondent is asked about the relevant symptoms of their control such that a clinical understanding of either is not required. The ITSQ is scored by transforming all items to a scale of 0–100 with the higher score indicating better treatment satisfaction. For each subscale, the sum score is divided by number of items. Missing values are imputed based on the mean of the non-missing items. The ITSQ was administered after 28 weeks of treatment.

- Diabetic treatment effect status assessed after 28 weeks of treatment:
  - HbA1c level
  - Hypoglycemic episodes classified as total number, by timing of event (classified as day or night), and by type of event (classified as symptomatic or not)
  - BMI classified as BMI group (< 25 as normal, between 25 and 29.9 as overweight, and ≥ 30 as obese) and by absolute value change (classified as improved or worsened)

Statistical strategy
According to an a priori statistical analysis plan, for each statistical test performed, the relationship between the factors of interest, overall satisfaction and satisfaction within each domain of the ITSQ was examined by linear regression with listwise entry (all variables entered as a block to test interaction with treatment satisfaction as the dependent variable). Statistical significance was considered to be achieved with a minimum P value of 0.05.

Examining the relationship between patient and disease characteristics and treatment satisfaction
First, in order to confirm successfully randomized cohorts, differences between treatment groups for baseline patient and disease characteristics were examined by ANOVA or chi square as appropriate for each characteristic.

Next, a listwise linear regression analysis was performed to examine baseline predictors of treatment satisfaction assessed at week 28. The model included all demographic
and diabetes history variables collected at baseline (age, gender, ethnicity, duration of diabetes diagnosis, BMI, and number/type of diabetes-related co-morbid conditions). Independent linear regression models were assessed for each of the ITSQ subscales and the Overall score.

The relationship of total number of co-morbid conditions and treatment satisfaction (each subscale and Overall) was examined by Pearson correlations. Additionally, the relationship between types of diabetes-related co-morbid conditions and treatment satisfaction was examined by independent linear regression (listwise) analyses of the diabetes-related co-morbid conditions found in this sample and treatment satisfaction.

**Examining the relationships between treatment outcomes and treatment satisfaction**

The relationship between the number of minor hypoglycemic events and treatment satisfaction was assessed by Pearson correlation (2-tailed). Type of event (symptomatic or not) and time of event (day or night), cumulative over the treatment period, and treatment satisfaction were examined by linear regression analyses independently for each subscale and Overall ITSQ score.

The relationship between BMI change during treatment and treatment satisfaction (Overall and all subscales) was first examined by performing an ANOVA, controlling for baseline weight, using change in BMI group from baseline to week 28. An independent ANOVA, controlling for baseline weight, examined the relationship between absolute changes in BMI, comparing those who had improved BMI to those whose BMI had worsened over the 28 weeks.

The relationship between HbA1c levels at week 28 and treatment satisfaction was examined by Pearson correlation coefficients.

The relationship between treatment group and treatment satisfaction was examined by ANOVAs for the total ITSQ score and each of its domains.

**Results**

**Sample description**

A total of 233 patients with Type 2 diabetes, a slight majority of which were male (52.9%) and Caucasian (51.7%) with an average age of 52.15 (± 10.34) years, were randomized into the INITIATE study. Two hundred and nine (209) of these subjects had treatment effect data and 197 of these completed the ITSQ and were included in these analyses. The average duration of diabetes diagnosis was 8.29 years (± 5.24), and mean baseline HbA1c was 9.71% (± 1.47%).

**Statistical findings**

**INITIATE trial clinical results**

The INITIATE trial demonstrated that patients receiving BIAsp 70/30 were significantly more likely to have achieved HbA1c targets vs. Glar (66% vs. 40% of patients to HbA1c < 7%, p < 0.01, and 42% vs. 28% of patients to HbA1c ≤ 6.5%, p < 0.05, respectively), and significantly improving total HbA1c reduction (-0.43%; p < 0.01 between arms). Post-prandial glucose excursions at lunch and supper were also significantly more tightly controlled (p < 0.05), though an increase in minor hypoglycemia and weight gain occurred (both p < 0.05) [27].

**Examining the relationship between patient and disease characteristics and treatment satisfaction**

There were no significant differences found between treatment groups for any of the baseline patient or disease characteristics, indicating that randomization of subjects to treatment groups used in the present analyses was successful.

As shown in Table 1, presence of diabetes-related co-morbid conditions at baseline was significantly predictive of overall treatment satisfaction as well as for Lifestyle Flexibility and in the Hypoglycemic and Glycemic Control subscales. Age was also predictive of treatment satisfaction related to Lifestyle Flexibility.

The number of total co-morbid conditions in the sample ranged from 0 to 7 with a mean of 1.70 (± 1.50) and a median of 1.00. The diabetes-related co-morbid conditions identified in this sample were retinopathy, nephropathy, neuropathy, and macro-angiopathy. There were no significant relationships between total number of co-morbid conditions and treatment satisfaction. However, neuropathy was a significant predictor of treatment satisfaction (overall and in all subscales), indicating that pain is broadly associated with satisfaction outcomes. One other noted predictor was retinopathy, which was significant for the Device Satisfaction subscale. The relationship between types of diabetes-related co-morbid condition and treatment satisfaction is shown in Table 2.
Examining the relationships between treatment outcomes on treatment satisfaction

A total of 1,073 hypoglycemic events, of which 1,072 were minor events, were reported over the 28 weeks of treatment. Hence, the analysis is based on minor hypoglycemic events only. The number of minor hypoglycemic events per patient was found to be significantly associated only with the Hypoglycemic Control subscale ($r = -0.263$, $p < 0.001$). However, as shown in Table 3, the time of a hypoglycemic event was found to be highly predictive of overall treatment satisfaction and for each subscale, with daytime events resulting in decreased treatment satisfaction. There were no significant impacts for the type of hypoglycemic event.

The majority of patients (81%) did not change BMI group (normal, overweight or obese) during the treatment period, and no significant relationships between BMI group change and treatment satisfaction were found. The absolute BMI change ranged from -2.57 (decrease) to +6.83 (mean $1.61 \pm 1.62$) with the majority of the population (85.5%) increasing in BMI levels. For those 14.5% of patients who had a decrease in BMI, there was a systematic improvement in treatment satisfaction across all scores, although only the difference for the Lifestyle Flexibility subscale was significant ($p < 0.05$). The lack of statistical significance may be due to the small number of patients who did change BMI group power (19%).

Pearson correlation coefficients showed a significant relationship between HbA1c and treatment satisfaction. While associations were low, the relationship of HbA1c level to treatment satisfaction was significant for the Overall score ($r = -0.16$, $p < 0.05$), Inconvenience of Regimen ($r = -0.15$, $p < 0.05$), Glycemic Control ($r = -0.23$, $p < 0.001$), and Device Satisfaction ($r = -0.17$, $p < 0.05$).

No significant differences were found in the total ITSQ score or any of the domains during comparisons between treatment groups, indicating in part the potential that an increase in HbA1c could also lead to a decrease in treatment satisfaction.

Table 1: Impact of Demographics and Diabetes History on Insulin-related Treatment Satisfaction

| Coefficient (t-ratio)                     | ITSQ Overall (n = 197) | ITSQ IR (n = 196) | ITSQ LF (n = 196) | ITSQ HC (n = 197) | ITSQ GC (n = 197) | ITSQ DD (n = 197) |
|-------------------------------------------|------------------------|-------------------|------------------|------------------|------------------|------------------|
| Intercept                                 | 72.42*** (6.54)        | 74.63*** (5.89)   | 68.90*** (4.95)  | 67.67*** (4.83)  | 82.53*** (6.32)  | 73.68*** (6.32)  |
| Age                                       | 0.21 (1.51)            | 0.23 (1.43)       | 0.35* (2.00)     | 0.22 (1.24)      | 0.17 (1.05)      | 0.14 (0.92)      |
| Gender (0 = M, 1 = F)                     | -2.30 (-0.87)          | 0.45 (0.15)       | 0.23 (0.07)      | -6.42 (-1.92)    | -4.43 (-1.43)    | -1.95 (-0.70)    |
| Ethnicity (1 = C, 0 = Other)              | -0.99 (-0.36)          | -0.19 (-0.06)     | -2.80 (-0.81)    | -1.44 (-0.42)    | -2.35 (-0.73)    | 0.90 (0.31)      |
| Length of time with diabetes (↑ longer)   | -0.17 (-0.66)          | -0.09 (-0.32)     | -0.06 (-0.17)    | -0.42 (-0.30)    | -0.13 (-0.42)    | -0.17 (-0.61)    |
| Weight-BMI                                | 0.17 (0.66)            | 0.01 (0.04)       | -0.23 (-0.70)    | 0.50 (1.53)      | 0.10 (0.31)      | 0.23 (0.83)      |
| Number of diabetes related co-morbid conditions | -3.93* (-2.15)   | -2.33 (-1.10)     | -5.68* (-2.46)   | -5.08* (-2.19)   | -5.48* (-2.54)   | -1.93 (-1.00)    |

| F = 1.195                                 | F = 0.538              | F = 1.812         | F = 1.918        | F = 1.497        | F = 0.534        |
| $R^2 = 0.036$                             | $R^2 = 0.016$          | $R^2 = 0.053$     | $R^2 = 0.056$    | $R^2 = 0.044$    | $R^2 = 0.016$    |
| Adj $R^2 = 0.006$                         | Adj $R^2 = -0.014$     | Adj $R^2 = 0.024$ | Adj $R^2 = 0.027$| Adj $R^2 = 0.015$| Adj $R^2 = -0.014$|

* p < 0.05, ** p < 0.01, *** p < 0.001
* IR = Inconvenience of Regime subscale, LF = Lifestyle Flexibility subscale, HC = Hypoglycemic Control subscale, GC = Glycemic Control subscale, DD = Device Delivery subscale

Table 2: Impact of Diabetic-related Co-morbid Conditions on Treatment Satisfaction

| Coefficient (t-ratio)                     | ITSQ Overall (n = 197) | ITSQ IR (n = 196) | ITSQ LF (n = 196) | ITSQ HC (n = 197) | ITSQ GC (n = 197) | ITSQ DD (n = 197) |
|-------------------------------------------|------------------------|-------------------|------------------|------------------|------------------|------------------|
| Intercept                                 | 84.11*** (56.89)       | 87.24*** (51.36)  | 78.85*** (41.72) | 81.75*** (42.67) | 86.43*** (49.64) | 84.79*** (54.58) |
| Retinopathy (N = 0, Y = 1)                | 7.29 (1.50)            | 8.94 (1.60)       | 0.95 (0.15)      | 2.31 (0.37)      | 10.17 (1.78)     | 10.90* (2.13)    |
| Nephropathy (N = 0, Y = 1)                | 2.08 (0.40)            | -0.61 (-0.10)     | 3.68 (0.55)      | 3.79 (0.56)      | -2.57 (-0.42)    | 3.94 (0.71)      |
| Neuropathy (N = 0, Y = 1)                 | -10.56*** (-3.53)      | -8.81* (-2.54)    | -12.42** (-3.22) | -10.80** (-2.79) | -13.43*** (-3.82) | -8.00* (-2.54)  |
| Macro Angiopathy (N = 0, Y = 1)           | -5.18 (-1.09)          | -1.16 (-0.21)     | -4.58 (-0.76)    | -5.63 (-0.92)    | -5.93 (-1.06)    | -6.44 (-1.29)    |
| F = 3.622**                               | F = 1.888              |                   |                   |                   |                   |                   |
| $R^2 = 0.069$                             | $R^2 = 0.037$          | $R^2 = 0.057$     | $R^2 = 0.044$    | $R^2 = 0.080$    | $R^2 = 0.054$    |                   |
| Adj $R^2 = 0.050$                         | Adj $R^2 = -0.018$     | Adj $R^2 = 0.038$ | Adj $R^2 = 0.025$| Adj $R^2 = 0.062$| Adj $R^2 = -0.035$|

* p < 0.05, ** p < 0.01, *** p < 0.001
* IR = Inconvenience of Regime subscale, LF = Lifestyle Flexibility subscale, HC = Hypoglycemic Control subscale, GC = Glycemic Control subscale, DD = Device Delivery subscale
increase in number of daily injections (BID vs. QD) may not negatively affect treatment satisfaction.

Examining the combined impact of significant factors

As shown in Table 4, when all previously identified significant relationships were examined together, neuropathy continued to have the broadest impact on treatment satisfaction (Overall and in all 5 subscales). Improved treatment efficacy (HbA1c) maintained the previously identified significant impacts on overall treatment satisfaction and in 3 subscales. The number of hypoglycemic events remained significant only for the Hypoglycemic Control subscale, however the timing of the hypoglycemic event significantly impacted overall satisfaction and 3 subscales. The number of hypoglycemic events such as patients’ activity levels are likely to be major events they may have a greater potential for serious health consequences. It may also be noteworthy that minor diurnal events may be more bothersome to daily functioning, and therefore more negatively influence treatment satisfaction. On the contrary, minor nocturnal events such as patients’ activity levels are likely to be higher during daytime hours. As there was only 1 severe hypoglycemic event during the study, we were not able to examine the relationship between major events and treatment satisfaction.

Also, contrary to common wisdom, no significant differences in treatment satisfaction were found between treatment groups despite the fact that BIAsp 70/30 was administered twice a day (with a pen device) and Glar was administered once a day (with a syringe) as a fixed characteristic of the treatment arm. This finding suggests that number of daily injections may not impact overall treatment satisfaction. However, it should be noted that it remains unclear if daytime hypoglycemic events are also more "feared" or problematic for patients, as nighttime events may be more difficult to identify or respond to and if they are major events they may have a greater potential for serious health consequences. It may also be noteworthy that minor diurnal events may be more bothersome to daily functioning, and therefore more negatively influence treatment satisfaction.

Discussion

Clinical wisdom and common sense would presume, and research has at times supported the idea, that there is a negative impact of unfavorable treatment characteristics, burden of treatment and side effects on patient-reported treatment satisfaction. However, we did not find a significant relationship between the number of minor hypoglycemic events, weight gain or treatments with 1 vs. 2 daily injections with overall treatment satisfaction. The relationship of weight gain to treatment satisfaction was restricted only to Lifestyle Flexibility. This significant finding may be due to the nature of the items in the Lifestyle Flexibility domain, as 2 of the 3 items in that subscale concern eating times and meals. The timing of the hypoglycemic event (minor hypoglycemic events that occurred during the day vs. nocturnal episodes) did have a significantly negative impact on overall treatment satisfaction. However, it should be noted that it remains unclear if daytime hypoglycemic events are also more "feared" or problematic for patients, as nighttime events may be more difficult to identify or respond to and if they are major events they may have a greater potential for serious health consequences. It may also be noteworthy that minor diurnal events may be more bothersome to daily functioning, and therefore more negatively influence treatment satisfaction. On the contrary, minor nocturnal events such as patients’ activity levels are likely to be higher during daytime hours. As there was only 1 severe hypoglycemic event during the study, we were not able to examine the relationship between major events and treatment satisfaction.
Regarding the belief that treatment efficacy is the primary driver of treatment satisfaction, BIAsp 70/30 was found to have superior efficacy (improvement in HbA1c) to Glar in the INITIATE trial, evidence that is supported in a subsequent clinical trial [33]. Although there is no data available to confirm that patients knew their HbA1c levels, doses were regularly titrated based on blood glucose levels and patients self-monitored their blood glucose levels twice daily. Further, HbA1c level was found to be significantly related to Overall Satisfaction suggesting that subjects were aware of treatment efficacy either by information from the physician or by how they felt. Thus, it is highly likely that the patients were aware of their HbA1c values, and able to use this factor when appraising their treatment satisfaction. Despite superior efficacy, treatment satisfaction was equivalent between the 2 treatment groups. This finding may be explained by the Decisional Balance Model of Treatment Satisfaction, whereby Overall treatment satisfaction is determined by the balance between the positive value of treatment and the negative harms and inconveniences of the medication [3]. Thus, the appraisal of treatment satisfaction for a treatment such as BIAsp 70/30, with superior efficacy when compared to Glar, but increased minor hypoglycemia and weight gain, may be equivalent. The Balance Theory model and our findings in the present analysis are supported by those in recent trials where competing insulin preparations which demonstrate disparate side effect and efficacy profiles have caused non-significant differences in treatment satisfaction [34]. The Balance Theory model may also help explain the lack of significant differences in overall treatment satisfaction in pen vs. syringe device studies, even when there is clearly stated preference for the pen device, if treatments have equivalent efficacy and safety [28].

It should be noted that the ITSQ overall score is not a truly independent rating of overall treatment satisfaction as it is the sum of the subscale items rather than a separate item assessing overall satisfaction. Post-hoc analyses with this study data have shown that the correlations between subscales and the overall score are all above 0.77 and a regression analysis of subscales on the Overall score, found all subscales were significantly associated with the Overall score at the 0.01 level, indicating a balance of the subscale weights contributing to the Overall score. We believe that in order to fully understand the impact of variables on treatment satisfaction, the impact on both subscales and Overall treatment satisfaction should be identified.

Understanding the relative weight of subscales in the overall assessment may be valuable as this can help identify subscales of greater importance given a particular treatment or patient characteristic. Future studies may find it helpful to include a truly independent measure of overall satisfaction to further understand the contribution of subscales to overall satisfaction.

In this study we have examined both “simple” regression models looking at similar types of factors (e.g. demographic and patient characteristics) as well as more “complex” multivariate models allowing us to test if significance of variables from the like factor model is maintained when "competing" against a wide spectrum of...
influences. This incremental approach to the importance of factors, provides a more compressive picture of these factors and allows use to refine our understanding of their influence treatment satisfaction. It should be noted that although statistically significant relationships were identified in the study, the amount of variance accounted for in some of the models was small to moderate. Further, as with all statistical significance, it is unclear what the relationship is between the statistical and clinical significance of the findings and it is often difficult to separate out the unique contributions of variables which may be related, such as age and co-morbidity. Older type 2 diabetes patients are more likely to have diabetes related co-morbid conditions and these conditions may negatively impact a patient’s ability to self-manage their disease, creating a barrier to positive outcomes.

Thus, it is important to understand the unique contributions of each of these factors to overall as well as domains of satisfaction. For example, although age did not have a significant impact on Overall satisfaction, neuropathy, a painful diabetes related condition, was the strongest predictor for decreased treatment satisfaction overall and across subscales. Although it may be difficult for physicians to reduce neuropathy-related pain in diabetes, lessening of pain has been found to play only a partial role in explaining treatment satisfaction for people with chronic pain. Several studies have found that provider-patient interaction factors such as confidence, trust and positive communications are also important for improving satisfaction [35-37]. Thus, improving provider-patient relationships for persons with diabetes and neuropathy may help improve their treatment satisfaction in general and lead to improved outcomes. This study did not examine the impact of non-diabetes related co-morbid conditions on treatment satisfaction and we suggest that this relationship be examined in future studies.

Diabetes related etinopathy was also a significant indicator for the Device Delivery subscale, indicating that visual impairments and the sensory experience should be taken into consideration in choosing an insulin delivery device for a given patient. When pen devices have been compared head-to-head (FlexPen vs. Humalog Pen vs. syringe), significant improvements in patient preferences have been found in favor of the devices which improve the readability of the dose scale and user confidence [28,38]. The auditory feature of pen devices has also been shown to affect the patient’s confidence in selecting the correct dose [39], suggesting that auditory impairments also be considered when exploring treatment options.

These findings may also help clinicians identify patients who are at risk for poor treatment satisfaction, so that the clinicians can develop targeted treatment plans for these patients. Certainly, patients with neuropathy or frequent daytime hypoglycemic events should be targeted for a discussion regarding their potential dissatisfaction with treatment. Additionally, their compliance and diabetes self-management behavior, critical factors for diabetes treatment success, should be assessed to identify if these behaviors have been negatively affected by their dissatisfaction. Finally, although cost of treatment was not examined in this study, it should be noted that biphasic insulin has been projected to be cost-effective therapy vs. basal insulin alone for long-term treatment [40,41]. Given the superior efficacy and equivalent patient-reported treatment satisfaction between these treatments, it may be reasonable for clinicians to consider the estimated improvements in quality-adjusted life expectancy along with potential cost savings associated with complication reductions.

Understanding the complex and multidimensional nature of treatment satisfaction as well as the interaction between factors that may impact how persons with diabetes appraise their treatments is still evolving. This study has begun to examine factors that may impact these relationships; considerably more research will be required to further unravel the myths and the realities of treatment satisfaction in diabetes. There are some shortcomings of this study that should be noted so that future research can address these issues. First, although satisfaction with previous treatments may significantly impact satisfaction with new treatments [42], the reality is that it is often difficult or unfeasible to assess. The lack of baseline treatment satisfaction also prevents the assessment of responsiveness of a measure to differences in treatment. These methodological challenges present limitations both in the broader treatment satisfaction literature and in this study. As the subjects were insulin naïve at baseline, previous insulin treatment satisfaction could not be measured. However, the absence of any differences in baseline demographic or health characteristics in the sample in combination with the study entry criteria requiring all subjects to have failed OADs, suggests that baseline differences in treatment satisfaction would not have been significant. Furthermore, a 4 week run-in period prior to the addition of the study insulin ensured that all randomized subjects were stabilized on an identical OAD regimen at baseline. It is also known that RCT populations are generally not representative of the entire population. Therefore, it would be very informative to examine these relationships in a more representative diabetic population such as in a clinic-based study. Additionally, although this study examined potential baseline patient/disease characteristics and key treatment outcomes that may influence treatment satisfaction, data on other important ongoing treatment factors such as burden and compliance were not collected. There may also be other diabetes as well as non
diabetes related co-morbid conditions not reported in this sample, which may impact treatment satisfaction. These additional influences should be considered when looking at the broader spectrum of variables that may or may not impact treatment satisfaction. Inclusion of these variables or other potential drivers of treatment satisfaction may have increased the amount of variance in predicting treatment satisfaction in this study.

Conclusion
Using baseline and end of study treatment effect data from a randomized controlled trial comparing the efficacy of BIAsp 70/30 (administered QD via pen device) and Glar (administered QD via syringe), and by correlating it to measured responses in the ITSQ, we examined the relationship between treatment satisfaction, patient/disease characteristics, treatment outcomes, and treatment groups. By assessing these factors independently, certain significant relationships were identified. However, not all of these significant findings survived when examined in combination with each other. Thus, in order to more accurately characterize the impact of patient/disease characteristics and treatment outcomes on treatment satisfaction, a more comprehensive approach at capturing data on all potentially relevant variables is necessary. This approach can enhance our understanding of factors that exert enduring or broader impacts on treatment satisfaction.

The perfect drug for diabetes would improve efficacy with no side effects, and the perfect patient with diabetes would be otherwise healthy. However, given the real world, clinicians will need to balance the advantages and disadvantages of various treatments and patient types, and aim to more fully understand the myths and realities of patient-reported treatment satisfaction to identify the optimal treatment for a given patient.

Competing interests
The study reported on within this article was sponsored by Novo Nordisk. Authors 2 and 3 are current employees of Novo Nordisk.

Authors’ contributions
MB, DB, MLT and PR participated in the study design and conceptualization of the project. MB, DB, DC participated in the data analysis planning and execution. All authors read and approved the manuscript.

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