Dear Editor,

Insulin glargine (iGla) and detemir (iDet) are the two most commonly used basal insulins in basal-bolus regimens in type 1 diabetes (T1D) [1]. While both these analogues have similar onset of action, their duration of effect has remained a matter of intense debate [1]. Under steady-state conditions, the mean duration of action of iGla is 24–25.6 hours and that of iDet is 21.5 hours [1]. Consequently, a higher proportion of patients are prescribed a twice-daily dosing of iDet as compared to iGla for a full basal coverage [1]. In addition to the concerns regarding the last few hours of the iDet action, twice-daily dosing also results in increased costs and hence its limited use in resource-poor settings. Unfortunately, most of the studies on the efficacy and action duration have compared either of the insulin analogues with NPH, and head-to-head iGla-iDet comparisons are limited [2-4]. In particular, it is unknown whether there is really a waning of effect of iDet during the last few hours of its action duration in daily clinical practice, which would justify its twice-daily dosing.

Self-monitored blood glucose (SMBG) records of the last three months in 22 randomly selected children with T1D (11 each using iDet and iGla in once-daily bedtime dosing) were compared. In addition, the mean HbA\textsubscript{1c}, mean daily total, and basal and mealtime bolus insulin doses were calculated. All patients were using either aspart or lispro as bolus insulin. We presumed that if the iDet effect really wanes, the required doses of evening insulin bolus might be higher in patients using iDet.

The mean age of the two groups of patients was similar (8.9 ± 4.7 vs. 8.3 ± 4.8 years, respectively). Five children in the iGla group and four in the iDet group were pubertal. The mean duration of diabetes was 3.2 ± 2.3 years. In the iGla group, four (36%) families belonged to lower, five (45%) to middle, and two (18%) to upper socioeconomic status (SES). The iDet group had three (27%), five (45%), and three (27%) families in the lower, middle, and upper SES, respectively. The parents’ educational levels were also similar in the two groups. The glycaemic parameters, episodes of hypoglycaemia, and the requirement of daily basal and bolus insulin doses were also similar in the two groups (Table I).

These results indicate a similar efficacy of both basal analogues when used in once-daily dosing in routine clinical practice and are consistent with our previous studies in children with T1D [2, 3]. The common belief amongst physicians that a twice-daily dosing is required in a substantial proportion of patients who are on iDet could not be ascertained in this small study [1]. The basis of such belief, however, is not supported by available literature. There are a limited number of studies on head-to-head comparison of iDet and iGla in patients with T1D [4, 5]. All the comparison studies conducted on patients with either T1D or type 2 diabetes (T2D) also used once-daily dosing of these two basal insulin analogues and demonstrated similar efficacy [5–7]. In the only available trial that compared once- versus twice-daily iDet in patients with T1D, the HbA\textsubscript{1c} at four months was 8.1 ± 0.9 vs. 8.0 ± 1.0% with once- and twice-daily iDet, respectively, with an adjusted between-group difference of 0.12%, showing non-inferiority for once-daily dosing [8]. This study concluded that the most suitable routine starting schedule for iDet is once-daily injection in a basal-bolus regimen for T1D [8]. The findings were the same in another study that compared once- and twice-daily iDet dosing in T2D [9]. A recent meta-analysis that included mainly observational studies generally considered closer to the “real-world” situation also did not indicate favouring a twice-daily administration of iDet [4].

A similar requirement of pre-dinner bolus insulin doses in both groups indicates that the effect of once daily iDet remained similar to iGla during the last few hours of its 24-hour action. This finding assumes importance in view of the recently described “dusk phenomenon” characterised by unexplained pre/post-dinner hyperglycaemia in patients with diabetes [10]. Although the underlying mechanisms are unknown, one of the ways suggested to overcome this phenomenon is to increase the basal insulin.
supply in patients using insulin pumps, indirectly indicating an increased requirement of the evening insulin [10]. This also would have meant an increased requirement of pre-dinner insulin in our patients using iDet if we presumed that the iDet effect wanes during last few hours of its daily profile. Additionally, in our extensive experience of using basal insulin analogues, we have never used iDet in twice-daily dosing [2, 3]. We suggest conducting larger studies comparing iGla with iDet as well as once- versus twice-daily dosing of iGla in real-life clinical practice. Until then, the treating physicians may consider using iDet as once-daily dosing only in children with T1D.

Table I. Comparison of glycaemic parameters, hypoglycaemic events, and insulin doses of patients treated with insulin glargine and insulin detemir

| Variable                        | Glargine group Mean (SD) | Detemir group Mean (SD) | p-value |
|---------------------------------|--------------------------|-------------------------|---------|
| Blood glucose                   |                          |                         |         |
| Before breakfast                | 148.40 (86.54)           | 129.84 (59.14)          | 0.56    |
| Before lunch                    | 156.80 (79.97)           | 170.10 (79.14)          | 0.69    |
| Before dinner                   | 163.72 (87.13)           | 167.72 (79.13)          | 0.91    |
| At Bedtime                      | 158.87 (92.35)           | 162.81 (76.04)          | 0.91    |
| Hypoglycaemic episodes per month|                          |                         |         |
| Asymptomatic                    | 1.81 (0.40)              | 1.63 (0.50)             | 0.36    |
| Symptomatic                     | 0.63 (0.50)              | 0.81 (0.60)             | 0.45    |
| Severe symptomatic              | 0.18 (0.40)              | 0.09 (0.30)             | 0.55    |
| Nocturnal                       | 0 (0)                    | 0 (0)                   |         |
| HbA1c                           | 7.88 (1.25)              | 7.57 (1.03)             | 0.53    |
| Basal insulin dose (U/kg/day)   | 0.231 (0.41)             | 0.371 (0.13)            | 0.29    |
| Bolus insulin dose (U/kg/day)   | 0.488 (0.51)             | 0.620 (0.28)            | 0.46    |
| Before breakfast bolus (U/kg/day)| 0.172 (0.40)             | 0.206 (0.09)            | 0.78    |
| Before lunch bolus (U/kg/day)   | 0.178 (0.29)             | 0.221 (0.10)            | 0.64    |
| Before dinner bolus (U/kg/day)  | 0.141(0.06)              | 0.190 (0.10)            | 0.17    |

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