Childhood Height Growth Rate Association With the Risk of Islet Autoimmunity and Development of Type 1 Diabetes

Zhiguo Li,1,10 Riitta Veijola,2 Eileen Koski,1 Vibha Anand,1 Frank Martin,3 Kathleen Waugh,4 Heikki Hyöty,5 Christiane Winkler,6,7,8 Michael B. Killian,9 Markus Lundgren,10,11 Kenney Ng,1 Marlena Maziarz,10,11 and Jorma Toppari12,12,*

1Center for Computational Health, IBM T.J. Watson Research Center, Yorktown Heights, 10598 NY, and Cambridge, 02142 MA, USA
2Department of Pediatrics, PEDEGO Research Unit, University of Oulu, 90014 Oulu, and Oulu University Hospital, 90029 Oulu, Finland
3JDRF, New York, 10281 NY, USA
4Barbara Davis Center for Diabetes, University of Colorado, Denver, 80440 CO, USA
5Department of Virology, Faculty of Medicine and Health Technology, Tampere University, and Fimlab Laboratories, Pirkanmaa Hospital District, 33520 Tampere, Finland
6Institute of Diabetes Research, Helmholtz Zentrum München, German Research Center for Environmental Health, 85764 Munich-Neuherberg, Germany
7Forschergruppe Diabetes e.V. at Helmholtz Zentrum, München, German Research Center for Environmental Health, 85764 Munich-Neuherberg, Germany
8Forschergruppe Diabetes, Technical University Munich, at Klinikum rechts der Isar, 80334 Munich, Germany
9Pacific Northwest Research Institute, Seattle, 98122 WA, USA
10Department of Clinical Sciences, Lund University Diabetes Center, 21428 Malmö, Sweden
11Department of Pediatrics, Kristianstad Hospital, 29133 Kristianstad, Sweden
12Institute of Biomedicine, Research Centre for Integrative Physiology and Pharmacology, and Centre for Population Health Research, University of Turku, and Department of Pediatrics, Turku University Hospital, 20520 Turku, Finland

Correspondence: Jorma Toppari, MD, PhD, Institute of Biomedicine, Research Centre for Integrative Physiology and Pharmacology, and Centre for Population Health Research, University of Turku, and Department of Pediatrics, Turku University Hospital, 20520 Turku, Finland. Email: jortop@utu.fi; or Zhiguo Li, PhD, Center for Computational Health, IBM T.J. Watson Research Center, Yorktown Heights, 10598 NY, USA. Email: lizh@us.ibm.com

*M.M. and J.T. contributed equally.

Abstract

Context: Rapid growth has been suggested to promote islet autoimmunity and progression to type 1 diabetes (T1D). Childhood growth has not been analyzed separately from the infant growth period in most previous studies, but it may have distinct features due to differences between the stages of development.

Objective: We aimed to analyze the association of childhood growth with development of islet autoimmunity and progression to T1D diagnosis in children 1 to 8 years of age.

Methods: Longitudinal data of childhood growth and development of islet autoimmunity and T1D were analyzed in a prospective cohort study including 10 145 children from Finland, Germany, Sweden, and the United States, 1-8 years of age with at least 3 height and weight measurements and at least 1 measurement of islet autoantibodies. The primary outcome was the appearance of islet autoimmunity and progression from islet autoimmunity to T1D.

Results: Rapid increase in height (cm/year) was associated with increased risk of seroconversion to glutamic acid decarboxylase autoantibody, insulin autoantibody, or insulinoma-like antigen-2 autoantibody (hazard ratio [HR] = 1.26 [95% CI = 1.05, 1.51] for 1-3 years of age and HR = 1.48 [95% CI = 1.28, 1.73] for >3 years of age). Furthermore, height rate was positively associated with development of T1D (HR = 1.80 [95% CI = 1.15, 2.81]) in the analyses from seroconversion with insulin autoantibody to diabetes.

Conclusion: Rapid height growth rate in childhood is associated with increased risk of islet autoimmunity and progression to T1D. Further work is needed to investigate the biological mechanism that may explain this association.

Key Words: type 1 diabetes, autoantibodies, prospective cohort, child growth

Abbreviations: GADA, glutamic acid decarboxylase autoantibodies; HLA, human leukocyte antigen; IA-2A, islet antigen 2 antibodies; IAA, insulin autoantibodies; IGF-1, insulin-like growth factor 1; T1DI, Type 1 Diabetes Intelligence study.

Rapid linear growth has been suggested as a risk factor for type 1 diabetes (1). According to the accelerator hypothesis (2-4), obesity and rapid weight gain could contribute to development of type 1 diabetes via insulin resistance (5). There are some results supporting this hypothesis (6-8). Rapid linear growth in infancy has been associated with increased risk of islet autoimmunity and type 1 diabetes. The ongoing TEDDY study (8) recently analyzed distinct infant and childhood growth phases in relation to the risk of type 1 diabetes and found that a lower height growth rate in infancy (hazard

Received: 8 November 2021. Editorial Decision: 1 March 2022. Corrected and Typeset: 26 March 2022
© The Author(s) 2022. Published by Oxford University Press on behalf of the Endocrine Society. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.
ratio [HR] = 0.79 [95% CI = 0.70, 0.90] per 1 cm/year) and higher rate in early childhood (after inflection point between infant and childhood growth periods that occurs around 7-10 months of age) [HR = 1.48 [95% CI = 1.22, 1.79] per 1 cm/year] was associated with increased risk of progression from islet autoimmunity to type 1 diabetes. This suggests that growth rates in distinct childhood growth phases might be associated differently with development of islet autoimmunity and progression to clinical type 1 diabetes. The infancy-childhood-puberty (ICP) growth model recognizes different regulatory mechanisms of growth in successive phases (9).

Taking these findings into account, our study focused specifically on the childhood growth phase, for which we have had a unique opportunity to examine the Type 1 Diabetes Intelligence study (T1DI) cohort (10), a large cohort of children prospectively followed for development of islet autoimmunity and diagnosis of type 1 diabetes. Longitudinal data in this cohort represents child growth patterns in the United States and Europe. Infants and toddlers in the T1DI cohort were systematically followed for development of islet autoantibodies starting at very young ages, often as early as 6 months of age. While previous work sheds light on development of islet autoimmunity in early childhood, including infancy, those analyses were restricted to the period up to 4 years of age (7). In this study, we analyze the association of rapid growth during the growth period from early childhood to pre-adolescence, which we define as the growth period from 1 to 8 years of age.

We hypothesized that rapid growth rate in childhood is associated with both development of islet autoimmunity and progression to type 1 diabetes. To test this hypothesis, we analyzed data on 10,145 children from 1 to 8 years of age to evaluate the association between 4 main predictors (height, weight, and rates of change in height and weight) and 2 main outcomes (seroconversion [with 4 types of seroconversion events] or type 1 diabetes) in 3 analysis timeframes (from 1 year of age to confirmed seroconversion to islet autoimmunity; from one year of age to onset of type 1 diabetes; and from seroconversion to onset of type 1 diabetes).

**Methods**

**Study Population**

The prospective T1DI study cohort consists of 24,662 children who have been followed in 5 natural history studies of type 1 diabetes including 2 in the United States, DAISY (11) and DEW-IT (12), and 3 in Europe, specifically, Germany (BABYDIAB) (13), Sweden (DiPiS) (14), and Finland (DIPP) (15). The focus of these ongoing studies is development of islet autoimmunity and progression to type 1 diabetes in children at high genetic or familial risk. All 5 studies have followed children at least up to 15 years of age or until diagnosis with type 1 diabetes, whichever occurred first.

Supplementary Figure S1 describes the formation of the present study cohort (16). From the full T1DI cohort, we previously analyzed (10) an infant-toddler subcohort restricted to children with first autoantibody measurement by 2.5 years of age (n = 16,709). From this infant-toddler subcohort, we selected children 1.0 to 8.0 years of age who had at least 3 height and weight measurements and at least 1 measurement of each of the 3 islet autoantibodies we considered (those against insulin [IAA], glutamic acid decarboxylase [GADA], and insulinoma-like antigen-2 [IA-2A]). Children diagnosed with type 1 diabetes before 1 year of age were excluded from our analyses. The final dataset, comprising 10,145 children, is hereafter referred to as the “childhood growth cohort.” Of these, 131 (1.3%) developed type 1 diabetes by 8 years of age during follow-up. Baseline data on sex, harmonized human leukocyte antigen (HLA) risk group (10), and study site were available for each subject. Since very few children in DEW-IT were recruited early enough to have the necessary growth and autoantibody data, they were not included in the analysis.

In all studies from which data were included in this analysis, height and weight were measured by trained staff such as clinicians, clinical staff, or research personnel. Measurements were taken using appropriate professional equipment, such as stadiometers or infantometers, and scales with a high degree of precision. In most cases, height was measured with children standing as soon as they were able to do so, which was generally between 1 and 2 years of age.

**Statistical Methods**

The HLA typing data was divided into 4 groups based on previously assessed risk of type 1 diabetes (17) and treated as a categorical variable with “HLA group A” being the highest risk group and “HLA group D” the lowest, with the latter treated as the reference in the analyses. Study site was treated as a categorical variable, and sex as binary (1 = male, 0 = female). The missing height (cm) and weight (kg) measurements were first imputed based on multiple imputation methodology using CDC reference which yielded low estimates of error (18). Imputations were then further refined with the best linear unbiased predictors for each subject estimated by linear mixed effects models with height or weight as the outcome, age and quadratic age as the fixed effects, with random intercepts and slopes, and adjusted for sex and study site (Supplementary Materials Supplement 2) (16). We compared the observed and estimated growth trajectories to evaluate the appropriateness of the imputation model (Supplementary Figures S2-S3) (16). Growth rate measurements, specifically, rates of change in height (cm/year) and weight (kg/year), were estimated using the first derivatives of the polynomials in the fitted models. Autoantibody status (1 = positive, 0 = negative) for IAA, GADA, and IA-2A was available from each study site using positivity thresholds appropriate for the assays used in each study. Missing autoantibody status information was imputed for some of the visits based on autoantibody status at previous or later visits in a given timeframe (Supplementary Materials Supplement 3) (16).

The outcomes of interest were events defined in detail below. An event was said to be observed (event = 1) if the outcome of interest was documented for a given subject within the analysis timeframe, otherwise, the observation was censored (event = 0) at the time of the last observed visit within the analysis timeframe or at 8 years of age. The main outcomes were seroconversion and type 1 diabetes, which were used in 3 sets of analyses differing by start time (t0) and outcome: (1) 1 year of age (t0) to seroconversion; (2) 1 year of age (t0) to type 1 diabetes diagnosis; and (3) seroconversion (t0) to type 1 diabetes diagnosis. The age at seroconversion with a given autoantibody (or autoantibodies) was defined as the age at the first of 2 consecutive samples positive for that autoantibody (or autoantibodies). In our analysis, we considered 4 seroconversion events: (A) seroconversion to the first of any of IAA, GADA, or IA-2A; (B)
Table 1. Characteristics of 9 models

| Model | Start (t₀) | Outcome | Islet Autoantibody seroconversion event | Adjustments | Subjects (events) (n) |
|-------|------------|---------|----------------------------------------|-------------|----------------------|
| 1A    | Age = 1 year | Seroconversion | Any (IAA, GADA, or IA-2A) | Sex, HLA | 10 008 (592) |
| 1B    | IAA        |          |                                        |             | 10 045 (361) |
| 1C    | GADA       |          |                                        |             | 10 086 (458) |
| 1D    | Multiple   |          |                                        |             | 10 070 (359) |
| 2     | Age = 1 year | Type 1 diabetes | n/a | Sex, HLA, islet autoantibody seroconversion event | 10 144 (313) |
| 3A    | Seroconversion | Type 1 diabetes | Any (IAA, GADA, or IA-2A) | Sex, HLA | 10 145 (131) |
| 3B    | IAA        |          |                                        |             | 10 145 (77) |
| 3C    | GADA       |          |                                        |             | 10 145 (70) |
| 3D    | Multiple   |          |                                        |             | 10 145 (92) |
Germany, and the United States. The majority of the children had the HLA genotype belonging to risk group B (n = 4941, 48.7%) while 2300 (22.7%) belonged to HLA group D, which served as the reference group in the analysis. The mean (SD) duration of follow-up was 6.9 (1.6) years. During that time the average number of height and weight measurements available per subject was 8.1 (5.1) and 8.2 (5.0), respectively. Additionally, on average, we imputed 1.2 (2.0) measurements for height and 1.1 (2.0) for weight.

One Year of Age (t0, in the analysis) to Seroconversion (Fig. 1)

Overall, in this part of the analyses, we noted a statistically significant positive association between each of the outcomes and rate of change in height over time in cm/year (height rate) for children in both the 1-3 years and 3-8 years age groups (Fig. 1A-D). Height was negatively associated with seroconversion with any of IAA, GADA, or IA-2A (Fig. 1A) as well as with GADA (Fig. 1C). The rate of change in weight over time in kg/year (weight rate) in children from 1 to 3 years of age tended to be negatively associated with seroconversion, as evidenced by the HR estimates being consistently < 1, albeit not statistically significantly. In all 4 analyses, we noted statistically significant positive associations between HLA group A (highest risk of type 1 diabetes) vs D (lowest risk) and all outcomes, as well as HLA group B vs D for 2 outcomes with multiple autoantibodies (Fig. 1A and 1C). Additional details on the main results from each of the 4 models (Fig. 1A-1D) are provided below.

Table 2. Characteristics of the T1DI Study growth rate analysis dataset (n = 10 145)

| Participants by study and country, n (%) | Number of events (n (%)) |
|-----------------------------------------|--------------------------|
| BABYDIAB, Germany                        | 1324 (13.1)              |
| DAISY, United States                     | 1373 (13.5)              |
| DiPiS, Sweden                            | 1233 (12.2)              |
| DIPP, Finland                            | 6215 (61.3)              |
| Duration of follow-up from 1 to 8 years of age, mean (SD) | 6.94 (1.61) |
| Number of growth measures available per subject, mean (SD) | 8.12 (5.05) |
| Height                                   | 8.24 (5.01)              |
| Weight                                   | 8.12 (5.05)              |
| Number of imputed growth measures per subject, mean (SD) | 1.24 (1.95) |
| Height                                   | 1.13 (1.96)              |
| Weight                                   | 1.24 (1.95)              |
| HLA group, n (%)                         | 592 (5.8)                |
| A                                       | 1369 (13.5)              |
| B                                       | 4941 (48.7)              |
| C                                       | 1535 (15.1)              |
| D                                       | 2300 (22.7)              |
| Seroconversion with IAA, GADA, or IA-2A  | 361 (3.6)                |
| Seroconversion with GADA                 | 458 (4.5)                |
| Seroconversion with multiple persistent autoantibodies | 359 (3.5) |
| Type 1 diabetes                          | 131 (1.3)                |

The Cox regression analysis from 1 year of age to seroconversion with the first of any of GADA, IAA, or IA-2A as the outcome was based on a sample size of 10 008 and 592 events (Fig. 1A). The estimated HR and the associated 95% CI of the association between height (cm) and the outcome was HR = 0.95 (95% CI = 0.93, 0.97), P < 0.001. The rate of change in height (cm/year) was associated with the outcome in children 1-3 years of age (HR = 1.26 [1.05, 1.51], P = 0.014) and in children 3-8 years of age (HR = 1.48 [1.28, 1.73], P < 0.001) (Fig. 1A).

The analysis from 1 year of age to seroconversion with IAA was based on n = 10 045 and 361 events (Fig. 1B). The rate of change in height (cm/year) was positively associated with seroconversion with IAA in children 1-3 years of age (HR = 1.32 [1.08, 1.63], P = 0.008), but not for children 3-8 years of age (HR = 1.13 [0.90, 1.43], P = 0.283) (Fig. 1B).

The analysis from 1 year of age to seroconversion with GADA as the outcome was based on n = 10 086 and 458 events (Fig. 1C). Seroconversion with GADA was negatively associated with height (cm) (HR = 0.97 [0.94, 0.99], P = 0.008), and positively associated with the rate of change in height (cm/year) in children 1-3 years of age (HR = 1.30 [1.05, 1.61], P = 0.015), and in children 3-8 years of age (HR = 1.52 [1.28, 1.81], P < 0.001) (Fig. 1C).

Lastly, the analysis from 1 year of age to seroconversion with multiple autoantibodies as the outcome was based on n = 10 070 and 359 events (Fig. 1D). The rate of change in height (cm/year) was positively associated with this outcome in children 1-3 years of age (HR = 1.47 [1.19, 1.82], P < 0.001), and in children 3-8 years of age (HR = 1.33 [1.07, 1.65], P = 0.009) (Fig. 1D).

One Year of Age (t0) to Type 1 Diabetes (Fig. 2)

The Cox regression analysis from 1 year of age to type 1 diabetes as the outcome was based on n = 10 144 subjects and 131 events (Fig. 2). The direction of the association between height (cm) and the outcome was positive, though not statistically significant (HR = 1.05 [1.00, 1.10], P = 0.072). The results for the height rate and the weight rate were consistent with those summarized above in “One Year of Age (t0 in the analysis) to Seroconversion” for seroconversion as the outcome. Specifically, the HR estimates for height rate (cm/year) for ages 1-3 and 3-8 were > 1, and those for the weight rate (kg/year) were < 1. However, none of these associations were found to be statistically significant. As before, HLA group A vs D was positively associated with the outcome. Additionally, a positive association between IAA positivity (vs negativity) and development of type 1 diabetes by age 8 years was observed (HR = 3.91 [2.41, 6.22], P < 0.001), and similarly between IA-2A positivity (vs negativity) and type 1 diabetes (HR = 39.5 [21.3, 73.3], P < 0.001) (Fig. 2).

Seroconversion With Seroconversion Event A-D (t0) to Type 1 Diabetes (Fig. 3)

The results in the analyses starting from age at seroconversion up to type 1 diabetes as the outcome were relatively consistent regardless of the type of seroconversion (Fig. 3). The HR estimate for the height rate was > 1 in all 4 analyses but was only marginally statistically significant. The rate of change in
weight (in kg/year) was negatively associated with risk of type 1 diabetes, especially for children older than 3.3 years from seroconversion, with HRs ranging from 0.21 to 0.26 in the 4 analyses. The association between a lower rate of weight gain and diagnosis of type 1 diabetes was estimated at HR = 0.21 (0.09, 0.5), \( P < 0.001 \) after 3.3 years after seroconversion with any of GADA, IAA, IA-2A; HR = 0.25 (0.09, 0.70), \( P = 0.009 \) after 3.3 years after seroconversion with IAA (Fig. 3B); HR = 0.25 (0.09, 0.72), \( P = 0.010 \) after 3.3 years after seroconversion with GADA (Fig. 3C); and HR = 0.26 (0.11, 0.63), \( P = 0.003 \) after 3.3 years after seroconversion with multiple autoantibodies (Fig. 3D). For all analyses, subjects positive for IA-2A were more likely to be diagnosed with type 1 diabetes compared with those who were negative (Fig. 3A-3D). Similar associations between IAA positivity and development of type 1 diabetes were also seen (Fig. 3A-C). In this analysis, the HLA risk group was not found to be associated with development of type 1 diabetes (Fig. 3).

**Discussion**

In this analysis, specifically focusing on the age range 1.0 to 8.0 years, rapid childhood growth was associated with both onset of islet autoimmunity and progression from that to clinical type 1 diabetes. Although the strength of the association varied slightly depending on the type of the first appearing autoantibody, rapid height rate consistently predicted islet autoimmunity, as well as development of diabetes from islet autoimmunity, although less significantly compared with the islet autoimmunity endpoint. The well-known association of
HLA class II genotype with the risk of islet autoimmunity and type 1 diabetes was also evident (21) and was taken into account in our analysis of the putative growth effect. Interestingly, height was inversely associated with the risk of autoimmunity, suggesting that subsequent catch-up growth is connected to the risk. This is also consistent with the findings in the TEDDY study, in which lower height rate in infancy and higher height rate in early childhood were associated with more rapid progression from autoimmunity to diagnosis of type 1 diabetes (8). In most previous studies, infant and childhood growth periods have been analyzed together rather than separately, which may explain many inconsistencies in the previous findings.

In contrast to height rate, weight rate was inversely associated with the rate of progression to type 1 diabetes. We hypothesize that this phenomenon is caused by the typical weight loss before the clinical diagnosis. Otherwise, we should expect weight to increase together with height. On the other hand, here the association of rapid linear growth with the disease process cannot be explained by an increase in body weight. It is typical that development of overweight is associated with slightly accelerated height rate, but here the children experienced weight loss due to deteriorating endogenous insulin secretion and gradual development of hyperglycemia and polyuria before the diagnosis. However, there was also an inverse association with weight rate and onset of islet autoimmunity, which cannot be explained by preclinical weight loss and needs further investigation. This was also different from the result of the German Fr1da study which showed that obesity but not overweight was associated with an increased risk of islet autoimmunity (22).

Regulation of growth is described by an infancy-childhood-puberty (ICP) growth model which refers to differential regulatory mechanisms in infancy, childhood, and puberty (9). Infant growth is often considered a continuation of fetal growth that rapidly decelerates during the first year of life, whereafter childhood growth continues until the end of puberty and the pubertal growth spurt overlaps it during the last few years before adult stature is achieved. Nutrition is the main driver of growth in infancy, while growth hormone is the most important factor affecting height development in childhood, and sex hormones induce the pubertal growth spurt (23). Most previous studies on the association of growth with the risk of islet autoimmunity and type 1 diabetes have analyzed infant and childhood growth together. This approach may have obscured features that were specific to these distinct growth periods. The present data strongly point to a link between childhood growth and risk of islet autoimmunity and diabetes. Our results are in agreement with those from the TEDDY study that showed a positive association of accelerated childhood growth with progression to diabetes (8). Interestingly, the same study showed an inverse association between rapid infant growth and progression to diabetes, emphasizing the importance of analyzing...
growth rates at distinct time periods. It is also in agreement with our finding of negative association of height and development of islet autoimmunity by age 8 years. In the TRIGR study, the risk of islet autoimmunity with multiple autoantibodies was positively associated with the child’s height velocity during the first 2 years of life (24). In the DAISY study, greater height growth velocity of children over 2 years of age was associated with both increased risk of islet autoimmunity and type 1 diabetes (2). Blom et al reported that linear growth was positively associated with diabetes risk in Swedish boys, while the association remained nonsignificant in girls albeit the trend was similar (1). In our analyses, male sex was associated with an earlier onset of islet autoimmunity, but growth rate association was independent of sex.

We do not know whether there is a causal relationship between a rapid growth rate and the risk of islet autoimmunity and further progression to type 1 diabetes. However, rapid growth usually comes with high growth hormone and
insulin-like growth factor 1 (IGF-1) levels, which may interact with insulin and glucose metabolism. The roles of growth hormone and IGF-1 are not similar in infancy and childhood, as infant growth is less dependent on growth hormone than childhood growth. We do not have data for IGF-1 in the T1DI cohort in order to evaluate whether there were any associations between IGF-1 levels and the risk of islet autoimmunity or type 1 diabetes. Beyerlein et al measured IGF-1 and insulin-like growth factor binding protein-3 (IGFBP-3) in a subset of children but found no association with the risk of islet autoimmunity. In their study, early age at infant peak body mass index (BMI) was inversely associated with the risk of islet autoimmunity, whereas height growth rate from infancy forward was not. In further analyses, rapid increase of BMI from infancy to 3 years of age was positively associated with an increased risk of islet autoimmunity, whereas a pattern of higher height SD score at birth followed by a decrease to average values after 3 years was associated with a reduced rate of islet autoimmunity. These latter findings are in agreement with our results, although the study combined data from infancy and childhood growth. In the T1DBIT and DIABIMMUNE studies, lower IGF-1 levels were reported in children with islet autoimmunity and type 1 diabetes than in autoantibody-negative children, which may be caused by decreased intraportal insulin levels leading to growth hormone resistance in the liver and thereby to lower IGF-1 production. This leads to increased growth hormone secretion due to declined negative feedback by IGF-1, and consequently growth hormone can stimulate paracrine secretion of IGF-1 in the growth plate to stimulate growth. While we speculate that this may be true in this setting, it is known that severe long-standing hyperglycemia and concomitant high levels of growth hormone may be associated with impaired growth, so-called Mauriac syndrome.

Rapid growth presents a high metabolic demand to the beta cells causing endoplasmic reticulum stress and oxidative stress, which have been hypothesized to originate beta cell demise and subsequent autoimmunity. This is difficult to measure in vivo and remains therefore hypothetical. However, the clear association between childhood growth rate and the risk of islet autoimmunity and progression to type 1 diabetes gives us an insight into further research needs. New biomarkers of beta cell stress and apoptosis need to be identified and validated.

The strengths of the study are prospective systematic collection of data, its large study population and representation of multiple nationalities, which makes the findings more generalizable. Our study also has some limitations. We were not able to analyze the infant growth phase, which limits comparability to earlier studies covering growth from birth onwards. The T1DI cohort comprises five studies, each of which had its own study protocol. Visit schedules varied, as well as inclusion criteria (HLA criteria or family history). Despite these differences, it was possible to combine and harmonize the data to make the analyses described in this manuscript possible.

In conclusion, rapid height growth rate in childhood is associated with increased risk of islet autoimmunity and, among subjects who seroconverted with IAA, with rapid progression to type 1 diabetes. Including rapid childhood growth in risk score models may improve the prediction accuracy of time to islet autoimmunity and type 1 diabetes. Further work is needed to investigate how much improvement in the risk score can be gained, as well as to elucidate the biological mechanism that may explain these associations.

Acknowledgments

The authors thank the participants of the DAISY, DiPiS, DIPP, DEW-IT, and BABYDIAB studies, and the T1DI Study Group that includes, in addition to the authors, BABYDIAB: Anette G. Ziegler, Ezio Bonifacio, Peter Achenbach; Forschergruppe Diabetes e.V. and Institute of Diabetes Research, Helmholtz Zentrum München, German Research Center for Environmental Health, Munich Neuherberg, Germany, der TU München, Munich, Germany; DAISY: Marian Rewers, Brigitte I. Frohnert, Jill Norris, Andrea Steck, Liping Yu; University of Colorado, Anschutz Medical Campus, Barbara Davis Center for Diabetes; DEW-IT: William A. Hapogian, Rachel Hervey; Pacific Northwest Research Institute. Seattle; DiPiS: Åke Lernmark, Helena Elding Larsson, Lampros Spiliopoulos, Josefín Jonsson; Department of Clinical Sciences Malmö, Lund University; DIPP: Jorma Ilonen, University of Turku and Turku University Hospital, Mikael Knip, Tampere University Hospital, and University of Helsinki; IBM: Mohamed Ghalwash, Bin Liu, Ying Li, B.C. Kwon, Harry Stravropoulos, Ashwani Malhotra, Shelley Moore, Jianying Hu; JDRF: Jessica Dunne, Olivia Lou.

Funding

This work was supported by funding from JDRF (IBM: 1-RSC-2017-368-I-X, 1-IND-2019-717-I-X), (DAISY: 1-SRA-2019-722-I-X, 1-RSC-2017-517-I-X, 1-ECR-2017-388-A-N), (DiPiS: 1-SRA-2019-720-I-X, 1-RSC-2017-526-I-X), (DIPP: 1-RSC-2018-555-I-X), (DEW-IT: 1-SRA-2019-719-I-X, 1-RSC-2017-516-I-X) as well as National Institutes of Health (NIH) (DAISY: DK032493, DK032083, DK104351; DEW-IT: UR6/CCU017247).

The DIPP study was funded by JDRF (grants 1-SRA-2016-342-M-R, 1-SRA-2019-732-M-B); European Union (grant BMH4-CT98-3314); Novo Nordisk Foundation; Academy of Finland (Decision No 292538 and Centre of Excellence in Molecular Systems Immunology and Physiology Research 2012-2017, Decision No. 250114); Special Research Funds for University Hospitals in Finland; Diabetes Research Foundation, Finland; and Sigrid Juselius Foundation, Finland.

The BABYDIAB study was funded by the German Federal Ministry of Education and Research to the German Center for Diabetes Research.

The DiPiS study was funded by Swedish Research Council (grant no. 14064), Swedish Childhood Diabetes Foundation, Swedish Diabetes Association, Nordisk Insulin Fund, SUS funds, Lion Club International, district 101-S, The Royal Physiographic Society, Skåne County Council Foundation for Research and Development, and LUDC-IRC/EXODIAB funding from the Swedish foundation for strategic research (Dnr IRC15-0067) and Swedish Research Council (Dnr 2009-1039).

Additional funding for DEW-IT was provided by the Hussman Foundation and by the Washington State Life Science Discovery Fund.

Disclosures

The authors have nothing to disclose.
Author Contributions
Z.L. performed the analytic work. Z.L., J.T., and M.M. contributed to the study design and interpretation of the results. Z.L., J.T., and M.M. contributed to the manuscript. E.K. reviewed and edited the manuscript. All authors contributed to discussion, review, and approval of the manuscript.

Guarantor
William Hagopian, Markus Lundgren, Marian Rewers, Riitta Veijola, and Anette Ziegler, as representatives of the data originating sites, are the overall guarantors of this project with responsibility for the integrity of the data. Zhiguo Li, as a representative of IBM, is the technical research lead for this project with responsibility for the accuracy of the data analysis. Frank Martin is a representative of JDRF, the convener and funder of the overall initiative.

Data Availability
The data that support the findings of this study are not publicly available because they were used under license for the current study only. Data are, however, available upon reasonable request from the originating sites, whose representatives are William Hagopian (DEW-IT), Markus Lundgren (DiPiS), Marian Rewers (DAISY), Riitta Veijola (DIPP), and Anette Ziegler (BABYDIAB). Zhiguo Li is responsible for the accuracy of the data analysis. The sources used included open-source software; however, the specific code is subject to proprietary constraints but can be made available upon reasonable request to the corresponding author.

References
1. Blom L, Persson L, Dahlquist G. A high linear growth is associated with an increased risk of childhood diabetes mellitus. Diabetologia. 1992;35(6):528-533.
2. Lamb MM, Yin X, Zerbe GO, et al. Height growth velocity, islet autoimmunity and type 1 diabetes development: the diabetes autoimmunity study in the young. Diabetologia. 2009;52(10):2064-2071.
3. Larsson HE, Hansson G, Carlsson A, et al; DiPiS Study Group. Children developing type 1 diabetes before 6 years of age have increased linear growth independent of HLA genotypes. Diabetologia. 2008;51(9):1623-1630.
4. Hyypönen E, Virtanen SM, Kenward MG, Knip M, Åkerblom HK. Obesity, increased linear growth, and risk of type 1 diabetes in children. Diabetes Care. 2000;23(12):1755-1760.
5. Wilkin TJ. The accelerator hypothesis: weight gain as the missing link between Type I and Type II diabetes. Diabetologia. 2001;44(7):914-922.
6. Couper JJ, Beresford S, Hrite C, et al. Weight gain in early life predicts risk of islet autoimmunity in children with a first-degree relative with type 1 diabetes. Diabetes Care. 2009;32(11):94-99.
7. Elding Larsson H, Vehik K, Haller MJ, et al. Growth and risk for islet autoimmunity and progression to type 1 diabetes in early childhood: the environmental determinants of diabetes in the young study. Diabetes. 2016;65(7):1988-1995.
8. Liu X, Vehik K, Huang Y, et al; the TEDDY Study Group. Distinct growth phases in early life associated with the risk of type 1 diabetes: the TEDDY study. Diabetes Care. 2020;43(3):556-562.
9. Karlberg J. A biologically-oriented mathematical model (ICP) for human growth. Acta Paediatr. 1989;78(s350):70-94.
10. Anand V, Li Y, Liu B, et al; T1DI Study Group. Islet autoimmunity and HLA markers of presymptomatic and Clinical Type 1 Diabetes: joint analyses of prospective cohort studies in Finland, Germany, Sweden, and the U.S. Diabetes Care. 2021;44(10):2269-2276.
11. Rewers M, Bugawan TL, Norris JM, et al. Newborn screening for HLA markers associated with IDDM: diabetes autoimmunity study in the young (DAISY). Diabetologia. 1996;39(7):807-812.
12. Wion E, Brantley M, Stevens J, et al. Population-wide infant screening for HLA-based type 1 diabetes risk via dried blood spots from the public health infrastructure. Ann N Y Acad Sci. 2003;1005:400-403.
13. Ziegler AG, Hummel M, Schenker M, Bonifacio E. Autoantibody appearance and risk for development of childhood diabetes in offspring of parents with type 1 diabetes: the 2-year analysis of the German BABYDIAB Study. Diabetes. 1999;48(3):460-468.
14. Larsson HE. A Swedish approach to the prevention of type 1 diabetes. Pediatr Diabetes. 2016;17(Suppl 22):73-77.
15. Kupila A, Muona P, Simell T, et al; Juvenile Diabetes Research Foundation Centre for the Prevention of Type 1 Diabetes in Finland. Feasibility of genetic and immunological prediction of type 1 diabetes in a population-based birth cohort. Diabetologia. 2001;44(3):290-297.
16. Li Z, Veijola R, Koski E, et al; T1DI Study Group. Supplemental material for Childhood Height Growth Rate Association with the Risk of Islet Autoimmunity and Development of Type 1 Diabetes. Posted February 2, 2022. https://doi.org/10.5281/zenodo.5948373
17. Erlich H, Valdes AM, Noble J, et al; for the Type 1 Diabetes Genetics Consortium. HLA DR-DQ Haplotypes and genotypes and type 1 diabetes risk: analysis of the type 1 diabetes genetics consortium families. Diabetes. 2008;57(4):1084-1092.
18. Li Z, Toppari J, Lundgren M, et al; T1DI study group. Imputing Longitudinal Growth Data in International Pediatric Studies: Does CDC Reference Suffice? AMIA Annu Symp Proc. 2022;2022:754-762.
19. American Diabetes Association. Diagnosis and classification of diabetes mellitus. Diabetes Care. 2013;37(Supplement 1):S81-S90.
20. Grambsch PM, Therneau TM. Proportional hazards tests and diagnostics based on weighted residuals. Biomетrika. 1994;81(3):515-526.
21. Ilonen J, Kiviniemi M, Lempainen J, et al; Finnish Pediatric Diabetes Register. Genetic susceptibility to type 1 diabetes in childhood - estimation of HLA class II associated disease risk and class II effect in various phases of islet autoimmununity. Pediatr Diabetes. 2016;17(Suppl 22):8-16.
22. Ziegler A-G, Kick K, Bonifacio E, et al; Forlida Study Group. Yield of a public health screening of children for islet autoantibodies in Bavaria, Germany. JAMA. 2020;323(4):339-351.
23. Tse WY, Hindmarsh PC, Brook CG. The infancy-childhood-puberty model of growth: clinical aspects. Acta Paediatr Scand Suppl. 1989;356:43-44; discussion 44.
24. Paccard D, Nucci AM, Cuthbertson D, et al; TRIGR investigators. Association between family history, early growth and the risk of beta cell autoimmunity in children at risk for type 1 diabetes. Diabetologia. 2021;64(1):119-128.
25. Beyerlein A, Thiering E, Pfueger M, et al. Early infant growth is associated with the risk of islet autoimmunity in genetically susceptible children. Pediatr Diabetes. 2014;15(7):534-542.
26. Yassouridis C, Leisch F, Winkler C, Ziegler A-G, Beyerlein A. Associations of growth patterns and islet autoimmunity in children with increased risk for type 1 diabetes: a functional analysis approach. Pediatr Diabetes. 2017;18(2):103-110.
27. Shapiro MR, Wasserfall CH, McGrail SM, et al. Insulin-like growth factor dysregulation both preceding and following type 1 diabetes diagnosis. Diabetes. 2020;69(3):413-423.
28. Peet A, Hamalainen A-M, Kool P, Ilonen J, Knip M, Tillmann V; DIABIMMUNE Study Group. Early postnatal growth in children with HLA-conferred susceptibility to type 1 diabetes. Diabetes Metab Res Rev. 2014;30(1):60-68.
29. Nambam B, Schatz D. Growth hormone and insulin-like growth factor-1 axis in type 1 diabetes. Growth Horm IGF Res. 2018;38:49-52.
30. Eizirik DL, Colli ML. Revisiting the role of inflammation in the loss of pancreatic β-cells in TIDM. Nat Rev Endocrinol. 2020;16(11):611-612.