

Adult height determinants in cystic fibrosis

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Prepuberal insulin secretory indices are long-term predictors of short adult stature in cystic fibrosis

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Abstract

Objective: Diabetes is a frequent comorbidity in cystic fibrosis (CF), related to multiple unfavorable outcomes. During the progression of β -cell dysfunction to diabetes, insulin deficiency could possibly reduce the anabolic support to grow even in the absence of significant glycemic derangements. To test this hypothesis, we evaluated whether prepuberal insulin secretory indices are independent predictors of adult height. *Design:* Observational cohort study.

Research design and methods: A longitudinal analysis of 66 CF patients (33 females) from an ongoing cohort received at prepuberal age (median age of 12 years) modified 3-h oral glucose tolerance tests with 30-min insulin and C-peptide sampling, modeling of insulin secretory and sensitivity parameters, anthropometric evaluation. The latter was repeated when adults after a median follow-up of 9 years.

Results: In alternative models, we found a positive association with either basal insulin secretion (mean 0.22, 95% CI 0.01, 0.44 z-scores) or prepuberal β -cell glucose sensitivity (mean 0.23, 95% CI 0.00, 0.46 z-scores) and adult height, while total insulin secretion was negatively related to adult height (mean -0.36, 95% CI -0.57, -0.15 z-scores or mean -0.42, 95% CI -0.69, -0.16 z-scores, respectively). The high total insulin secretion of low adult height patients was mainly due to late (>60 min) secretion and was associated with a worse glucose response during OGTT.

Conclusions: Abnormal insulin secretion associated with high glucose response during OGTT predicts a decrease in adult height z-score. Our results suggest that insulin secretory defects in CF affect growth prior to the development of fasting hyperglycemia.

Key Words

- cystic fibrosis
- oral glucose tolerance test
- ▶ insulin secretion
- ▶ beta cell glucose sensitivity

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Introduction

Insulin secretory defects in cystic fibrosis (CF) were initially shown in small-sized studies that employed various sophisticated techniques such as the hyperglycemic clamp (1, 2, 3) and intravenous glucose tolerance test (IVGTT) (4, 5, 6, 7, 8). In larger studies based on the analysis of insulin and C-peptide responses during oral glucose tolerance tests (OGTTs). These secretory defects were found to be frequent, also in normotolerant CF patients who compensate with an increased insulin sensitivity (9, 10, 11, 12, 13) and to worsen with advancing age,





as we found in the population followed at the CF Center in Milan (14), in which nomograms for OGTTderived insulin secretory parameters were produced to explore their clinical associations and significance in prospective studies.

The progressive deterioration in insulin secretion eventually leads to CF-related diabetes (CFRD), whose prevalence climbs toward 50% in adult patients (15). CFRD, in turn, has been associated with other multiple unfavorable outcomes, ranging from deterioration of nutritional status, pulmonary function and increased mortality (15). Specifically, CFRD prevents patients from achieving their full height genetic potential (16), which is a severe consequence per se, being related to mortality (17). However, linear growth reduction precedes CFRD (16, 18) and cannot be reversed with intensive insulin treatment once diabetes is diagnosed (19), suggesting that insulin secretory defects may play a negative role.

The concept that subclinical insulin secretory defects may affect linear growth independently from hyperglycemia has been suggested by a small-sized study in CF (20), but further evidence arises from studies in type 1 diabetes and in healthy subjects. According to mechanistic studies, insulin is one of the main regulators of growth hormone (GH)/insulin-like growth factor 1 (IGF-1) axis (21), frequently involved in growth failure associated with several chronic conditions including CF. In the past, subjects with uncontrolled type 1 diabetes may have suffered dwarfism associated with Mauriac Syndrome (22). Currently, the degree of glycemic control may still impact linear growth (23, 24), but an independent protective effect of residual β cell function has been documented (25). Furthermore, an earlier work in healthy adult males (26) showed that insulin responses to glucose are associated with height.

Evidence mentioned above supports the idea that the slowly progressing β cell dysfunction in CF possibly reduces the anabolic support to grow even in the absence of significant glycemic derangements. To test this hypothesis, we evaluated whether prepuberal insulin secretory indices measured during OGTT are independent predictors of adult height, measured approximately 10 years later.

Research design and methods

Design

This was a prospective cohort study with a mean follow-up of 9 years.

Setting

Between 2003 and 2013, the CF Center of Milan offered participation in this study to all CF patients at prepuberal age and regularly followed up with clinical and laboratory assessment, including an annual OGTT. The study was approved by the Milano Area 2 ethical committee (n. 452, 556 and 777). Patients were informed about the purpose of the study and gave permission to include their clinical and laboratory data in this research.

Patient inclusion criteria

OGTTs were performed between the ages of 8.5 and 12.9 years in CF patients of both sexes who were prepuberal (according to menarche age and or Tanner Stage), clinically stable in the previous 3 weeks (absence of major clinical events including pulmonary exacerbations, no change in their habitual treatment regimen including the introduction of antibiotics or steroids) and had not previously received a CFRD diagnosis or treatment with insulin or oral hypoglycemic agents in the previous 6 months. Patients who had received or were listed for a lung or liver transplant began taking GH replacement, modulator therapy, or corticosteroids treatment at any time prior to the end of follow-up were also excluded. OGTTs were performed between 2003 and 2013. All patients who reached 18 years of age and had their height recorded at the time of the study were included; the last available height was used and the end of follow-up was set at that date. Sixty-six patients complied with these criteria and were included in this analysis.

Assessment at baseline

All subjects received a 3-h OGTT (1.75 g/kg, maximum 75 g) sampling at baseline and at 30-min intervals the plasma glucose, serum insulin and C-peptide concentrations. Pulmonary function (FEV1% and FVC) and nutritional status (height, weight and BMI) were also measured, as well as parents' height from which mid-parental height was calculated.

Analytical methods

Plasma glucose was measured on fluoride plasma samples (Gluco-quant; Roche/Hitachi analyzer; Roche Diagnostics), and the other analytes were measured by commercial assays (ECLIACobas C6000; Roche Diagnostics).





OGTT analysis

Based on the plasma glucose concentrations, patients were assigned to one of the following categories of glucose tolerance (15): normal glucose tolerance, normal glucose tolerance with impaired fasting glucose, indeterminate glucose tolerance, impaired glucose tolerance, cystic fibrosis-related diabetes without fasting hyperglycemia and cystic fibrosis-related diabetes with fasting hyperglycemia. Insulin secretory and insulin sensitivity parameters were derived from the modeling of glucose, insulin and C-peptide profiles as we previously described (14, 27, 28).

Specifically, insulin secretion was calculated from glucose and C-peptide concentrations by regularized least squares (20). Regularization involves the choice of smoothing factors that were selected to obtain glucose and C-peptide model residuals with SDs close to the expected measurement error (~1% for glucose and ~4% for C-peptide). Insulin secretion rates were calculated from the model every 5 min. The integral of insulin secretion during the 3-h OGTT was denoted as total insulin output. β -cell glucose sensitivity represents the responsiveness of insulin secretion to glucose.

Insulin sensitivity was determined in the basal state using the homeostasis model assessment from basal levels (HOMA) index and during the OGTT with the 3-h oral glucose tolerance test-based index of insulin sensitivity (OGIS) index (29).

Insulin clearance was calculated in the fasting state as the ratio between fasting insulin secretion and fasting insulin concentration and during the OGTT as the ratio between the integral of insulin secretion and that of insulin concentration.

The above parameters were also evaluated according to reference values for sex and age recently developed in the CF Center of Milan (14).

Statistical analysis

Continuous variables are reported as median (25th, 75th percentile), and categorical variables are reported as count (percentages). Comparisons across groups were carried out using the Wilcoxon rank-sum test with continuity correction (for continuous variables) or Pearson's chi-squared test or Fisher's exact test (for categorical variables).

Ordinary least square linear models were used to describe the relationship between indices of insulin and glucose response to OGTT and adult height z-score. Models were adjusted for sex (categorical; 0=female, 1=male), pancreatic insufficiency (defined by fecal elastase (30))

(dichotomous; 0=pancreatic sufficiency, 1=pancreatic insufficiency), target height z-score. Different models were fitted, each including a single index of insulin and glucose response to OGTT as predictors: β -cell glucose sensitivity, integral of total insulin secretion (adjusted for basal insulin secretion), OGTT insulin clearance (adjusted for basal insulin clearance), HOMA index and OGIS index, and OGTT glucose AUC (adjusted for basal glucose). A final complete model including all the indices was also fitted. The association between OGTT-derived indices and adult height was expressed as average predicted changes in height z-score for one interquartile increase of the index.

Statistical analyses were performed in R 4.1.0 (31).

Results

Baseline (prepuberal) patient characteristics are reported in Table 1. Of the subjects included, 57 (86%) were pancreatic insufficient (29 females and 28 males) and all were consuming pancreatic enzymes replacement therapy. None of the patients were assuming CFTR modulator therapy. Z-scores of weight, height and BMI were not significantly different between sexes, but they were reduced when compared to the general population. Most patients (68%) had normal glucose tolerance but β cell glucose sensitivity and insulin secretion were lower in males (although differences in β cell glucose sensitivity did not reach statistical significance).

Table 2 reports differences in anthropometric parameters and respiratory status evaluated at prepuberal and adult ages. Median follow-up was 9 years, during which 9/66 (14%) patients became diabetic and were on insulin at the end of follow-up (all of them were among pancreatic insufficient patients, 9/57 (16%)). Adult height z-score was lower than target height z-score by 0.22 z-scores (CI: 0.04, 0.40; P = 0.017) and 11 (17%) patients were underweight at the end of follow-up. Adult FEV1 (% of predicted) was -13% (CI: -20, -6.8; P < 0.001) compared to prepuberal age.

Table 3 shows results from the predictive models of adult height z-score. No significant differences were found between sexes and patients with and without pancreatic insufficiency. The model testing the association between glucose response and adult height showed a negative association for OGTT glucose AUC adjusted for basal glucose, although the association did not reach statistical significance. The insulin secretion model showed a positive association of adult height z-score with basal insulin secretion and a negative association with total insulin

(†)



Table 1Baseline patient characteristics by sex.

Characteristic	Overall, <i>n</i> = 66	Female, <i>n</i> = 33	Male, <i>n</i> = 33	P-value
Age (years)	11.95 (10.92, 12.49)	11.51 (10.67, 12.45)	12.25 (11.25, 12.59)	0.054
CFTR mutation				0.7
F508del homozygous	25 (38%)	11 (33%)	14 (42%)	
F508del heterozygous	27 (41%)	15 (45%)	12 (36%)	
Other	14 (21%)	7 (21%)	7 (21%)	
Meconium ileus	11 (17%)	5 (15%)	6 (18%)	0.7
Pancreatic insufficient	57 (86%)	29 (88%)	28 (85%)	>0.9
Liver disease	24 (36%)	10 (30%)	14 (42%)	0.3
Pseudomonas aeruginosa				0.8
No	19 (29%)	11 (33%)	8 (24%)	
Intermittent	43 (65%)	20 (61%)	23 (70%)	
Chronic	4 (6.1%)	2 (6.1%)	2 (6.1%)	
Burkholderia cepacia	1 (1.5%)	0 (0%)	1 (3.0%)	>0.9
Height z-score	-0.39 (-0.62, -0.17)	-0.47 (-0.69, -0.16)	-0.33 (-0.53, -0.19)	0.3
Weight z-score	-0.48 (-0.69, -0.19)	-0.51 (-0.73, 0.28)	-0.37 (-0.61, -0.13)	0.3
Body mass index z-score	-0.42 (-0.62, -0.12)	-0.46 (-0.68, -0.13)	-0.37 (-0.58, -0.11)	0.5
Body mass index category				>0.9
Underweight	0 (0%)	0 (0%)	0 (0%)	
Normal weight	66 (100%)	33 (100%)	33 (100%)	
Overweight	0 (0%)	0 (0%)	0 (0%)	
Obesity	0 (0%)	0 (0%)	0 (0%)	
FEV1 (% of predicted)	87 (81, 95)	87 (82, 95)	87 (81, 97)	>0.9
FVC (% of predicted)	95 (87, 101)	95 (86, 100)	95 (87, 101)	0.7
HbA1C (%)	5.90 (5.50, 6.20)	5.80 (5.60, 6.60)	5.90 (5.40, 6.10)	0.4
Glucose tolerance				0.039
Normal glucose tolerance	45 (68%)	18 (55%)	27 (82%)	
Normal glucose tolerance with	0 (0%)	0 (0%)	0 (0%)	
impaired fasting glucose				
Indeterminate glucose tolerance	10 (15%)	8 (24%)	2 (6.1%)	
Impaired glucose tolerance	8 (12%)	4 (12%)	4 (12%)	
Cystic fibrosis related diabetes	3 (4.5%)	3 (9.1%)	0 (0%)	
without fasting hyperglycemia				
Cystic fibrosis related diabetes with fasting hyperglycemia	0 (0%)	0 (0%)	0 (0%)	
Homeostatic model assessment for insulin resistance	1.09 (0.62, 1.56)	1.10 (0.51, 1.46)	1.09 (0.64, 1.58)	0.7
Oral glucose insulin sensitivity index (mL min ⁻¹ m ⁻²)	539 (497, 621)	560 (491, 621)	529 (507, 621)	0.9
β cell glucose sensitivity (pmol×min ⁻¹ ×m ⁻² ×mM ⁻¹)	79 (50, 106)	89 (52, 124)	71 (45, 94)	0.10
Basal insulin secretion (pmol min ⁻¹ m ⁻²)	82 (59, 96)	90 (67, 105)	72 (49, 90)	0.007
Integral of OGTT insulin secretion (nmol m ⁻²)	55 (43, 68)	57 (50, 70)	46 (37, 64)	0.013
Basal insulin clearance (L min ^{-1} m ^{-2})	2.22 (1.79, 2.95)	2.43 (2.03, 3.64)	2.19 (1.64, 2.46)	0.014
OGTT insulin clearance (L min ^{-1} m ^{-2})	1.66 (1.30, 1.97)	1.75 (1.44, 2.51)	1.61 (1.27, 1.85)	0.10
Basal glucose (mg dL)	80 (72, 89)	80 (76, 89)	81 (71, 89)	0.7
OGTT glucose AUC (mg dL min ⁻¹)	20,452 (18,375, 24,896)	21,615 (18,825, 27,510)	20,400 (18,120, 22,755)	0.2
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Bold indicates statistical significance.

secretion, both statistically significant. In the model including all OGTT parameters, β -cell glucose sensitivity was positively associated with adult height, while total insulin secretion maintained the negative association.

Figure 1 shows insulin secretion profiles for patients with an adult height below the lower tertile and those above the upper tertile. Patients with a low adult height

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reached peak insulin secretory rate significantly later than

patients above the upper tertile of adult height (mean peak

minute 70 vs 47; mean difference 22 min; 95% CI 8, 37;

P = 0.009), and secreted more insulin in the second and

third hour of the test (total insulin secretion AUC between

60 and 180 OGTT minutes 41458 vs 26154; mean difference

15305; 95% CI 7316, 23292; $P \le 0.001$).



Table 2 Differences for age, nutritional and pulmonary status between follow-up (adult age) and baseline (prepuberal age).Median follow-up of 9 years.

Characteristic	Baseline, <i>n</i> = 66	Follow-up, <i>n</i> = 66	Difference	95% CI	<i>P</i> -value
Height z-score	-0.39 (-0.62, -0.17)	-0.30 (-0.78, 0.10)	0.09	0.01, 0.20	0.078
Weight z-score	-0.48 (-0.69, -0.19)	-0.31 (-0.59, 0.08)	0.20	0.09, 0.30	<0.001
Body mass index z-score	-0.42 (-0.62, -0.12)	-0.13 (-0.42, 0.14)	0.25	0.15, 0.35	<0.001
FEV1 (% of predicted)	87 (79, 101)	79 (55, 96)	-13	-20, -6.8	<0.001

Bold indicates statistical significance.

Discussion

The main finding of this study is that insulin secretory indices and possibly OGTT glucose response measured at prepuberal age represent long-term predictors of adult height in patients with CF.

We selected a subgroup of patients followed at the CF center of Milan, in a narrow age range and all in the prepubertal stage. The OGTTs were performed when CFTR modulators were not yet introduced in clinical practice. We have previously shown that the distribution of insulin secretory parameters measured by OGTT is shifted toward lower values, compared to healthy control subjects, also in normotolerant CF patients (14).

Patients' height and BMI were generally reduced already at the prepuberal age (Table 2), as previously described in the Italian CF population (32). We evaluated the effect of the disease on adult height adjusting for each patient's target height (mid-parental height), to approximately account for maximum adult height potential. We found a median interquartile range (IQR) difference of -0.22(-0.51, 0.09) z-scores between target height and achieved adult height, with no significant difference between sexes (P = 0.6). Overall, our analysis supports the idea that insulin secretory indices are related to defective growth and are predictive of a reduced adult height.

Several factors may concur to explain this relationship. In our study, adult height was positively related to β cell glucose sensitivity, but it was negatively related to total insulin secretion. In the context of CF, β -cell glucose sensitivity is an indicator of a healthy endocrine pancreas that could have favored optimal growth. On the other hand, the association with total insulin secretion was negative, although with a wide CI. While limitations in the sample size surely contributed to this uncertainty, in part it may be due to modeling the relationship between insulin secretion and adult height as linear (another limitation due to the limited sample size). It is plausible, and it was our original hypothesis, that higher insulin should promote growth through the GH/IGF-1 axis, but our data suggest that this is limited to higher insulin secreted early during the OGTT. A physiological response to an OGTT sees a timely and early elevation of insulin levels that peaks between 30 and 60 OGTT min (33). In our sample, the highest total insulin secretion levels were not obtained through an elevation of insulin secretion in the early stages of OGTT, but through a delayed response that compensated for increased glucose concentration in the late part of the OGTT.

We did not provide a mechanistic explanation for how a delayed and lately increased insulin secretion may have negatively affected growth. To date, the available studies are not conclusive but may support several speculations. The GH/IGF-1 axis is one of the main regulators of linear growth in children (34). GH promotes growth indirectly by inducing the release of insulin-like growth factor 1 (IGF-1) from the liver and the local secretion of IGF-1 from cartilage cells in the growth plates of the long bones, but it also displays a direct, IGF-1-independent, effect (35). Sustained insulin levels stimulate IGF-1 secretion (36), but may also lead to the inhibition of GH secretion (37) and signaling (38), playing an overall detrimental effect on growth. It is therefore plausible that a pulsatile rather than continuous insulin secretion may be required for optimal growth.

The negative association between adult height and total insulin secretion was consistent with what we previously noted (13), i.e. that hyperglycemia can stimulate total insulin output (and insulin and C-peptide AUCs) to levels similar or even greater than in healthy subjects. This finding has practical implications, because the absolute insulin secretion rates or the insulin or C-peptide concentrations or AUCs may not accurately describe β -cell function defects.

Indeed, we showed in Fig. 1 that the main determinant of a high total insulin secretion in subjects below the lower tertile of adult height z-score was a late (>60 min) insulin secretion. Our data suggest that an increased total insulin secretion that is delayed because of a failing betacell sensitivity to glucose, not only is associated with a deteriorated glucose tolerance but is also associated with growth failure.

Our data do not exclude an adverse role of hyperglycemia on linear growth. High glucose levels in





	Ċ	Glucose model	del	-	Insulin model	_	βcell	β cell sensitivity model	model	Insulin	Insulin resistance model	model	Insuli	Insulin clearance model	model	0	Complete model	del
Characteristic	Beta	95% CI	P-value	Beta	95% CI	P-value	Beta	95% CI	P-value	Beta	95% CI	P-value	Beta	95% CI	P-value	Beta	95% CI	P-value
(Intercept)	0.05	-0.81, 0.91	>0.9	0.23	-0.33, 0.79	0.4	-0.43	-1.0, 0.11	0.12	-0.59	-1.6, 0.39	0.2	-0.05	-0.46, 0.36	0.8	-0.20	-2.2, 1.8	0.8
Jex Female	ı	I	ı	ı	I	ı	ı	ı	ı	ı	I	ı	ı	ı	ı	ı	ı	ı
Male	-0.07	-0.33, 0.18	0.6	-0.11	-0.36, 0.14	0.4	0.01	-0.24, 0.27	>0.9	-0.03	-0.29, 0.22	0.8	-0.07	-0.34, 0.21	0.6	-0.16	-0.45, 0.12	0.3
Pancreatic insufficiency	0.01	-0.38, 0.39	>0.9	-0.09	-0.44, 0.26	0.6	0.01	-0.39, 0.40	>0.9	-0.08	-0.46, 0.30	0.7	-0.07	-0.45, 0.32	0.7	0.05	-0.34, 0.43	0.8
Target height	0.63	0.34, 0.93	<0.001	0.66	0.38, 0.93	<0.001	0.64	0.35, 0.94	<0.001	0.64	0.34, 0.94	<0.001	0.65	0.33, 1.0	<0.001	0.65	0.35, 0.95	<0.001
z-score Basal glucose (IQR	0.06	-0.11, 0.22	0.5													-0.01	-0.22, 0.21	>0.9
increase)	016		0,060													010	100110	0
UGLI Blucose AUC (IQR increase)	-0.10	-0.32, 0.01	000.0													0.10	-0.11, 0.31	0. 4.
Basal insulin secretion (IQR increase)				0.22	0.01, 0.44	0.039										0.06	-0.25, 0.37	0.7
Integral of OGTT insulin secretion				-0.36	-0.57, -0.15	0.001										-0.42	-0.69, -0.16	0.002
uck increase) β-cell glucose sensitivity (IQR increase)							0.12	-0.05, 0.28	0.2							0.23	0.00, 0.46	0.046
Homeostatic										0.07	-0.07, 0.22	0.3				0.12	-0.09, 0.33	0.2
assessment for insulin resistance																		
(IQR increase) Oral glucose										0.08	-0.11, 0.26	0.4				0.03	-0.20, 0.26	0.8
insulin sensitivity index (IQR increase)																		
Basal insulin clearance (IQR													0.00	0.00, 0.00	0.4	0.00	0.00, 0.00	0.5
increase)															c 1 0			с С
clearance (IQR increase)													-0.04	-0.09, 0.01	01.0	000	-0.00, 0.02	7.0

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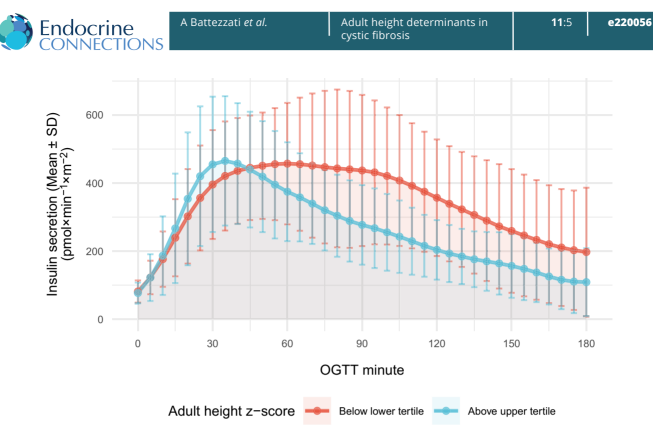


Figure 1

Prepuberal insulin secretion profiles of the patients in the extreme tertiles of adult height.

the late part of OGTT are probably the driver for this late insulin secretion and in the simple model of glucose response, OGTT glucose response, but not basal glucose, was negatively associated with adult height, at a nearly significant level. It is already known that CFRD has a large effect on nutritional status (39) and that hyperglycemia, consequent to insulin secretory defects and insulin resistance, could be a causative factor as it has been associated with defective linear growth (23). In theory, hyperglycemia could be responsible for either nutrient renal losses or reduced immune defenses especially in the respiratory tract (40), even though hyperglycemia is generally not accompanied by reduced linear growth in the absence of insulin secretory defects (as in type 2 diabetes).

We could not determine if the negative association with adult height was due to the abnormal insulin response or due to hyperglycemia. Our complete model seems to favor the differences in insulin secretion as an explanatory variable for differences in adult height, while glucose response could not explain an additional amount of variance in the outcome (although this may also be due to sample size limitations).

Insulin therapy is usually effective in glucose control, but only approximate insulin secretion from the pancreas; the fact that it cannot completely normalize suboptimal growth, both in CF (21) and type 1 diabetes

(41), may suggest that an optimal insulin secretion is more important that glucose control for growth. Furthermore, several findings stand against the idea that hyperglycemia of diabetes has driven the observed relationship of insulin secretory indices with adult height. In agreement with what has been already described in our group (42), future diabetics were already in shorter stature at prepubertal age prior to the development of CFRD, which suggests that overt hyperglycemia is not the cause of short stature. In addition, in this study, when accounting for insulin secretory parameters, neither insulin sensitivity nor OGTT glucose response was associated with height.

Finally, more severe genetic defects may promote a greater inflammatory phenotype which may be associated with reduced linear growth and even growth arrest (43). Greater inflammation would cause greater insulin resistance (44), further exacerbated by the use of steroid drugs. Against this hypothesis, we did not find a negative association between height with insulin sensitivity. This trait is highly variable over time in CF (45), thus it is very difficult for insulin resistance to be a long-term predictor of adult height. Furthermore, we have shown that insulin sensitivity and β cell function may be inversely related to CF (13). Also for this reason it is suggested that insulin secretory defects are related to linear growth independently from the inflammatory phenotype in CF patients.





If confirmed, this association is likely to be clinically relevant. First, height is an independent predictor of mortality in CF (17). Secondly, insulin secretory defects can be treated (intervention studies remain to be done) but they are presently not until the diagnosis of CFRD (15). Several trials have tested the effects of insulin administration in the absence of hyperglycemia, but the evidence is inconclusive (46). Moran (47) and Minicucci (48) performed controlled studies and observed effects on BMI, however for their relatively short duration and small size they were not powered to detect effects on height. Finally, this is a natural history study that may set a standard to compare the effect of emerging CFTR modulator therapies.

The main strengths of this study are the inclusion of patients in a strict age range prior to puberty, the measurement of parameters of β-cell function ranging from the simplest ones (insulin concentrations) to more sophisticated ones (*β*-cell glucose sensitivity, based on c-peptide and unaffected by hepatic insulin clearance), and in the prospective design that allowed a long-term follow-up. The main weakness of this study relates to the limited sample size of patients followed at a single center. The number of patients who reached 18 years of age and had their height recorded at the time of the study resulted in the sample size, and no a priori calculation of sample size was performed. While this study may provide an early signal of the association of prepuberal insulin secretory indices with short adult stature in cystic fibrosis, these findings should be confirmed in larger studies. Furthermore, the GH/IGF-1 axis was not explored.

In conclusion, we have followed for almost 10 years a cohort of CF patients who had received an OGTT and measured glucose and insulin secretory responses and insulin sensitivity. Our data indicate that final height is compromised in Cystic Fibrosis. Reduced height is not related to a reduction in the absolute amount of insulin secreted in response to glucose. Nonetheless, reduced height is related to a reduced beta-cell glucose sensitivity, a defect in the quality of insulin secretion that causes a delayed and lately sustained insulin response to glucose. This study does not provide a mechanistic explanation but suggests that altered insulin dynamics may play a role in the reduction of adult height. The possible interference of altered insulin secretory dynamics in the complex regulation of the GH/IGF-1 axis remains to be clarified.

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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Author contribution statement

A Ba wrote the manuscript. A Bif researched data. A Bis researched data and reviewed/edited the manuscript. A F performed statistical analysis and reviewed/edited the manuscript. C C reviewed/edited the manuscript. E N researched data. G A reviewed statistical analysis and reviewed/edited the manuscript. G B reviewed statistical analysis and reviewed/edited the manuscript. G D C researched data. S B reviewed/edited the manuscript.

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