Sirolimus: Efficacy and Complications in Children With Hyperinsulinemic Hypoglycemia: A 5-Year Follow-Up Study

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Introduction: Sirolimus, a mammalian target of rapamycin inhibitor, has been used in congenital hyperinsulinism (CHI) unresponsive to diazoxide and octreotide. Reported response to sirolimus is variable, with high incidence of adverse effects. To the best of our knowledge, we report the largest group of CHI patients treated with sirolimus followed for the longest period to date.

Methods: Retrospective study of CHI patients treated with sirolimus in a tertiary service and review of the 15 publications reporting CHI patients treated with mammalian target of rapamycin inhibitors. Comparison was made between the findings of this study with those previously published.

Results: Twenty-two CHI patients treated with sirolimus were included in this study. Twenty showed partial response, one showed complete response, and one was unresponsive. Five of the partially/fully responsive patients had compound heterozygous \textit{ABCC8} mutations and five had heterozygous \textit{ABCC8} mutations. A total of 86.4% (19/22) developed complications, with infection being the most frequent (17/22), of which 11 were of bacterial etiology, followed by persistent diarrhea (3/22) and hyperglycemia (2/22). Seventeen patients stopped sirolimus: 13 from infections; 2 from hyperglycemia; and 2 from alternative treatment (lanreotide) response. Compared with data previously published, our study identified a higher number of partially sirolimus-responsive CHI cases, although the high rate of complications while on this medication limited its potential usefulness.

Conclusion: Sirolimus candidates must be carefully selected given its frequent and potentially life-threatening side effects. Its use as a short-term, last-resort therapy until normoglycemia is achieved with other agents such as lanreotide could avoid pancreatectomy. Further studies evaluating the use of sirolimus in patients with CHI are required.

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Freeform/Key Words: sirolimus, mTOR inhibitors, hyperinsulinemic hypoglycemia, \textit{ABCC8} gene, side effects

Abbreviations: BWS, Beckwith-Wiedemann syndrome; CHI, congenital hyperinsulinaemic hypoglycaemia; DM, diabetes mellitus; HI, hyperinsulinemic hypoglycemia; PET, positron emission tomography; PI3K, phosphatidylinositol 3-kinase; mTOR, mammalian target of rapamycin; mTORC, mammalian target of rapamycin complex.

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Congenital hyperinsulinemic hypoglycemia (CHI) is characterized by the inappropriate, excessive secretion of insulin from the pancreatic β-cell and is the most frequent cause of persistent hypoglycemia in infancy [1]. CHI is a heterogeneous disorder with respect to clinical presentation, imaging, histology, and genetics. To date, mutations in at least 10 different genes (ABCC8, KCNJ11, GLUD1, GCK, HADH, SLC16A1, HNF4A, HNF1A, PMM2, PGM1) have been reported in the genetic etiology of CHI [2–5]. Mutations in ABCC8 and KCNJ11 are associated with severe CHI that is unresponsive to medical treatment with diazoxide and octreotide. There are three main histological types of CHI, each with a specific genetic etiology: focal, diffuse, and atypical [6]. In the past, patients with medically unresponsive forms of CHI were treated with subtotal (95% to 98%) pancreatectomy. In 2014, Senniappan et al. [7] reported the use of the mammalian target of rapamycin (mTOR) inhibitors, such as sirolimus, as an alternative therapeutic strategy in CHI patients unresponsive to therapy with diazoxide or octreotide.

Sirolimus (formerly known as rapamycin) inhibits the mTOR pathway, potentially limiting the production of insulin from β cells [8]. mTOR is a serine/threonine kinase regulated by phosphatidylinositol 3-kinase (PI3K) that integrates a complex cascade of signals that regulate cell growth, metabolism, proliferation, and survival in response to extracellular nutritional and environmental signals [9]. Although the exact underlying mechanisms are unclear, mTOR-signaling activation has been implicated in the regulation of β-cell mass, cell proliferation, and size. Integration of genomic, proteomic, and systems biology analysis suggests that the mTOR pathway shows a variable but constant activation and overexpression in acinar cells of patients with CHI [10] and in insulin-producing islet cells [11]. There are >300 interactions between mTOR-related molecules and 8 CHI genes (GLUD1, SLC16A1, HADH, KCNJ11, ABCC8, HNF1A, GCK, HNF4A), which were identified by the use of a CHI network [12]. Sirolimus has been shown to have the following actions: (i) inhibition of mTOR complex 1 (mTORC1), which inhibits β-cell mass; (ii) inactivation of mTORC2 (with prolonged exposure), which induces chronic insulin resistance [13]; (iii) possible reduction in the number of insulin receptors via mTOR inhibition on pancreatic β cells, leading to decreased insulin secretion [14]; (iv) in vivo, sirolimus alters the glucose-stimulated secretion of insulin [15]; (v) in cultures, sirolimus reduces the number of ductal cells, a potential source of islet cells [15]; and (vi) in mice, sirolimus induces a 50% decrease in β-cell mass by apoptosis [16].

Conversely, two publications dismiss the use of mTOR inhibitors in CHI: (i) in islets of two children with diffuse CHI, cell proliferation remained high with no reduction after mTOR inhibitor treatment [17]. The caveat of this study is that both patients had previously not responded to sirolimus and the study was not performed immediately after stopping sirolimus. (ii) An in silico transcriptome array study showed no substantial association between mTOR signaling and the focal form of CHI [17].

Adverse effects of sirolimus include stomatitis [18], increased risk of infection, immunosuppression, abnormalities in renal function, fatigue, pneumonitis, episodes of transient elevation of aminotransferase concentrations [17], and elevation of triglyceride concentrations [7]. These adverse effects are reversible with dose reduction. Studies in kidney transplant recipients [19] have suggested that sirolimus used in the long term may be diabetogenic; this complication was recently published in a patient with CHI [20]. The mechanisms by which sirolimus induces diabetogenic effects may include: (i) impaired β-cell proliferation by impairing AKT activation and signaling (IGF-1/mTOR/Akt pathway) [13]; (ii) direct β-cell toxicity; (iii) negative modulation of insulin action in peripheral tissues including skeletal muscle, liver, and white fat; (iv) direct insulin resistance from ectopic triglyceride deposition [18]; and (v) glucose intolerance by upregulation of liver gluconeogenesis [21].

Growing bodies of evidence from CHI patients treated with sirolimus suggest that sirolimus use meets with limited success and carries a greater risk of side effects than any of the standard medications for CHI [17]. Therefore, it must be used with caution in children with severe CHI, and only after discussion with families about its side effects and potential complications [17].
The aim of this study was to assess the efficacy of mTOR inhibitors (glycemic control, fasting tolerance, and weaning of supportive medical therapy) and to identify adverse reactions/complications (frequency and severity) related to sirolimus use in a cohort of patients with severe forms of CHI.

1. Methods

A. Study Design and Participants

The current study is a single-center retrospective analysis of data from 22 children with severe CHI, unresponsive to maximum doses of diazoxide and octreotide, who were treated with sirolimus between the years 2013 and 2018. Diagnosis and management in all cases was conducted within the Endocrinology Department of Great Ormond Street Hospital for Children NHS Trust, London, United Kingdom. This series of patients includes the progress of seven patients previously reported in the literature with CHI managed on sirolimus therapy [7, 20, 22, 23]. Other mTOR inhibitors were not used in this study.

Ethical approval had been obtained before the trial of sirolimus in 2014 by Senniappan et al. [7]. Patients were only begun on sirolimus therapy after informed consent was obtained from parents. The protocol for sirolimus use in CHI patients in our center had been elaborated in combination with our pharmacy. The dose of sirolimus was gradually increased every 4 to 5 days based on blood glucose concentrations, sirolimus plasma concentrations, and side effects. If blood glucose concentrations were stable, the dose of sirolimus was not increased (even if plasma concentrations were below the target range) to avoid possible complications. Complete response to sirolimus was defined as glycemic control achieved exclusively with sirolimus, partial response as glycemic stability obtained with sirolimus concomitantly with another CHI agent, and no response as absent glycemic improvement despite combination of sirolimus with other CHI medications. Glycemic control in individuals with CHI is defined as plasma glucose concentrations maintained between 3.5 and 7.0 mmol/L with avoidance of hypoglycemia (<3.5 mmol/L) episodes, along with the capacity to extend the fasting time (approaching or achieving age-appropriate fasting length) without developing hypoglycemia while generating an increase in ketone bodies and free fatty acids. Patients did not receive any live vaccinations while on sirolimus.

The data collection included the following aspects: characteristics of patients with CHI (presentation of CHI), onset of sirolimus therapy (concomitant management and sirolimus dose and plasma concentrations, duration of treatment), side effects while on sirolimus therapy (type of complication, dose, and plasma concentrations of sirolimus at the time), and eventual patient management (medication ± feed regimen). Data are presented as median (range), where range includes maximum to minimum values. Pathology images were obtained from the Histopathology Department of Great Ormond Street Hospital for Children NHS Trust after pancreatectomy in children who had previously been treated with sirolimus.

Previous publications on the use of mTOR inhibitors in children with CHI were identified in PubMed (last search performed 9 August 2018); results were summarized in this study.

2. Results

Fifteen publications of cases of CHI where mTOR inhibitors had been used were identified. These include a total of 31 children treated in countries in Europe, Middle East, and South East Asia. Nineteen of 31 (61.3%) patients have been reported to have a positive glycemic response to mTOR inhibitors in monotherapy or combined with another CHI medication. Any complication of any nature and severity while on mTOR inhibitor therapy has been reported in 26 of 31 (83.8%). Table 1 outlines these publications and characteristics related to the use of mTOR inhibitors.

The characteristics of the 22 CHI patients included in our study are summarized in Table 2. A total of 90.9% of patients (20/22) were diagnosed with CHI within the first month of
| Reference | No. of Cases | Age at Start | HI Genes, PET, Histology | Prior Unresponsiveness to Other CHI Medications, Doses (Response to Medication Specified) | Max Dose mTOR Inhibitor, mg/m²/d | mTOR Plasma Level, ng/mL | Feeds, IV Fluids, Other Medication During mTOR Trial | mTOR Follow-Up Duration | Complications While on mTOR Inhibitor | Glycemic Control; Mono or Combination | Fasting Length on mTOR Inhibitor | mTORStopped, Reason |
|-----------|--------------|--------------|--------------------------|------------------------------------------------------------------------------------------------|-------------------------------|--------------------------|-----------------------------------------------|--------------------------|--------------------------------------|-------------------------------|---------------------------------|-----------------------------|-----------------------------|
| Soniappan [7] | N = 4, 8-16 wk | 2: maternal het ARCC8 1: no mutations 1: Hom ARCC8 PET: 3 diffuse | Diaz (20 mg/kg/d) + oct (35 μg/kg/h) + ggc (5 μg/kg/h) + IV | NS, 1.5-2.5 mg/m²/d | NS, aimed 5-15 | IV ggc, oct stopped in 3 pt; 1 pt remained on oct | 36-45 wk | 2 pt: mild lipid elevation. 1 pt: transient elevation AST 1 pt: none | Yes | 6-8 h | No (stopped for 3 d in 1 case, reason NS) |
| Shah [22] | N = 1, 9 wk | Comp het ARCC8 | Diaz (15 mg/kg/d) + oct (30 μg/kg/h) + ggc (5 μg/kg/h) + IV | 1.6 | 2.5-12.8 | IV, ggc discontinued; oct decreased | 7 wk | None | Yes; combined with oct (20 μg/kg/d) | 4 h | No, ongoing at age 16 wk |
| Amato LA [24] | N = 1, 7 mo | De novo het ARCC8 | Diaz (20 mg/kg/d) + oct (27 μg/kg/h) + frequent feeds | 6 | NS, aimed 5-15 | Decreased hypercaloric feeds | 3 mo | Growth failure | Yes; combined with oct (16 μg/kg/d) | NS | No, ongoing at age 12 mo |
| Abraham [25] | N = 1, 3 mo | Hom ARCC8 PET: diffuse Histology: diffuse | Diaz (20 mg/kg/d) + oct (60 mg/kg/h) + ggc (5 μg/kg/h) + IV + lanr 10 mg/m + pancreatostomy | 5 | NS, aimed 5-15 | IV, medications discontinued except for lanr maintained until 7 mo of age | 10 mo | C. jejuni gastroenteritis | Yes; during first 4 mo with lanr. Following 6 mo mono | Yes, age 13 mo as CHI resolved |
| M¨eder [8] | N = 1, 8 wk | Comp het ARCC8 PET: diffuse | Diaz (20 mg/kg/d) + oct (35 mg/kg/h) + nif + IV | NS | 9.62 | IV, medications discontinued | 10 mo | NEC, persistent mild elevation of LFTs and lipids Oral aphthosis, mild acne. Transient hematemesis | Yes | Mono | 6 h | No, ongoing at age 1 y |
| Minute [26] | N = 1, 8 YOL | Biallelic het ARCC8 PET: diffuse | Diaz (30 mg/kg/d) + 3 hourly feeds + IM ggc inj | 7mg/m²/d | 12.1 | Diaz reduced to 4 mg/kg/d. No need for ggc | 6 m | Yes; combined with diaz | NS | No, ongoing at age 8 y, 6 mo |
| Kara [27] | N = 2 | Pt 1: hom KCNJ11 Pt 2: hom ARCC8 | Max doses of diaz + oct + ggc + IV; pt 2 had previous 80% pancreatostomy | Pt 1: 2 Pt 1:27.9 | NS | Pt 1: IV, medications discontinued Pt 2: Oct decreased | NS | Mid elevation liver enzymes (pt 2); renal and hepatic failure (pt 1); sepsis (pt 1) | Yes; mono in pt 1; combined with oct (6.6 μg/kg/d) in pt 2 | 10 h in pt 2 NS | Yes, in pt 1 from side effects; ongoing in pt 2 |
| G¨uemes [23] | N = 1, 3 mo | UPD 11p15 (BWS) | Diaz (15 mg/kg/d) + oct (30 μg/kg/h) + ggc (5 μg/kg/h) + IV | 11 | Max 24.7 | IV, ggc discontinued; oct decreased | 19 mo | None | Yes, combined with oct (27 μg/kg/d) | 4 h | No, ongoing at age 22 mo |
| ¨Unal S [28] | N = 1, 35 DOL | Hom KCNJ11 | Diaz (65 mg/kg/d) + oct (40 μg/kg/h) + ggc (10 μg/kg/h) + nif (1 mg/kg/d) | 0.5 | NS | Diaz, nif, ggc stopped; oct decreased to 5 μg/kg/h | 4 mo | None | Yes; combined with oct (5 μg/kg/d) | NS | No, ongoing at age 5 mo |
| Loke KY [29] | N = 1, 15 mo | Paternal het ARCC8 PET: diffuse | Partially responsive to diaz (12.6 mg/kg/d) | 4.4 | 25-30 | Diaz stopped | 15 mo | Oral ulcers, elevation of ALT and CK | Yes; mono | Yes, age 3 y, 3 mo as CHI resolved |

(Continued)
| Reference | No. of Cases; Age at mTOR Started | HI Genetics, PET, Histology | CHI Medications, Doses (Response to Medication Specified) | Max Dose mTOR Inhibitor, mg/m²/d | mTOR Plasma Level, ng/mL | Feeds, IV Fluids, Other Medication During mTOR Trial | mTOR Follow-Up Duration | Complications While on mTOR Inhibitor | Glucose Control Mono or Combination | Fasting Length on mTOR Inhibitor | mTOR Stopped, Reason |
|-----------|----------------------------------|-----------------------------|----------------------------------------------------------|----------------------------------|------------------------|--------------------------------------------------|----------------------|-------------------------------------|----------------------------------|-----------------------------|-------------------|
| Szymanowski [17] | N = 10; 0–57 mo | 1: Hom ABCC8 | 5: Diaz (6–11 mg/kg/d) + SST analogs (23–66 µg/kg/d) | Sirolimus, 6.6–14.1 | 5 pt showed whole or partial glycemic response | 5–16 mo | Triglyceride elevation (5), anemia (1), sepsis (2), varicella zoster (1), exocrine pancreatic insufficiency (2) | Yes; 3 pt (30%) | NS | Yes, in 7 from ineffectiveness in glycemic control; 3 pt still on mTOR after 4, 15, and 16 mo |
| Haliloglu [30] | N = 2; 17–21 DOL | 2: Hom ABCC8 | Pt 1: diaz (15 mg/kg/d) + oct (90 µg/kg/d) | Sirolimus, 3.1 | Pt 1: elevation of AST and ALT | 1–2 mo | Yes; combined with oct (8 µg/kg/d in pt 1 and 23 µg/kg/d in pt 2) | NS | Yes, in both because of hepatitis |
| Al-Balwi [31] | N = 3; 4–5 mo | 2: Hom KCNJ11 | Diaz (20 mg/kg/d) + oct (40 µg/kg/d, 1 pt had a previous pancreatectomy) | 9–14 | No modification of other meds | 47–60 d | Mild increase LDL (2. mild leucosis) | No | NS | Yes, in the 3 pt as no response |
| Dastamani [30] | N = 1; 16.6 YOL | Autosomal dominant ABCC8 | Responsive to diaz (15.9 mg/kg/d) | 4.25 | Diaz dose reduced | 1 mo | Facial cellulitis, abscess, and DM | NS | Yes, at 17.6 years because of DM |
| Korula [32] | N = 1; 29 DOL | Hom KCNJ11 PET: diffuse | Diaz (20 mg/kg/d) + oct (40 µg/kg/d) + nif and IV | 4 | IV fluids, diaz, nif, and oct discontinued | 13 mo | None | Yes; combined with cornstarch | 5 h | No, ongoing at 14 mo of age |

Abbreviations: comp, compound; diaz, diazoxide; DOL, days of life; gcg, glucagon; het, heterozygous; HI, hyperinsulinaemic hypoglycaemia; hom, homozygous; inj, injection; lanr, lanreotide; LFT, liver function test; max, maximum; mono, monotherapy; NEC, necrotizing enterocolitis; nif, nifedipine; NS, not specified; oct, octreotide; pt, patient; SST, somatostatin; UPD, paternal uniparental disomy; YOL, y of life.

aNEC was attributed to octreotide as patient responded well to its discontinuation rather than a sirolimus effect.

b5 pt: whole or partial response, fast increased to 3 to 10 h; 5 patients: no response.

cBoth patients were also detected to have congenital hypothyroidism with normally positioned gland in the neonatal period.
life, with 2 cases diagnosed during the second year of life. An \( \text{ABCC8} \) mutation was identified in 68.2% (15/22) of the patients, whereas 13.6% (3/22) had other genetic findings associated with CHI. Only 18.2% (4/22) had no genetic findings related to CHI. 18F-fluoro-L-dihydroxyphenylalanine positron emission tomography (PET) scan revealed diffuse forms of CHI in 63.6% (14/22) and focal forms in 9.1% (2/22), whereas no imaging was performed in 27.3% (6/22). Three patients had a pancreatectomy before commencing sirolimus (one had negative CHI genetics, one was a compound \( \text{ABCC8} \) heterozygote, and one had a \( \text{GCK} \) mutation).

The onset of management with sirolimus is shown in Table 3. At the time sirolimus was commenced, 1 patient was on no CHI medications, another was on cornstarch, 10 were receiving 1 CHI medication, and 10 were on \( \geq 2 \) CHI medications. Partial/total response was observed in 95.5% (21/22) patients in our cohort, of which 1 showed total response and 20 partial response. A total of 4.5% (1/22) were unresponsive; 23.8% (5/21) of the partially/fully responsive patients had compound heterozygous \( \text{ABCC8} \) mutations and 23.8% (5/21) had heterozygous \( \text{ABCC8} \) mutation.

Complications detected while on sirolimus are summarized in Table 4. In those who had an infection, a bacterial etiology was identified in 64.7% (11/17) of cases. Of note, a 16-year-old female with CHI resulting from a dominant \( \text{ABCC8} \) gene mutation developed diabetes mellitus (DM) when clarithromycin was added to sirolimus [20]. Her dominant \( \text{ABCC8} \) mutation predisposes to DM (her mother with the same mutation had also developed DM), and there could have been a drug–drug interaction that potentiated the diabetogenic effect. The eventual management strategy required to achieve normoglycemia, whether on sirolimus or other medication/surgery, is shown in Table 5. Five of 22 patients (22.7%) remain on weaning doses of sirolimus requiring intensive feeding regimens (4 are on overnight continuous feeds, with added 10% carbohydrate content). The three patients that had a pancreatectomy before commencing sirolimus therapy all showed partial responses, and are currently managed on diazoxide (n = 2) and octreotide (n = 1), combined with continuous overnight feeds. Of note, four patients received sirolimus for a range of 13 months to 2.5 years,

### Table 2. Characteristics of Patients With CHI In This Study

| CHI Patient Characteristics | Values |
|-----------------------------|--------|
| **Sex**                     | 14 Females, 8 Males |
| **Weeks of gestation, median (range)** | 38 (33–40.5) |
| **Birth weight, median (range)** | 4050 g (2138–5300) |
| **Birth weight, median (range), SDS** | 1.71 SDS (−0.65 to +3.81) |
| **Perinatal CHI risk factors** | Gestational diabetes (2) |
| **Age at presentation** | Within first 2 d of life (19) |
| **Genetics** | 15 \text{ABCC8} mutations: 4 homozygous \text{ABCC8}, 5 Compound heterozygous \text{ABCC8}, 6 Autosomal dominant heterozygous \text{ABCC8} (2 maternal, 3 paternal, 1 \textit{de novo}) |
| **Syndrome** | 1 BWS |
| **PET scan report** | Diffuse (14) |

Abbreviation: SDS, standard deviation score.
were then switched to lanreotide, and currently remain with good glycemic control having avoided a pancreatectomy; in 2 of these cases, the reason for stopping sirolimus was re-
sponsiveness to lanreotide, whereas in the rest it was due to infection. A fifth patient remains on sirolimus and lanreotide, having also avoided surgery. Two patients with compound heterozygous \textit{ABCC8} mutations, classified as partially responsive to sirolimus, were initially able to fast for 8 hours; over time, their fasting tolerance dropped to 2 to 3 hours, at which point sirolimus was stopped.

Histopathology of the pancreas in four patients after the use of sirolimus is shown in Figure 1. The histopathology of the excised pancreatic tissue showed some variability in the amount of fibrous tissue within pancreas, but no greater than in the normal pancreas. Two patients had focal disease and two had diffuse disease. There were no other specific features identified. Pancreatectomy took place 4 to 24 months after sirolimus treatment. The reasons for stopping sirolimus and performing surgery were lesionectomy of focal form in one patient, no response to sirolimus in one, and side effects from sirolimus in two.

3. Discussion

To the best of our knowledge, this study reports the largest cohort of patients with CHI treated with sirolimus at a single center, representing a 5-year experience of this therapy. Given the controversies surrounding the usefulness, variable therapeutic success, and risks of this medication for CHI, previous case publications were reviewed. Overall, published cases show glycemic improvement, but at the expense of frequent side effects. The caution exerted when comparing our manuscript with previous case reports relies on the diverse nature of such case reports, given the rarity of CHI and, moreover, the use of mTOR inhibitors for this indication. These reports include small number of subjects, of different ages, with
various genetic makeup findings, the use of diverse concomitant treatments at different
doses, the definition of response to medication, and documentation of effects. The caveat is
that most case reports do not clearly state a definition of positive glycemic response to mTOR
inhibitors, but rather indicate if each patient was able to come off IV glucose and discharged
home on monotherapy or combination of CHI medications maintaining stable glucose con-
centrations. Critique could also be made to a certain extent on the preciseness of the criteria
for responsiveness in the current study; however, the rarity of this challenging disorder
combined with numerous management variables (IV dextrose infusion, continuous gastric
feeding, glucagon infusion) render it difficult to establish more rigorous targets. Publication
bias should be taken into account, because it is possible that trials with an unimpressive
outcome to this drug may have been unpublished.

mTOR inhibitors have proven their efficacy to control excessive insulin release in insu-
linomas [33], where the mTOR pathway is abnormally activated. In an attempt to avoid
extensive pancreatectomy in children with severe diffuse CHI, which eventually renders
children diabetic with pancreatic exocrine deficiency, and is, in some cases, only partially
curative with an ongoing need for medication [34], sirolimus was promoted as a promising
therapeutic agent by Senniappan et al. [7].

The patients reported in this study, in concordance with those in the literature, had
extremely severe forms of CHI that had already failed to respond to medical treatment with
maximum doses of diazoxide and octreotide and, in some, that was also unresponsive to
glucagon and nifedipine. This highlights the challenges inherent in the management of these
patients. The mTOR pathway has been shown to have a variable but constant activation and
overexpression in acinar cells of patients with CHI [10] and in insulin-producing islet cells
Conversely, a study analyzed pancreatic tissue from two sirolimus-unresponsive patients with CHI in which there was no reduction in cell proliferation [17]. An interpretation to this could be that the resistance to sirolimus might be due to the fact that the cell proliferation could be mTOR-independent. The histopathology of four pancreatic tissue samples examined in this study did not show important changes after the use of sirolimus; however, one possible criticism of the study may lie in the diverse time postsirolimus use and the variability in the pathological analysis of tissue.

Since the first publication of sirolimus use for CHI, 15 case reports have followed that are quite diverse in terms of their characteristics. Sirolimus has been mainly used in infants but also in an 8-year-old boy [26]. Most children harbor ABCC8 or KCNJ11 mutations in homozygosity or compound heterozygosity, but sirolimus has also been successfully used in a patient with Beckwith-Wiedemann syndrome (BWS) [23]. The majority of patients in case studies received sirolimus, but one received everolimus [17], which has a different half-life and plasma level range. Comparison between studies with surgically naïve patients and other patients with previous pancreatectomy [25, 27, 31] can also add a further layer of complexity.

### Table 5. Eventual Management of Patients With CHI

| Eventual Management | Management Details |
|---------------------|--------------------|
| Pancreatectomy       | After sirolimus start (3 permanently stopped sirolimus, 1 restarted sirolimus after operation): 4 |
|                      | Prior sirolimus start: 3 |
|                      | No pancreatectomy: 15 |
| Twenty-one patients that stopped sirolimus responded to: | Pancreatectomy + diazoxide (3.8–10.2 mg/kg/d) + feeds on-demand: 2 |
|                      | Pancreatectomy + octreotide (20–39.3 μg/kg/d) + continuous overnight feeds 10%–16% CHO: 2 |
|                      | Pancreatectomy + octreotide (13 μg/kg/d): 1 |
|                      | Pancreatectomy + lanreotide (30 mg/7 wk) + on-demand feeds: 1 |
|                      | Pancreatectomy + on-demand feeds: 1 |
|                      | Octreotide (40 μg/kg/d) + continuous overnight feeds 10%–16% CHO: 2 |
|                      | Octreotide (36 μg/kg/d) + continuous overnight feeds 10% CHO: 1 |
|                      | Octreotide (32 μg/kg/d) + continuous overnight feeds 18% CHO: 1 |
|                      | Octreotide (24.6 μg/kg/d) + on-demand feeds: 1 |
|                      | Octreotide (20 μg/kg/d) + continuous overnight feeds 10% CHO: 1 |
|                      | Octreotide (16 μg/kg/d) + continuous overnight feeds 10% CHO: 1 |
|                      | Lanreotide (30 mg/4 wk) + on-demand feeds: 1 |
|                      | Lanreotide (60 mg/3 wk) + continuous overnight feeds: 1 |
|                      | Lanreotide (60 mg/4 wk) + 3 hh feeds: 1 |
|                      | Overnight continuous feeds: 1 |
|                      | Frequent feeds: 1 |
|                      | Metformin because of DM: 1
|                      | Fasting capacity while on sirolimus (h): median (range): 8 (3–18) |
| Twenty-one patients that stopped sirolimus responded to: | No episodes of hypoglycemia at all: 2 |
|                      | Genetics: |
|                      | Autosomal dominant heterozygous ABCC8: 6 |
|                      | ABCC8 compound heterozygous: 5 |
|                      | ABCC8 homozygous: 4 |
|                      | Negative genetics for known CHI genes: 4 |
|                      | GCK: 1 |
|                      | BWS: 1 |
|                      | Rubinstein-Taybi syndrome: 1 |
| One patient that remains on sirolimus | Lanreotide (60 mg/4 wk), sirolimus (3.2 mg/m²/d; level 2 ng/L) + continuous overnight feeds 10% CHO: 1 |
|                      | Fasting capacity while on sirolimus (h): 6 |
|                      | Genetics: ABCC8 homozygous: 1 |

**Abbreviation:** CHO, carbohydrate content.

*aAs published in Dastamani A et al., [20]"
Although sirolimus has been mainly used in diffuse forms of CHI, in our study, this dual mTORC1/mTORC2 inhibitor agent was also used in two patients with 18F-fluoro-L-dihydroxyphenylalaninePET/CT scans compatible with focal disease. Both patients were heterozygous for a paternally inherited ABCC8 mutation; one did respond to sirolimus and had no side effects, and the other is currently still on sirolimus and octreotide at the age of 15 months with the complication of frequent skin infections around the gastrostomy site.

In our study, as in most case publications, sirolimus was started in combination with other CHI medications to counter the variable but constant activation of the mTOR pathway in CHI [12]. Despite the wide range of dose used, aiming to maintain medication blood concentrations within a therapeutic window and minimizing side effects, the median dose range is nevertheless higher than those reported in the literature. The higher doses may perhaps have influenced our higher rate of success, because partial/total response was observed in 95.5% of our cohort (the majority showing a partial response). This is a striking difference from the results published by Szymanowski et al. [17], in which the mTOR inhibitor was found to be clinically useful in 30% of patients in whom it was used. These authors also suggest that patients with dominantly inherited mutations in ABCC8 have a higher rate of response [17], concurring with the findings from another study [11]. In our study, heterozygous mutations and compound heterozygous mutations in ABCC8 were more likely to have a partial/total

Figure 1. Histopathology of pancreas after sirolimus use. (A) Patient with diffuse CHI resulting from a homozygous ABCC8 mutation, 4 mo after sirolimus treatment. High-power view of an islet showing scattered enlarged islet-cell nuclei. This change was present in islets throughout the tissue, in keeping with diffuse disease. (B) Patient with focal CHI resulting from paternal compound heterozygous ABCC8 mutation, 12 mo after sirolimus. There is an increase in islet-cell tissue with strong proinsulin staining. There was no diffuse islet-cell nuclear enlargement. The appearances suggest an atypical focal form of the disease. (C) Patient with focal CHI resulting from a paternal compound heterozygous ABCC8 mutation, 21 mo after sirolimus use. The image shows enlarged and clustered islets that stain strongly for proinsulin, in keeping with focal disease. (D) Patient with diffuse CHI resulting from homozygous ABCC8 mutation, 24 mo after sirolimus. There were scattered enlarged islet-cell nuclei throughout the tissue, indicating diffuse disease. Images taken with Nikon Eclipse E600 microscope; (A, D) Stained with hematoxylin and eosin; (B, C) stained with antibody to proinsulin. Original magnification: (A) ×400, (B) ×20, (C) ×100, (D) ×400.
response, with one case with *de novo* heterozygous *ABCC8* mutation achieving glycemic control on sirolimus monotherapy and avoiding surgery. We speculate that it seems unlikely that compound heterozygosity *per se* is affecting the response, but rather that the good outcome observed with compound heterozygosity may reflect the functional impact of the two mutations. Our patient with BWS previously reported as sirolimus-responsive in combination with octreotide [23] was switched at age 23 months to lanreotide, to which a good glycemic response was obtained.

Some of the published CHI cases on sirolimus have been shown to fast for up to 8 hours without developing hypoglycemia [7]. In our study, children on sirolimus showed improvement in their fast tolerance at the start of the treatment, and fasting time ranged from 3 to 8 hours on their latest fast provocation test. Two patients seemed to show a lower fasting capacity over time, perhaps suggesting that it may eventually become less effective in terms of glycemic control. Additionally, one publication reported increased quality of life since sirolimus helped to stabilize glycemia in a patient [26]. Of interesting, of the 17 patients that stopped sirolimus, only 4 required pancreatectomy; the rest were managed medically. This could indicate a use for sirolimus for short periods, thereby avoiding near-total pancreatectomy and its long-term adverse effects. Nevertheless, with such an approach, we recommend close monitoring of sirolimus trough concentrations and its side effects.

Cellular signaling studies in dermal fibroblasts from one patient with severe nonketotic hypoinsulinemic hypoglycemia resulting from mosaic genetic activation of class 1A PI3K (insulin signaling pathway) showed that sirolimus potently suppressed the increased basal AKT phosphorylation at very low doses (1.1 to 2.2 ng/mL). This offers mTOR inhibitors (or potentially PI3K inhibitors) a future therapeutic role in this rare form of hypoketotic hypoglycemia [35].

As in most studies using sirolimus in CHI patients, we also aimed for trough concentrations of 5 to 15 ng/mL. Interestingly, a case report stated that in a patient with CHI, the target was increased to that used in transplant patients, without any major side effects, although no glycemic response was achieved [31]. Postrenal transplant children seem to tolerate well doses of sirolimus of around 6 mg/m\(^2\)/d (target concentration 10 to 20 ng/mL) [36]. In our experience, sirolimus plasma concentrations were of limited use to titrate drug dose because their reproducibility was low given their dependence on sirolimus enteral absorption rate. This is why dose titration was based on maintenance of euglycemia (even if sirolimus concentrations were below the cutoff value) with avoidance of side effects. Reducing the dose, as observed in other studies, reverses side effects. Oral mucositis is frequent and can be minimized by rinsing the mouth with water or juice after the dose. However, a report on three children with CHI who received sirolimus doses as high as 35 mg/m\(^2\)/d reported no major side effects or infections [31].

Our study, with a wide range of trough concentrations of sirolimus, has the same rate of complications as collected in the case reports from the literature (86.4% vs 83.8%), which is not to be taken lightly. These complications can appear early, or after many months, and mainly involve infections of varying degrees of severity, in agreement with the literature. Infection is the main cause of discontinuation of sirolimus. Autosomal dominant *ABCC8* gene mutations causing CHI in infancy are known, in some cases, to be linked to DM at a later stage of life. A published case illustrates how sirolimus (combined with clarithromycin) may have precipitated the onset of diabetes [20].

Because the natural history of the condition is to get milder over time with slow progressive β-cell apoptosis, in some cases sirolimus would only be required for a short period, avoiding possible long-term consequences. For other cases, sirolimus could keep some children stable until they are older and are then switched to another treatment (e.g., lanreotide) to avoid pancreatectomy. This strategy worked for four of our patients. Similarly, Haliloglu et al. [30] had reported 2 patients responsive to sirolimus combined with octreotide, in which the former drug was discontinued because hepatitis, and both were managed on octreotide-LAR after the age of 7 months. Sirolimus may be a hazardous drug, however; even
after a 95% pancreatectomy, 60% of patients will remain with profound hypoglycemia [37] and therefore remain at risk for severe brain injury. Discussion with parents must therefore include the variable response and potential side effects of sirolimus as well as their preferred choice of treatment.

It is difficult to confirm causality of complications for every case, given that these infants need central access, admitted in the hospital for long periods, usually on a combination of medications, and may have other associated comorbidities; it is not uncommon for them to develop infections and other side effects. Because complications mainly developed while combined with other medications, the latter may also be held responsible of some of these. For instance, the CHI case published by Korula et al. [32] had a nonfermenting gram-negative bacilli sepsis before sirolimus treatment and had not had any infections or side effects in the 14 months of treatment on sirolimus. Further studies analyzing the long-term use of sirolimus in children with CHI are needed. Also, the response of children with other CHI genetic mutations to sirolimus needs to be investigated.

Although the potential role for mTOR inhibitors in certain CHI cases cannot be ignored, a thoughtful decision-making process regarding the optimal therapy for patients with severe medically unresponsive CHI must be undertaken, balancing potential benefit vs risks. The latter are common and potentially life-threatening; in view of these concerns, we would suggest the use of mTOR inhibitors as a last-resort therapy for extremely severe CHI cases unresponsive to conventional CHI medications, for curtailed periods with close monitoring, with the caveat that plasma concentrations are of limited use given their variability.

4. Research in Context

A. Evidence Before This Study

mTOR inhibitors, in particular sirolimus, have been used in the past 5 years in children with severe forms of congenital hyperinsulinism (CHI) that do not respond to maximal doses of diazoxide and octreotide. Because CHI is a rare disease, a PubMed search (in English, Spanish, and Greek) was performed, and characteristics of the 15 publications reporting cases in which mTOR inhibitors were used for this condition were analyzed. Reported response to sirolimus in the literature is variable, but with a high rate of adverse effects.

B. Added Value of This Study

Our study, to the best of our knowledge, reports the largest series of CHI patients treated with sirolimus, with the longest follow-up duration. We have compared our findings with those previously published. Our study describes a higher number of CHI cases that are partially sirolimus-responsive and a similar high rate of complications while on this medication.

C. Implications of All the Available Evidence

Given the frequent and potentially severe complications while on this medication, candidates for this therapy should be meticulously selected. Transient use of sirolimus, until achievement of euglycemia on therapies such as lanreotide, could be a future approach to avoid pancreatectomy.

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