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1 Increasing knowledge in *IGF1R* defects: lessons from 35 new patients.

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51 ABSTRACT

52 (248words)

Background: The IGF1R is a keystone of fetal growth regulation by mediating the effects of IGF-I and IGF-II. Recently, a cohort of patients carrying an *IGF1R* defect was described, from which a clinical score was established for diagnosis. We assessed this score in a large cohort of patients with identified *IGF1R* defects, as no external validation was available. Furthermore, we aimed to develop a functional test to allow the classification of variants of unknown significance (VUS) *in vitro*.

Methods: DNA was tested for either deletions or single nucleotide variant (SNV) and the phosphorylation
of downstream pathways studied after stimulation with IGF-I by western blotting of fibroblast of nine
patients.

Results: We detected 21 *IGF1R* defects in 35 patients, including eight deletions and 10 heterozygous, one homozygous, and one compound-heterozygous SNVs. The main clinical characteristics of these patients were being born small for gestational age (90.9%), short stature (88.2%), and microcephaly (74.1%). Feeding difficulties and varying degrees of developmental delay were highly prevalent (54.5%). There were no differences in phenotypes between patients with deletions and SNVs of *IGF1R*. Functional studies showed that the six missense SNVs tested were associated with decreased AKT phosphorylation.

67 Conclusion: We report eight new pathogenic variants of *IGF1R* and an original case with a homozygous
68 SNV. We found the recently proposed clinical score to be accurate for the diagnosis of *IGF1R* defects
69 with a sensitivity of 95.2%. We developed an efficient functional test to assess the pathogenicity of SNVs,
70 which is useful, especially for VUS.

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72

74 INTRODUCTION

75 Insulin-like growth factors IGF-I and IGF-II are major factors which stimulate fetal growth. Both bind to the type 1 IGF receptor (IGF1R). Binding of IGFs to this receptor leads to autophosphorylation of 76 77 intracellular tyrosine residues, which in turn leads to activation of the phosphatidyl-inositide 3-kinase 78 (PI3K)/AKT and mitogen-activated protein kinase (MAPK)/ERK signaling pathways, resulting in cellular 79 proliferation and growth [1]. IGF1R is located at chromosome 15q26, contains 21 exons, and leads to the 80 expression of a dimeric transmembrane tyrosine-kinase receptor (figure 1A. and 1C.) [2]. IGF1R and the insulin receptor (IR) share more than 50% homology and hybrid dimers can be generated, the function of 81 82 which is still unclear [3].

83 The first description of the involvement of *IGF1R* defects in pre and post-natal growth failure was made by Pasquali et al. in the late seventies [4]. The authors described patients with intra-uterine growth 84 85 retardation (IUGR), post-natal growth failure, and microcephaly due to a 15q26 terminal deletion that 86 included *IGF1R* (OMIM #612626). Butler et al. then described a similar phenotype in patients with ring 87 chromosome 15 with *IGF1R* deleted [5]. The phenotype-genotype correlation was unclear because of the difference in gene content of these large deletions. In 2013, Abuzzahab et al. reported a loss of function of 88 the IGF1R in two children with either compound heterozygous pathogenic missense variants or a 89 90 heterozygous pathogenic nonsense variant of IGF1R (OMIM #270450) [6]. Since then, many others have 91 reported pathogenic variants in *IGF1R*, mainly in the heterozygous state and rarely in the compound 92 heterozygous state (n = 3) [7 and references within]. Finally, in 2012, Gannagé-Yared *et al.* reported the 93 first patient with a homozygous pathogenic variant [8]. Most of the reported cases were born small for 94 gestational age (SGA) with no or poor catch-up growth, but the final heights are widely variable and can 95 be in the normal range. Indeed variable phenotypic expression has already been reported, even in relatives 96 carrying the same molecular defect, which makes the diagnosis of IGF1R defects difficult [9,10]. 97 Furthermore, some authors have highlighted that the phenotype of patients with *IGF1R* defects overlaps 98 with that of either Silver-Russell syndrome (SRS, OMIM #180860) or SHORT syndrome (OMIM

99 #269880), leading to the late diagnosis of *IGF1R* defects [11,12]. However, the presence of microcephaly 100 appears to be highly specific for IGF1R defects. Thus, the recent international consensus about SRS 101 mentioned *IGF1R* defects as a differential diagnosis of SRS in case of absence of relative macrocephaly, 102 meaning that head circumference should be assessed in a patient born SGA with poor catch-up growth to 103 distinguish between these etiologies [13]. From the first descriptive cohort of 25 patients with IGF1R 104 defects, Walenkamp et al. proposed a clinical score to drive molecular investigations [14]. This score 105 combines the following four items: birth length or weight < -1 standard deviation score (SDS), head 106 circumference < -2 SDS at first presentation, height at first presentation < -2.5 SDS, and plasma IGF-I 107 levels above the mean for age and gender. Molecular testing for IGF1R should be proposed if three or 108 more items are present, with a sensitivity of 76% in their cohort [14]. Recently, Janchevska et al. 109 identified two anomalies of IGF1R in a cohort of 64 patients born SGA with no catch-up growth, 110 supporting the hypothesis that the prevalence of these defects is high enough to search for them in this 111 particular group of patients [15]. Concerning the treatment of postnatal short stature, the efficiency of 112 recombinant growth hormone (rGH) therapy in patients with *IGF1R* defects is still controversial and only 113 isolated cases with variable age at onset, duration, and dose of treatment have been reported [7,14].

A few functional studies in either fibroblasts or cell lines have been reported, generally showing the inability of the mutated receptor to activate downstream pathways, especially phosphorylation of the receptor itself and/or AKT and rarely ERK [15–18]. In 2009, Fang *et al.* demonstrated *IGF1R* haploinsufficiency due to a mRNA decay phenomenon in a nonsense variant in exon 18 [10]. Most groups have not observed any effect on the expression of the transmembrane IGF1R in patients with missense variants [6,10,19,20].

We report here a large cohort of 21 *IGF1R* defects, including eight previously unreported pathogenic variants. Furthermore, we established phenotype-genotype correlations and assessed the efficiency of rGH therapy in these patients. Finally, we developed a reproductive functional test to assess the responsibility of variants of unknown significance (VUS) in the phenotype.

124 MATERIALS AND METHODS

125 **Patients**

Patients were either followed in our clinic or referred by other clinical centers for molecular analysis. A clinical file, including comprehensive clinical and biological data, growth charts, and treatment was completed for all patients. Each patient had been examined by a geneticist and/or a pediatric endocrinologist. Molecular diagnosis strategy following clinical presentation is depicted in supplementary figure 1. Written informed consent for participation was received either from the patients themselves or their parents, in accordance with French national ethics rules for patients recruited in France (Assistance Publique – Hôpitaux de Paris authorization n°681).

133 Auxologic methods

Length, weight, and head circumference at birth are expressed as SDS according to Usher and McLean
charts [21]. Post-natal growth parameters are expressed as SDS according to Sempé charts [22]. The age
of puberty onset (breast development for girls and testis enlargement (≥ 4mL) for boys) was considered to
be normal from 8 to13 years for girls and 9 to 14 years for boys.

138 Molecular analysis

All molecular diagnosis of the *IGF1R* defects was performed in the same laboratory of molecular genetics.
DNA was extracted from blood leukocytes using an in-house protocol after cell lysis by a salting out
procedure, as previously described [23]. DNA was quantified using a NanoDrop ND-1000
Spectrophotometer (Invitrogen, France).

The main known molecular causes of SRS (loss of methylation at 11p15, maternal uniparental disomy of
chromosome 7) or Temple syndrome (OMIM#616222) at the 14q32.2 locus were ruled out by methylation
analysis, as detailed in a previous study [23].

IGF1R deletions were assessed by multiplex ligation-dependent probe amplification (MLPA) using the
 SALSA MLPA P217 IGF1R probe mix (MRC Holland, Amsterdam, Netherlands), following the

manufacturer's instructions. MLPA data were analyzed using the Novel Software Coffalyser.NETprovided by MRC-Holland.

For single nucleotide polymorphism (SNP) microarray analysis, samples were processed using cytoSNP-12, or HumanOmniExpress-24 microarrays (Illumina, San Diego, CA, USA). Automated Illumina microarray experiments were performed according to the manufacturer's instructions. Images were acquired using an iScan System (Illumina). Image analysis and automated CNV calling were performed using GenomeStudio v.2011.1 and CNVPartition v.3.1.6. SNP profiles were analyzed by examination of signal intensity (Log R ratio, i.e. ln (sample copy number/reference copy number)) and allelic composition (BAF, i.e. B Allele Frequency).

For the detection of *IGF1R* SNV, DNA was amplified and sequenced by direct Sanger sequencing procedures, using the ABI PRISM Big Dye Terminator v3.0 Cycle Sequencing Kit and an ABI 3100 Genetic Analyzer (Life Technologies, Courtaboeuf, France). Sequences were then analyzed with SeqScape v2.6 (Life Technologies).

Variants are described in accordance with the recommendations of the Human Genome Variation Society.
All the new variants were recorded in the ClinVar database (https://www.ncbi.nlm.nih.gov/clinvar).

Variant interpretation was performed following the American College of Medical Genetics and Genomics and the Association for Molecular Pathology (ACMG/AMP) classification of variants [24]. Six main categories are evaluated according to these guidelines: population data (prevalence of the variant in control populations), computational in silico predictive data, functional characterization, segregation, de novo data and allelic data (e.g. variant detected in *trans* with a pathogenic variant for a recessive disorder). Depending on these data, variants are classified as benign or likely benign, pathogenic or likely pathogenic and some stay of uncertain significance.

Functional test

171 Patient fibroblasts, obtained from skin biopsies after informed consent and control fibroblasts obtained from the Coriell Institute of Medical Research (Camden, N.J.), were cultured in Dulbecco's Modified 172 173 Eagle Medium enriched with glutamate, sodium pyruvate, penicillin, streptomycin, and 10% fetal-calf 174 serum at 37°C. After 24 h of serum-free culture in six-well plates, cells were stimulated with [50 ng/ml] 175 IGF-I (Peprotech, US) for 10 min before lysis. We found these stimulation conditions to be the most 176 accurate to assess both AKT and ERK phosphorylation in controls. For both non-stimulated and stimulated conditions, 4.2 to 12.1 µg of protein was deposited in a NuPAGETM 4-12% Bis-Tris Gel 177 (Thermo Fischer Scientific, US). Electrophoresis was performed on an XCell SureLock™ Mini-Cell 178 179 Electrophoresis system (Thermo Fischer Scientific, US). Membranes were incubated with polyclonal 180 antibodies against either phospho-AKT (Ser473, Cell Signaling, US, 1:2000), pan-AKT (Cell Signaling, 1:1000), phospho-ERK1/2 (Tyr204, Cliniscience, France, 1:800), ERK1/2 (Cell Signaling, 1:1000), or 181 GAPDH for normalization (Cell Signaling, 1:2000). Then, membranes were incubated with an HRP-182 183 conjugated secondary antirabbit antibody (1:3,000), revealed with ChemiDoc[™] XRS+ System (Bio-Rad, US), and analyzed with Quantity One v4.6.6 software. Immunoblot images were quantified using ImageJ 184 185 1.50 software (https://imagej.nih.gov).

186

187 **mRNA** quantification

Total mRNA was extracted from non-stimulated cells using NucleoSpin miRNA® (Macherey-Nagel) and
cDNA obtained by reverse-transcriptase polymerase chain reaction (RT-PCR, Superscript II, Invitrogen,
France). cDNA was then amplified and quantified on a QuantStudio 7 Flex Real-Time PCR system
(Thermo Fischer) using primers localized in exons 7-8 by SYBR Green technology (Applied Biosystem,
US).

193 **Biological assays**

194 IGF-I serum concentrations were determined by different techniques, as patients were followed in 195 different centers. However, IGF-I levels were collected along with the normal values (NV) for most patients. Thus, IGF-I levels were considered as high if >1 SDS according to the technique used. We
express IGF-I levels as SDS according to age and gender from control matched references [25].

198 Statistical analysis

Characteristics of the population are described as percentages for qualitative variables or as SDS and mean
(range) for continuous variables. For statistical analysis, Pearson's test was used for correlations, Fisher's
test for dichotomous variables, and the t-test for continuous variables.

202 RESULTS

203 Genetic results

Between 2006 and 2018, 111 samples of DNA were tested for IGF1R mutations/deletions. We identified 204 205 IGF1R defects in 35 patients from 20 different families. Aside from the 20 index cases, we identified 206 IGF1R defects in 15 relatives, including three siblings, seven fathers, and five mothers. Among the 20 207 index cases, molecular analysis was prescribed for 13 patients for a clinical suspicion of an IGF1R defect 208 and for seven for a clinical suspicion of SRS (Supplementary figure 1). The molecular diagnosis of the 209 IGF1R defect was made at 9.2 years of age (0.8 to 18.1) for the index cases. Eight patients carried a 210 heterozygous deletion (figure 1B.). Eleven carried a single nucleotide variant (SNV): 8 missense, two 211 nonsense, and one insertion at the boundary of intron 5-exon 6. Sequencing of the cDNA of the latter 212 variant obtained from lymphocytes confirmed that the inserted guanine was present in the cDNA, leading 213 to a frameshift and a premature stop codon (N417Efs*52, Supplementary figure 2). One patient carried 214 two missense SNVs. Among the 13 SNVs identified, 10 patients had a heterozygous SNV, one patient 215 had compound heterozygous missense SNV, and one carried a homozygous missense SNV (figure 1). 216 Parental DNA samples were available for 13 patients. Three inherited the anomaly from their mother, five 217 from their father, two from both parents, and the anomaly arose *de novo* for three patients. Among the 13 218 variants, we identified eight new pathogenic or likely pathogenic variants (Table 1). Five deletions 219 included the entire *IGF1R* gene, one interstitial and four terminal lengthening from 3.13 to 5.01Mb (figure 1B., Del1 to 5), whereas three included only part of *IGF1R* with length from 19kb to 234kb (figure 1B.,
Del6 to 8).

222 Clinical features

223 The intragenic deletions (exon 2, Del6, n = 3) and *IGF1R* terminal deletions (Del7, n = 2 and Del8, n = 1) 224 did not include other disease-causing OMIM genes and were thus analyzed together with the SNVs for the 225 clinical study. Clinical characteristics are shown in Table 2. There was no statistical difference in clinical 226 presentation between patients with large deletions and pathogenic variants of *IGF1R*. We calculated the 227 clinical score recently proposed by Walenkamp et al. for 21 patients for whom clinical data required for 228 this scoring system were fully available (birth weight or length < -1 SDS, height at presentation < -2.5229 SDS, head circumference at presentation < -2 SDS (microcephaly) and IGF-I level > 0 SDS) [14]. Twenty 230 patients (95.2%) met at least three of the four criteria and 11 (52.4%) fulfilled all four. Among them, all 231 had a birth weight or length < -1 SDS, 17 (81.0%) had a height at presentation below -2.5 SDS, and 19 232 (90.5%) had microcephaly. All 21 patients scored positive for elevated IGF-1 levels if considered at the 233 different endpoints (including during rGH treatment). However, five patients (23.8%) would have not met 234 this criterion if IGF-I levels were considered only prior to the initiation of rGH treatment. One patient 235 (carrying Del6) did not achieve a positive clinical score, with only two items [being born with a height or weight < -1 SDS and high levels of IGF-I (during rGH treatment only)]. 236

Given the clinical overlap between SRS and *IGF1R* defects, the Netchine-Harbison clinical scoring
system (with a positive clinical diagnosis of SRS for a score of at least 4/6) was assessable for 10 patients
and only one scored 4/6 [13,26] (lacking relative macrocephaly at birth and body asymmetry items).
Nevertheless, most patients scored 3 out of 6, comprising the following items: being born SGA, post-natal
growth retardation, and feeding difficulties.

					ACMP/AMP					
cDNA nomenclature NM_000875.4	Reference	Amino-acid substitution NP_000866.1	ClinVar	GnomAD	Detailed staging	Variant classification#				
c.118C>T	This study	R40C	SCV000926288	Not reported	PM1 PM2 PP2 PP3	Likely pathogenic				
c.384T>C	This study	F112L	SCV000926289	Not reported	PS4 PM1 PM2 PP1 PP2 PP3 PS3	Pathogenic				
c.904G>T	This study	E302*	SCV000926290	Not reported	PVS1 PM2 PP3 PP4	Pathogenic				
c.995G>A	This study	C332Y	SCV000926291	Not reported	PM1 PM2 PP2 PP3 BS4	Likely pathogenic				
c.1247+1-1247+2insG	This study	N417Efs*52	SCV000926292	Not reported	PS3 PM2 PM4 PP1 PP4	Pathogenic				
c.3162G>A	This study	M1054I	SCV000926293	Not reported	PM1 PM2 PP2 PP3 PP4	Likely pathogenic				
c.3454G>A	This study	G1152R	SCV000926294	Not reported	PS4 PM1 PM2 PP1 PP2 PP3 PP4	Pathogenic				
c.3539C>A	This study	S1180Y	SCV000926295	Not reported	PS4 PM1 PM2 PP2 PP1 PP3	Pathogenic				
c.2629C>T	[30]	R877*	rs150221450	AF: 3.977.10 ⁻⁶	PVS1 PM2 PP3 PP4	Pathogenic				
c.3530G>A	[14]	R1177H	SCV000926296	Not reported§	PM1 PM2 PP2 PP3	Likely pathogenic				
c.3595G>A	EGL Genetic Diagnostics	G1199R	rs886044448	Not reported	PS4 PM1 PM2 PP1 PP2 PP3 PP4	Pathogenic				
c.4055G>T	dbSNP	G1352V	rs759808066	AF: 1.607.10 ⁻⁵	PM2 PP2 PP3 PS3 BS4	Likely pathogenic [¤]				
c.4066G>A	[14]	E1356K	rs746562843	AF: 642.10 ⁻⁵	PM2 PP2 BS4 PS3	Likely pathogenic ^x				

Table 1

		All n =	= 35		Deletions n	n = 6	SNVs n = 29			
	Mean	Range	n (%)	Mean	Range	n (%)	Mean	Range	n (%)	-
Sex (Female/Male)			15/20			2/4			13/16	0.68
Birth parameters:										
Term (WA)	37.7	31.0;41.5	23	38.4	37.0;40.0	5	37.5	31.0;41.5	18	0.82
Preterm (< 37 WA)			5/23 (21.7)			0/5 (0)			5/18 (27.8)	0.55
Weight (SDS)	-2.5	-3.8;-1	23	-2.3	-3.2;-1.4	5	-2.5	-3.8;-1.0	18	0.58
SGA #(weight <- 2 SDS)			17/23 (73.9)			3/5 (60.0)			14/18 (77.8)	1
Length (SDS)	-3.3	-5.2;-1.4	22	-3.0	-4.2;-2	5	-3.4	-5.2;-1.4	17	0.46
SGA # (length < -2 SDS)			20/22 (90.9)			4/5 (80.0)			16/17 (94.1)	1
Head circumference (SDS)	-2.6	-3.9;-0.3	17	-2.4	-3.0;-1.6	5	-2.7	-3.9;-0.3	12	0.63
Microcephaly (HC< -2 SDS)			14/17 (82.4)			4/5 (80.0)			10/12 (83.3)	1
Relative macrocephaly#			3/17 (17.6)			1/5 (20.0)			2/12 (16.7)	1
Clinical features:										
Microcephaly			20/27 (74.1)			4/4 (100)			16/23 (69.6)	0.70
Feeding difficulties#			12/22 (54.5)			4/5 (80.0)			8/17 (47.1)	0.69
Anorexia/thinness			11/14 (78.5)			4/4 (100.0)			7/10 (70.0)	1.0
Enteral nutrition			5/14 (35.7)			2/4 (50.0)			3/10 (30.0)	1
Developmental delay			12/22 (54.5)			4/5 (80.0)			8/17 (47.1)	0.69
Language			6/22 (27.3)			3/5 (60.0)			3/17 (17.6)	0.31
Motor			6/22 (27.3)			1/5 (20.0)			5/17 (29.4)	1
Cognitive			12/22 (54.5)			4/5 (80.0)			8/17 (47.1)	0.69
Assistance in school			14/23 (60.9)			4/5 (80.0)			10/18 (55.6)	0.70
Clinical score ([14]):										
Birth weight or length < -1 SDS			21/21 (100)			4/4 (100)			17/17 (100)	
Microcephaly (HC< -2 SDS)			19/21(90.5)			4/4 (100)			15/17 (88.2)	
Height # (< -2.5 SDS)			17/21(81.0)			4/4 (100)			13/17 (76.5)	
IGF-I >0 SDS			21/21 (100)			4/4 (100)			17/17 (100)	
Final height (SDS):	-2.5	-4.2;1.0	25	-2.9	-4.1;-2.1	5	-2.3	-4.2;1.0	20	0.30
Without rGH	-2.2	-4.2;1.0	13	-3.2	-4.1;-2.1	2	-2.0	-4.2;1.0	11	0.42
With rGH	-2.8	-4.2;-1.0	12	-2.8	-3.8;-2.1	3	-2.8	-4.2;-1.0	9	0.99
Short stature (< -2 SDS)			19/25 (76.0)			5/5 (100)			14/19 (73.7)	0.72

Table2

245 No deafness was reported in our cohort. One girl (carrying a missense SNV) had a slightly delayed onset 246 of puberty (onset at 13.3 years), whereas the onset of puberty of the other 16 patients (10 boys) occurred at 247 the normal age. Three patients were treated with GnRH analogs together with rGH at the onset of puberty 248 to preserve the duration of growth due to a low predicted final height, despite the onset of puberty at a 249 normal age. Four patients carrying a missense SNV had attention deficit hyperactivity disorder (ADHD), 250 which required medication. Three patients developed obesity in childhood with metabolic syndrome for 251 one as a young adult. One patient (father of two affected children) had early type 2 diabetes and one 252 patient had episodes of hypoglycemia in infancy. Noticeably, four patients (two with IGF1R deletion, two 253 with a missense SNV) had cardiac defects, including one case of transient inter-auricular communication 254 (IAC), one of IAC and rhythmic troubles, one of patent foramen ovale, and one of severe cardiac 255 insufficiency, which led to heart transplantation (carrying a missense SNV).

256 Only two cases of homozygous pathogenic variants have yet been reported [8,12]. Thus, the pedigree and 257 growth curves of the girl with the F112L homozygous pathogenic variant are shown in figure 2. Although 258 the girl with the homozygous pathogenic variant (II.4) showed severe growth retardation of approximately 259 -4 SDS with tremendously elevated IGF-I [621 ng/mL (NV 20-300) at 1.6 years] and IGFBP-3 [5605 260 ng/mL (NV 800-3700)], both parents (I.1 and I.2) and one older sister (II.3) with the heterozygous pathogenic variant showed impaired postnatal growth of approximately -2 SDS, with a final height in the 261 262 lower range of normal curves. The unaffected younger siblings (II.1 and II.2) showed normal growth around the mean. Furthermore, the homozygous carrier (II.4) had a patent foramen ovale, severe oeso-263 264 gastric reflux, anorexia requiring enteral support for one year (naso-gastric tube), and psychomotor delay 265 with learning disability, whereas no other member of the family presented with such clinical features.

All clinical data are available for each patient in Supplementary Table 1.

267 rGH therapy

268 Eighteen patients received rGH treatment, starting at an age of 7.5 years (1.5;15.3) under the SGA European Medicines Agency (EMA) indication, with a mean height at the start of therapy of -3.8 SDS (-269 270 5.6; -1.6). The starting dose was 46.6 μ g/kg/day (35.0;85.5) and was significantly increased for only five 271 of the 15 patients for whom data on the dose evolution was available. For most patients (60.0%), the dose 272 of rGH was not raised because of high serum levels of IGF-I. Among the 12 patients that completed rGH 273 treatment and reached their final height, the mean height gain was 1.0 SDS (0.2;2.5), which positively 274 correlated with the duration of treatment ($\rho = 0.76$, p = 0.004) and negatively correlated with the age at the 275 start of rGH ($\rho = -0.68$, p = 0.01). IGF-I serum levels were high (over 1 SDS) for 11 (47.8%) patients 276 before any treatment, with a mean of 1.9 SDS (-2.0;7.1), which rose to 3.3 SDS (0.3;9.5) under rGH 277 therapy.

278 IGF1R functional test

We performed functional analysis on fibroblasts for seven index cases, two affected parents and four 279 280 controls (figure 3). The six missense SNVs all showed a decrease in phosphorylated AKT, although the results for the S1180Y variant did not reach statistical significance (p = 0.065). Both G1352V and 281 282 E1356K were predicted as VUS after in silico analysis and showed a significant decrease in AKT 283 phosphorylation *in vitro* (p = 0.009 and p = 0.002, respectively), suggesting that these variants are likely 284 pathogenic. There were no alterations of AKT phosphorylation for two patients carrying either a nonsense 285 SNV or a chromosome 15q26.6 deletion that included the entire IGF1R gene. The results concerning ERK 286 phosphorylation were highly variable and we observed no significant modifications in this pathway (Supplementary figure 3). All but one patient showed normal *IGF1R* expression. This patient, who carries 287 288 a 15q26.6 heterozygous deletion, including IGF1R, showed expression of 37.7% of controls 289 (Supplementary figure 4).

290 **DISCUSSION**

We report a large cohort of patients carrying various *IGF1R* defects and describe eight new pathogenic
variants. Furthermore, we developed an *in vitro* functional test to assess the pathogenic impact of VUS.

As previously described, *IGF1R* defects are mainly present in the heterozygous state. Nevertheless, we found two patients with missense SNVs on both alleles, including one patient with compound heterozygous pathogenic variants and one with a homozygous one. The patient carrying the compound heterozygous SNVs did not phenotypically differ from the other patients although both variants were pathogenic. On the other hand, the homozygous pathogenic variant was associated with a more severe phenotype in terms of growth, microcephaly, and mental retardation relative to that of her relatives who carry the same variant in the heterozygous state.

300 With the advent of next-generation sequencing (including exome sequencing or a gene panel of growth 301 disorders, microcephaly, or cognitive impairment, which can include *IGF1R*), the identification of SNVs 302 will increase in the future. Thus, the description and registration of new SNVs with a precise phenotypic 303 description is necessary to distinguish between those that are benign and those that are pathogenic. 304 Furthermore, we demonstrated that functional characterization of such SNVs is sometimes necessary. In 305 our cohort, such experiments were helpful for the classification of two SNVs reported as SNPs with a very 306 low allele frequency and classified as VUS based on the ACMG/AMP recommendations because of 307 incomplete penetrance (E1356K and G1352V) [24]. However, the definition of "unaffected" carrier was 308 only based on the reported final heights of the two fathers who carried the variants, as other criteria were 309 not available (birth parameters, head circumference and IGF-I levels). Those two variants were finally 310 classified as likely to be pathogenic after demonstration of their functional consequences.

The *in vitro* studies showed impairment in the ability to activate downstream pathways for the receptors affected by missense SNVs, especially the AKT pathway. We were unable to demonstrate any significant functional consequences of deletions or nonsense SNV, unlike previous studies [10,27,28]. It is possible that discrepancies between our results on deletions and those of previous studies may be due to different 315 IGF-I concentrations used for stimulation. Indeed, Choi et al. showed a progressive increase in AKT 316 phosphorylation in fibroblasts from a patient with an IGF1R deletion in response to increasing IGF-I 317 concentrations from 1 to 400 ng/mL [28]. Ester et al. reported the same pattern with lower concentrations 318 (5 to 20 ng/mL) [18]. Thus, it is possible that the IGF-I concentration we used (50 ng/mL) did not allow 319 proper discrimination of AKT phosphorylation between deletions and controls [29]. Nevertheless, the aim 320 of this functional study was to assess pathogenic impact of SNVs of unknown significance, and we found 321 IGF-I concentration of 50 ng/mL to be effective. Unlike missense pathogenic variants, which may lead to 322 a dominant-negative effect, deletions or nonsense variants may lead to haploinsufficiency. Although we 323 could not quantify membrane IGF1R, we demonstrated that IGF1R mRNA levels were low in fibroblasts 324 from one patient with a deletion, favoring haploinsufficiency [10,28,30].

325 Very recently, a scoring system has been proposed for a clinical suspicion of an *IGF1R* defect [14]. This 326 clinical score showed 95.2% sensitivity for our cohort. All patients were born with weight or length < -1327 SDS and microcephaly was almost always present. However, post-natal short stature (with a threshold set 328 at -2.5 SDS) was inconstantly observed in our cohort. However, height at first evaluation was usually 329 below -2 SDS. The clinical scoring system could be adapted for this item, so as not to miss patients with IGF1R defects for whom height is not severely affected but this will result in an increased number of 330 patients that should be tested and thus, to a reduced specificity. Elevated circulating IGF-I levels were 331 332 absent prior to rGH therapy for 23.8% of the patients in our cohort but IGF-I levels rose markedly after initiating rGH treatment. The absence of high IGF-I levels prior to rGH therapy can be explained by the 333 previously described feeding difficulties of some patients with IGF1R defects, which can lead to 334 335 nutritional deficiency and low basal levels of IGF-I [31]. This pattern of low IGF-I levels which increase 336 rapidly after initiating rGH therapy, should alert clinicians to the possibility of an *IGF1R* defect in a child 337 born SGA, especially with the presence of microcephaly. However, the high sensitivity of this clinical 338 score favors its use in routine diagnosis to drive genetic tests. The specificity of this clinical score should 339 be assessed in large cohorts of SGA patients with the help of molecular studies, as well as in patients with idiopathic short stature since fetal growth restriction, although highly prevalent, is not constant in patientscarrying *IGF1R* defect.

342 This cohort allowed us to better characterize the phenotype of patients with an IGF1R defect. As previously described, fetal and post-natal growth retardation, microcephaly, and elevated IGF-I serum 343 344 levels were highly prevalent in our cohort [7,14]. As in previously reported cases, we identified several 345 cardiac anomalies in these patients, mostly benign. However, one patient underwent heart transplantation 346 because of severe cardiac failure. These findings are in accordance with previous observations and argue 347 in favor of a systematic cardiac ultrasound evaluation when an IGF1R defect is identified [8,12,19,32]. 348 Another interesting feature of our cohort is the presence of ADHD in several patients, which was only 349 been previously reported for one case [29]. Furthermore, as reported in the Dutch cohort, we found a high 350 prevalence of feeding difficulties, sometimes requiring nutritional support, [9,11,14]. This latter feature may have misled some clinicians to consider a clinical diagnosis of SRS at first evaluation. Indeed patients 351 352 with SRS or IGF1R defects share several symptoms, including being born SGA, post-natal growth 353 retardation, and high circulating levels of IGF-I [33,34]. However patients with IGF1R defects usually 354 present with microcephaly, which distinguishes them from SRS patients, for whom head circumference is 355 relatively preserved at birth [13,26]. The recent international consensus on the diagnosis and management of SRS stated that *IGF1R* defects represents a differential diagnosis and may be considered easily after the 356 357 major molecular defects of SRS are ruled out, especially for those patients with no relative macrocephaly 358 [13].

The efficiency of rGH therapy in this cohort is difficult to ascertain, as this was a retrospective and multicentric analysis with varying management in terms of the age at onset, the initial dose, dose adaptation, and discontinuation of treatment. Since the duration and age at the start of treatment significantly correlate with height gain, rGH treatment should be considered for patients with no catch-up growth at four years of age, under the EMA SGA indication. However, both clinicians and patients (or parents) should be aware of this unpredictable response to rGH therapy. Our *in vitro* experiments,

accounting for the functional consequences of the variants, were unable to distinguish between those patients who responded well or poorly to rGH therapy. It would be of interest however to set up such a prognostic tool. The high baseline IGF-I levels do not reflect the biological effect of IGF-I since the IGF1R signalization is impaired. These high IGF-I levels raised concern on potentially negative long term effects. Nevertheless, the pathophysiological comprehension of such elevated circulating IGF-I levels allows us to consider these levels only as a consequence of IGF-I resistance. Thus, IGF-I levels should not be interpreted and used in patients with *IGF1R* defects in the same manner as in unaffected patients.

In conclusion, we provide extensive clinical data on a large cohort of patients carrying *IGF1R* defects. We identified eight new pathological variants, including one homozygous pathogenic variant. We validated the clinical scoring system that has been recently proposed for patients with *IGF1R* defects. Finally, we developed a functional test to assess IGF1R activity *in vitro* that is useful for sorting VUS, which is of particular importance, especially for accurate genetic counseling.

377

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495

496 TABLES

497 Table 1. Description of the identified single nucleotide variants in the cohort and predictions of the

498 pathological consequences. ACMP/AMP: American College of Medical Genetics and Genomics and the

Association for Molecular Pathology classification of variants [24]. **PS3** corresponds to a pathogenic

effect in functional test. AF: allele frequency. # Classification performed using the InterVar classification
system; § First reported in Walenkamp *et al.* [14]. ¤ Classified as "variant of unknown significance"
before the functional test results.

Table 2. Clinical features of the patients with *IGF1R* defects. SNV: Single nucleotide variant. WA: weeks of amenorrhea; SDS: standard deviation score; SGA: born small for gestational age; HC: head circumference; rGH: recombinant growth hormone. # Items included in the Netchine-Harbison clinical scoring system for Silver-Russell syndrome diagnosis.

507 FIGURES

Figure 1. A. Schematic representation of the position of *IGF1R* on chromosome 15. B. Representation of
the eight identified deletions using the UCSC (University of California Santa Cruz) software. C.
Representation of the identified single nucleotide variants (SNV) from exons 1 to 21. Arrows indicate the
SNVs identified in the cohort. The corresponding functional domains of the protein are shown to the right.
L1 and 2: leucine-rich repeat domains; CR: cysteine-rich region; FN1 to 3: fibronectine type III domains;
TM: trans-membrane region; TK: tyrosine kinase domain; CT: C-terminal segment.

Figure 2. Growth curves and pedigree of the family of the patient carrying the homozygous variant
F112L/F112L. SDS: standard score deviation; NGT: naso-gastric tube.

516 Figure 3. A. Western blot showing phosphorylated-AKT (P-AKT), total AKT, and GAPDH for patients

and controls. **B.** Quantification of AKT phosphorylation calculated as: *R* after IGF-I stimulation / *R* not

518 stimulated, with R = [(P-AKTpatient/AKTpatient)/GAPDHpatient]/ [(P-

519 AKTcontrol/AKTcontrol)/GAPDHcontrol]. *p < 0.05; **p < 0.01; WT: wild type allele. Experiments

- 520 were repeated from 3 to 6 times for each individual. Error bars represent the standard error of the mean.
- 521 Del ex1-21 corresponds to Del4 in figure 1 and Del ex20-21 to Del7.

522 STATEMENTS

523 Competing interests: The authors have nothing to disclose.

525 Eloïse Giabicani: wrote the manuscript, performed the experiments, collected and analyzed the data and 526 revised the manuscript. Marjolaine Willems: collected and analyzed the data and revised the manuscript Virginie Steunou: performed the experiments. Sandra Chantot-Bastaraud: performed the experiments, 527 528 analyzed the data and helped in revising the manuscript. Nathalie Thibaud: performed the experiments 529 Walid Abi Habib: analyzed the data and helped in revising the manuscript. Salah Azzi : performed the 530 experiments, analyzed the data and helped in revising the manuscript. Bich Lam: performed the experiments, analyzed the data and helped in revising the manuscript. Laurence Bérard: collected data and 531 532 helped to revise the manuscript. Hélène Bony-Trifunovic: collected data and helped to revise the 533 manuscript. Cécile Brachet: collected data and helped to revise the manuscript. Elise Brischoux-Boucher: 534 collected data and helped to revise the manuscript. Emmanuelle Caldagues: collected data and helped to revise the manuscript. Régis Coutant: collected data and helped to revise the manuscript. Marie-Laure 535 536 Cuvelier: collected data and helped to revise the manuscript. Georges Gelwane: collected data and helped 537 to revise the manuscript. Isabelle Guemas: collected data and helped to revise the manuscript. Muriel 538 Houang: collected data and helped to revise the manuscript. Bertrand Isidor: collected data and helped to 539 revise the manuscript. Claire Jeandel: collected data and helped to revise the manuscript. James Lespinasse: collected data and helped to revise the manuscript. Catherine Naud-Saudreau: collected data 540 541 and helped to revise the manuscript. Monique Jesuran-Perelroizen: collected data and helped to revise the manuscript. Laurence Perrin-Sabourin: collected data and helped to revise the manuscript. Juliette Piard: 542 collected data and helped to revise the manuscript. Claire Sechter: collected data and helped to revise the 543 544 manuscript. Pierre-François Souchon: collected data and helped to revise the manuscript. Caroline Storey: 545 collected data and helped to revise the manuscript. Domitille Thomas: performed the experiments, 546 collected data and helped to revise the manuscript. Yves Le Bouc: collected data and helped to revise the 547 manuscript. Sylvie Rossignol: collected data and helped to revise the manuscript. Irène Netchine: analyzed the data and revised the manuscript. Frédéric Brioude: analyzed the data, wrote and revised themanuscript.



















	PATIEN	ΓS	G	FAMILY						
	NAME	SEX	NOMENCLATURE	PROTEIN	ALLELIC	SEGREGATION	MOTHER	FATHER	TARGET	TERM
ADHD: attention deficit	natient1	М	15a26 3(98954957 102461162)x1	NA	HT7	mother	-4 1	-0.5	-2 1	40
hyperactivity disorder	relative1	F	$15\alpha 26 3(98954957 \ 102461162)x1$	NA	HT7	unknown	-0.6	-1 7	-1 3	NA
BL: birth length	patient2	M	15g26.3(98610818 102397836)x1	NA	HTZ	de novo	1.9	0.8	1.4	39
BW: birth weight	natient3	M	15g26 2g26 3(98426952 102461162)x 1	NA	HT7	unknown	-2.4	1 4	-0.3	37
DVT: developmental	natient4	M	15q26 2q26 3(97450781 102461162)x1	NA	HT7	unknown	<u>_</u>). 0	2.4	13	39
GnRH: gonadotropin	patient5	F	15a26.2a26.3(98464591 101599113)x1	NA	HT7	de novo	-1.6	0.8	-0.5	37
HC: head circumference	natient6	M	15g26 3(99492046 99511073)x1	NA	HT7	unknown	NA	NA	NA	NA
HTZ cp: compound HTZ	relative6	M	15q26 3(99492046 99511073)x1	NA	HT7	father	-1.6	-3 5	-2.8	38.3
HTZ: heterozygous	patient7	F	15q26.3(99496341_99730813)x1	NA	HTZ	de novo	-1.5	-1.7	-1.7	35
IAC: interatrial communication	patient8	F	15g26.3(99229330_99290406)x1	NA	HT7	father	-1.5	-1.2	-1.5	32
min: minimum	relative8a	M	15q26 3(99229330 99290406)x1	NA	HT7	father	-1 5	-1 2	-1 2	31
NA: not available	relative8b	M	15q26.3(99229330_99290406)x1	NA	HTZ	unknown	-3.8	0	-1.7	41
POF: patent foramen ovale	patient9	M	c.118C>T	R40C	HT7	unknown	-0.9	-3.3	-2	41
rGH: recombinant	patient10	F	c.384T>C	F112L	HM7	father + mother	-2.4	-2.5	-2.6	34
growth hormone	relative10a	F	c.384T>C	F112L	HT7	unknown	0.3	-1	-0.5	NA
SDS: standard deviation score	relative10b	F	c.384T>C	F112L	HT7	unknown	-2.4	-2.5	-2.6	NA
SRS: Silver-Russell syndrome	relative10c	M	c.384T>C	F112L	HT7	unknown	-2.4	0	-1	NA
WA: weeks of amenorrhea	patient11	F	c.904G>T	F302*	HT7	unknown	-2.2	-2.5	-2.5	34
	patient12	F	c.995G>A	C332Y	HT7	father	-1.8	-1.3	-1.7	40
	relative12	M	c.995G>A	C332Y	HT7	unknown	NA	NA	NA	NA
	patient13	M	c.1247+1-1247+2insG	N417FfsX52	HTZ	mother	-2.8	-0.5	-1.4	41.5
	relative13	F	c 1247+1-1247+2insG	N417FfsX52	HT7	unknown	-2.9	-7	-2.2	NA
	natient14	M	c 2629C>T	R877*	HT7	unknown	-1 3	-1	-1	37
	natient15	M	c 3162G>A	M1054I	HT7	unknown	-0.6	-0.8	-0.6	38 5
	patient16	F	c 3454G>A	G1152R	HT7	father	-2 9	-4 1	-3.8	40
	relative16a	M	c.3454G>A	G1152R	HTZ	unknown	-2.4	-1.7	-1.8	NA
	relative16b	M	c.3454G>A	G1152R	HT7	father	-2.9	-4.1	-3.3	39
	patient17	M	c.3530G>A	R1177H	HT7	unknown	-3.2	-2.2	-2.5	38
	patient18	M	c.3539C>A/c.4055G>T	S1180Y/G1352V	HTZ comp	father + mother	-2.4	_,_ 1	-0.7	40
	relative18a	M	c.4055G>T	G1352V	HT7	unknown	NA	_ NA	NA	NA
	relative18b	F	c.3539C>A	S1180Y	HTZ	unknown	NA	NA	NA	NA
	natient19	F	c 3595G>A	G1199R	HT7	mother	-2.9	0	-1.6	38 5
	relative19	F	c.3595G>A	G1199R	HTZ	unknown	NA	NA	_,;; NA	NA
	patient20	F	c.4066G>A	E1356K	HTZ	father	-1.1	0.5	-0.4	37
	relative20	M	c.4066G>A	E1356K	HTZ	unknown	NA	NA	NA	NA
						-				

BIRTH PAR	AMETERS				rGH THER	APY		GR	OWTH			DEVELO	OPMENT				
WEIGHT	LENGTH	HC		AGE AT	HEIGHT AT		HEIGHT	HEIGHT	FINAL	ASSISTED	DVT	LANGUAGE	MOTOR	COGNITIVE	HC	FEEDING	
(SDS)	(SDS)	(SDS)	rGH	ONSET	ONSET (SDS)	DURATION	GAIN (SDS)	min (SDS)	HEIGHT (SDS)	SCHOOL	DELAY	DELAY	DELAY	DELAY	<-2 SDS	DIFFICULTIES	
-1,9	-2	-3	yes	4,1	-3,0	10,9	1,4	-3,3	-2,4	1	1	1	0	1	1	1	
NA	NA	NA	no	-	-	-	-	NA	-4,1	NA	NA	NA	NA	NA	NA	NA	
-1,4	-2,3	-1,6	no	-	-	-	-	-2,1	-2,2	1	1	1	0	1	NA	0	
-2,3	-4,2	-2,33	yes	2,9	-5,6	on going	on going	-5,7	NA	1	1	1	1	1	1	1	
-3,2	-3,1	-3	yes	1,5	-4,6	14,7	2,5	-4,6	-2,1	0	0	0	0	0	1	1	
-2,6	-3,4	-2,3	yes	3,0	-5,1	3,5	1,5	-5,3	-3,8	1	1	0	0	1	1	1	
NA	NA	NA	no	-	-	-	-	NA	NA	NA	NA	NA	NA	NA	NA	NA	
-2,1	-3,1	NA	yes	6,4	-3,1	on going	on going	-3,2	-3,5	1	1	0	0	1	1	1	
-2,5	-3,8	-2,8	no	-	-	-	-	-4,8	NA	0	0	0	0	1	1	0	
-3,5	-3,8	-2,6	yes	2,9	-3,5	11,2	2,1	-3	-1,8	1	1	1	1	1	1	1	
-2	-2,5	NA	yes	7,2	-1,6	7,2	0,7	-2,4	-1	0	1	0	1	0	0	0	
-1,4	NA	NA	no	-	-	-	-	NA	-1,2	0	0	0	0	0	0	NA	
-2,3	-2,5	NA	no	-	-	-	-	-2,7	-2,7	1	0	0	0	0	1	0	
-3,8	-4	-3,3	yes	4,1	-4,6	on going	on going	-4,5	NA	1	1	1	1	1	1	1	
NA	NA	NA	no	-	-	-	-	NA	-2,4	NA	NA	NA	NA	NA	0	NA	
NA	NA	NA	yes	11,3	-2,8	on going	on going	-2,2	NA	1	NA	NA	NA	NA	0	NA	
NA	NA	NA	no	-	-	-	-	NA	-2,5	NA	NA	NA	NA	NA	0	NA	
-1	-1,4	-0,3	yes	12,8	-4,4	2,8	0,4	-4,4	-4,2	1	1	0	0	1	1	0	
-3,2	-4,8	-3,9	yes	9,5	-2,7	on going	on going	-2,8	NA	1	1	0	1	1	1	1	
NA	NA	NA	no	-	-	-	-	NA	-1,3	NA	NA	NA	NA	NA	NA	NA	
-2,8	-3,9	NA	yes	10,4	-3,3	5,8	0,1	-3,3	-3,2	0	0	0	0	0	1	0	
NA	NA	NA	no	-	-	-	-	NA	-2,8	NA	NA	NA	NA	NA	NA	NA	
-1,8	-3,1	-2,8	no	-	-	-	-	-3	NA	NA	0	0	0	0	1	0	
-2,5	-2,9	-0,3	yes	15,3	-2,7	0,3	0,2	-2,6	-2,5	1	0	0	0	0	1	0	
-2,7	-3,3	NA	yes	4,2	-4,8	4,3	0,1	-4,7	NA	1	1	0	0	1	1	1	
NA	NA	NA	no	-	-	-	-	NA	-4,2	0	NA	NA	NA	NA	NA	NA	
-3,7	-5,2	-3,5	yes	11,8	-4,5	6,2	0,4	-4,6	-4,1	0	0	0	0	0	1	1	
-2,6	-4,2	-2,9	yes	5,5	-3,4	10,1	1,7	-3,4	-3	0	0	0	0	0	1	0	
-2,2	-2,7	-3	no	-	-	-	-	-2,5	NA	1	1	1	1	1	1	1	
NA	NA	NA	no	-	-	-	-	NA	1	NA	NA	NA	NA	NA	0	NA	
NA	NA	NA	yes	7,5	NA		NA	NA	-2,1	NA	NA	NA	NA	NA	1	NA	
-2,1	-3,7	-3,7	yes	11,7	-4,3	4,1	1,4	-4,4	-2,9	0	0	0	0	0	1	0	
NA	NA	NA	no	-	-	-	-	NA	-2,9	NA	NA	NA	NA	NA	NA	NA	
-2,8	-2,6	-3,6	no	-	-	-	-	NA	NA	NA	NA	NA	NA	NA	0	1	
NA	NA	NA	no	-	-	-	-	NA	0,5	NA	NA	NA	NA	NA	NA	NA	

CLINICS	LAS	T EVALUAT	ION	IG	F-I BEFORE	TREATM	IENT	IGF-	I DURING	TREAT	MENT	CLIN	2019)			
OTHERS		HEIGHT	HC		IGF1	IGF-I	IGF1	ACE	IGF1	IGF-I	IGF1	SCORE	BW/BL	HEIGHT	HC	IGF1
UTHERS	AGE	(SDS)	(SDS)	AGE	(ng/mL)	SDS	>0 SDS	AGE	(ng/mL)	SDS	>0 SDS	SCORE	<-1 SDS	<-2,5 SDS	<-2 SDS	>0 SDS
Hypoglycaemia	15,8	-1,6	-1,5	3,7	279	5,5	1	5,1	521	8,8	1	4	1	1	1	1
	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	1	NA	NA
Knees luxation	18,9	-2,2	NA	8,2	460	3,6	1	12,1	704	2,9	1	2/3	1	0	NA	1
IAC	12,8	-2,5	-2,8	2,9	95	0,6	1	4,2	135	1,1	1	4	1	1	1	1
Overweight	18,0	-2,1	NA	1,5	25	-2,0	0	2,5	169	2,9	1	4	1	1	1	1
GnRH analogs	15,5	-3,6	NA	2,6	138	1,1	1	4,2	306	3,7	1	4	1	1	1	1
	NA	-3,5	NA	56,2	224	1,5	1	NA	NA	NA	NA	NA	NA	NA	NA	1
ADHD	13,5	-3,1	NA	NA	NA	NA	NA	6,7	460	6,0	1	4	1	1	1	1
IAC	10,4	-4,5	-5,1	10,3	414	0,6	1	NA	NA	NA		4	1	1	1	1
	14,0	-1,4	NA	1,9	112	0,9	1	5,9	220	1,3	1	4	1	1	1	1
ADHD, cryptorchidism	19,1	-0,9	NA	7,2	142	-0,3	0	9,2	556	3,9	1	2	1	0	0	1
	49,5	-1,2	-1,6	NA	NA	NA	NA	NA	NA	NA	NA	NA	1	NA	0	NA
	17,0	-2,5	na	14,7	968	4,5	1	NA	NA	NA	NA	3	1	0	1	1
POF, hypothyroidism	5,2	-3,7	-4,9	4,0	314	4,3	1	5,2	710	9,8	1	4	1	1	1	1
	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	0	NA
ADHD	13,7	-2,2	NA	11,3	269,3	-1,2	0	11,7	582	1,6	1	1/3	NA	0	0	1
	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	1	0	NA
GnRH analogs	15,3	-4	NA	12,8	947	4,1	1	13,8	1470	7,9	1	4	1	1	1	1
ADHD, GnRH analogs	11,3	-2,2	NA	9,3	353	0,5	1	10,3	510	1,4	1	4	1	1	1	1
	NA	-1,3	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
	16,6	-3,2	NA	10,0	252	-0,2	0	11,6	449	1,3	1	4	1	1	1	1
	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	1	NA	NA
	7,1	-2,2	NA	5,4	273	3,5	1	NA	NA	NA	NA	4	1	1	1	1
Obesity	21,5	-2,5	-2,7	15,2	1260	7,1	1	16,1	908	4,4	1	3	1	0	1	1
	12,6	-4,7	na	4,2	226	2,1	1	8,1	964	7,4	1	4	1	1	1	1
Type II diabetes	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	1	NA	NA
	18,1	-4,1	-4,1	11,8	300	-0,1	0	15,7	412	0,2	1	4	1	1	1	1
	15,1	-1,7	NA	5,6	310	4,3	1	6,6	653	9,5	1	4	1	1	1	1
Pyelic dilatation	6,6	-1,6	-4,3	5,4	304	4,2	1	NA	NA	NA	NA	3	1	0	1	1
	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	0	NA
	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	1	NA
	20,8	-2,9	NA	12,9	383	-0,2	0	14,6	790	3,0	1	4	1	1	1	1
	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	1	NA	NA
Heart failure	6,2	-1,1	-1,8	0,7	151	2,0	1	NA	NA	NA	NA	3	1	1	0	1
	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

SUPPLEMENTARY DATA

Table 1. General data of the whole cohort of patients carrying an IGF1R defect.

Figure 1. Molecular strategy adopted for *IGF1R* defects diagnosis in the laboratory. SGA: small for gestational age, SRS: Silver-Russell syndrome, MLPA: multiplex ligation-dependent probe amplification. *IGF1R* defect suspicion was based on the presence of microcephaly and/or elevated IGF-I levels.

Figure 2. cDNA sequencing for the patient carrying the NM_000875.4:c.1247+1_1247+2insG variant, responsible for a one-base-pair insertion, leading to a frameshift and premature stop codon (N417EfsX52).

Figure 3. Quantification of ERK phosphorylation calculated as:

[(P-ERKpatient/ERKpatient)/GAPDHpatient]/ [(P-ERKcontrol/ERKcontrol)/GAPDHcontrol]. WT: wildtype allele. Experiments were repeated from 3 to 6 times for each individual. Error bars represent the standard error of the mean. Del ex1-21 corresponds to Del4 in figure 1 and Del ex20-21 to Del7.

Figure 4: IGF1R cDNA expression in fibroblasts. Primers used:

Igf1R-219-F: ACAGGGATCTCATCAGCTTCAC and Igf1R-219-R: TCCACCATGTTCCAGCTGTT. The amplicon length was 109 bp, spanning exons 7 and 8. Del ex1-21 corresponds to Del4 in figure 1 and Del ex20-21 to Del7.