Mosaicism for GNAS methylation defects associated with pseudohypoparathyroidism type 1B arose in early post-zygotic phases

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Abstract

Background: Pseudohypoparathyroidism type 1B (PHP1B; MIM #603233) is a rare imprinting disorder (ID), associated with the GNAS locus, characterized by parathyroid hormone (PTH) resistance in the absence of other endocrine or physical abnormalities. Sporadic PHP1B cases, with no known underlying primary genetic lesions, could represent true stochastic errors in early embryonic maintenance of methylation. Previous data confirmed the existence of different degrees of methylation defects associated with PHP1B and suggested the presence of mosaicism, a phenomenon already described in the context of other IDs.

Results: With respect to mosaic conditions, the study of multiple tissues is a necessary approach; thus, we investigated somatic cell lines (peripheral blood and buccal epithelium and cells from the urine sediment) descending from different germ layers from 19 PHP patients (11 spor-PHP1B, 4 GNAS mutated PHP1A, and 4 PHP with no GNAS (epi)genetic defects) and 5 healthy controls. We identified 11 patients with epigenetic defects, further subdivided in groups with complete or partial methylation defects. The recurrence of specific patterns of partial methylation defects limited to specific CpGs was confirmed by checking methylation profiles of spor-PHP1B patients diagnosed in our lab (n = 56). Underlying primary genetic defects, such as uniparental disomy or deletion, potentially causative for the detected partial methylation were excluded in all samples.

Conclusions: Our data showed no differences of methylation levels between organs and tissues from the same patient, so we concluded that the epimutation occurred in early post-zygotic phases and that the partial defects were mosaics. The number of patients with no detectable (epi)genetic GNAS defects was too small to exclude epimutations occurring in later post-zygotic phases, affecting only selected tissues different from blood, thus leading to underdiagnosis during routine molecular diagnosis. Finally, we found no correlation between methylation ratios, representing the proportion of epimutated cells, and the clinical presentation, further confirming the hypothesis of a threshold effect of the GNAS loss of imprinting leading to an "all-or-none" phenotype.

Keywords: GNAS, Imprinting, Mosaicism, Pseudohypoparathyroidism, Albright hereditary osteodystrophy, PTH resistance, Epigenetics, Methylation defects

Background

Pseudohypoparathyroidism type 1B (PHP1B; MIM #603233) is a rare imprinting disorder (ID) characterized by renal resistance to parathyroid hormone (PTH) in the absence of other endocrine or physical abnormalities. Renal resistance to PTH, the hallmark of PHP, was documented by markedly blunted or absent urinary phosphate and cAMP responses to PTH injection. More recently, resistance to the action of thyroid-stimulating hormone (TSH) has also been described in a large subset of patients, while growth hormone (GH) secretion appears to be conserved [1–3].

In 2000, the PHP1B phenotype was associated with a loss of imprinting (LOI), at one or more GNAS differentially methylated regions (DMRs), which disrupts the parental-specific expression of its transcripts (the isolated, at the A/B DMR or the broad, at the NESP DMR associated with the simultaneous LOM at AS, XL.
and A/B DMRs) [2]. The autosomal dominant maternally transmitted form of the disorder (AD-PHP1B) is caused by microdeletions of long-range imprinting control elements (ICRs) that regulate GNAS imprinting [4–9]. Conversely, sporadic PHP1B cases (spor-PHP1B) present a broad GNAS LOI with no known underlying genetic lesion; thus, they could represent true stochastic errors in early embryonic maintenance of methylation [10]. In a small subset of PHP1B patients, the broad GNAS LOI resulted to be secondary to a complete or segmental uniparental disomy (UPD) of chromosome 20 [11, 12].

The DNA methylation undergoes a process of erasure, acquisition, and maintenance during gametogenesis, then it is maintained during the epigenetic reprogramming in the pre-implantation embryo, and disruptions in any of these steps may lead to IDs, resulting in the aberrant development of embryogenesis, placentation, and postnatal growth [13–15].

The examination of DNA methylation patterns in embryos and in corresponding sperm samples demonstrated no methylation errors at tested imprinted loci in sperm, suggesting that the paternal transmission of epigenetic abnormalities was unlikely [16]. The protection of imprint marks should mainly rely on maternal proteins stored in the oocyte before fertilization (i.e., DPPA3, ZFP57, TRIM28, and DNMT1 gene products), because both the oocyte and the embryo are transcriptionally silent until the zygotic gene activation (ZGA), which takes place at the late four-cell stage in humans. This hypothesis was further supported by experiments in mouse embryos showing that ovarian stimulation disrupted maternal-effect gene products required for imprint maintenance [17].

The phenomenon of epigenetic mosaicism has been already described in the context of other IDs, such as Angelman syndrome (AS), Silver–Russell syndrome (SRS), Beckwith–Wiedemann syndrome (BWS), and transient neonatal diabetes (TND), and it was proposed that the absence of maternal gene products could cause complex chimeras with variable degrees of normally and aberrantly imprinted cells. Relaxation of IGF2 imprinting was previously found in BWS and in children with non-syndromic somatic overgrowth. In particular, the extent and the location of (epi)genetic mosaicism may be responsible for the phenotype heterogeneity. Additionally, in most cases of TND, the hypomethylation was mosaic and more marked in non-leukocyte than in leukocyte samples [18–20].

The description of rare cases of monozygotic twins discordant for the methylation status of specific genes, which represent examples of post-zygotic epimutations arose after the twinning event, confirmed the existence of somatic mosaicism also for epigenetic defects [21, 22]. Single-cell DNA methylation analysis revealed the presence of epigenetic chimerism in pre-implantation embryos, emerging during the early pre-implantation phase of development, which translated into incomplete methylation patterns in whole embryos. However, further studies are needed to conclusively determine the mechanisms through which such chimerism is established [23].

Previously published data confirmed the existence of different degrees of methylation defects associated with PHP1B and suggested the presence of somatic mosaicism, with the contribution in a given tissue of both normal cells and cells with epigenetic defects [10, 24, 25]. Up to now, investigations were conducted on PHP patients’ DNA samples obtained from a single tissue, mainly peripheral blood, and no specific study has been performed to confirm and characterize the mosaicism associated with partial GNAS imprinting defects.

With respect to mosaic conditions, the study of multiple tissues is a necessary approach, so we collected the genetic material of somatic cell lines descending from different germ layers (ectoderm, mesoderm, and endoderm) that eventually gives rise to tissues and organs through the process of organogenesis, and we extensively studied their genetic and epigenetic status at the GNAS locus in both PHP patients and healthy controls.

Results

We investigated genomic DNA samples obtained from different tissues of 5 healthy controls, 1 patient with PHP1A due to a large GNAS deletion, considered as a positive reference for methylation studies, and 19 PHP patients (11 spor-PHP1B, 4 GNAS-mutated PHP1A, and 4 PHP with no GNAS (epi)genetic defects).

Sanger sequencing analysis allowed to detect a point mutation in Gsa-coding exons in 4 patients (patients 12–15) and to exclude small genetic defects in the 15 other patients and 5 healthy controls. Methylation analysis identified 11 patients with broad GNAS LOI (raw methylation ratios in the Additional file 1: Table S1).

The comparison of raw methylation ratios confirmed the reproducibility and reliability of the methylation specific–multiplex ligand-dependent probe amplification (MS-MLPA) assay, allowing data validation for subsequent statistical analysis. Data obtained from the serial dilution of a sample with a GNAS maternal deletion confirmed the ability to detect different levels of methylation (Fig. 1).

After the analysis of genetic markers upstream and downstream the GNAS locus to determine whether UPD could be the primary genetic cause of the LOI, all cases resulted having at least one heterozygous VNTR in the 20q region (Additional file 1: Table S2). In patients 2 and 11, we were also able to exclude uniparental heterodisomy as parent’s DNA was available. The exclusion of
uniparental isodisomy and aneuploidies as causative genetic aberration further supported that the detection of partial LOI is derived from the presence of an epigenetic mosaicism rather than a genetic mosaicism.

Single CpG methylation ratios in the three tissues (peripheral blood and buccal epithelium and cells from the urine sediment) were the same for each subject, and in the subsequent statistical analysis, we considered mean values (Table 1 and Fig. 2).

As expected, in healthy samples and in PHP patients affected by GNAS point mutations, the NESP DMR was hemimethylated on the paternal allele, while AS, XL, and A/B DMRs on the maternal allele.

The investigation of different tissues of four patients with no GNAS (epi)genetic defects failed to find an imprinting defect in cell lineages different from leukocytes, which could explain the clinical phenotype in the absence of anomalies in blood samples.

Because of the variability among different CpG methylation ratios belonging to the same DMR in spor-PHP1B with respect to negative and positive references (NESP DMR = 0.97 ± 0.17 (mean of three CpGs), AS DMR = 0.11 ± 0.11 (mean of three CpGs), XL DMR = 0.10 ± 0.11 (mean of four CpGs), and A/B DMR = 0.05 ± 0.10 (mean of two CpGs)), we considered single CpGs instead of whole DMRs in the downstream analysis. This CpG-specific analysis (raw data in the Additional file 1: Table S1) allowed subdivision of spor-PHP1B patients in two major groups, one with a “complete” methylation defect (n = 6) and another with a “partial” methylation defect (n = 5) (Fig. 3).

The “partial” cluster could be further subdivided for the presence of a partial methylation defect limited to specific CpGs. Only one patient (patient 2) showed a partial methylation defect in all CpGs, with methylation ratios significantly different (P values < 0.01 or < 0.001) from the positive deleted control. Conversely, the other four patients displayed a partial LOM limited to the AS DMR, the XL DMR, or both the AS and the XL DMRs. In particular, patient 5 had a partial LOM at the AS DMR (AS1 = 0.20 ± 0.03 and AS3 = 0.17 ± 0.01), while patients 3 and 4 had a partial LOM at both AS and XL DMRs (AS1 = 0.21 ± 0.04/0.17 ± 0.02, AS3 = 0.21 ± 0.06/0.15 ± 0.01, XL1 = 0.20 ± 0.03/0.17 ± 0.01, XL2 = 0.22 ± 0.03/0.13 ± 0.01, XL3 = 0.27 ± 0.22/0.13 ± 0.02, and XL4 = 0.24 ± 0.24/0.09 ± 0.01) (Additional file 1: Table S1).

The methylation status of the A/B DMR of partial patients, with the exception of patient PHP2, resulted more severely affected by the LOM when compared with
| Sample     | Tissue | NESP1 Mean SD | NESP2 Mean SD | NESP3 Mean SD | AS1 Mean SD | AS2 Mean SD | AS3 Mean SD | XL1 Mean SD | XL2 Mean SD | XL3 Mean SD | XL4 Mean SD | A/B1 Mean SD | A/B2 Mean SD |
|------------|--------|---------------|---------------|---------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|--------------|--------------|
|Ctrls B (5) | 0.55   | 0.05          | 0.45          | 0.05          | 0.49        | 0.05        | 0.56        | 0.06        | 0.47        | 0.03        | 0.55        | 0.04         | 0.48         |
|Ctrls S (5) | 0.58   | 0.01          | 0.48          | 0.02          | 0.50        | 0.04        | 0.54        | 0.04        | 0.50        | 0.06        | 0.57        | 0.02         | 0.46         |
|Ctrls U (5) | 0.58   | 0.07          | 0.49          | 0.09          | 0.52        | 0.06        | 0.56        | 0.09        | 0.50        | 0.13        | 0.60        | 0.05         | 0.48         |
|Mut pts B (4) | 0.56 | 0.02          | 0.51          | 0.05          | 0.54        | 0.03        | 0.57        | 0.04        | 0.46        | 0.03        | 0.59        | 0.03         | 0.54         |
|Mut pts S (3) | 0.55 | 0.08          | 0.48          | 0.04          | 0.49        | 0.06        | 0.55        | 0.11        | 0.48        | 0.05        | 0.57        | 0.02         | 0.53         |
|Mut pts U (3) | 0.57 | 0.03          | 0.49          | 0.03          | 0.55        | 0.09        | 0.55        | 0.04        | 0.56        | 0.04        | 0.59        | 0.05         | 0.48         |
|No def pts B (4) | 0.56 | 0.06          | 0.48          | 0.06          | 0.52        | 0.04        | 0.54        | 0.08        | 0.48        | 0.06        | 0.57        | 0.03         | 0.54         |
|No def pts S (4) | 0.57 | 0.06          | 0.49          | 0.05          | 0.49        | 0.06        | 0.61        | 0.06        | 0.45        | 0.09        | 0.61        | 0.02         | 0.51         |
|No def pts U (4) | 0.58 | 0.05          | 0.47          | 0.04          | 0.46        | 0.09        | 0.60        | 0.08        | 0.55        | 0.01        | 0.61        | 0.03         | 0.56         |
|Del pt B (1)  | 1.05  | 0.08          | 0.90          | 0.02          | 1.04        | 0.15        | 0.00        | 0.00        | 0.00        | 0.00        | 0.00        | 0.07         | 0.00         |
|PHP1B pts B (11) | 1.05 | 0.13         | 0.89          | 0.11          | 0.96        | 0.12        | 0.15        | 0.07        | 0.03        | 0.06        | 0.13        | 0.08         | 0.12         |
|PHP1B pts S (10) | 1.08 | 0.16         | 0.92          | 0.13          | 0.90        | 0.12        | 0.15        | 0.07        | 0.02        | 0.05        | 0.14        | 0.09         | 0.08         |
|PHP1B pts U (11) | 1.08 | 0.19         | 0.97          | 0.22          | 0.98        | 0.16        | 0.13        | 0.07        | 0.03        | 0.06        | 0.12        | 0.11         | 0.07         |

Data in parenthesis are the number of samples reported.
B = blood, S = saliva, U = urine.
AS and XL DMRs, with the exclusion of the AS CpG number 2 whose methylation ratio was nearly 0 in all samples. This observation further supported the hypothesis of the existence of a second independent ICR regulating their epigenetic status, independently from the A/B DMR. Moreover, the gain of methylation (GOM) at the NESP DMR was complete in all partial samples, except in patient PHP2.

To confirm our findings about “partial LOM” subclusters, we checked GNAS methylation profiles obtained from genomic DNA extracted from peripheral blood samples in our series of spor-PHP1B patients diagnosed in our lab \((n = 56)\), and we observed that these patterns of partial LOM were recurrent. In particular, if we consider the whole cohort of spor-PHP1B patients \((n = 67)\), we found that the prevalence of cases with partial epigenetic anomalies was about 42%: 5 patients had a partial LOM limited to the AS DMR, 12 patients had a partial LOM limited to the XL DMR, 5 patients had partial LOMs at AS and XL DMRs, and 7 patients had partial methylation defects at all 4 DMRs (Fig. 4).

Finally, we searched for an explanation of the phenotypic variability, which means the number or severity of hormonal resistances and the presence of signs of Albright hereditary osteodystrophy (AHO), according to the methylation ratio and/or the specific affected CpGs, but we failed to find any significant difference among patients with or without AHO rather than with one or more endocrine alterations.

**Discussion**

In the present work, we investigated GNAS imprinting in DNA samples obtained from different tissues of a small series of PHP patients, as previously published data confirmed the existence of different degrees of methylation defects associated with PHP1B and suggested the presence of epigenetic mosaicism [10, 19].

The broad GNAS LOI was previously associated with complete or segmental 20qUPD in small subsets of patients, but this hypothesis was unlikely in our case series as all patients were heterozygous for at least one genetic marker, further supporting the presence of epigenetic mosaicism in subjects with partial methylation defects [11, 12].

To study mosaic conditions, the experimental approach relies on the analysis of multiple tissues from the same subject; thus, we collected the genetic material of cell lines descending from different germ layers (ectoderm, mesoderm, and endoderm). We did not observe significant differences in methylation levels in different tissues from the same patient, so we concluded that the partial LOI was a mosaic condition due to an epimutation which occurred in early post-zygotic phases, before the germ layers’ formation. In particular, in patients with partial LOI, we determined methylation ratios of 0.15–0.25, consistent with an error in methylation that occurred at the four- or eight-cell phase, i.e., the phase during which the post-zygotic genome-wide passive demethylation takes place [13–17].
Following fertilization, parental genomes undergo a global demethylation to facilitate the remodeling from two distinct differentiated gamete-specific states to a pluripotent embryonic state, with the exception of imprinted loci whose differential methylation is preserved by maternal zygotic effect genes accumulated during the oocyte growth. Maternal-effect proteins that protect imprinted methylation sites during pre-implantation development identified up to now include developmental pluripotency-associated 3 (DPPA3; also called STELLA/PGC7), zinc finger protein 57 (ZFP57), tripartite motif-containing 28 protein (TRIM28; also called KAP1/TIF1b) and DNA methyltransferase 1 (DNMT1) [16, 17]. Mutations in ZFP57 were associated with TND, the first heritable global ID compatible with life described in humans, and perhaps additional methylation anomalies of imprinted loci are not yet described [26].

Although the underlying molecular cause of spor-PHP1B is not clear, the mosaicism in cases with partial GNAS methylation defects is consistent with a failure of methylation maintenance in the early zygote, and a maternal-effect mutation might be postulated. On the other hand, in patients affected by complete broad methylation defects, an imprinting defect originating in gametes, resulting in unstable methylation of the ICRs afterwards, could be considered, but further studies to detect factors associated with the GNAS imprinting maintenance are needed. Conversely, defects affecting cells from a specific tissue would derive from molecular errors during later phases of the embryo development, but unfortunately, the number of investigated patients with no detectable GNAS (epi)genetic defects was too small to fully exclude epimutations occurring in later post-zygotic phases.

Studies in the mouse model performed to investigate structure and regulation of the GNAS-imprinted cluster identified 2 candidate ICRs with a different mode of action, one at exon A/B, acting as an insulator model, and one covering AS and XL promoters, acting through a mechanism involving antisense RNAs [27].

![Graphs showing the methylation ratio (mean blood/saliva/urine replicates ± SD) at each investigated CpG for two representative spor-PHP1B patients with partial (PHP2) and severe (PHP8) loss of imprinting (LOI) at GNAS DMRs compared to values obtained in healthy controls (negative ctrls) and the patient with the GNAS locus deletion (positive ctrl). All methylation ratios observed in spor-PHP1B patients were significantly different from those in wild type (*P < 0.001). Methylation ratios determined in spor-PHP1B patients could be not significantly different (no symbol above the column) or significantly different (*P < 0.05; **P < 0.01; ***P < 0.001) from those due to ablation of the maternal allele.](image-url)
of different patterns of mild epigenetic anomalies in our patients, in particular, partial LOMs limited to the AS DMR, the XL DMR, or both AS and XL DMRs associated with a complete LOM at the A/B DMR (18%, 12 of 67 pts), partial LOM limited to the AS DMR (7%, 5 of 67 pts), partial LOM limited to AS and XL DMRs (7%, 5 of 67 pts), and partial LOI at all 4 DMRs (10%, 7 of 67 pts).

Some DMRs, defined as germline or primary DMRs, acquire their allelic methylation during gametogenesis and are stably maintained throughout the post-zygotic development. Conversely, somatic or secondary DMRs acquire methylation during development and are regulated by nearby germline DMRs [28]. Previous studies found that AS, XL, and A/B DMRs are germline DMRs and that the AS DMR affects the somatic NESP DMR methylation status, as its antisense transcript could act in a similar manner to Air [29]. As a matter of fact, the isolated LOM at the A/B DMR, both in sporadic and inherited patients, does not affect the imprinting of the NESP DMR [30]. Our observations further supported this hypothesis because the GOM at the NESP DMR was complete also in partial patients, except for patient 2, who displayed a complete LOM at the CpG AS2, raising the possibility that this region contains a cis-acting sequence involved in the NESP DMR methylation establishment.

Finally, in our previous work, conducted in 63 patients with PHP type 1 and GNAS imprinting defects, we found no correlation between the degree of methylation defects and the severity of clinical parameters [25]. The absence of correlation between methylation ratios and the phenotype observed in the present study further confirms our previous findings and can be explained by a threshold effect leading to an “all-or-none” phenotype.

Conclusions
To conclude, further studies to unravel the mechanisms underlying GNAS LOI are needed and, in particular, it will be crucial to determine whether GNAS LOI in spor- PHP1B patients is associated with a primary or secondary epigenetic defect, with the final aim to correctly evaluate the recurrence risk of this imprinting syndrome.

Methods
The present study included 5 healthy controls, 1 subject affected by a whole GNAS locus deletion (g.56′657′752_59′228_953del, previously described in reference 34), used as positive reference for methylation defects at GNAS DMRs, and 19 PHP patients (11 with methylation defects at the GNAS locus, 4 with point mutations in GNAS Gsa-coding exons, and 4 with no GNAS/PDE4D/PRKAR1A (epi)genetic defects and no deletions of the long arm of chromosome 2). Some of these PHP patients (1–6, 9–10, 12–14, and 16–17) were presented in our previous works [25, 28, 29, 31]. The clinical diagnosis was based upon the presence or absence of specific clinical and biochemical signs, i.e., hypocalcemia, hyperphosphatemia, and raised serum PTH levels in the absence of vitamin D deficiency and/or typical AHO manifestations, considered major and minor criteria for the diagnosis of PHP [32–35]. Clinical details, including some additional patient-specific features, and molecular diagnosis are summarized in Table 2. Informed consent for genetic and epigenetic studies was obtained from all subjects involved in the study.

Genomic DNA was extracted from samples of the peripheral blood (Nucleon BACC2 genomic DNA purification kit, cod.RPN8502, GE Healthcare), saliva, and urine (Puregene Core kit, cod. 158622, Qiagen), in order to obtain the genetic material from somatic cell lines descending from different germ layers. Buccal epithelial cells are of ectodermal derivation and urine sediment cells are derived from the mesoderm, while leukocytes are from the endoderm. Mutation analysis was performed by direct sequencing (BigDye™Terminator v3.1 Cycle Sequencing Kit, cod.4337456, Applied
Biosystems) of GNAS 1–13 exons and flanking intronic sequences (ENSBMBL ID: ENSG00000087460). Informative genetic markers in the 20q region were evaluated to exclude a possible misdiagnosis of spor- PHP1B in the presence of aneuploidies (monosomy or trisomy) or uniparental disomy (UPD). According to sample availability, the analysis was performed in blood, saliva, and urine samples. For variable number tandem repeat (VNTR) genotyping, we set up a cost-effective three-primer approach based on the simultaneous use of a couple of sequence-specific primers, the reverse one containing a poliA tail at the 5’ end to allow easier allele scoring, associated with a fluorescently labeled universal forward M13-tailed FAM oligonucleotide. All primers and experimental conditions are available upon request. Copy number variant (CNV) analysis of STX16 and GNAS loci and methylation analysis of GNAS DMRs were performed by MS-MLPA (cod. ME031-100R+ EK1-FAM, MRC-Holland) (methylation-specific probe location, GR37/hg19, summarized in Additional file 1: Figure S1). MS-MLPA data analysis was performed using the Coffalyser.Net (MRC-Holland). The detection of expected MS-MLPA methylation ratios of about 0.5 reflected a normal hemimethylated status in healthy controls (considered as a negative reference), while finding of MS-MLPA methylation ratios of about 1 and 0 reflected the apparent LOI in the GNAS-locus-deleted subject (considered as positive reference). At least two technical replicates were performed for each biological sample. A serial dilution, in triplicate, of GNAS-deleted DNA (0, 12.5, 25, 37.5, 50, 62.5, 75, 87.5, and 100%) in wild type DNA was created to define the MS-MLPA detection range of different percentages of epigenetic mosaics. Statistical analysis was performed using GraphPad Prism version 5 for Windows (www.graphpad.com). The exploratory data analysis included means ± SD, median, and interquartile ranges for values measured in the blood, saliva, and urine. Based on data distribution, means ± SD were used for further comparisons. One-way ANOVA with Bonferroni’s Multiple Comparison test was used to evaluate the intercluster variability at single CpGs. After Bonferroni correction, a P value < 0.05 was deemed to indicate statistical significance. Continuous variables, i.e., hormonal and biochemical parameters, were reported as means ± SD, whereas dichotomous variables, i.e., the presence of AHO signs, were expressed as proportions. Differences between means and proportions were checked by

### Table 2: Clinical characteristics and molecular analysis of patients included in the present study

| pt ID | GNAS point mut | GNAS meth def | Sex | PTH | TSH | SS | Ob | RF | Br | MR/BD | OS | Additional features |
|-------|----------------|---------------|-----|-----|-----|----|----|----|----|-------|----|---------------------|
| 1     | No             | Yes           | F   | 183 | 2.7 |    |    |    |    |       |    | Long QT syndrome    |
| 2     | No             | Yes           | M   | 899 | 2.6 | Yes|    |    |    |       |    | Hypertension, Fahr syndrome, and renal damage |
| 3     | No             | Yes           | M   | 258 | 4.1 |    |    |    |    |       |    |                     |
| 4     | No             | Yes           | F   | 215 | 2.5 |    |    |    |    |       |    |                     |
| 5     | No             | Yes           | M   | 454 | 4.9 |    |    |    |    |       |    |                     |
| 6     | No             | Yes           | M   | 114 | 5.4 | Yes |    |    |    |       |    | Subclinical hypothyroidism |
| 7     | No             | Yes           | M   | X   |     | Yes|    |    |    |       |    | Long QT syndrome |
| 8     | No             | Yes           | F   | 152 | 0.29| Yes|    |    |    |       |    | Leukopenia, normocytotic microcromic anemia, autoimmune primary hypothyroidism, and osteopenia |
| 9     | No             | Yes           | M   | 288 | 7   |    |    |    |    |       |    | Asperger syndrome |
| 10    | No             | Yes           | F   | 615 | 1.8 |    |    |    |    |       |    | Asthenia associated with lipothyria, genu valgo, and hypertension |
| 11    | No             | Yes           | M   | 338 | 2.1 | Yes|    |    |    |       |    | LGA at birth |
| 12    | Yes            | No            | F   | 373 | 8.34| Yes|    |    |    |       |    | Polandes cyst, aortic stenosis, rGHrH, and genu valgo |
| 13    | Yes            | No            | F   | 236 | 1.49| Yes| Yes|    |    |       |    | Irregular menses and hypertension |
| 14    | Yes            | No            | F   | 6    | 4.48| Yes|    |    |    |       |    |                     |
| 15    | Yes            | No            | F   | 138 | 34  | Yes|    |    |    |       |    | Oligohydramnios, congenital hyperthyroidism, rickets, spastic diplegia, and amenorrhea |
| 16    | No             | No            | F   | 106 | 7.9 | Yes| Yes| Yes| Yes|       |    | Cholelithiasis, hypoadrenalism |
| 17    | No             | No            | M   | 77  | 4   | Yes|    |    |    |       |    | Congenital adrenal hyperplasia |
| 18    | No             | No            | F   | 77.3| 1.65| Yes|    |    |    |       |    | Hashimoto thyroiditis, favism, and multiple fractures |
| 19    | No             | No            | F   | 37  | 1   | Yes|    |    |    |       |    | Oligomenorrhea |

Age (years at diagnosis)
PHT PTH serum levels, normal range 10–65 pg/mL, TSH TSH serum levels, normal range 0.4–3.9 mU/L, X elevated, but value not available, Ob obesity, RF round face, MR/BD mental retardation and/or behavioral defects, Br brachydactyly, OS ectopic ossifications, SS short stature, LGA large for gestational age
Student’s t and chi-squared tests, respectively. In particular, for the epigenotype–phenotype correlation investigation, we considered the following clinical parameters: the precocity of the disease (defined as the age at diagnosis), the severity of hormonal resistances (defined as PTH, calcium, phosphorus, and TSH levels at diagnosis), and the severity of AHO, defined as the number of AHO signs.

**Additional file**

**Additional file 1: Figure S1.** Methylation-specific probes location, GR37/hg19. Table S1. Raw methylation ratios. Table S2. Variable number tandem repeats (VNTRs) in the 20q region analysed to exclude uniparental disomy (UPD). (DOXC 378 kb)

**Abbreviations**

AHO: Albright hereditary osteodystrophy; AS: Angelman syndrome; BWS: Beckwith–Wiedemann syndrome; cAMP: Cyclic adenosine monophosphate; DMRs: Differentially methylated regions; Gαi: Growth hormone; GOM: Gain of methylation; Gsα: Alpha subunit of the stimulatory G protein; ICRs: Imprinting control elements; ID: Imprinting disorder; LOI: Loss of imprinting; LOM: Loss of methylation; MS-MLPA: Methylation-specific-multiplex ligand-dependent probe amplification; PHP: Pseudohypoparathyroidism; PTH: Parathyroid hormone; SD: Standard deviation; SRS: Silver–Russell syndrome; TND: Transient neonatal diabetes; TSH: Thyroid-stimulating hormone; UPD: Uniparental disomy; VNTR: Variable number tandem repeat; ZGA: Zygotic gene activation

**Acknowledgements**

The authors are members of and acknowledge the EuroPHP network (EuroPHP) and the EUCID.net (COST action BM1208 on imprinting disorders; www.imprinting-disorders.eu).

**Competing interests**

The authors declare that they have no competing interests.

**Ethics approval and consent to participate**

Informed consent was obtained from all patients (or legal guardians for minors) and relatives included in the present study. All procedures were performed in compliance with relevant legislation and institutional guidelines.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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**Received:** 19 August 2017 **Accepted:** 30 January 2018

**Published online:** 06 February 2018

**References**

1. Barret D, Breslau NA, Wax MB, Molinoff PB, Downs RW Jr. New form of pseudohypoparathyroidism with abnormal catalytic adenyl cyclase. Am J Phys. 1989;257(2 Pt 1):E277–83.
2. Liu J, Litman DM, Rosenberg MJ, Yu S, Bieseker LG, Weinstein LS; A GNAS1 imprinting defect in pseudohypoparathyroidism type IB. J Clin Invest. 2000; 106(9):1167–74.
3. Mantovani G, Bondioni S, Linglart A, Maghnie M, Cistermino M, Corbetta S, Lania AG, Beck-Peccoz P, Spada A. Genetic analysis and evaluation of resistance to thyrotropin and growth hormone-releasing hormone in pseudohypoparathyroidism type IB. J Clin Endocrinol Metab. 2007;92: 3738–42.
4. Bastepe M, Frohlich LF, Hendy GN, Indridason OS, Josse RG, Kohiyama H, Korkko J, Nakamoto JM, Rosenblloom AL, Syper AH, et al. Autosomal dominant pseudohypoparathyroidism type Ib is associated with a heterogeneous microdeletion that likely disrupts a putative imprinting control element of GNAS. J Clin Invest. 2003;112(8):1253–63.
5. Bastepe M, Frohlich LF, Linglart A, Abu-Zara HS, Tojo K, Ward LM, Juppner H. Deletion of the NESP55 differentially methylated region causes loss of maternal GNAS imprints and pseudohypoparathyroidism type IB. Nat Genet. 2005;37(1):25–7.
6. Linglart A, Gensure RC, Olney RC, Juppner H, Bastepe M. A novel STX16 deletion in autosomal dominant pseudohypoparathyroidism type IB redefines the boundaries of a cis-acting imprinting control element of GNAS. Am J Hum Genet. 2005;76(3):804–14.
7. Chilliambhi S, Turan S, Hwang D, Chen H, Juppner H, Bastepe M. Deletion of the noncoding GNAS antisense transcript causes pseudohypoparathyroidism type IB and biparental defects of GNAS methylation in cis. J Clin Endocrinol Metab. 2010;95(8):3993–4002. https://doi.org/10.1210/jc.2009-2205.
8. Richard N, Abeguillé G, Coudray N, Mifttie H, Gruchy N, Andrieux J, Cathebras P. Epigenetic reprogramming in mammals. Hum Mol Genet. 2005;14:R47–53.
9. Morgan HD, Santos F, Green K, Dean W, Reik W. Epigenetic reprogramming in mammals. Hum Mol Genet. 2005;14:R47–58.
10. Shi L, Wu J. Epigenetic regulation in mammalian preimplantation embryo development. Reprod Biol Endocrinol. 2009;7. https://doi.org/10.1186/1477-7827-7-59.
11. Shi X, Chen S, Zheng H, Wang L, Wu Y. Abnormal DNA methylation of imprinted loci in human preimplantation embryos. Reprod Sci. 2014; 21(8):978–83.

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17. Denomme MM, Mann MR. Maternal control of genomic imprint maintenance. Reprod BioMed Online. 2013;27(6):629–36. https://doi.org/10.1016/j.rbmo.2013.06.004.

18. Okaïmoto K, Morison IM, Taniguchi T, Reeve AE. Epigenetic changes at the insulin-like growth factor II/H19 locus in nephron development is an early event in Wilms tumorigenesis. Proc Natl Acad Sci U S A. 1997;94(10):5367–71.

19. Itoh N, Becroft DM, Reeve AE, Morison IM. Proportion of cells with paternal 11p15 uniparental disomy correlates with organ enlargement in Wiedemann-Beckwith syndrome. Am J Med Genet. 2000;92(2):111–6.

20. Mackay DJ, Boonen SE, Clayton-Smith J, Goodship J, Hahnenmann JM, Kant SG, Njølstad PR, Robin NH, Robinson DO, Siebert R, et al. A maternal hypomethylation syndrome presenting as transient neonatal diabetes mellitus. Hum Genet. 2006;120(2):262–9.

21. Galetzka D, Hansmann T, El Hajj N, Weis E, Irscher B, Ludwig M, Schneider-Ratzke B, Kohlschmidt N, Beyer V, Bartsch O, et al. Monozygotic twins discordant for constitutive BRCA1 promoter methylation, childhood cancer and secondary cancer. Epigenetics. 2012;7(1):47–54. https://doi.org/10.4161/epi.7.1.18814.

22. Bliek J, Elders M, Maas SM, Oostra RJ, Mackay DM, van der Lip K, Callaway JL, Brooks A, van’t Padde S, Westerveld A, et al. Lessons from BWS twins: complex maternal and paternal hypomethylation and a common source of haematopoietic stem cells. Eur J Hum Genet. 2009;17(12):1625–34. https://doi.org/10.1038/ejhg.2009.77.

23. Lorthongpanich C, Cheow LF, Balu S, Quake SR, Knowles BB, Burkholder WF, Solter D, Messerschmidt DM. Single-cell DNA-methylation analysis reveals epigenetic chimerism in preimplantation embryos. Science. 2013;341(6150):1110–2. https://doi.org/10.1126/science.1234617.

24. Maupertii-Mehouas S, Marot V, Reynes C, Bertrand G, Feillet F, Carel JC, Simon D, Bihan H, Gajdos V, Devouge E, et al. Quantification of the methylation at the GNAS locus identifies subtypes of sporadic pseudohypoparathyroidism type Ib. J Med Genet. 2014;51(1):55–63. https://doi.org/10.1136/jmg.2010.081356.

25. Elli FM, de Sanctis L, Bollati V, Tarantini L, Filopanti M, Barbieri AM, Peverelli E, Beck-Peccoz P, Spada A, Mantovani G. Quantitative analysis of methylation defects and correlation with clinical characteristics in patients with pseudohypoparathyroidism type I and GNAS epigenetic alterations. J Clin Endocrinol Metab. 2014;99(3):E508–17. https://doi.org/10.1210/jc.2013-3086.

26. Mackay DJ, Callaway JL, Marks SM, White HE, Acerini CL, Boonen SE, Dayanikli P, Firth HV, Goodship JA, Haeeners AP, et al. Hypomethylation of multiple imprinted loci in individuals with transient neonatal diabetes is associated with mutations in ZPFP. Nat Genet. 2008;40(9):949–51. https://doi.org/10.1038/ng.187.

27. Williamson CM, Ball ST, Nottingham WT, Skinner JA, Plagge A, Turner MD, Powles N, Hough T, Papworth D, Fraser WD, et al. A cis-acting control region is required exclusively for the tissue-specific imprinting of Gnas. Nat Genet. 2004;36(8):894–9.

28. Court F, Tayama C, Romanelli V, Martin-Trujillo A, Iglesias-Platas I, Okamura K, Sugahara N, Simon C, Moore H, Harness JV, et al. Genome-wide parent-of-origin DNA methylation analysis reveals the intricacies of human imprinting and suggests a gemline methylation-independent mechanism of establishment. Genome Res. 2014;24(4):554–69. https://doi.org/10.1101/gr.164913.113.

29. Haywood BE, Bonthron DT. An imprinted antisense transcript at the human GNAS1 locus. Hum Mol Genet. 2000;9(5):835–41.

30. Elli FM, Linglart A, Garin I, de Sanctis L, Bordogna P, Grybek V, Pereda A, Giachero F, Vennia E, Hanna P, Mantovani G, Perez de Sanclares G. The prevalence of GNAS deficiency-related diseases in a large cohort of patients characterized by the EuroPh phosphosensor. J Clin Endocrinol Metab. 2016;101(10):3657–68.

31. Garin I, Elli FM, Linglart A, Silve C, de Sanctis L, Bordogna P, Pereda A, Clarke JT, Kannengisser C, Coutant R, Tenebaum-Rakover Y, Admoni O, de Sanclares GP, Mantovani G. Novel microdeletions affecting the GNAS locus in pseudohypoparathyroidism: characterization of the underlying mechanisms. J Clin Endocrinol Metab. 2015;100(4):E681–7. https://doi.org/10.1210/jc.2014-3098.

32. Elli FM, de Sanctis L, Cecolini B, Barbieri AM, Bordogna P, Beck-Peccoz P, Spada A, Mantovani G. Pseudohypoparathyroidism type Ia and pseudopseudohypoparathyroidism: the growing spectrum of GNAS inactivating mutations. Hum Mutat. 2013;34(3):411–6.

33. Elli FM, Bordogna P, de Sanctis L, Giachero F, Vennia E, Segni M, Mazzanti L, Boldrin V, Toromanovic A, Spada A, et al. Screening of PRKAR1A and PDE4D in a large Italian series of patients clinically diagnosed with Albright hereditary osteodystrophy and/or pseudohypoparathyroidism. J Bone Miner Res. 2016;31(6):1215–24. https://doi.org/10.1002/jbmr.2785.

34. Mantovani G, Spada A, Elli FM. Pseudohypoparathyroidism and Gao-cAMP-linked disorders: current view and open issues. Nat Rev Endocrinol. 2016;12(6):347–56. https://doi.org/10.1038/nrendo.2016.52.

35. Thiele S, Mantovani G, Barlier A, Boldrin V, Bordogna P, De Sanctis L, Elli FM, Freson K, Garin I, Grybek V, et al. From pseudohypoparathyroidism to inactivating PTH/PTHrP signalling disorder (iPPSD), a novel classification proposed by the EuroPhP network. Eur J Endocrinol. 2016;175(6):P1–P17.