

ARTICLE

A mild form of Mucopolysaccharidosis IIIB diagnosed with targeted next-generation sequencing of linked genomic regions

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Next-generation sequencing (NGS) techniques have already shown their potential in the identification of mutations underlying rare inherited disorders. We report here the application of linkage analysis in combination with targeted DNA capture and NGS to a Norwegian family affected by an undiagnosed mental retardation disorder with an autosomal recessive inheritance pattern. Linkage analysis identified two loci on chromosomes 9 and 17 which were subject to target enrichment by hybridization to a custom microarray. NGS achieved 20-fold or greater sequence coverage of 83% of all protein-coding exons in the target regions. This led to the identification of compound heterozygous mutations in *NAGLU*, compatible with the diagnosis of Mucopolysaccharidosis IIIB (MPS IIIB or Sanfilippo Syndrome type B). This diagnosis was confirmed by demonstrating elevated levels of heparan sulphate in urine and low activity of α -N-acetyl-glucosaminidase in cultured fibroblasts. Our findings describe a mild form of MPS IIIB and illustrate the diagnostic potential of targeted NGS in Mendelian disease with unknown aetiology. *European Journal of Human Genetics* (2012) 20, 58–63; doi:10.1038/ejhg.2011.126; published online 29 June 2011

Keywords: next-generation sequencing; DNA target enrichment; sequence capture; Sanfilippo Syndrome type B; mucopolysaccharidosis; *NAGLU*

INTRODUCTION

The advent of next-generation sequencing (NGS) techniques is expected to accelerate the identification of mutations causing Mendelian disorders¹ and both whole-genome and whole-exome approaches have proven successful in recent studies.^{2–6} In each case, massively parallel sequencing has produced a large amount of sequence information and identified large numbers of novel genetic variants of unknown significance, which makes the identification of the causal mutation challenging. So far, the application of NGS for diagnostic purposes is already established, but the choice of approach used to identify disease causing mutation(s) remains a challenge. This report illustrates the potential and power of using family data.

Mucopolysaccharidosis IIIB (MPS IIIB) (Sanfilippo Syndrome type B; OMIM 252920) is an autosomal recessive disorder caused by an enzyme deficiency of α -N-acetyl-glucosaminidase due to mutations in the *NAGLU* gene (NM_000263).⁷ This enzyme is involved in the degradation of the glycosaminoglycan compound heparan sulphate and impaired function leads to the accumulation of partially degraded mucopolysaccharides in tissues, and increased excretion of these compounds in urine. The disorder is clinically heterogeneous and > 100 different *NAGLU* mutations have been reported.⁸ The clinical picture varies, but is usually characterised by developmental delay,

behavioural abnormalities and sleep disturbance in early childhood. Progressive dementia and coarsening of facial features evolve slowly. As initial symptoms and signs of MPS IIIB might be subtle and non-specific during childhood, the diagnosis may be easily overlooked in its early years by the child neurologist. When these patients are subsequently transferred to the care of adult medicine, it is our experience that the search for an aetiological diagnosis is less ambitious and that such patients may remain undiagnosed despite neurological deterioration.

The Norwegian family presented in this paper demonstrated an unusually mild form of MPS IIIB. Urine analysis performed during the primary investigations of the family 10 years before reinvestigation failed to detect abnormal amounts of glycosaminoglycans. As the family remained undiagnosed on reinvestigation, linkage analysis in combination with targeted DNA capture and NGS was applied to obtain the diagnosis. The results are presented here.

PATIENTS AND METHODS

Patients

The subject family contained eight siblings, of whom four (three males and one female) were affected by an unknown genetic disorder, later diagnosed as MPS IIIB as described below (Figure 1). Medical data on affected individuals were

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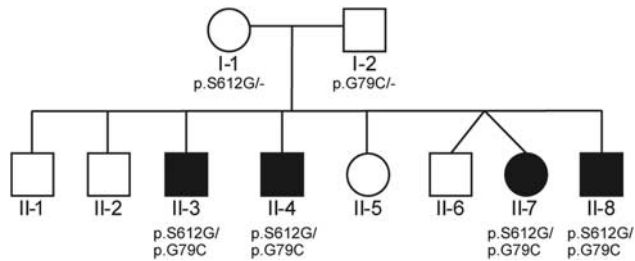


Figure 1 Pedigree of the family.

compiled from several different sources: Medical history taken in conversation with the parents, healthy siblings and other relatives, notes on numerous visits to the local nursery, medical charts from hospital admissions, reports from ophthalmological examination with funduscopy, results on cerebral CT and MRI examinations, and clinical neurological examination performed on all siblings.

Linkage analysis

All 10 members of the family were genotyped using the Affymetrix GeneChip Human Mapping 10K 2.0 array,⁹ following the suppliers protocol. Washing and staining were performed with the Affymetrix Fluidics Station 450 (Affymetrix Inc., Santa Clara, CA, USA) and the signal intensities detected with the GeneChip Scanner 3000 7G (Affymetrix Inc., Santa Clara, CA, USA). The results from the Affymetrix arrays were analysed in GeneChip Genotyping Analysis Software (GTYPE)¹⁰ and GeneChip Operating Software (GCOS). Data handling, Mendelian error control and statistical analyses were done using Progeny Lab software (version 5, Progeny Software, LLC, South Bend, IN, USA), PEDSTATS¹¹ and MERLIN.¹²

Target capture by DNA microarray hybridisation

Based on the linkage results, exons defined in Ensembl release 48 were selected from three adjacent regions on chromosome 17 (NCBI Build 36.1/hg18 chromosome coordinates 28 813 215–28 942 222; 29 615 779–31 041 457; 36 815 436–39 115 918). Exon target regions were extended to include flanking sequence such that each target was no shorter than 500 bp. Based on this target list, a custom Sequence Capture 385K human array was designed and manufactured by Roche Nimblegen Inc. (Madison, WI, USA). Due to the presence of repetitive elements, not all desired regions (hereafter referred to as 'targets') could be represented on the array (represented regions hereafter are referred to as 'tiled regions'). The array contained oligonucleotide probes of length 60–90 nucleotides with 3 bp spacing. A total of 526 210 bp were tiled. The tiled region covered 86% of all target exon sequences (90% of all targeted coding exons). Target and tiled regions are available in BED file format upon request. Similar target regions from within a second linkage peak on chromosome 9 were tiled on a second array (details available upon request).

Twenty micrograms of genomic DNA from individual II-4 (Figure 1) was sonicated using a Bioruptor sonicicator (Diagenode s.a., Liege, Belgium) to an average size of 400 bp. DNA ends were blunted by incubation with Klenow polymerase before ligation to oligonucleotide linker (previously prepared by annealing the following two oligonucleotides: LM-PCR-A 5' P-GAGGATCCAG AATTCTCGAGTT and LM-PCR-B 5' CTCGAGAATTCTGGATCCTC). Linker dimers were removed by purifying the ligated DNA using AMPure beads (Beckman Coulter Inc., Brea, CA, USA). DNA was hybridised to the arrays on a MAUI hybridisation station (Biomicro Systems, Salt Lake City, UT, USA) according to Roche Nimblegen instructions for 65 h. Arrays were washed according to Roche Nimblegen instructions. Captured DNA was eluted from arrays by incubating at 95 °C for 5 min with 500 μ l water under a coverslip with gasket in a microarray hybridisation chamber (Agilent Technologies Inc., Santa Clara, CA, USA). Eluted DNA was concentrated using a YM-100 microcon concentrator (Millipore, Billerica, MA, USA) before LM-PCR amplification using the above LM-PCR-B oligonucleotide. Enrichment efficiency was verified using selected PCR amplifiers chosen to represent exons targeted for enrichment (data not shown). Before sequencing, the majority of linker sequence was removed by *Bam*HI digestion.

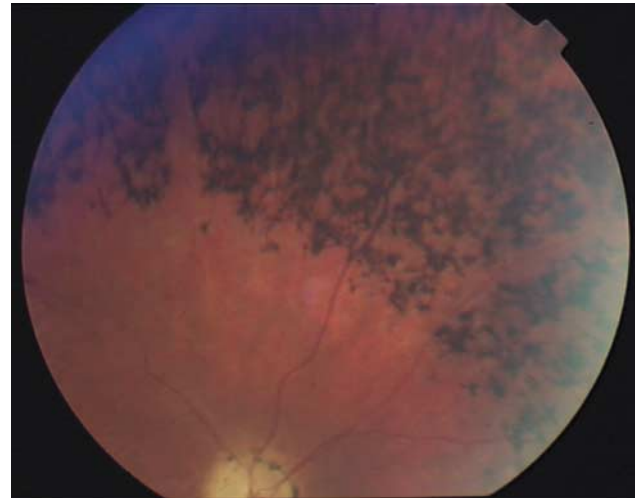


Figure 2 Fundus photography of individual II-4 illustrating retinitis pigmentosa with dark pigment deposits, optic disc pallor and thinning of the arteries.

Next-generation sequencing

Two micrograms of enriched DNA was used to generate a library and run on a single lane of an Illumina GAIIX sequencer (Illumina Inc., San Diego, CA, USA), with paired-end reads of 50 bp, following the manufacturer's instructions. Image analysis and base calling were performed using Illumina's RTA software version 2.4 (Illumina Inc.) and pipeline software version 1.4 (Illumina Inc.). Reads were filtered to remove those with low base call quality using Illumina's default chastity criteria. We used MAQ¹³ to map the sequence reads to chromosome 17, assemble alignments and call single-nucleotide polymorphisms (SNPs). SNPs were characterised (presence in dbSNP build 129, non-coding/coding, non-synonymous/synonymous) using SNPnexus.¹⁴ Custom Perl scripts were used to identify regions with zero coverage, and prepare BED files for viewing data on the University of California at Santa Cruz (UCSC) genome browser (<http://genome.ucsc.edu>). Fold enrichment of the regions targeted on the entire array was calculated using the formula used by Volpi *et al*¹⁵: $(\Sigma R_{EMT_{Tm}}/S_{Tm})/(\Sigma R_{MG}/S_G)$, where $\Sigma R_{EMT_{Tm}}$ is the number of reads mapping to the repeat-masked target region, S_{Tm} is the size of the repeat-masked target region, ΣR_{MG} is the number of reads mapping uniquely to the human genome and S_G is the size of the human genome.

Sanger sequencing

In addition, both the GC-rich first exon of *NAGLU* and the mutation identified by NGS in the sixth exon of *NAGLU* were verified by Sanger sequencing using PCR amplification and an Applied Biosystems (Foster City, CA, USA) 3730 DNA Analyzer. The GC-rich first exon of *NAGLU* was amplified using AmpliTaq Gold 360 Master Mix (Applied Biosystems). Primers are available upon request.

RESULTS

Clinical data

The four affected individuals went to special schools, but acquired skills sufficient to handle their own money, run errands in the grocery store and keep a simple job under supervision. From their mid-twenties their cognitive and motor functions started to deteriorate. Night vision became increasingly impaired due to retinitis pigmentosa (Figure 2). Ambulation became gradually restricted not only because of retinitis pigmentosa, but also due to cerebellar and sensory ataxia, and three of the four affected siblings had to use a wheelchair. Speech was slowly lost as they developed repetitive and stereotypic behaviour with sudden aggressive outbursts. A tendency for incontinence for urine and faeces was present from childhood. Cerebral CT showed

gross cortical atrophy and marked thickening of the cranial bones. Cerebral MRI demonstrated gross cortical atrophy, dilatation of the ventricles and mild cerebellar atrophy (Figure 3). Further details of the clinical phenotype of each patient can be found in Table 1.

Linkage results

Linkage analysis, assuming a fully penetrant recessive mode of inheritance and a frequency of 0.0001 of the disease allele, identified two loci with the maximum expected LOD score of 2.3 on chromosomes 9 and 17 (Figure 4). Maximum LOD score results on the remaining chromosomes were all negative. The intervals 9p21.3-p22.3 and 17q11.2-q12 contained nearly 22 Mb of sequence and 450 genes in total according to the Ensembl genome browser (<http://www.ensembl.org>).

Targeted DNA capture and NGS

Targeted high-throughput resequencing was performed for selected exons within both the chromosome 9 and 17 linkage peaks on DNA from a single patient (see Materials and methods). Chromosome 9 regions were captured on a custom microarray and sequenced on half a PicoTiterPlate of a Roche-454 GS FLX Titanium instrument; however, no candidate mutations likely to explain the patients' phenotype were discovered (data not shown). Selected exons from within the chromosome 17 linkage peak were similarly enriched on a custom microarray and sequenced using a single lane of Illumina sequencing. These regions encompassed 126 known and predicted genes, of which 109 were protein coding. A total of 1115 exons were targeted, of which 1023 were finally represented by 930 tiled regions totalling 526 210 bp. Protein-coding bases totalled 140 202 bp. Performance of the capture array and Illumina sequencing are detailed in

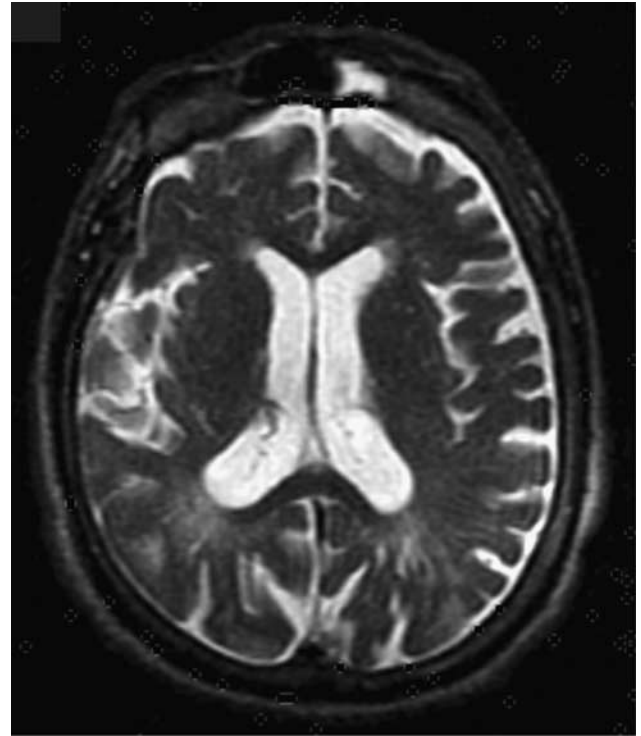


Figure 3 Cerebral axial T2-weighted MRI of individual II-4 demonstrating cortical atrophy and widening of the lateral ventricles. Note hypo-intensive foci in periventricular white matter, possibly representing small cysts or dilated perivascular spaces filled with mucopolysaccharides or cerebrospinal fluid.

Table 1 Clinical information on affected individuals

	II-3	II-4	II-7	II-8
Sex	Male	Male	Female	Male
Year of birth	1949	1952	1956	1963
Walked at age	Before 18 months	18 months	Normal	Normal
Language development	Slow	Sentences at 4 years	Normal	Slow
Education/employment	Regular primary school at 7 years, special school from 12 years. Manual work until 35 years	Regular primary school at 7 years, special school from 10 years. No schooling after 13 years. Manual work until 35 years	Regular primary school at 7 years, later transferred to special school. Never employed	Special school from 9 years. Manual work until 40 years
Tonsillitis	Numerous tonsillectomy at 8 years	Numerous tonsillectomy at 6 years	NA	Numerous
Vision/eye findings	Impaired dark vision	RP at 39 years (Figure 2)	Optic disc pallor and loss of pigment in the periphery of the retina	RP identical to II-4. Impaired dark vision at 20 years
Behaviour	Sleep disturbance and aggressive behaviour recorded at 40 years. Stereotypic and compulsive walking ^a	Anxious, restless and aggressive with stereotypic movements recorded at 39 years	Restless with compulsive walking. Aggressive in adult life	Emotionally unstable, psychotic episodes. Sleeping difficulties. Rapid cognitive decline last 2 years
Neuroimaging (CT or MRI)	NA	Cortical and central atrophy with periventricular white matter cysts (MRI) (Figure 3)	Cortical and central atrophy (CT)	Cortical and central atrophy with cysts in corpus callosum (MRI)
Other neurological findings	Gait ataxia, hyperreflexia and inverted plantar responses	Gait ataxia; dorsal column and peripheral sensory component. Plantar responses inverted	Gait ataxia. Deep tendon reflexes were hyperactive and plantar responses inverted	Gait ataxia. Deep tendon reflexes were hyperactive and plantar responses inverted
Other clinical findings	NA	NA	Epilepsy	NA

Abbreviations: NA, not available; RP, retinitis pigmentosa; CT, computed tomography; MRI, magnetic resonance imaging.

^aFor the past 20 years, he has been walking the same circle 12 h per day, touching doorframes that need a new coat of paint every second year.

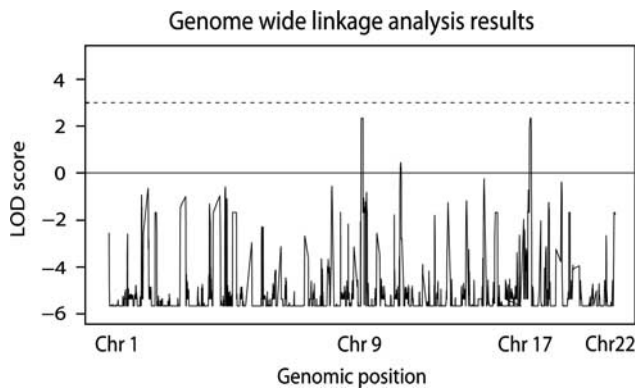


Figure 4 LOD score results from the genome-wide linkage analysis. Two regions on chromosomes 9 and 17 reached the maximum estimated LOD score of 2.3.

Table 2 Performance of the capture array and the Illumina sequencing for the chromosome 17 linked region

<i>Sequencing output (entire array)</i>	
Number reads passing filters	22 173 516
Reads mapping uniquely to human genome	20 203 662 (91%)
Reads mapping multiple times to human genome	1 113 480 (5%)
Reads failed to map to human genome	856 374 (4%)
<i>Capture array performance (entire array)</i>	
Number unique reads mapping to tiles	14 260 070 (70.5%)
<i>Sequence depth (chr17 tiled regions only)^a</i>	
Number unique reads mapping to tiles	5 076 025
Sequence generated over tiled regions	210 442 925 bp
Tile median coverage depth	424-fold
Tiled coding exon median coverage depth	356-fold
Tiled base coverage ≥ 20 -fold	485 835 of 526 210 (92.3%)
Tiled bases with zero coverage	22 283 of 526 210 (4.2%)
Coding exon bases coverage ≥ 20 -fold	131 118 of 140 202 (93.5%)
Coding exon bases with zero coverage	4 729 of 140 202 (3.4%)

^aNote that the array also contained chr14 sequences not relevant to this particular family, and that the sequence depth figures given here correspond only to the chr17 sequences referred to in this article.

Table 2, and compares favourably with similar published experiments with $\sim 70\%$ reads on target and 92% of tiled regions covered by ≥ 20 -fold sequencing depth. Applying the formula for calculating enrichment,¹⁵ the array achieved 1744-fold enrichment of target regions over the input genomic DNA (see Materials and methods). In summary, of all protein-coding exons within the linkage peak, 83% were captured and sequenced to ≥ 20 -fold sequencing depth. A total of 13.1% had no sequence coverage, either being unsuitable for tiling due to repetitive elements (10%) or due to technical limitations of the capture and sequencing technologies (3.1%).

Within the tiled regions, a total of 1168 single-nucleotide variants (SNVs) were identified, each with a coverage depth of 10-fold or greater. Initially, we removed all SNVs that overlapped with known SNPs present in the dbSNP build 129 database, after which 575 SNVs remained (Supplementary File 1). Of 75 SNVs affecting coding sequence, 47 were synonymous variants. Of the remaining novel missense or non-sense variants, only a single SNV appeared in a homozygous state. However, this homozygous SNV caused a conservative valine-glycine shift in an unconserved amino-acid position of the *NLE1* gene, and as a result was not considered a likely causative

mutation. The search for mutations was, therefore, extended to include further possibilities such as splice site variations and compound heterozygous variants. While considering the possibility that the *NAGLU* gene could be a potential causative gene in a compound heterozygous state due to the presence of two missense mutations in exon 6, it was noticed that the first exon contained a 460-bp sequence coverage gap in a region of high GC content. The *NAGLU* gene was considered a prime candidate based on phenotype descriptions in OMIM (MIM ID #252920), so this gap was sequenced by traditional Sanger sequencing and a third missense SNV was detected. Sanger sequencing of exon 1 and parts of exon 6 in the parents allowed identification of the parental source of each mutation. Two missense mutations are detailed as follows in the reference sequence NM_000263 (one of the SNVs seen in the Illumina data was not replicated by traditional Sanger sequencing):

Paternal: c.235G>T/p.G79C (in exon 1)

Maternal: c.1834A>G/p.S612G (in exon 6)

Crucially, both of these mutations have been previously described as causing MPS IIIB /Sanfilippo Syndrome type B. Sequence depth over the *NAGLU* gene, including the gap region where the paternal mutation was later found by Sanger sequencing is depicted in Figure 5. Sequence traces of Sanger sequencing used to confirm the Illumina data and cover the gap are included as insets, with their locations on the gene indicated.

We examined the effect of GC content and capture tile size on sequence coverage, and observed that tiles with extremes of low or high GC content suffered from reduced sequence coverage. In addition, tile size was an important factor determining coverage, with small tile size (≤ 100 bp) associated with low coverage depth (Figure 6a). When focusing on tiles containing sequencing gaps, it is clear that small tiles can be poorly covered irrespective of GC content, but when tile GC content exceeds 60%, coverage in larger tiles also deteriorates (Figure 6b).

Biochemical verification of MPS IIIB diagnosis

To verify the diagnosis of Sanfilippo Syndrome type B, urine was analysed from all affected family members. Increased levels of secreted glycosaminoglycans were observed by the dimethylmethylene blue test, and the presence of excess heparan sulphate was confirmed by thin layer chromatography (data not shown). In addition, α -N-acetylglucosaminidase activity in cultured fibroblasts derived from individual II-4 was found to be 3% of normal control levels (data not shown), thus confirming the diagnosis.

DISCUSSION

The affected individuals described in this paper were diagnosed with MPS IIIB by genome-wide linkage analysis and subsequent NGS of the linked regions. The lack of NGS sequence coverage observed in the first exon of the *NAGLU* gene, which prevented diagnosis based on NGS data alone, is typical of GC-rich first exons, as has been observed before.¹⁶ Regions of low or zero sequence coverage typically corresponded to GC-rich genomic areas or small target regions. These regions limit capture (poor hybridisation to small fragments) and sequencing (GC-rich regions remain challenging for sequencing-by-synthesis technology). GC-rich regions may also elute less efficiently from arrays, further contributing to their low sequence coverage. Developments in sequencing technology such as direct single molecule sequencing may remove much of this bias in the future.¹⁷ However, in the time elapsed since the experiments presented here were performed, some improvements in coverage have been obtained simply by limiting the number of PCR cycles employed during the procedure.¹⁸

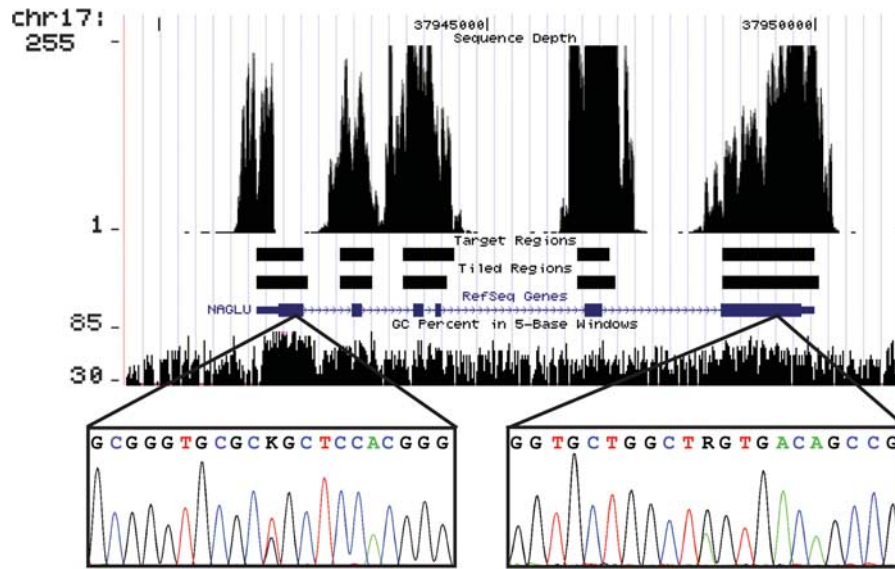


Figure 5 Screen shot from UCSC genome browser showing the *NAGLU* gene structure in blue (<http://genome.ucsc.edu/>) and GC content from 30 to 85%. Custom tracks were added to show sequence coverage (capped at 255-fold for ease of display), targeted exons and tiled regions covered on the sequence-capture array. Insets show Sanger sequencing traces from the regions containing the paternal (on the left; c.235G>T) and maternal (on the right; c.1834A>G) mutations. Note the coincidence of high GC content with zero NGS sequence coverage.

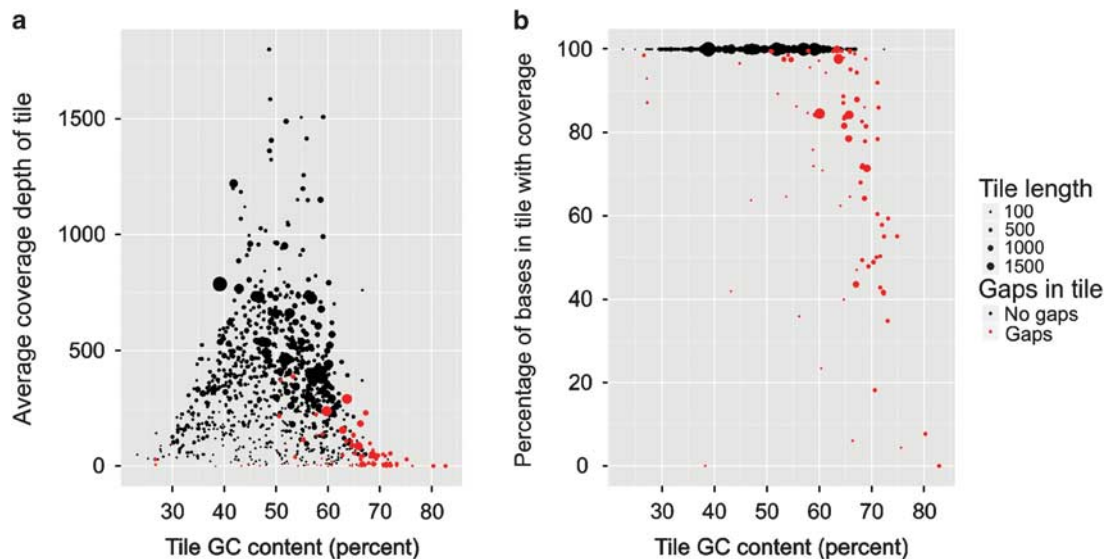


Figure 6 Effect of tile GC content and size on sequence coverage depth of all tiled regions. (a) Effect of tile length and GC content on tile average coverage depth. (b) Effect of tile length and GC content on percent of tile bases with ≥ 1 -fold coverage.

We identified two compound heterozygous mutations in the *NAGLU* gene, the paternal c.235G>T/p.G79C and maternal c.1834A>G/p.S612G. Both mutations have previously been identified in MPS IIIB patients. The p.S612G mutation has been identified in both compound heterozygous and homozygous states, all producing attenuated phenotypes, whereas the p.G79C mutation has been identified in a homozygous state in a severely affected individual.^{19–21} The patients reported here are mildly affected and this supports the hypothesis that particular mutations, including p.S612G, seem to reduce the clinical severity of the disease.^{21,22} Even in the presence of the p.G79C mutation known to produce a severe phenotype, the allele with the p.S612G mutation seems to be able to provide a residual enzyme activity sufficient to mitigate the course of the disease.

The patients described here range in age from 47 to 61 years. The clinical course of the disease was quite similar in all four, being

remarkably mild the first two decades of life with stable, but reduced intellectual ability. In their third decade of life, however, they all showed a marked decline both cognitively and physically. Three of them were diagnosed with retinitis pigmentosa. The phenotype resembles previously reported MPS IIIB patients with an attenuated form, in particular the ones known to have at least one allele with the p.S612G mutation.²¹ Since the paternal allele p.G79C has previously been associated with the severe phenotypic form of MPS IIIB, it is likely that the attenuated form seen in the patients described here is due to residual enzyme activity derived largely from the maternal p.S612G allele. This is consistent with the observations of Valstar *et al*,²¹ who reported four patients carrying this allele in compound heterozygote and one homozygous carrier, all of whom had the attenuated MPS IIIB disease form. In the patient's cells examined in this study, α -N-acetyl-glucosaminidase activity was only 3% of normal

levels. This activity was probably produced by the p.S612G allele and may be of clinical relevance. Like most MPS IIIB mutations, p.S612G does not affect the active site of the enzyme, but is rather expected to affect the folding and/or stability of the enzyme.²³ As such, the attenuated form of the disease may be the most responsive to therapies based on 'chemical chaperones' which may hold promise if toxicity issues can be resolved.²³

Clinical and allelic heterogeneity are common features in many Mendelian diseases. A selection bias of severely affected individuals and families in genetic studies leads to disease descriptions and diagnosis criteria which are appropriate for the more severe forms of the disease spectrum. The recent paper by Valstar *et al*²¹ suggests that this is also the case in MPS IIIB. In an overview of the clinical history and molecular basis of an unbiased cohort of 44 Dutch patients with MPS IIIB, they demonstrate that as much as 79% had the attenuated form, even though the general notion has been that the severe form is the most frequent. Such a selection bias in medical genetic research makes the clinical and genetic diagnostics of the milder phenotypes challenging. However, with the advent of several genome-wide techniques, in particular NGS and exome sequencing, genetic diagnostics will no longer have to rely on the candidate gene approach, facilitating the diagnosis of a greater proportion of mildly affected individuals with genetic disorders with unspecific manifestations. The approach presented here clearly demonstrates the valuable diagnostic potential of NGS and also the utility and power of additional family information.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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