







METHODS

Beyond copy number: A new, rapid, and versatile method for sequencing the entire *SMN2* gene in SMA patients

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Abstract

Spinal muscular atrophy (SMA) is caused by bi-allelic loss or pathogenic variants in the *SMN1* gene. *SMN2*, the highly homologous copy of *SMN1*, is considered the major phenotypic modifier of the disease. Determination of *SMN2* copy number is essential to establish robust genotype–phenotype correlations and predict disease evolution, to stratify patients for clinical trials, as well as to define those eligible for treatment. Discordant genotype–phenotype correlations are not uncommon in SMA, some of which are due to intragenic *SMN2* variants that may influence the amount of complete *SMN* transcripts and, therefore, of full-length *SMN* protein. Detection of these variants is crucial to predict SMA phenotypes in the present scenario of therapeutic advances and with the perspective of SMA neonatal screening and early diagnosis to start treatments. Here, we present a novel, affordable, and versatile method for complete sequencing of the *SMN2* gene based on long-range polymerase chain reaction and next-generation sequencing. The method was validated by analyzing samples from 53 SMA patients who lack *SMN1*, allowing to characterize paralogous, rare variants, and single-nucleotide polymorphisms of *SMN2* as well as *SMN2*–*SMN1* hybrid genes. The method identifies partial deletions and can be adapted to determine rare pathogenic variants in patients with at least one *SMN1* copy.

KEYWORDS

next-generation sequencing, paralogous variants, phenotype–genotype correlations, *SMN2* copies, spinal muscular atrophy

1 | INTRODUCTION

Spinal muscular atrophy (SMA) is the second most common recessive genetic disease of infancy and early childhood, with an incidence of 1 in 5000–10,000 live births and a worldwide carrier frequency of 1:51

(Sugarman et al., 2012). SMA patients are classified into different clinical groups based on the age of onset, clinical severity, and achieved motor milestones. In the most severe form, type I SMA, patients are never able to sit and generally die of respiratory failure before the age of 2 years. Intermediate type II SMA patients are able to sit but never walk, thus

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being confined to wheel-chair. Type III patients walk unassisted but may lose this ability during infancy or adolescence (Wang et al., 2007; Zerres & Rudnik-Schöneborn, 1995).

Bi-allelic absence or pathogenic variants of the *Survival of Motor Neuron 1* (*SMN1*) gene cause SMA (Lefebvre et al., 1995). A centromeric and nearly identical paralog, *SMN2*, encodes in principle the same protein as *SMN1* (Lefebvre et al., 1995; Monani et al., 1999; Rochette et al., 2001). However, a silent transition within exon 7 of the *SMN2* gene causes exon skipping and results in a truncated, nonfunctional variant (*SMN-Δ7*) (Lorson et al., 1999). It has been estimated that each *SMN2* copy can produce only around 10% to 15% of functional SMN protein, depending on the cells and tissues studied (Boza-Morán et al., 2015; Soler-Botija et al., 2005; Wirth et al., 2013).

The number of *SMN2* copies and the presence of intragenic *SMN2* variants are known modifiers of SMA disease severity (Bernal et al., 2010; Prior et al., 2009; Ruhno et al., 2019). Indeed, numerous studies show that the higher the *SMN2* copy number, producing larger amount of full-length SMN protein, the milder the associated SMA phenotype and vice versa. However, this inverse correlation is not absolute (Calucho et al., 2018). Whereas the determination of *SMN2* copy number is widely implemented to study SMA patients, the actual structures and genomic sequences of *SMN2* copies are usually not included in the characterization of SMA patients.

The current scenario of SMA therapy is rapidly evolving due to the approval in the last years of nusinersen/Spinraza, an antisense-tailored therapy (Finkel et al., 2017), AVXS101/Zolgensma, an adeno-associated viral-based gene therapy (Mendell et al., 2017), as well as the recent approval of the first oral drug to treat SMA, risdiplam/Evrysdi (<http://www.fda.gov>). However, these disease-modifying therapies are expensive treatments, and their efficacy needs to be periodically assessed. Although responses to treatment vary in SMA patients, it is not yet known whether specific features of *SMN2* are correlated with these responses (Cuscó et al., 2020). Thus, it becomes crucial to investigate genomic *SMN2* data to better characterize SMA patients and accurately predict disease evolution.

Here, we report a novel method for sequencing the whole *SMN2* gene based on long-range polymerase chain reaction (PCR) and next-generation sequencing (NGS). The method allows determining all variants described so far as disease modifiers in SMA patients without *SMN1* as well as to identify new variants and structural changes. Furthermore, the technique can be adapted to determine rare pathogenic variants in heterozygous patients with at least one *SMN1* copy. Inclusion of this technique in the routine diagnosis of SMA patients is expected to improve individual genotype-phenotype correlations and, therefore, to help predict more accurately the evolution of the disease.

2 | MATERIALS AND METHODS

2.1 | Patients

We studied 53 genetically confirmed SMA patients with homozygous absence of *SMN1*. The vast majority of studied patients had three

SMN2 gene copies ($n = 51$), but the cohort included one patient each with two and four *SMN2* copies, respectively. In addition, samples from three patients who carry one *SMN1* copy were studied to assess the versatility of the method to detect *SMN1*-specific pathogenic variants. These patients were: patient SMA54, who had a heterozygous deletion of *SMN1*, the pathogenic variant c.399_402del (p.(Glu134Serfs*14)) in the other *SMN1* allele, and three *SMN2* copies; patient SMA55, who had a heterozygous deletion of *SMN1*, the pathogenic variant c.815A>G (p.(Tyr272Cys)) in the other *SMN1* allele, one complete *SMN2* copy, and two partial *SMN* copies comprising exons 1 to 6 (also known as *SMN1/2Δ7-8* deletion, Arkblad et al., 2006), and SMA55F, father of SMA55 harboring 2 *SMN1* copies (one with the variant c.815A>G), and two *SMN2* copies.

Genetic confirmation of SMA by bi-allelic defects in *SMN1* (Alías et al., 2009) as well as *SMN2* copy number determination by multiplex ligation-dependent probe amplification (MLPA) were carried out as previously described (Alías et al., 2011). Patients were classified as I, II, or III according to their severity and motor milestone achievements. One patient presenting with type 0 (congenital) SMA was also studied. All patients were unrelated with the exception of the SMA55/55F pair mentioned above (a child with type 0 SMA and his father) and two pairs of siblings, SMA17/SMA18 (both type II) and SMA27/SMA51 (types II and III, respectively). DNA samples were obtained from peripheral blood. All participants or their legal guardians signed written informed consent. The study was approved by the Ethical Committee of our Hospital (PR(AG)229/2018).

2.2 | PCR design and library preparation

We studied the complete genomic *SMN2* sequence, including promoter, 5'-UTR and 3'-UTR regions. To this end, we designed three overlapping PCRs (~12-kb each) to amplify a target region of approximately 31.5 Kb (chr5:69,342,511-69,374,064). These long-range PCRs were successfully set up using TaKaRa LA Taq[®] DNA polymerase (#RR002A; Takara Bio). Given the high homology between *SMN1* and *SMN2* sequences, primers were not specific to *SMN2*, but the 53 patients analyzed have zero *SMN1* copies, allowing in principle an *SMN2* exclusive analysis. In addition, samples from three patients with at least one *SMN1* copy were analyzed. Primer sequences and PCR conditions are given in Table S1.

After amplification, the concentration of the three PCR products was measured using Qubit 2.0 Fluorometer (Thermo Fisher Scientific) and mixed equimolarly. One thousand nanogram of the obtained mixture was fragmented with NEBNext[®] dsDNA Fragmentase[®] (New England Biolabs) to generate DNA fragments of ~200 bp. Then, the NEBNext Ultra DNA library prep kit for Illumina and NEBNext[®] MultiplexOligos for Illumina[®] Dual Index Primers Set 1 (New England Biolabs) were used to generate the libraries. The necessary purifications and size selections were performed using AMPure XP beads (Beckman Coulter).

The quality and size of the libraries were assessed using QIAxcel (Qiagen) and were quantified with Qubit. Finally, the

libraries of all patients were equimolarly mixed and sequenced using a 500-cycle MiSeq reagent kit v2 with a paired-end run of 2×251 bp reads in a MiSeq instrument (Illumina). The number of patients included in each run was calculated to ensure a minimum coverage of $\times 400$. All procedures were performed following the manufacturer's instructions.

In samples harboring at least one copy of *SMN1*, pathogenic variants were ascribed to *SMN1* using a long-range PCR and Sanger sequencing of *SMN1* as previously described (Kubo et al., 2015), with slight modifications.

2.3 | Bioinformatics analysis

The data analysis pipeline included the quality trimming of Illumina sequences using Trimmomatic (Bolger et al., 2014). Sequences were mapped to an artificial genome reference that contains only the *SMN2* coordinates, based on the reference genome (UCSC hg 19 version, build 37.1). This strategy avoids arbitrarily reporting only one of the possible alignments produced by most mapping algorithms and therefore avoids dispersion of the read-depth signal from all *SMN* genes. This approach increases the power to detect small changes of coverage and variants with lower AB ratios, which are characteristic of heterozygous variants in multicopy regions. Results show a single location per each read that corresponds to any of the *SMN* genes. Mapping was performed using burrows-wheeler aligner (BWA)-align and BWA-sample with default parameters. Variant calling was performed with Genome Analysis Toolkit (GATK) Unified Genotyper and Haplotype Caller (McKenna et al., 2010), and variant annotation with ANNOVAR (Codina-Solà et al., 2016; Wang et al., 2010).

2.4 | Genetic variant types

We discriminated the genetic variants identified in three categories: (1) paralogous sequence variants (PSVs), which are positions differing between duplicated genes (in this context, variants differing between *SMN1* and *SMN2*); (2) rare single-nucleotide variants (SNVs), which are those present in less than 1% of the population or never described; and (3) single-nucleotide polymorphisms (SNPs), which are variations in a single position present in over 1% of the population. Determining whether the variants found are SNPs, SNVs, or PSVs allowed us to evaluate the specificity of the method in the detection of *SMN1* versus *SMN2* and to identify hybrid structures in the samples. This classification is not related to a possible effect of variants on *SMN2* expression and phenotype, a topic out of the scope of this article.

PSV ratios were computed as quotients of the number of reads belonging to a given functional copy and the total coverage.

Mutation nomenclature refers to GenBank NC_000005.9, RefSeq NM_000344.3 for *SMN1*, and RefSeq NM_017411.4 for *SMN2*.

3 | RESULTS

In all analyzed samples, a mean depth of coverage of $1720 \times$ ($435 \times$ – $7478 \times$) was obtained. This high coverage allows to accurately determine the allelic frequency of the detected variants in each sample (AB ratio). Given that the copy number of *SMN2* genes had been already studied by MLPA, we could use the AB ratio to calculate the number of copies in which the detected variant was present.

To assess the specificity of our method to identify genetic variants in the 5q13 complex region, we first analyzed PSVs that are specific for the *SMN2* gene. To this end, a list of *SMN1* and *SMN2* PSVs was created based initially on a literature review (Monani et al., 1999), and expanded with the information obtained from a basic local alignment search tool-like alignment tool (BLAT) between the two genes using the reference genome Hg19. The detailed information obtained from sequencing the *SMN2* gene in samples from 53 patients was used to determine and eventually confirm which positions should be considered real PSVs or SNPs or SNVs. Based on previous reports and BLAT data, we initially considered 22 PSVs (Table 1). Four of these variants exhibited high variability between samples, indicating that they should not be considered genuine PSVs (by definition). Two other variants, including the candidate position g.69371981A/C previously described as a PSV (Monani et al., 1999) and g.69367553G/A were not found neither in *SMN1* nor *SMN2* genes from our samples and were considered as very infrequent variants. The remaining 16 changes are genuine PSVs between *SMN1* and *SMN2* genes, being c.835-1606C/T in intron 6 considered here for the first time as a PSV (Table 1). These PSVs were present in all the reads and in more than 95% of the samples, confirming that all these positions were specific for *SMN2*. However, in two patients (SMA04 and SMA39) we observed some discrepancies in the PSV ratio compatible with the presence of hybrid genes. Both patients presented six *SMN1* PSVs all located in intron 6 (Chr5: 69370451- 69370895). The PSV ratio indicates that in SMA04, two of the three *SMN* copies are *SMN2*-*SMN1* hybrids, while in SMA39, only one of the three copies is an *SMN2*-*SMN1* hybrid (Figure 1).

We also detected variants previously described as beneficial for *SMN2* function (Bernal et al., 2010; Prior et al., 2009; Ruhno et al., 2019; Wu et al., 2017). Using the AB ratio data, we were able to confirm the presence of the c.859G>C (p.(Gly287Arg)) variant in patient SMA52 with two *SMN2* copies in the heterozygous state and of c.835-44A>G (g.69372304A>G, commonly known as A-44G) in one of the three *SMN2* copies of patient SMA21 (Figure 2 and Table S2). Furthermore, in two sisters with different phenotypes (SMA27, type II and SMA51, type III) and three *SMN2* copies, we found the PSV conversion c.835-1897C>T (g.69370451C>T) in one of their three alleles. In our series, we did not identify any of the other more recently published variants considered as candidates to be modifiers (Ruhno et al., 2019; Wadman et al., 2020).

Finally, to verify whether our technique could also be extended to identify *SMN1* variants, we studied samples from three SMA patients harboring at least one *SMN1* copy. The PSV genotype in all three cases was in agreement with the copy numbers previously determined by MLPA. SMA54 (1_ *SMN1*/3_ *SMN2* MLPA genotype) showed the *SMN1* PSVs at a frequency of $\sim 25\%$ (one of the four *SMN* copies of the

TABLE 1 The 22 candidate positions for paralogous sequence variants (PSVs) between *SMN1* and *SMN2* are shown

SMN1 position	SMN2 position	Gene location	Ref SMN1	Ref SMN2	Categorization	Conversion nomenclature SMN2>SMN1	Source of information
70231509	69356085	Intron 1	G	A	SNP ^a	c.82-3157A>G	BLAT
70240028	69364605	Intron 4	G	A	SNP ^a	c.628-457A>G	BLAT
70242435	69367010	Intron 6	T	C	SNP ^a	c.834+432C>T	BLAT
70242978	69367553	Intron 6	A	G	SNV ^b	c.834+975G>A	BLAT
70244142	69368717	Intron 6	A	G	SNP ^a	c.834+2139G>A	Monani et al. /BLAT
70245876	69370451	Intron 6	T	C	PSV	c.835-1897C>T	Monani et al. /BLAT
70246016	69370591	Intron 6	G	A	PSV	c.835-1757A>G	Monani et al. /BLAT
70246019	69370594	Intron 6	T	C	PSV	c.835-1754C>T	Monani et al. /BLAT
70246156	69370731	Intron 6	G	A	PSV	c.835-1617A>G	Monani et al. /BLAT
70246167	69370742	Intron 6	T	C	PSV	c.835-1606C>T	BLAT
70246320	69370895	Intron 6	G	A	PSV	c.835-1453A>G	Monani et al. /BLAT
70246793	69371368	Intron 6	G	A	PSV^c	c.835-980A>G	Monani et al. /BLAT
70246872	69371448	Intron 6	-	AGGCA	PSV^c	c.835-900_835-896del	Monani et al. /BLAT
70246919	69371499	Intron 6	A	C	PSV^c	c.835-849C>A	Monani et al. /BLAT
70247219	69371799	Intron 6	G	A	PSV^c	c.835-549A>G	Monani et al. /BLAT
70247290	69371870	Intron 6	T	C	PSV	c.835-478C>T	Monani et al. /BLAT
70247401	69371981	Intron 6	C	A	SNV ^{b,c}	c.835-367A>C	Monani et al. /BLAT
70247724	69372304	Intron 6	G	A	PSV^c	c.835-44A>G	Monani et al. /BLAT
70247773	69372353	Exon 7	C	T	PSV^c	c.840T>C	Monani et al. /BLAT
70247921	69372501	Intron 7	A	G	PSV^c	c.*3+100G>A	Monani et al. /BLAT
70248036	69372616	Intron 7	A	G	PSV^c	c.*3+215G>A	Monani et al. /BLAT
70248501	69373081	Exon 8	G	A	PSV^c	c.*239A>G	Monani et al. /BLAT

Note: These positions were obtained from a previous bibliographic compilation (Monani et al., 1999) and were complemented with a BLAT between the two genes, as deposited in the reference genome Hg19. Repetitive regions (polyA, polyT, and polyGT) were discarded. The candidate positions were genotyped in the patients studied in this study ($n = 53$) and in samples with at least one *SMN1* ($n = 3$) to check for consistency. From the 22 candidates, six were discarded as PSVs, four of them are listed as SNPs instead, and the remaining two as rare SNV. Therefore, a total of 16 nucleotides (in bold in the Table) differentiate *SMN1* and *SMN2* genes, 10 of which had been previously described and validated, 5 had been described but not validated, and 1 is considered here as a PSV for the first time.

Abbreviations: BLAT, BLAST-like alignment tool; SNP, single-nucleotide polymorphism; SNV, single-nucleotide variant.

^aThese positions show high variability between samples and were therefore classified as SNPs (including position c.835-367C/A previously validated by Monani et al., 1999).

^bIn these positions, the same nucleotide has always been found in both *SMN1* and *SMN2*. Thus, G>A and A>C exchanges appears to be very rare SNV found in the reference genome.

^cThese 11 positions were previously classified as bona fide PSVs, after being tested in a control population of 15 individuals (Monani et al., 1999).

patient). The *SMN1* PSV frequency in SMA55 and SMA55F was ~50% in line with their genotypes (1_*SMN1*/1_*SMN2* and 2_*SMN1*/2_*SMN2*, respectively). A similar consistency was obtained with *SMN1* pathogenic variants. Thus, c.399_402del (p.(Glu134Serfs*14)) in SMA54 showed a frequency of ~25%, while c.815A>G (p.(Tyr272Cys)) showed a frequency of ~50% in SMA55 and of 25% in SMA55F. *SMN1*-specific Sanger sequencing (Kubo et al., 2015) revealed that patients SMA54 and SMA55 present this variant in hemizygous state, whereas in SMA55F it was detected in heterozygous state (Figure 3).

In addition, in patient SMA55, it was also possible to corroborate the presence of the common partial *SMN1*/2Δ7-8 deletion of the 3' region previously detected by MLPA (Figure 4). Indeed, analysis of the AB ratios of all SNVs revealed a discrepancy between the 5' and 3' *SMN* region, showing ratios compatibles with four copies (25%–75%) in the 5' region (from promoter to exon 6), but compatible with two copies (50%–100%) in the 3' region confirming the presence of the two alleles with the *SMN1*/2Δ7-8 deletion (further explanation in Figure 4).

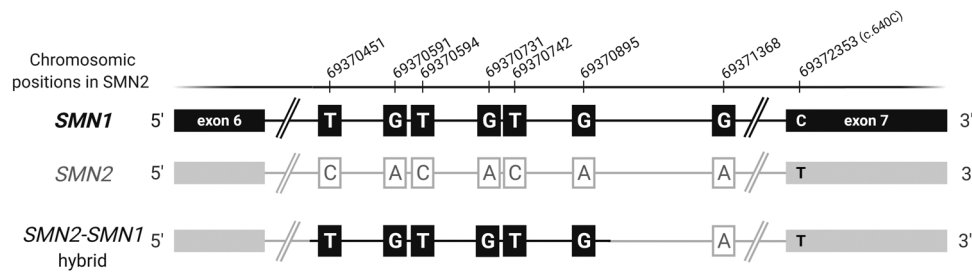


FIGURE 1 Structure of the *SMN2-SMN1* hybrid detected in patients SMA4 and SMA39. PSVs were genotyped in all patients. In SMA4 and SMA39, six *SMN1* PSVs located in intron 6 (Chr5: 69370451-69370895) were detected, which indicates the presence of hybrid genes. The AB ratio indicates that in SMA04, two of the three copies are *SMN2-SMN1* hybrids (*SMN1* PSVs in 66%) while in SMA39, only one of the three copies is an *SMN2-SMN1* hybrid (*SMN1* PSVs in 33%). PSV, paralogous sequence variant

4 | DISCUSSION

We have developed a novel, practical method for the genomic analysis of both *SMN1* and *SMN2* regions. The technique focuses on the genomic characterization of each *SMN2* copy in SMA patients regardless of the gene copy number and can also be applied to detect pathogenic variants in heterozygous SMA patients with at least one *SMN1* copy.

Currently published methods to study the complete sequence of the *SMN2* gene include whole genome sequencing (WGS) and multiplexed direct genomic selection (MDiGS) sequencing (Chen et al., 2020; Ruhno et al., 2019). WGS is a rather expensive and laborious technique and usually needs a complex bioinformatics analysis. In MDiGS, whole DNA is prepared in libraries, target regions are captured by bacterial artificial chromosomes (BAC) probes of *SMN* (not specific for *SMN1* or *SMN2*), as well as of *CFTR* and *PLS3*, for quantitative comparison. Thus, the hybridization step is rather large and complex. By contrast, the new method described here based on long-range PCR and NGS can be easily implemented in any genetics laboratory performing NGS applications.

The main advantages of our method, specifically designed to deal with the complexity of the SMA region, include accessible cost, relative simplicity, and speedy results (usually obtained in around 3 days). This practical protocol can easily genotype both PSVs and rare variants already described in SMA patients, but it might also help to identify new variants and *SMN2-SMN1* hybrid genes as part of more investigative and personalized approaches. The high coverage obtained at the nucleotide level with the new method allows the determination of the number of copies in which a specific variant is present, which, in turn, is essential to characterize the genomic architecture of each *SMN2* copy. Indeed, although the method does not directly quantify *SMN2* copy numbers, calculation of the number of copies and the presence of partial *SMN* genes is straightforward using the allelic frequencies of the variants and might be used to, for example, confirm results reported by other methods, such as MLPA.

Employing this method to samples from SMA patients, we elaborated an updated list of 16 PSVs between *SMN1* and *SMN2*, including one previously never described. PSV genotyping would help to detect hybrid genes and to discover new, potentially relevant conversions that

	<i>SMN2</i> status	NGS result (IGV image)	AB Ratio Expected (Mut/Total)
SMA52	<i>mut</i> AAGAACGAAGG <i>wt</i> AAGAAGGAAGG	 C = 352 (56%) G = 276 (44%)	1/2 = 0.5
SMA21	<i>mut</i> TCTATGTCTAT <i>wt</i> TCTATATCTAT <i>wt</i> TCTATATCTAT	 G = 278 (36%) A = 503 (64%)	1/3 = 0.33

FIGURE 2 Utility of AB ratios to calculate the number of the copies in which variants are present. Patient SMA52 has two *SMN2* copies (determined by MLPA) and the variant NM_017411.4:c.859G>C (p.(Gly287Arg)) was detected with a frequency of 56%, in agreement with the AB ratio expected for the variant in one over two alleles. Patient SMA21 has three *SMN2* copies (determined by MLPA), and the variant c.835-44A>G (NC_000005.9: g.69372304A>G) was detected with a frequency of 36%, in agreement with the AB ratio expected for the variant in one over three alleles. MLPA, multiplex ligation-dependent probe amplification

Patient	SMA55	SMA55F
AB ratio expected	Total: 1/2 = 0.5 <i>SMN1</i> : 1/1 = 1	Total: 1/4 = 0.25 <i>SMN1</i> : 1/2 = 0.5
<i>SMN1</i> - <i>SMN2</i> status	<i>SMN1</i> GCTGTCA <i>SMN2</i> GCTATCA	GCTGTCA <i>SMN1</i> GCTATCA <i>SMN1</i> GCTATCA <i>SMN2</i> GCTATCA <i>SMN2</i>
<i>SMN1</i> - <i>SMN2</i> NGS (IGV image)	G= 3725 (52%) A= 3483 (48%) G C T A T C A	G= 1338 (25%) A= 3915 (74%) G C T A T C A
<i>SMN1</i> specific PCR (Sanger electropherogram)	Hemizygous state	Heterozygous state

FIGURE 3 Detection of the pathogenic variant NM_017411.4:c.815A>G in samples SMA55 and SMA55F. Patient SMA55 has one *SMN1* copy (with the variant c.815A>G) and one *SMN2* copy, while his father (SMA55F) has two *SMN1* (one copy with the variant c.815A>G) and two *SMN2* copies (determined by MLPA and Sanger). The pathogenic variant c.815A>G was detected in SMA55 and SMA55F through NGS with a frequency of 52% and 25%, respectively. The *SMN1*-specific PCR performed confirms that the pathogenic variant c.815A>G is present in *SMN1* since we observed the variant in hemizygous status in SMA55 and in heterozygous status in SMA55F. MLPA, multiplex ligation-dependent probe amplification; NGS, next-generation sequencing; PCR, polymerase chain reaction

might act as disease modifiers (Ruhno et al., 2019; Wu et al., 2017). Our method is highly specific given that it allows detection of positive modifiers, such as c.859G>C, c.835-44A>G (g.69372304A>G), and c.835-1897C>T (g.69370451C>T). The method also determines the number of *SMN2* copies carrying modifier variants through analysis of allelic frequencies (Figure 2).

Based on the analysis of the entire genomic *SMN2* region and according to the PSV genotyping, we also identified specific *SMN2*-*SMN1* hybrid structures previously undetected by MLPA. The presence of

hybrid *SMN* genes has been previously described based mainly on the analysis of exons 7 and 8 (Cuscó et al., 2001; Hahnen et al., 1996). This phenomenon occurs because the complex 5q13 region contains segmental duplications prone to nonallelic homologous recombinations, deletions, duplications, and gene conversion events. Usually, hybrid genes are detected because of the homozygous absence of *SMN1* exon 7 coupled to the presence of *SMN1* exon 8. Thus, hybrids upstream exon 7 cannot be detected by common methods of diagnosis, including MLPA. In contrast, with our long PCR-based approach, we were able to detect

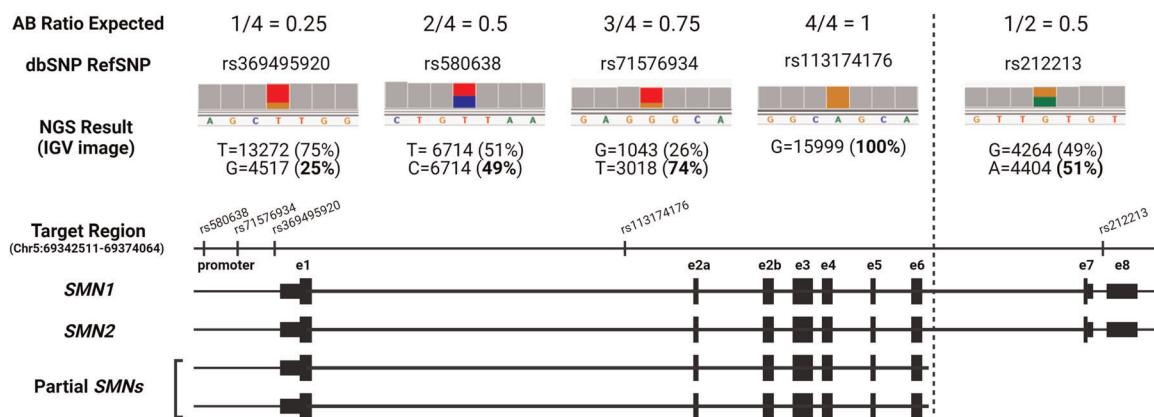


FIGURE 4 Description of the utility of AB ratios to determine the presence of two partial *SMN* genes (*SMN1*/2Δ7/8). The patient (SMA55) has one *SMN1* (with a pathogenic variant in exon 6*), one *SMN2*, and two partial *SMNs* from promoter to exon 6 (determined by MLPA). In the 5' region (promoter-ex6), the patient has a total of four *SMN* copies; consequently, we detected different SNPs with an allelic frequency of 25%, 50%, 75%, and 100%. By contrast, in the 3' region (in6-ex8), the patient has two *SMN* copies, and only variants with an allelic frequency of 50% are detected. Note that PSVs are located in the 3' region; therefore, it is not possible to determine whether these partial genes are derived from *SMN1* or *SMN2*. MLPA, multiplex ligation-dependent probe amplification; PSV, paralogous sequence variant; SNP, single-nucleotide polymorphisms

novel hybrid genes consisting of a *SMN2* gene with a fragment of intron 6 derived from *SMN1* (Figure 1). Although additional studies are needed to characterize the function of these two hybrid genes, their detection might provide clues about possible functional differences between *SMN2* genes.

The method described here also allows a complete genomic *SMN1* analysis. In fact, we were able to detect the pathogenic variants c.399_402del (p.(Glu134Serfs*14)) and c.815A>G (p.(Tyr272-Cys)) in patients and carriers, and to determine their frequencies (e.g., homozygous vs. heterozygous cases). Further confirmation that a certain variant is located in *SMN1* can be achieved with a specific PCR of the gene (Kubo et al., 2015). Given that in our method, we analyze the whole *SMN1*, including promoter, 5' and 3' regions as well as all introns, it emerges as potentially useful to study complex SMA cases in which only one allele alteration (deletion or point mutation) has been detected using conventional techniques (Alías et al., 2009).

Finally, we were able to confirm the presence of partial *SMN* copies (*SMN1/2Δ7-8*) using the AB ratios. The *SMN1/2Δ7-8* deletion has been widely described in the literature (Arkblad et al., 2006; Calucho et al., 2018; Chen et al., 2020; Ruhno et al., 2019; Vijzelaar et al., 2019) as a variant relatively common in general population (Europeans 15.7%, Vijzelaar et al., 2019). In almost all cases, the breakpoints are consistent (3643 bp before exon 7 and 1587 bp after exon 8 according to Ruhno et al., 2019), so it is considered that this deletion is the result of a single deletion event (Vijzelaar et al., 2019).

The number of *SMN2* copies and specific variants of the gene have been established as the main disease modifiers of the SMA phenotype. However, to date, studies in discordant siblings do not support the hypothesis that the intra-familial variability is due to variants in the *SMN2* locus (Calucho et al., 2018; Cuscó et al., 2006; Ruhno et al., 2019) as is the case of our sisters sharing the modifier variant c.835-1897C>T (see Table S2). Although *SMN2* is the target for splicing modifiers in the current therapeutic scenario, thorough sequencing of the gene is almost never performed in genetic diagnostic laboratories. Application of our method and thus the availability of detailed *SMN2* sequences of SMA patients would help to solve discrepancies in genotype–phenotype correlations, as well as deepen the study of intra-familial variability, in a prospective manner, for the analysis of presymptomatic cases detected in newborn screening. Indeed, being the method so specific and fast, it is envisaged that it could be accommodated within the time frame for therapeutic decisions in SMA newborns. It is also essential to unveil possible linkages between specific *SMN2* variants, factors involved in *SMN2* splicing, and responses to therapies. SMA treatments are very expensive, and proof of their efficacy is periodically assessed in SMA patients. In particular, nusinersen is an 18-mer oligonucleotide that binds the ISNN1 region of the intron 7, and to date, none of the modifier variants described is located in this region (Ruhno et al., 2019; Wadman et al., 2020; this study). However, in addition to the known modifier variants, other features of their *SMN2* genes may be correlated with the level of responsiveness and effectiveness, an issue that warrants further investigation. Thus,

the discovery and validation of positive and negative *SMN2* variants in each patient remain a crucial issue in SMA diagnosis and research. An additional benefit of implementing the new method besides the characterization of *SMN2* sequences in patients with a homozygous deletion of *SMN1* is its application to study *SMN1* in SMA patients retaining at least one *SMN1* copy. Therefore, the versatile method described here is a useful tool to approach *SMN1* and *SMN2* deep sequencing, which can be easily implemented in most SMA diagnostic laboratories.

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CONFLICT OF INTERESTS

A patent is in preparation for this methodology and pipeline (I.C. and E. F. T.). E.F.T has received grant support to conduct clinical trials on SMA from Ionis/Biogen and serves as a consultant to AveXis, Novartis, Biogen, Biologix, Cytokinetics, Roche. The remaining authors declare that there are no conflict of interests.

AUTHOR CONTRIBUTIONS

The enclosed manuscript has been seen and approved by all the authors, and they have taken care to ensure the integrity of the work.

WEB RESOURCES

The following web sources have been used to carry out this study:

<http://www.fda.gov>,
<https://genome-euro.ucsc.edu/>,
<http://www.hgmd.cf.ac.uk/ac/index.php>,
<https://mutalyzer.nl/>,
<https://gnomad.broadinstitute.org/>,
<https://arxiv.org/abs/1303.3997v2>.

DATA AVAILABILITY STATEMENT

All data and scripts used to generate the analyses of this paper are available upon request unless that the type of request compromises ethical standards or legal requirements. All variants reported in this article are now listed in the ClinVar database (<https://www.ncbi.nlm.nih.gov/clinvar/>).

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