

# Cryptic Splice-altering Variants in MYBPC3 Are a Prevalent Cause of Hypertrophic Cardiomyopathy

**Running title:** *Lopes et al.; Cryptic splicing variants in MYBPC3 cause HCM*

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## Nonstandard Abbreviations and Acronyms

HCM: hypertrophic cardiomyopathy

NGS: next-generation sequencing

WES: whole-exome sequencing

The yield of genetic testing in hypertrophic cardiomyopathy (HCM) is only 40%, even in patients with family histories of the disease. This may be caused by a high prevalence of non-genetic phenocopies or complex genetic mechanisms, but may also reflect the inability of conventional diagnostic sequencing to detect intronic variants distant from the essential donor/acceptor dinucleotides with the potential to disrupt splicing (also known as cryptic splice mutations). The availability of whole genomes from the genome aggregation database (gnomAD; <https://gnomad.broadinstitute.org>) and novel prediction tools, has improved the capacity for analyzing and interpreting deep intronic sequences.

In this study we performed large-scale unbiased screening of intronic variants in *MYBPC3* in 1644 unrelated and consecutive patients with HCM (49.5±15.6 years old at diagnosis, 1103 (67.1%) male, 1000 white (60.8%), 156 Asian (9.5%) and 75 black (4.6%)). All patients gave written informed consent and the study was approved by the regional ethics committee (15/LO/0549). Eight hundred and seventy-four probands were screened with next-generation sequencing (NGS) of the whole-genomic region of 41 genes and 770 probands were screened using whole-exome sequencing (WES) including a ~ 100bp intronic region beyond the intron-exon boundaries. Sequencing, variant calling, filtering and annotation were as previously

described<sup>1,2</sup>. To prioritize intronic variants with an impact on splicing, a deep-learning approach, SpliceAIv1.3, was applied (<https://github.com/Illumina/SpliceAI>)<sup>3</sup> with a very stringent threshold ( $\geq 0.9$ ). To predict branchpoint disruptions, LabBranchoR (<https://github.com/jpaggi/labbranchor>) was run. To evaluate the functional consequences of a novel variant, expression of mRNA was analyzed using blood samples, according to published methods<sup>4</sup>. The data that support the findings of this study are available from the corresponding author upon reasonable request.

Five hundred and forty-six patients (33.2%) had coding (or canonical splicing) variants with minor allele frequency (MAF) $<1 \times 10^{-4}$  in *MYH7*, *MYBPC3*, *TNNT2*, *TNNI3*, *MYL2*, *MYL3*, *TPMI*, and *ACTC1*. Four variants with a very high ( $\geq 0.90$ ) SpliceAI score were detected: 3 located in intron 13 [c.1224-52G>A (0.99; N=18), c.1224-80G>A (0.94; N=3) and c.1224-21A>G (0.90; N=1)] and another, c.906-36G>A (0.91; N=2), in intron 9 (Figure 1 panel A). These 4 variants were present in 24/1644 (2.2% of otherwise mutation-negative patients). All are predicted to cause a cryptic splice site and an expansion to a larger subsequent (micro)exon, in turn leading to a frameshift and stop codon, as previously shown for c.1224-52G>A, c.1224-80G>A and c.906-36G>A with RNA assays<sup>5-7</sup>.

Eighteen patients had c.1224-52G>A (1.1% compared to MAF in GnomAD of  $2.8 \times 10^{-5}$ , (OR=197.9 (CI 95%: 66.9-585.6;  $p < 0.0001$ )). The variants c.1224-80G>A, c.906-36G>A and c.1224-21A>G were not found in GnomAD. Co-segregation for c.1224-52G>A was demonstrated in three families (figure 1 panel B).

Of the 24 index patients carrying these 4 variants, age of diagnosis was 10 to 72 years, 16 (67%) were male and 6 were Asian (25% vs 9.5% in the overall cohort,  $p=0.009$ ); the remainder were white. Nineteen (79%) had a family history of sudden death and/or HCM. Maximal wall

thickness varied between 13-30mm and 6 (25%) had left ventricular outflow tract obstruction. Seven were deemed to be at high sudden death risk or already had an implantable cardioverter-defibrillator at baseline.

Finally, a novel variant, c.1898-23A>G with a low SpliceAI score (0.04, GnomAD MAF 0.000005) segregated with the phenotype in two “mutation-negative” families (Figure 1 panel C). These families were enrolled and studied with whole-exome sequencing with the initial aim of novel gene discovery and this variant was the only suitable candidate found. We have then searched for this variant in the first cohort of 874 patients and found it in an additional proband. As shown in Figure 1 panel D, the *MYBPC3* RT-PCR amplification of total RNA showed an additional longer 473 bp band detected in the probands’ RNA. Sanger sequencing revealed that these longer bands contained the complete sequence of intron 19. This mis-spliced transcript introduces a premature termination codon in intron 19 and is expected to cause nonsense-mediated mRNA decay, leading to haploinsufficiency. Given the low SpliceAI score for this variant, we sought to understand the mechanism that could explain mis-splicing; we have found a high probability of branchpoint disruption (probability of position being a branchpoint reduced by 60%).

Two recent studies of small cohorts of mutation-negative cases (46<sup>5</sup> and 93<sup>6</sup> probands) described 8 novel cryptic splice-altering variants in *MYBPC3*, with a prevalence of 9% and 6.5%, respectively. In our study, the prevalence was only 2.2% in otherwise mutation-negative patients. The difference may be explained by the very stringent criteria employed and the fact that whole-exome sequencing coverage is limited to ~ 100 bp from the exon-intron boundary, meaning that deeper intronic variants might have been missed. None of our candidate variants were described in a previous analysis of *MYBPC3* splice variants at non-canonical splice sites,

that used both a splice prediction tool and a mini-gene assay to identify variants that alter *MYBPC3* splicing<sup>8</sup>.

If the variant calling in our cohort was limited to conventional splice-sites, all the families with cryptic splice-altering variants would have been considered mutation-negative. One variant in particular (c.1224-52G>A) was unusually common (1.1%). The cause for the high frequency is uncertain, but seems unlikely to be a recent founder effect, considering the heterogeneous geographic origin of the patients. As a comparison, the causal HCM variant considered as the most common to date (p.Arg502Trp) has an estimated prevalence of 1.4-2.0% (95% CI)<sup>9</sup>. Patients with these particular cryptic splicing intronic variants in *MYBPC3* showed a higher prevalence of Asian ethnicity; this finding needs to be replicated in other populations.

Sequencing of deep intronic regions of *MYBPC3* increases the yield of genetic testing and thus improves counselling and evaluation of families with HCM. New phenotype-modifying genetic therapies tailored for splicing altering variants are being tested in animal models; as such this increase in yield might in the future be translated into tailored therapy.

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## Figure Legend:

**Figure Panel A.** MYBPC3 deep intronic variants c.1224-52G>A and c.1224-80G>A generate cryptic splice sites within intron 13 with expansion of exon 14 (A). c.906-36G>A generates a cryptic splice site within intron 9 with expansion of exon 10. Score refers to spliceAI score.

**Panel B.** Pedigrees demonstrating co-segregation of the MYBPC3 c.1224-52G>A variant with HCM phenotype in 3 families. Circles: women; squares: men. +: mutation-positive; -: mutation-negative; colored symbols: affected (HCM).

**Panel C.** Pedigrees demonstrating co-segregation of the MYBPC3 c.1898-23A>G variant with HCM phenotype in 2 previously “mutation-negative” families. Circles: women; squares: men. +: mutation-positive; -: mutation-negative; colored symbols: affected (HCM).

**Panel D.** RT-PCR analysis of the ectopic expression of *MYBPC3* in the blood of controls (lanes 1-2) and index patients carrying the c.1898-23A>G variant (lane 3: H25, lane 4: H26). Negative controls were loaded in lane 5 (-RT) and 6 (non-template). L- 100 bp DNA ladder. The lower bands (368 bp, MYBPC3) correspond to the normal mRNA, and the longer bands (473 bp, IR19-MYBPC3) correspond to a mis-spliced intron 19-retained transcript.