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Targeted panel sequencing in adult patients with left ventricular non-compaction reveals a large genetic heterogeneity

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For Review Only

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3 **Targeted panel sequencing in adult patients**
4 **with left ventricular non-compaction reveals a large genetic**
5 **heterogeneity**
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8 **Running title: Genetic complexity of adult left ventricle non compaction**
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3 and systems dissection of mechanisms underlying dilated cardiomyopathy"; PROMEX
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10 **ABSTRACT**

11 Left ventricular non-compaction (LVNC) is a cardiomyopathy that may be of genetic origin,
12 however few data are available about the yield of mutation, the spectrum of genes and allelic
13 variations. The aim of this study was to better characterize the genetic spectrum of isolated
14 LVNC in a prospective cohort of 95 unrelated adult patients through the molecular
15 investigation of 107 genes involved in cardiomyopathies and arrhythmias.
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22 Fifty-two pathogenic or probably pathogenic variants were identified in 40 patients (42%)
23 including 31 patients (32.5%) with single variant and 9 patients with complex genotypes
24 (9.5%). Mutated patients tended to have younger age at diagnosis than patients with no
25 identified mutation. The most prevalent genes were *TTN*, then *HCN4*, *MYH7*, and *RYR2*. The
26 distribution includes 13 genes previously reported in LVNC and 10 additional candidate
27 genes.
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35 Our results show that LVNC is basically a genetic disease and support genetic counseling
36 and cardiac screening in relatives. There is a large genetic heterogeneity, with predominant
37 *TTN* null mutations and frequent complex genotypes. The gene spectrum is close to the one
38 observed in dilated cardiomyopathy but with specific genes such as *HCN4*. We also identified
39 new candidate genes that could be involved in this sub-phenotype of cardiomyopathy.
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49 KEY WORDS: Left ventricular non compaction, cardiomyopathy, molecular genetic, next
50 generation sequencing
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INTRODUCTION

Left ventricular non-compaction (LVNC, OMIM300183) is a relatively rare cardiomyopathy, with or without LV dysfunction, characterized by excessively prominent trabeculations and associated deep recesses that communicate with the ventricular cavity¹. LVNC is part of unclassified cardiomyopathies according to the European Society of Cardiology² and to genetic cardiomyopathies by the American Heart Association³.

The prevalence of LVNC was estimated at 0.014% to 1.3% depending on the age of patients^{4,5}. Multiple imaging techniques are usually useful for the diagnosis of LVNC, with variable echocardiographic or magnetic resonance imaging diagnostic criteria but no clear consensus so that the positive diagnosis may be challenging⁶. The phenotypic expression and evolution of isolated LVNC is highly variable, and clinical features can range from asymptomatic to symptomatic, with a relatively stable course over several years or an evolution towards severe complications including congestive heart failure, ventricular arrhythmia and sudden cardiac death, atrial arrhythmias and systemic embolic events⁶.

LVNC is supposed to be related to a premature arrest of compaction of the loose myocardial meshwork during fetal embryogenesis, with persistent trabeculated myocardium, but the precise pathophysiology remains poorly understood. A family history is noticed in a significant proportion of patients and predominant mode of inheritance is autosomal dominant, with some cases with an X-linked transmission⁷.

Several genes have been identified as LVNC disease causing. The first reported genetic cause of isolated LVNC was described by Bleyl et al. with mutations in the X-linked *TAZ* gene, also responsible for Barth syndrome⁸. The sarcomere-encoding genes (*MYH7*, *ACTC1*, *TNNT2*, *MYBPC3*, *TMP1*, and *TNNI3*) appear to account for 17 to 30% of LVNC^{9,10} but other genes such as *DTNA* (α -dystrobrevin), *NKX-2.5*, Z-line protein-encoding *ZASP/LDB3*, lamin A/C (*LMNA*) genes have been also associated with LVNC¹¹. Recently, the *TTN* gene was also reported as involved in this disease¹²⁻¹⁴, with a highly heterogeneous

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3 prevalence, as high as 19% and the first responsible gene for LVNC in a German study of 68
4 index cases¹³ to 7% in adults but 0% in children from a Dutch cohort of 327 patients¹⁴.
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7 Several studies have analyzed the spectrum of genes in LVNC⁹⁻¹⁶ but with heterogeneous
8 strategies (Sanger or Next-generation sequencing), usually in retrospective cohorts without
9 imaging core-lab, and usually with a relatively small panel of genes (45 genes in the Dutch
10 study¹⁴, exome sequencing in the recent German study¹³. Therefore, the exact spectrum of
11 LVNC-causing variants, their prevalence and their impact in genetic counseling remain poorly
12 understood. Furthermore, the unique specificity of LVNC as an independent nosology entity
13 has been questioned and LVNC has been suggested as an overlapping phenotype with
14 hypertrophic or dilated cardiomyopathy⁹. To explore these issues, a prospective French
15 national research program was launched and focused on consecutive adult patients with a
16 recent diagnosis of isolated LVNC. The general aim was to better characterize the allelic and
17 genetic spectrum of LVNC through a large panel of genes previously reported in the various
18 cardiac hereditary diseases. The objective was also to identify new potential candidate genes
19 that could be involved in the phenotype of LVNC.
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34 **METHODS**

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37 The present study was conducted as part of the Programme Hospitalier de Recherche
38 Clinique (PHRC Ref: 2011-A - 00987-34, coordinator Pr. G. Habib, Marseille) aimed at
39 describing the clinical spectrum of LVNC and at characterizing the genetic spectrum of LVNC
40 through a next-generation sequencing (NGS) strategy in a new prospective cohort.
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45 **Patients, inclusion criteria**

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48 The study included unrelated patients with a minimal age at inclusion of 18 years old,
49 enrolled between 2012 and 2013 in 13 French centers for inherited cardiac diseases.
50 Collected data included clinical data (initial presentation, first symptoms, and data from
51 cardiac and neurologic examination), family history, and tests including ECG,
52 echocardiography, MRI, CT scan, Holter monitoring when available, as well as follow-up
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3 data. Only patients with a recent diagnosis of isolated LVNC (maximum in the last 6 months
4 before inclusion) were enrolled. All echocardiographic documents were sent and reviewed by
5 a core lab (Marseille) to confirm the diagnosis. Diagnosis of isolated LVNC was considered
6 definite when several criteria were present in left ventricle (LV): (i) multiple trabeculations with
7 deep endomyocardial recesses, (ii) two-layer myocardial structure with a thin compacted (C)
8 and a thick non compacted (NC) layer, (iii) color Doppler evidence of perfused intertrabecular
9 recesses, (iv) systolic NC / C ratio > 2 (parasternal short-axis view); (v) no associated heart
10 disease¹⁷. Cardiac MRI was also frequently performed in our series, with a NC/C ratio of >2.3
11 in diastole as the recommended threshold for the diagnosis of LVNC using this technique¹⁸.
12 However, for the purpose of the current study, only the echocardiographic criteria were used
13 as inclusion criteria. Only patients with a diagnosis validated by an imaging core lab
14 confirming a definite diagnosis of LVNC were included. Informed consent, blood samples,
15 and clinical evaluations were obtained from all patients, with a protocol approved by the
16 Ethics Committee of AP-HM (Assistance Publique- Hôpitaux de Marseille).
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33 **Genetic analysis**

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36 Targeted gene enrichment, high-throughput sequencing: Patients' DNAs were extracted from
37 peripheral blood with **QIAasymphony**® (Qiagen, Hilden, Germany) and qualitatively checked
38 using Tape Station DNA genomic array (Agilent, Santa Clara, USA). Custom targeted gene
39 enrichment and DNA library preparation were performed using the **Nimblegen** EZ choice
40 probes® and Kappa HTP Library preparation kit® according to the manufacturer's
41 instructions (Nimblegen®, Roche Diagnostics, Madison, USA). The targeted regions include
42 all coding exons and **+/- 50 base pairs** of flanking intronic regions of 107 genes known to be
43 involved in cardiomyopathies (77 genes) and arrhythmias (30 genes) (**Suppl. Table 1**). The
44 targeted regions were sequenced using the Illumina MiSeq platform on a 500 cycle Flow Cell
45 (Illumina, Santa Cruz, USA) and MiSeq Software generates FASTQ format files after
46 demultiplexing patients' sequences.
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3 Bioinformatics pipeline: In presence of overlapping paired-end reads, these were merged
4 with Flash¹⁹. Merged single reads and paired-end reads were then aligned on Hg19 human
5 reference genome using BWA-MEM²⁰. This was further followed by a local realignment
6 around insertion and/or deletions and a quality base recalibration by using of the GATK
7 program²¹. PCR and optical duplicates were highlighted with the MarkDuplicates Picard tool
8 (<http://broadinstitute.github.io/picard>) and were further removed using samtools²². Resulting
9 .bam outputs from merged single reads and properly paired-end reads were then combined
10 into a unique .bam file. Variant calling was performed using the GATK Haplotype Caller
11 program²¹ simultaneously on all sequenced samples. Detected variants were then annotated
12 using ANNOVAR²³ and CADD²⁴ tools. Coverage statistics were produced using the HsMetrics
13 Picard tool. Detected variants with sequencing depth greater than 30X and with at least 20%
14 of reads supporting the alternative allele were kept for analysis. Detection of copy number
15 variation (CNV) was performed after coverage normalization, by computing the ratio of a
16 target's coverage of a given individual over the mean coverage of this target across all
17 patients of the same sequencing run.

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33 Variants interpretation: Pathogenicity of variants was determined according to current ACMG
34 guidelines²⁵ that recommend classifying variants into 5 categories: pathogenic, likely
35 pathogenic, unknown significance, likely benign and benign. A recent publication dedicated
36 to cardiomyopathies recommended the use of a frequency threshold of 0.01%²⁶. Variants
37 were filtered out according to their allele frequency as reported in the GnomAD database
38 (<http://gnomad.broadinstitute.org/>). We then evaluated each variant considering a careful
39 review of the literature, the location of the variant in the gene and the resulting corresponding
40 protein, the *in silico* prediction tools (Polyphen2, SIFT, GVG D and Mutation Taster for
41 missense variants and SpliceSiteFinder like®, MaxEntScan®, NNSPLICE®, GeneSplicer®
42 and Human Splicing Finder® for splicing variants) and functional studies when available.
43 Additionally, we looked at a local database of pathogenic variants related to our experience
44 on the molecular diagnosis of cardiomyopathies. In practice, we considered as “pathogenic”
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(class 5), a variant with confirmed pathogenicity criteria and already proved as responsible for cardiomyopathies or a novel nonsense variant with a frequency below 0.01%. We considered as “likely pathogenic” (class 4), unpublished variants with a frequency below 0.01% and unknown in our database, located in a functional domain of the protein and with pathogenicity prediction tools mainly (at least 3 out of 4 tools) in favor to a strong effect.

“Variants of unknown significance” were new variants with no evidence for predicted deleteriousness and published variants with a frequency over 0.01%. Such variants were not considered in this work until proof of pathogenicity but are presented in supplemental data (Suppl. Table 2). In *TTN* gene, only null variants (consensus splice sites, stop codons, insertions and deletions leading to a shift in the reading frame) were considered as pathogenic according to a recent publication on *TTN* mutations in cardiomyopathies²⁷, and we excluded other variants, especially *TTN* missense variants.

All variants considered as pathogenic and probably pathogenic have been confirmed by a second independent method (Sanger sequencing or MLPA)

Statistical analyses were performed with Fisher Exact test or chi2, for binary variables and Student t tests for continuous variables, when appropriate.

RESULTS

Patients

Ninety-five unrelated patients ranging from 19 to 81 years-old were included in the study with a mean age at diagnosis/inclusion of 46.3 (± 15) years old. This cohort was composed of 56 males and 39 females. At the time of inclusion, 46 patients had a NYHA score >1 , mean ejection fraction at inclusion was 42.5% (± 14.5) and mean heart rate was 68.5 bpm (± 15.6).

Performance of the custom panel

The Miseq sequencing run yielded an output of 1.8 to 2.2Gb per sample, with a mean sequencing depth per sample of 265 reads (SD:35.3). On average, 99.7% of selected targets (1740/1745) were covered over 30X and 98.5% (1720/1745) over more than 100X.

Allelic Spectrum

Cohort analysis led to the identification of 52 confirmed or highly suspected pathogenic variants (class 5 or 4) including 42 novel ones located in 23 different genes. Among these mutations, 50 were found in 22 cardiomyopathies related genes (*ACTC1*, *BAG3*, *DSC2*, *DSP*, *FLNC*, *HCN4*, *HEY2*, *LDB3*, *LMNA*, *MYBPC3*, *MYH6*, *MYH7*, *MYLK2*, *MYPN*, *NEXN*, *NKX2.5*, *PDLIM3*, *PKP2*, *RBM20*, *RYR2*, *TMEM43* and *TTN*) and 2 were observed in *ANK2* gene, known to be responsible for long QT Syndrome (Table 1).

Among the 22 cardiomyopathy genes, the most prevalent ones were *TTN* (19%, 10 variants), followed by *HCN4* and *MYH7* genes (10 %, 5 variants each), followed by *RYR2* (8%, 4 variants) then *MYH6* and *ACTC1* (6%, 3 variants each), then *MYBPC3*, *LDB3*, *MYLK2* and *NEXN* (4%, 2 variants). The 12 other genes were found mutated only once. Among the arrhythmias genes, *ANK2* was mutated in 2 patients (4%, 2 variants) (Fig. 1).

The 10 *TTN* truncating variants included 8 variants located in the A-band (80%) and two located at the end of the I-band at the junction with A-band. In the 22 remaining genes, 42 mutations were identified including 35 missense variants, 1 in-frame deletion, 5 null mutations (3 frame shifts, 1 splice and 1 non-sense mutations), and a CNV consisting in a complete deletion of *RYR2* exon 3 (Table 2).

In the second prevalent gene *HCN4*, 5 different missense variants were found including one already published in LVNC. These variants were located in transmembrane domain 4 (c.1123C>T, p.Arg375Cys), in the pore (c.1403C>T, p.Ala468Val, c.1438G>T, p.Gly480Cys and c.1444G>A, p.Gly482Arg) before the transmembrane domain 5 (c.1231C>G, p.Leu411Val). (Table 2)

In order to classify genes and variants according to their function in the cardiomyocytes, we defined 5 cellular “compartments” (Fig.1). The distribution of the 52 variants in the 23 genes showed that 52% of variants (N=27) were located in sarcomeric genes, 21% (N=11) were in ion channel or related genes, 8% (N=4) were in genes involved in the cellular structure, 6%

(N=3) were located in desmosome genes. In addition, 12% of variants (N=6) were found in transcription factors genes or genes involved in other structures or functions (eg. *NKX2-5*).

Finally, upon the 52 identified variants, 40 were located in the 13 already known LVNC genes (77%) and 12 were located in the 10 additional candidate genes (23%)(Tables 1 & 2).

Multiple mutations in patients

In the cohort, 9 patients (9.5%) presented a complex genotype feature with the presence of more than one pathogenic variant. Seven patients harbored two disease-causing variants in cardiomyopathy genes (Fig.2) and 2 patients carried at least 3 pathogenic variants: one with *BAG3*, *MYH7*, and *NKX2-5* variants and the second with *ACTC1*, *ANK2*, *LDB3* and *MYLK2*. Regarding the *TTN* gene, 8 patients were carrying a unique *TTN* variant and 2 patients carried a *TTN* mutation associated with another gene variant (*MYH6*, *NEXN*).

Mutations, patients and phenotype

According to our variant selection criteria, a pathogenic variant was identified in 40 patients of the cohort (20 males and 20 females; 50%), including 31 patients (32.5%) with single variant and 9 patients with complex genotypes (9.5%). *TTN* mutations are predominant and identified in 10 patients of the cohort (10.5%). In 55 patients (58%), no genetic cause was identified. Fifteen patients had a known family history of LVNC, 55 patients were sporadic cases and family history was not available for 25 patients. Among these groups, the mutation rate is 53%, 46% and 28% respectively. The difference between familial cases and sporadic cases is not significant (p-value : 0.77).

An analysis was performed regarding the age at diagnosis, ejection fraction, presence of dyspnea (NYHA>1) and heart rate comparing the groups of mutated patients vs patients with no identified genetic cause. Mutated patients tended to be younger at diagnosis (43.0 vs 48.7 years, p=0.07) but systolic dysfunction showed no significative difference between groups

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3 (Table 3). Interestingly, the mutation yield was higher in youngest patients <65 years old
4 (38/84, 45%) as compared to oldest patients >65 years (2/11, 18.2%, p-value: 0.11).
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7 Patients with complex genotypes (≥ 2 mutations), as compared to patients with single
8 variants, tended to be younger at diagnosis and to have a decreased ejection fraction
9 although differences were not significant (Table 3). Finally, in patients carrying a single
10 variant, we observed that the LV mean ejection fraction in patient with a mutation in
11 sarcomeric genes (N=18) was lower than in patients mutated in non-sarcomeric genes
12 (N=13) (43.8% vs 51.6%, p-value: 0.26).
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19 DISCUSSION

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22 We present here the results of the genetic analysis of a cohort of 95 independent
23 patients (index cases) with LVNC in order to evaluate the yield of mutation screening and to
24 assess the allelic and genetic spectrum of the disease. The design of our study has some
25 characteristics that may differ from previous studies since our project was a prospective
26 study performed in newly diagnosed consecutive patients (diagnosis less than six months)
27 with a validation of the cardiac diagnosis by an expert centralized imaging core-lab. This
28 design was conceived to limit potential inclusion bias and strengthen the representativeness
29 of the cohort. We also focused on isolated LVNC, without associated congenital heart
30 defects, in adult patients in order to have a more homogeneous population. Next generation
31 sequencing was performed with a panel of 107 genes involved in cardiomyopathies and
32 some arrhythmias, which represents the most comprehensive genetic analysis published
33 (exome sequencing was performed by Sedagast-Hamedani *et al* but [only selected genes
34 known to be involved in the phenotype were reported](#))¹³.
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49 When considering pathogenic or likely pathogenic variants, we identified a mutation in
50 42% of the patients, which represents a proportion slightly higher than previously reported in
51 this disease^{9,10,13,14,16}, but relatively similar to features observed in other cardiomyopathies²⁸.
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55 The distribution of genes revealed a high degree of genetic heterogeneity with putative or
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2 confirmed pathogenic mutations identified in 23 different genes. The distribution includes 13
3 genes previously published as associated with the phenotype of LVNC (77% of variants) and
4 10 additional candidate genes (9 cardiomyopathy and 1 arrhythmia genes) that were never
5 reported before as associated with LVNC (Table 1). Despite the stringent selection criteria of
6 variants, *TTN* represents the most prevalent gene in the cohort (19% of variants or 10.5% of
7 patients) including 8 variants located in the A-band (80%). As previously observed in patients
8 with DCM, *TTN* truncating mutations in the A-band region of the protein were over
9 represented²⁷.
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19 The following most prevalent genes were *HCN4* and *MYH7*, followed by *MYH6*, *RYR2*
20 and *ACTC1*. Considering others published reports^{13,14}, some discrepancies were observed in
21 the gene distribution especially regarding *TTN*, *MYH7*, *HCN4* and *LMNA*. Differences in
22 distribution may be related in part to the variable characteristics of the cohorts (isolated
23 LVNC or not, age at diagnosis, incident or prevalent cases). We observed a relatively high
24 proportion of patients with *HCN4* pathogenic variants as we found 5 different variants,
25 located in S4, S5 and pore domains of the protein (Table 2). Among these patients 3/5
26 presented bradycardia (one patient was implanted by a pace maker) but no valvular disease
27 has been reported in any of them^{29,30}. This suggests that this recently published gene²⁹
28 constitutes an important disease-causing gene in LVNC. The prevalence of *HCN4* did not
29 appear as such in previously published cohorts, possibly due to the absence of this gene in
30 some studied panels^{9-11,14,15} or a difference in the cohort recruitment^{13,16}. For other genes, a
31 higher rate of *MYH7* variants and a lower rate of *LMNA* variants were found in our cohort as
32 compared with the study of Sedaghat-Hamedani *et al.*¹³ while frequency of *TTN* and
33 *MYBPC3* variants were consistent. In the work of van Waning *et al.*¹⁴, the proportion of
34 *MYH7* and *MYBPC3* in the group of adult patients were consistent with us but a lower rate of
35 *TTN* variants was observed. In older publications^{9,10} in which only sarcomeric genes were
36 analyzed, the spectrum of genes showed that *MYH7* was the most prevalent gene then
37 *TNNT2*, *TMP1*, *TNNI3*. These last three genes were not found mutated in our cohort, which
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3 could be due to the fact that our cohort is composed by patients with an adult onset of the
4 disease. Interestingly, our results also strengthen the involvement of recently published
5 genes such as *HCN4*^{29,30} and *RYR2*³¹ and help to better estimate their prevalence. As a
6 whole, our finding about the large genetic and allelic spectrum is helpful in refining the genes
7 of interest for routine molecular diagnostic of patients with LVNC.
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13 Additionally, we tried to determine if patients reporting a familial history of LVNC, were
14 more frequently found with a mutation than patients presenting as sporadic cases. The
15 cohort includes 15 familial form and 55 sporadic cases, the 25 remaining patients had no
16 information's about their relatives. Quite the same rate of mutation identification was found in
17 the group of familial forms and sporadic cases (53 % and 46% respectively, difference not
18 significant)"
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25 Apart from the report of mutations in genes previously associated with LVNC, an
26 important finding of our study is that we identified mutations in 10 genes known to be
27 involved in cardiac inherited diseases but not described until now as associated with this
28 specific phenotype. Among these genes, 9 were previously reported as associated with other
29 sub-types of cardiomyopathy such as HCM, DCM and ARVC (Table 1), suggesting an
30 overlap between the various cardiomyopathies. Among these 10 genes, *MYLK2* and *NEXN*
31 were identified each in 2 patients and 7 genes (*BAG3*, *FLNC*, *HEY2*, *MYPN*, *PDLIM3*,
32 *TMEM43* and *DSC2*) were involved each in only one patient. Although these genes could be
33 good candidates for being pathogenic, the definitive role of these genes for causing LVNC
34 will require confirmation in further studies, especially through segregation analyses in
35 families or functional studies. Similarly, in 2 patients with no particular ECG abnormalities, we
36 identified variants in *ANK2* gene known to be involved in long QT syndrome. As *ANK2* was
37 never reported before in LVNC, cautious is necessary before any conclusion about the
38 causal role of these variants. However, *HCN4* was initially described in channelopathies and
39 then involved in LVNC^{29,30}, with a significant proportion in our cohort, illustrating the fact that
40 a given gene involved in arrhythmias does not preclude the potential role in LVNC.
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3 Another observation emerging from this study is the high level of patients (9.5%)
4 presenting complex genotypes with causative variants in two (or more) different genes. This
5 feature about double mutated patients was not previously described as so high in adult
6 patients with LVNC. In patients with complex genotypes, cumulative effect of variants have
7 been associated with a higher severity of the disease in one study¹⁴ but was not analyzed in
8 details in most of other studies. [The hypothesis of a gene-dose- effect can be suspected as](#)
9 [well as in other sub- morphotypes of cardiomyopathies, particularly in HCM.](#) In the present
10 cohort, patients with complex genotypes tended to have a same age at diagnosis (43.1 vs.
11 43.5 years old) but more symptoms (dyspnea > NYHA1: 67% vs 53%) and a lower ejection
12 fraction (36% vs. 47%). However these differences were not statistically significant and must
13 be confirmed in larger cohorts.
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25 The global analysis of the distribution of genes observed in our cohort of adult
26 patients with LVNC also provides useful information regarding the debated issue of whether
27 or not LVNC is an independent nosological entity or a phenotype overlapping with other
28 cardiomyopathy sub-types such as HCM or DCM^{26, 32-34}. The biggest cohorts published so far
29 about HCM patients (including 3267 HCM patients sequenced on 16 genes and 874 patients
30 sequenced on 20 genes) reported sarcomeric genes (especially *MYBPC3*) as the major
31 genes^{26,32}. In patients with DCM, the *TTN* gene has been consistently reported as the most
32 frequent mutated gene³³. In a study of 639 DCM patients sequenced on 84 genes, the
33 highest prevalence observed was for *TTN* (13%), *PKP2*, *MYBPC3*, *DSP*, *RYR2*, *DSC2* and
34 *SCN5A* genes³⁴. In the present cohort, we observed that the most prevalent genes are *TTN*,
35 then *MYH7*, *HCN4*, *MYH6*, and *RYR2*. This distribution, and the fact that *TTN* is by far the
36 most frequent gene we observed in LVNC, as well as the high level of complex genotypes,
37 suggests that the [genetic](#) profile of LVNC patients is relatively similar to patients with DCM
38 but not similar to patients with HCM^{26,28,32-37}. However, the distribution of LVNC patients
39 presents some specific findings, such as the relatively high rate of *HCN4* gene mutations,
40 which favor the possible specific role of some particular genes in this disorder.
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3 **Limitations.** Our results were derived from a cohort of adult-onset patients with
4 isolated LVNC. Therefore, results may not be applicable to a pediatric population or a
5 population with syndromic LVNC. Even though probably pathogenic variants completed all
6 the criteria for pathogenicity, we don't provide family segregation and functional analysis for
7 now.
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13 **In conclusion,** molecular analysis of 107 genes in 95 adult patients with isolated
14 LVNC shows a mutation detection rate of 42%. These data, coming from the most
15 comprehensive study available until now in terms of genes that were analyzed, show that
16 LVNC is basically a genetic disease in most cases, with a large genetic heterogeneity. The
17 global distribution of genes appears quite close to the profile observed in DCM patients, with
18 *TTN* as the most frequent mutated gene, but with some specific genes such as *HCN4*. We
19 found 9.5% of patients presenting a complex genotype with a disease causing variant in two
20 different genes located on different chromosomes. This observation could explain part of
21 intra-familial variable expressivity in case of bi-lineal inheritance, as some relative should be
22 carrier of a single variant with a moderate phenotype. We also described mutations in 10
23 genes not described until now as associated with LVNC. Although these genes are putative
24 good candidates for causing LVNC, the definitive causal role of these genes in this
25 phenotype will require confirmation in further studies.
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Table 1: List of genes published in LVNC cardiomyopathy and potentially new genes identified in the present work.

Cellular structure	Gene (NM)	Protein	Phenotype	Ref. in LVNC	This Cohort
Sarcomere					
	MYH7 (NM_000257.2)	Myosin heavy chain	HCM, DMC, LVNC	9-10	yes
	TNNT2 (NM_001001430.1)	Troponin T2	HCM, DMC, LVNC	9-10	no
	ACTC1 (NM_005159.4)	Cardiac Actin	HCM, DMC, LVNC	9-10	yes
	MYBPC3 (NM_000256.3)	Cardiac C protein	HCM, DMC, LVNC	9-10	yes
	TPM1 (NM_001018005.1)	Alpha-tropomyosin	HCM, DMC, LVNC	9-10	no
	TNNI3 (NM_000363.4)	Troponin I3	HCM, DMC, LVNC	9-10	no
	DTNA (NM_001390.4)	Alpha-Dystrobrevin	DCM, LVNC	11-15	no
	MYH6 (NM_002471.3)	Myosin light chain	DCM, HCM, LVNC	15	yes
	ACTN2 (NM_001103.2)	Actinin	HCM, LVNC	32	no
	TTN (NM_001256850.1)	Titin	DMC, HCM, LVNC	12	yes
	MYLK2 (NM_033118.3)	Myosin Light chain kinase	DCM	This work	yes
	MYPN (NM_032578.2)	Myopalladin	DCM	This work	yes
	NEXN (NM_144573.3)	Nexilin	HCM, DCM	This work	yes
Structure					
	FLNC (NM_001458.4)	Filamin-C	HCM, DCM	This work	yes
	LDB3 (NM_007078.2)	LIM Domain Binding 3	DCM, LVNC	11-15	yes
	LMNA (NM_170707.2)	Lamine A/C	DMC, LVNC	11-15	yes
Ion channel and related genes					
	ANK2 (NM_001148.4)	Ankyrin 2	LQT	This work	yes
	HCN4 (NM_005477.2)	Hyperpolarization Activated Cyclic Nucleotide Gated Potassium Channel 4	ARVC, LVNC	29,30	yes
	RYR2 (NM_001035.2)	Ryanodin receptor 2	ARVC, LVNC, CPVT	31	yes
	CASCQ2 (NM_001232.3)	Calsequestrin 2	QTL, LVNC	38	no
	SCN5A (NM_198056.2)	Sodium channel, voltage-gated, type V, alpha subunit	LQT, Brugada, DCM, LVNC	39	no

Other					
	NKx2.5 (NM_004387.3)	NK2 Homeobox 5	DCM, LVNC	11-15	yes
	TAZ (NM_000116.3)	Taffazin	Barth Syndrom, LVNC	8	no
	FBN1 (NM_000138.4)	Fibrillin	Marfan Syndrom, LVNC	40	no
	ABCC9 (NM_020297)	ATP Binding Cassette Subfamily C Member 9	DCM, LVNC	35	no
	PDRM16 (NM_022114)	PR/SET Domain 16	LVNC	41	no
	BAG3 (NM_004281.3)	BCL2 Associated Athanogene 3	DCM	This work	yes
	HEY2 (NM_012259)	Hairy-Related Transcription Factor 2	DCM	This work	yes
	PDLIM3 (NM_014476.4)	PDZ And LIM Domain 3	ARVC, HCM	This work	yes
	RBM20 (NM_001134363.1)	RNA Binding Motif Protein 20	DCM	13	yes
	TMEM43 (NM_024334.2)	TransmembraneProtein 43	ARVC	This work	yes
Desmosome					
	PKP2 (NM_004572.3)	Plakophilin 2	ARVC, LVNC	42	yes
	DSP (NM_004415.2)	Desmoplakin	ARVC, LVNC, DCM	43	yes
	DSC2 (NM_024422.3)	Desmocollin	ARVC	This work	yes

Table 2: List of pathogenic and probably pathogenic variants identified in the cohort.

Position c., cDNA position; Position p., protein effect; Published: No or Yes; Associated phenotype for published variants; GnomAD correspond to the allelic frequency, and Htz corresponds to the allele count in GnomAD in all populations; GVG, SIFT, Mutation taster and polyphen are algorithms corresponding to *in silico* Predictive Algorithms used for evaluation of missense variants. Range of scores for each are indicated in the title column; "Type" indicated the nature of the variant; MS: missense, NS; Nonsense, Del; deletion, Splice; mutation affecting splicing site. Column "interpretation", indicates conclusions about the pathogenicity of the variant: class 5 ; certainly pathogenic, Class 4; probably pathogenic. NA: not applicable

Gene	Position c.	Position p.	Published No/Yes	Associated phenotype	GnomAD, Htz	GVGD (C65-C0)	SIFT (0-1)	Mutation Taster (1-0)	Polyphen (1-0)	Type	Interpretation
ACTC1	c.670G>T	p.Asp224Tyr	N	NA	/	65	0	1	0.994	MS	Class 4
ACTC1	c.281A>G	p.Asn94Ser	N	NA	4.061e-6, 1	45	0	1	0.615	MS	Class 4
ACTC1	c.623G>A	p.Arg208His	N	NA	4.061e-5, 10	25	0	1	0,01	MS	Class 4
ANK2	c.11150T>A	p.Ile3717Asn	N	NA	1.083e-5, 3	45	0	0,995	0.865	MS	Class 4
ANK2	c.9145C>T	p.Arg3049Trp	N	NA	8.155e-6, 2	65	0	0,93	0.999	MS	Class 4
BAG3	c.465_466insGCG	p.Ala155delinsAlaAla	N	NA	/	NA	NA	NA	NA	MS	Class 4
DSC2	c.1448A>T	p.Asn483Ile	N	NA	/	15	0	NA	0.905	MS	Class 4
DSP	c.3035delA	p.Asp1012fs	N	NA	/	NA	NA	NA	NA	Del	Class 5
FLNC	c.1933_1935del	p.645del	N	NA	3.969e-5, 11	NA	NA	NA	NA	Del	Class 4
HCN4	c.1403C>T	p.Ala468Val	N	NA	4.065e-6, 1	65	0	1	0,95	MS	Class 4
HCN4	c.1123C>T	p.Arg375Cys	N	NA	4.061e-6, 1	65	0	1	0,99	MS	Class 4
HCN4	c.1231C>G	p.Leu411Val	N	NA	/	25	0	1	0,99	MS	Class 4
HCN4	c.1444G>A	p.Gly482Arg	Y	NCVG	/	65	0	1	1	MS	Class 5
HCN4	c.1438G>T	p.Gly480Cys	N	NA	/	65	0	1	1	MS	Class 4
HEY2	c.683C>T	p.Thr228Met	N	NA	3.231e-5, 2	0	0,05	1	0,45	MS	Class 4
LDB3	c.625G>C	p.Glu209Gln	N	NA	/	25	0	1	0,94	MS	Class 4
LDB3	c.608C>T	p.Ser203Leu	Y	CMD	2.538e-5, 7	65	0	1	0,88	MS	Class 5

LMNA	c.738delG	p.Gln246fs	N	NA	/	NA	NA	NA	NA	NA	Del	Class 5
MYBPC3	c.532G>A	p.Val178Met	Y	HCM	/	0	0,01	1	0,992	MS	Class 5	
MYBPC3	c.1504C>T	p.Arg502Trp	Y	HCM	5.411e-5, 15	65	0	1	0,484	MS	Class 5	
MYH6	c.1793dupA	p.Asn598fs	N	NA	8.122e-6, 2	NA	NA	NA	NA	Dup	Class 5	
MYH6	c.4828C>T	p.Arg1610Cys	N	NA	3.247e-5, 1	0	0	1	0,988	MS	Class 4	
MYH6	c.50G>T	p.Arg17Leu	Y	Cardiac septal defect	/	0	0	1	0,55	MS	Class 5	
MYH7	c.379C>A	p.Pro127Thr	N	NA	/	0	0	1	0,98	MS	Class 4	
MYH7	c.3830G>C	p.Arg1277Pro	N	NA	/	35	0	1	0,842	MS	Class 4	
MYH7	c.3586C>T	p.His1196Tyr	N	NA	/	15	0	1	0,613	MS	Class 4	
MYH7	c.2419C>G	p.Arg807Gly	N	NA	/	25	0,02	1	0,85	MS/Splice	Class 4	
MYH7	c.4588C>T	p.Arg1530X	N	NA	/	NA	NA	NA	NA	NS	Class 4	
MYLK2	c.1754T>A	p.Ile585Asn	N	NA	/	45	0	1	0,921	MS	Class 4	
MYLK2	c.1658G>A	p.Arg553His	N	NA	1.276e-5, 3	0	0,15	0,92	0,004	MS	Class 4	
MYPN	c.3457G>A	p.Gly1153Arg	N	NA	1.219e-5, 3	65	0	1	1	MS	Class 4	
NEXN	c.2012T>C	p.Ile671Thr	N	NA	2.541e-5, 7	0	0	1	0,936	MS	Class 4	
NEXN	c.1396A>C	p.Lys466Gln	N	NA	/	0	0	1	0,996	MS	Class 4	
NKX2-5	c.604C>G	p.Leu202Val	N	NA	/	25	0	1	0,07	MS	Class 4	
PDLIM3	c.742C>T	p.Arg248Cys	N	NA	8.153e-6, 2	0	0	1	1	MS	Class 4	
PKP2	c.2018G>A	p.Gly673Asp	Y	ARVC	/	65	0	1	1	MS	Class 5	
RBM20	c.1907G>A	p.Arg636His	Y	DCM	/	0	0	1	0,99	MS	Class 5	
RYR2	c.13936G>C	p.Asp4646His	N	NA	/	0	0	1	0,99	MS	Class 4	
RYR2	c.6180G>T	p.Gln2060His	N	NA	/	15	0	1	0,89	MS	Class 4	
RYR2	c.878A>C	p.Gln293Pro	N	NA	/	65	0	1	0,756	MS	Class 4	
RYR2	c.169- ? c.273+?del ?	/	Y		/	NA	NA	NA	NA	Del	Class 5	
TMEM43	c.317A>G	p.Tyr106Cys	N	NA	2.031e-5, 5	65	0	1	1	MS	Class 4	
TTN	c.93376delA	p.Arg31126fs	N	NA	/	NA	NA	NA	NA	Del	Class 4	
TTN	c.93376_93377de l	p.Arg31126fs	N	NA	/	NA	NA	NA	NA	FS	Class 4	

TTN	c.82724delA	p.Asn27575fs	N	NA	/	NA	NA	NA	NA	FS	Class 4
TTN	c.98039_98040insTCAA	p.Asn32680fs	N	NA	/	NA	NA	NA	NA	Ins	Class 4
TTN	c.64100_64101insTTGA	p.Asp21368X	N	NA	/	NA	NA	NA	NA	Ins	Class 4
TTN	c.53947C>T	p.Arg17983X	N	NA	/	NA	NA	NA	NA	NS	Class 4
TTN	c.61961G>A	p.Trp20654X	N	NA	/	NA	NA	NA	NA	NS	Class 4
TTN	c.44248C>T	p.Arg14750X	N	NA	/	NA	NA	NA	NA	NS	Class 4
TTN	c.80845C>T	p.Arg26949X	Y	CMD	/	NA	NA	NA	NA	NS	Class 4
TTN	c.43360C>T	p.Arg14454X	Y	CMD	/	NA	NA	NA	NA	NS	Class 4

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Table 3: Summary of clinical data according to the genetic status

	Mutated Patients (N=40)	Not mutated patients (N=55)		Single mutation (N=31)	Complex genotype (N=9)		Sarcomeric gene (N=18)	Non Sarcomeric gene (N=13)	
Mean Age (years)	43.0±15.5	48.7±15.2	p=0.07	43.5±14.5	43.1±15.3	p=0.94	42.1±15.7	45.5±12.9	p=0.78
NYHA>1 (%)	48	47	p=1.00	41	77	p=0.12	44	31	p=0.48
Mean Heart rate (bpm)	71±18	66±13	p=0.18	70.5±20	73±11	p=0.60	72±17	67±24	p=0.50
Patients with Ejection Fraction <50% (%)	57	71	p=0.19	52	67	p=0.47	61	46	p=0.48

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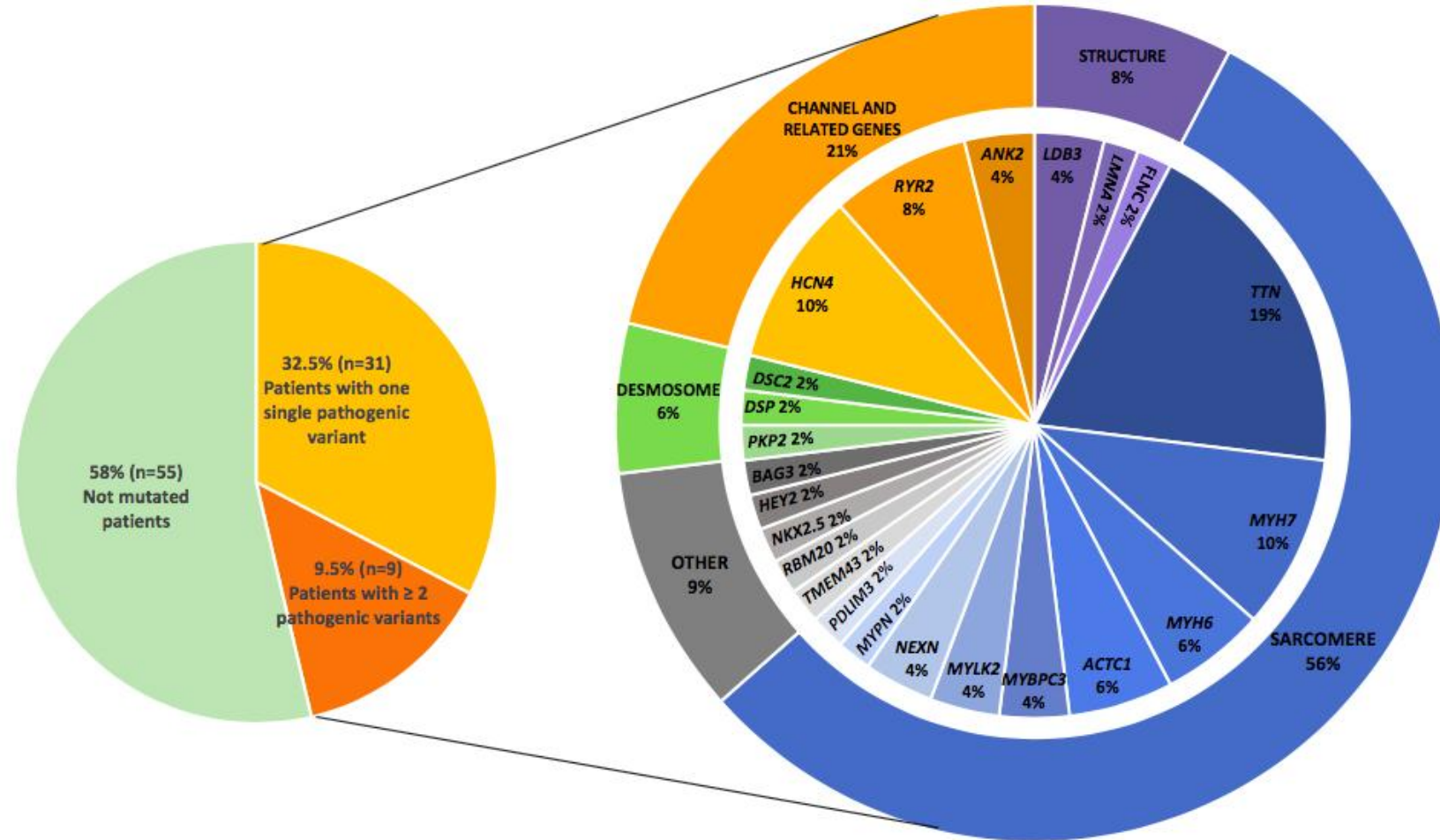
FIGURE LEGENDS

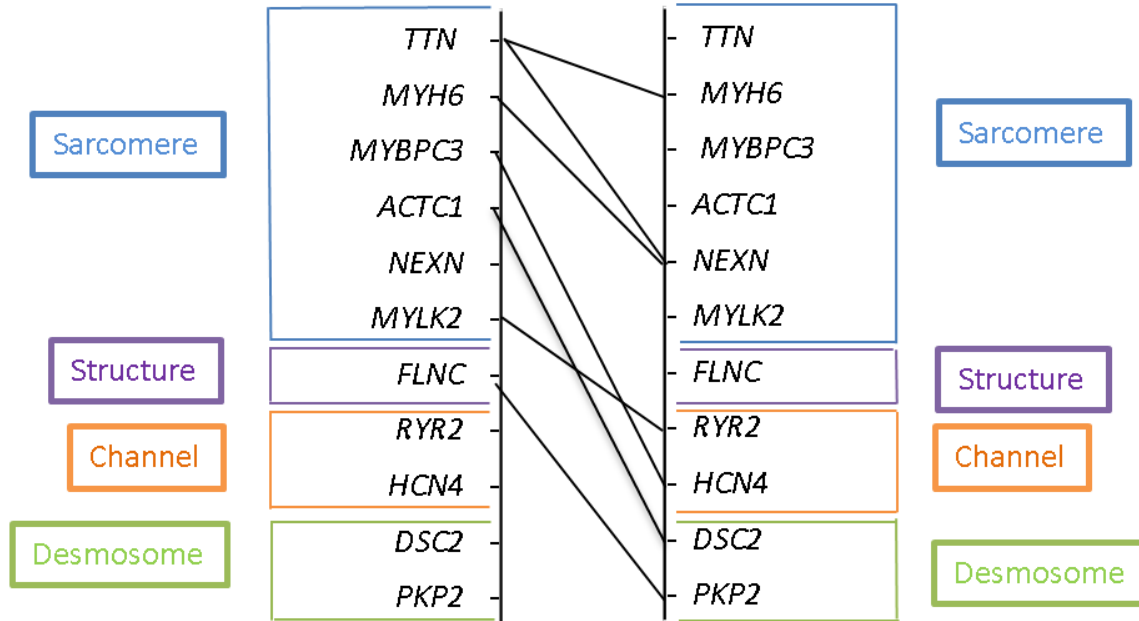
Figure 1: Distribution of genes according to their number of identified pathogenic variants and visualization of their cellular location and function.

Figure 2: Representation of genes association in patients carrying two pathogenic variants. Gene symbol were indicated on the right and left scales. For each of the seven patients carrying 2 mutations, the two mutated genes are connected by a straight line.

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Suppl Data- Table 1: List of genes analyzed in this cohort.

Gene	Reference Sequence	Chromosome
<i>AARS2</i>	NM_020745.2	chr6
<i>ABCC9</i>	NM_020297	chr12
<i>ACAD9</i>	NM_014049	chr3
<i>ACTA1</i>	NM_001100_	chr1
<i>ACTC1</i>	NM_005159.4	chr15
<i>ACTN2</i>	NM_001103.2	chr1
<i>AGK</i>	NM_018238.3	chr7
<i>AKAP9</i>	NM_005751.4	chr7
<i>ANK2</i>	NM_001148.4	chr4
<i>ANKRD1</i>	NM_014391.2	chr10
<i>BAG3</i>	NM_004281.3	chr10
<i>C2orf64(COA5)</i>	NM_001008215.1	chr2
<i>CACNA1B</i>	NM_000718.2	chr9
<i>CACNA1C</i>	NM199460.2	chr12
<i>CACNA2D1</i>	NM_000722.2	chr7
<i>CACNB2</i>	NM_201596.2	chr10
<i>CALR3</i>	NM_145046.3	chr19
<i>CASQ2</i>	NM_001232.3	chr1
<i>CAV3</i>	MN033337.2	chr3
<i>COX10</i>	NM_001303.3	chr17
<i>COX15</i>	NM_078470.4	chr10
<i>CSRP3</i>	NM_003476.3	chr11
<i>CTNNA3</i>	NM_013266.2	chr10
<i>DES</i>	NM_001927.3	chr2
<i>DSC2</i>	NM_024422.3	chr18
<i>DSG2</i>	NM_001943.3	chr18
<i>DSP</i>	NM_004415.2	chr6
<i>DTNA</i>	NM_001390.4	chr18
<i>EMD</i>	NM_000117.2	chrX
<i>EYA4</i>	NM_004100.4	chr6
<i>FBN1</i>	NM_000138.4	chr15
<i>FHL1</i>	NM_001159702	chrX
<i>FLNC</i>	NM_001458.4	chr7
<i>GAA</i>	NM_000152.3	chr17
<i>GJA5</i>	NM_005266.5	chr1
<i>GLA</i>	NM_000169.2	chrX
<i>GPD1L</i>	MN_015141.3	chr3
<i>HCN4</i>	NM_005477.2	chr15
<i>HEY2</i>	NM_012259	chr6
<i>JPH2</i>	NM_020433.4	chr20
<i>JUP</i>	NM_002230.2	chr17

<i>KCNA5</i>	NM_002234.2	chr12
<i>KCND3</i>	NM_004980.4	chr1
<i>KCNE1</i>	NM_000219.3	chr21
<i>KCNE1L</i>	NM_012282.2	chrX
<i>KCNE2</i>	NM_172201.1	chr21
<i>KCNE3</i>	NM_005472.4	chr11
<i>KCNH2</i>	NM000238.2	chr7
<i>KCNJ2</i>	NM000891.2	chr17
<i>KCNJ5</i>	NM_000890.3	chr11
<i>KCNJ8</i>	NM_004982.2	chr12
<i>KCNQ1</i>	NM_000218.2	chr11
<i>KRAS</i>	NM_004985.3	chr12
<i>LAMP2</i>	NM_002294.2	chrX
<i>LDB3</i>	NM_007078.2	chr10
<i>LMNA</i>	NM_170707.2	chr1
<i>MRPL44</i>	NM_022915.3	chr2
<i>MYBPC3</i>	NM_000256.3	chr11
<i>MYH6</i>	NM_002471.3	chr14
<i>MYH7</i>	NM_000257.2	chr14
<i>MYL2</i>	NM_000432.3	chr12
<i>MYL3</i>	NM_000258.2	chr3
<i>MYLK2</i>	NM_033118.3	chr20
<i>MYOM1</i>	NM_003803.3	chr18
<i>MYOZ2</i>	NM_016599.3	chr4
<i>MYPN</i>	NM_032578.2	chr10
<i>NEBL</i>	NM_006393.2	chr10
<i>NEXN</i>	NM_144573.3	chr1
<i>NKX2-5</i>	NM_004387.3	chr5
<i>NPPA</i>	NM_006172	chr1
<i>PDLIM3</i>	NM_014476.4	chr4
<i>PKP2</i>	NM_004572.3	chr12
<i>PLN</i>	NM_002667.3	chr6
<i>PRDM16</i>	NM_022114	chr1
<i>PRKAG2</i>	NM_016203.3	chr7
<i>PSEN1</i>	NM_000021.3	chr14
<i>PSEN2</i>	NM_000447.2	chr1
<i>PTPN11</i>	NM_002834.3	chr12
<i>RAF1</i>	NM_002880.3	chr3
<i>RANGRF</i>	NM_016492.4	chr17
<i>RBM20</i>	NM_001134363.1	chr10
<i>RYR2</i>	NM_001035.2	chr1
<i>SCN1B</i>	NM_199037.3	chr19
<i>SCN2B</i>	NM_004588.4	chr11
<i>SCN3B</i>	NM018400.3	chr11
<i>SCN4B</i>	NM174934.3	chr11

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SCN5A	NM_198056.2	chr3
SCO2	NM_001169109.1	chr22
SDHA	NM_004168.2	chr5
SGCD1	NM_000337.5	chr5
SLC25A4 (ANT1)	NM_001151.3	chr4
SNTA1	NM_003098.2	chr20
SOS1	NM_005633.3	chr2
SYNPO2	NM_133477.2	chr4
TAZ	NM_000116.3	chrX
TCAP	NM_003673.3	chr17
TGFB3	NM_003239.2	chr14
TMEM43	NM_024334.2	chr3
TMEM70	NM_017866.5	chr8
TMPO	NM_003276.2	chr12
TNNC1	NM_003280.2	chr3
TNNI3	NM_000363.4	chr19
TNNT2	NM_001001430.1	chr1
TPM1	NM_001018005.1	chr15
TTN	NM_001256850.1	chr2
TTR	NM_000371.3	chr18
VCL	NM_014000.2	chr10

Suppl. Data- Table 2: list of variants interpreted as VUS found in the cohort.

Gene name	HGVSc.	HGVSp.
AARS2	c.1752+13C>T	---
AARS2	c.44C>G	p.Ala15Gly
ABCC9	c.2424+10A>G	---
ABCC9	c.4212-31T>G	---
ABCC9	c.1981C>T	p.Arg661Cys
ABCC9	c.1165-7_1165-6delTT	---
ABCC9	c.-38C>A	---
ACAD9	c.244+7A>G	---
ACTN2	c.2057T>A	p.Ile686Asn
AGK	c.21G>A	p.Thr7Thr
AKAP9	c.11546+11G>C	---
AKAP9	c.11229G>A	p.Met3743Ile
AKAP9	c.11384A>G	p.Asn3795Ser
AKAP9	c.4246-4G>T	---
AKAP9	c.7212C>T	p.Thr2404Thr
ANK2	c.11032+45_11032+50delGTGTGT	---
ANK2	c.1288-40C>A	---
ANK2	c.2179-11A>G	---
ANK2	c.2024C>G	p.Thr675Arg
ANK2	c.5072A>G	p.Gln1691Arg
ANK2	c.7915C>G	p.His2639Asp
ANK2	c.2662C>A	p.Arg888Arg
ANK2	c.4710C>T	p.Thr1570Thr
ANK2	c.7161T>C	p.Ala2387Ala
CACNA1B	c.2268-23dupG	---
CACNA1B	c.3413+22T>C	---
CACNA1B	c.3711-41A>T	---
CACNA1B	c.4473+20G>C	---
CACNA1B	c.5777+34G>A	---
CACNA1B	c.274A>G	p.Thr92Ala
CACNA1B	c.282G>T	p.Trp94Cys
CACNA1B	c.4497C>T	p.Tyr1499Tyr
CACNA1B	c.5052C>T	p.Ala1684Ala
CACNA1B	c.6936C>T	p.Asn2312Asn
CACNA1C	c.1218-48C>T	---
CACNA1C	c.2531-39G>T	---
CACNA1C	c.5823-16G>A	---
CACNA1C	c.5930-33G>T	---
CACNA1C	c.3280A>G	p.Ile1094Val
CACNA1C	c.5519A>G	p.Glu1840Gly
CACNA1C	c.2460G>C	p.Lys820Asn
CACNA2D1	c.1516-10delT	---
CACNA2D1	c.728+29A>G	---
CACNB2	c.121_122insTTTTTT	p.Gln40_Ser41insPhePhe
CACNB2	c.1302+51_1302+52insCTTTTTTTTTTT	---
CACNB2	c.886-36dupT	---
CACNB2	c.1550A>C	p.Glu517Ala
CACNB2	c.1880G>A	p.Arg627His
CACNB2	c.1650C>T	p.Ser550Ser
COX15	c.507C>T	p.Tyr169Tyr
COX15	c.876C>G	p.Ser292Ser
COX15	c.999A>G	p.Ser333Ser
DES	c.1245-39G>A	---
DES	c.736-19G>A	---
DSC2	c.630+45G>A	---
DSC2	c.943-27A>G	---
DSC2	c.1448A>T	p.Asn483Ile
DSG2	c.1173C>A	p.Ser391Arg
DTNA	c.1138G>A	p.Ala380Thr
DTNA	c.549A>G	p.Glu183Glu
EMD	c.399+50C>T	---
FBN1	c.1148-33G>C	---
FBN1	c.1571C>T	p.Thr524Met
FBN1	c.3026C>T	p.Pro1009Leu
FBN1	c.165-7G>A	---
FBN1	c.1530G>A	p.Ser510Ser
FBN1	c.6402C>T	p.Pro2134Pro
FLNC	c.1048-11C>T	---
FLNC	c.1210+14delC	---
FLNC	c.4951+45T>C	---
FLNC	c.4301G>T	p.Arg1434Leu
FLNC	c.7546G>A	p.Glu2516Lys
FLNC	c.2733G>A	p.Lys911Lys
FLNC	c.492G>T	p.Arg164Arg
GAA	c.546+23C>G	---
GAA	c.2786G>A	p.Ser929Asn
GAA	c.2482-1G>T	---
GLA	c.427G>A	p.Ala143Thr
GLA	c.639+31C>G	---
HCN4	c.3081C>T	p.Pro1027Pro
HEY2	c.565T>A	p.Phe189Ile
HEY2	c.438G>T	p.Ser146Ser
HEY2	c.843C>G	p.Ser281Ser
JPH2	c.1836C>A	p.Pro612Pro
JPH2	c.1963C>A	p.Arg655Arg
KCNA5	c.1573C>T	p.Arg525Trp
KCND3	c.1269+18G>A	---
KCND3	c.1372-6dupT	---
KCNH2	c.431A>T	p.Asp144Val

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3	KCNH2	c.2592+3G>A	---
4	KCNH2	c.1263G>A	p.Thr421Thr
5	KCNH2	c.3258T>C	p.Pro1086Pro
6	KCNH2	c.3258T>C	p.Pro1086Pro
7	KCNJ8	c.-214C>A	---
8	KCNQ1	c.160_168dupATCGCGCCCC	p.Ile54_Pro56dup
9	KCNQ1	c.576C>A	p.Arg192Arg
10	LDB3	c.399_407delAGGCACCCC	p.Gly134_Pro136del
11	LDB3	c.668C>T	p.Ser223Leu
12	MRPL44	c.828-35A>G	---
13	MRPL44	c.792C>T	p.Thr264Thr
14	MYBPC3	c.2068-47C>G	---
15	MYBPC3	c.2994+37G>T	---
16	MYBPC3	c.2320G>A	p.Ala774Thr
17	MYBPC3	c.2602G>A	p.Gly868Ser
18	MYH6	c.1892-35A>G	---
19	MYH6	c.4828C>T	p.Arg1610Cys
20	MYH6	c.3978G>C	p.Lys1326Asn
21	MYH6	c.3979-9_3979-8delCC	---
22	MYH6	c.90C>T	p.Pro30Pro
23	MYH7	c.2419C>G	p.Arg807Gly
24	MYH7	c.5150A>T	p.His1717Leu
25	MYH7	c.571G>A	p.Val191Ile
26	MYH7	c.3246-3C>A	---
27	MYLK2	c.1658G>A	p.Arg553His
28	MYLK2	c.1754T>A	p.Ile585Asn
29	MYOM1	c.2795-26C>T	---
30	MYOM1	c.1615A>G	p.Ser539Gly
31	MYOM1	c.3283A>G	p.Ile1095Val
32	MYOM1	c.2274G>A	p.Ser758Ser
33	MYPN	c.2246G>A	p.Ser749Asn
34	MYPN	c.1899C>T	p.Asn633Asn
35	NEBL	c.2518+20C>A	---
36	NEBL	c.886A>G	p.Ser296Gly
37	PDLIM3	c.500C>T	p.Ala167Val
38	PDLIM3	c.742C>T	p.Arg248Cys
39	PKP2	c.634C>T	p.Arg212Cys
40	PKP2	c.2019C>T	p.Gly673Gly
41	PKP2	c.918C>T	p.Pro306Pro
42	PRDM16	c.3109+38_3109+41dupACAC	---
43	PRDM16	c.677-42G>A	---
44	PRDM16	c.885-14C>T	---
45	PRDM16	c.3091G>A	p.Glu1031Lys
46	PRDM16	c.561G>C	p.Gln187His
47	PRDM16	c.387+7G>A	---
48	PSEN1	c.549-15_549-14delGT	---
49	RAF1	c.771G>C	p.Ser257Ser
50	RYR2	c.11558-40T>C	---
51	RYR2	c.1292+39A>G	---
52	RYR2	c.1477-11delT	---
53	RYR2	c.2823-45C>T	---
54	RYR2	c.576+42A>C	---
55	RYR2	c.1454G>A	p.Arg485Gln
56	RYR2	c.13936G>C	p.Asp4646His
57	RYR2	c.14731C>A	p.Gln4911Lys
58	RYR2	c.4010A>G	p.Tyr1337Cys
59	RYR2	c.14808+7A>G	---
60	RYR2	c.11922G>A	p.Leu3974Leu
61	RYR2	c.12027C>T	p.Asn4009Asn
62	RYR2	c.3321G>A	p.Thr1107Thr
63	SCN2B	c.30T>C	p.Pro10Pro
64	SCN5A	c.1881G>A	p.Pro627Pro
65	SCN5A	c.4527C>T	p.Pro1509Pro
66	SDHA	c.-15G>A	---
67	SDHA	c.1894G>T	p.Val632Phe
68	SDHA	c.825C>T	p.Asp275Asp
69	SGCD	c.91C>T	p.Arg31Trp
70	SNTA1	c.160G>C	p.Gly54Arg
71	SOS1	c.3392-44T>C	---
72	SYNPO2	c.1070-13_1070-12dupTT	---
73	SYNPO2	c.1805C>T	p.Pro602Leu
74	SYNPO2	c.2818T>C	p.Ser940Pro
75	TAZ	c.331C>T	p.His111Tyr
76	TAZ	c.584-7delT	---
77	TGFB3	c.353-34C>G	---
78	TGFB3	c.517-47delG	---
79	TGFB3	c.755-28A>T	---
80	TMPO	c.280-20T>C	---
81	TMPO	c.1165G>A	p.Gly389Arg
82	TMPO	c.1235T>C	p.Ile412Thr
83	TMPO	c.1405C>G	p.Leu469Val
84	TMPO	c.1750A>G	p.Thr584Ala
85	TMPO	c.330A>G	p.Leu110Leu
86	TNNI3	c.116C>T	p.Ser39Phe
87	TNNT2	c.382-50G>A	---
88	TPM1	c.808A>G	p.Ile270Val
89	TTN	c.614_619delAGACAA	p.Lys205_Thr206del
90	TTN	c.10045A>G	p.Thr3349Ala
91	TTN	c.102116T>C	p.Phe34039Ser
92	TTN	c.107687C>T	p.Pro35896Leu
93	TTN	c.16985G>A	p.Gly5662Asp
94	TTN	c.18655G>A	p.Glu6219Lys
95	TTN	c.3295G>A	p.Val1099Met
96	TTN	c.343G>A	p.Val115Met

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3	TTN	c.39289C>G	p.Pro13097Ala
4	TTN	c.42524T>G	p.Phe14175Cys
5	TTN	c.49664C>G	p.Pro16555Arg
6	TTN	c.52373T>C	p.Val17458Ala
7	TTN	c.55000T>C	p.Cys18334Arg
8	TTN	c.60607C>T	p.Pro20203Ser
9	TTN	c.60934G>A	p.Glu20312Lys
10	TTN	c.64165C>A	p.Pro21389Thr
11	TTN	c.84203G>C	p.Ser28068Thr
12	TTN	c.86887T>C	p.Trp28963Arg
13	TTN	c.90594T>A	p.His30198Gln
14	TTN	c.95410T>C	p.Ser31804Pro
15	TTN	c.98021G>A	p.Arg32674His
16	TTN	c.27608A>G	p.Glu9203Gly
17	TTN	c.106531+6T>C	---
18	TTN	c.104913C>T	p.Ala34971Ala
19	TTN	c.12780G>T	p.Ala4260Ala
20	TTN	c.52878C>T	p.Val17626Val
21	TTN	c.71340C>T	p.Thr23780Thr
22	TTN	c.98061T>C	p.Ala32687Ala
23	VCL	c.239+23T>C	---
24	VCL	c.1223T>C	p.Ile408Thr
25	VCL	c.2655C>T	p.Phe885Phe
26	VCL	c.2760C>T	p.Ala920Ala
27	VCL	c.804A>G	p.Arg268Arg
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NB: For recessive gene variants were retain only if the patient is carrier of another variant in the same gene.