**FLNC missense variants in familial noncompaction cardiomyopathy**

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**Abstract**

The majority of familial noncompaction cardiomyopathy (NCCM) is explained by pathogenic variants in the same sarcomeric genes that are associated with hypertrophic (HCM) and dilated (DCM) cardiomyopathy. Pathogenic variants in the filamin C gene (FLNC) have been linked to HCM and DCM. We expand the spectrum of FLNC related cardiomyopathies by presenting two families with likely pathogenic FLNC variants showing familial segregation of NCCM and concurrent coarctation of the aorta and/or mitral valve abnormalities.

**Introduction**

Noncompaction cardiomyopathy (NCCM) is characterized by excessive trabeculation of the left ventricle (LV) with a noncompacted to compacted ratio of more than 2 according to current echocardiographic criteria, or 2.3 on CMR.1,2 Approximately 10% of patients diagnosed with NCCM have concurrent congenital heart defects (CHD).3,4 In 30-40% of cases diagnosed with NCCM a pathogenic variant can be identified. Around 80% of these pathogenic variants involve the same sarcomere genes, that are the major causes for hypertrophic cardiomyopathy (HCM) and dilated cardiomyopathy (DCM), in particular MYH7, MYBPC3 and TTN.5,6 Filamin C (FLNC) plays a central role in muscle functioning by maintaining the structural integrity of the muscle fibers. Pathogenic variants in FLNC were found to be associated with a wide spectrum of myopathies ranging from cardiomyopathies to distal skeletal myopathies. Truncating FLNC variants were previously associated with dilated cardiomyopathy7,8 and missense variants were identified in familial HCM and restrictive cardiomyopathy. FLNC has not been associated with NCCM or CHD before.

We present two Dutch families where familial NCCM with CHD were linked to rare FLNC missense variants. These observations suggest that the spectrum of clinical manifestations of FLNC variants include familial NCCM with CHD.

**Case Reports**

**Family A**

In this family (Figure 1A), a 52-year-old woman (II:3) was first diagnosed with NCCM when she underwent cardiologic examination for a suspected perimyocarditis. Echocardiography showed pericardial effusion and normal LV dimensions without LV dysfunction. The LV walls showed hypertrabeculation with end-systolic noncompacted/compacted (NC/C) ratio >2. Electrocardiographically, inferolateral repolarization abnormalities were observed. Cardiac magnetic resonance imaging (CMR) confirmed the diagnosis of NCCM with diastolic NC/C ratio >2.3 in the LV inferoseptal wall. She had elevated CK levels of 1234 U/L [ref <200 U/L]. No signs for neuromuscular disease were detected at neurologic examination. After seven years of follow-up, she remained cardiologically asymptomatic (NYHA class I).

Family screening revealed NCCM in two relatives. A niece (III:3), was diagnosed with NCCM at age 21 and had surgery at age seven for coarctation of the aorta (CoA). She fulfilled both the echocardiography and CMR diagnostic criteria for NCCM and had excessive long chordae of the anterior mitral valve leaflet (Figure 2A and B). The noncompaction had not been recognized in the past on echocardiography. Cardiologic screening of an asymptomatic brother (II:4) at age 54 years showed that he also had NCCM, with a NC/C ratio of 2.3 on echocardiography and a ratio of 2.9 on CMR. No previous cardiac imaging had been performed. He had elevated CK-levels of 265 U/L without neuromuscular signs. Ten years after the diagnosis NCCM he had an episode of atrial fibrillation that required electric cardioversion. CMR from the brother (II:1) and the son (III:1) of the proband were performed at age 57 and 15 years, respectively, showing borderline NC/C ratios of respectively 2.1 and 2.2 on MRI, i.e. just below the diagnostic criteria. Proband III-2 did not participate in the family screening.

**Family B**

In family B (Figure 1B) the diagnosis NCCM in a 17-year-old boy (III:1) was made by echocardiography. He was referred because of multiple unexplained episodes of syncope. He also had a ventricular septal defect (VSD) and a mild mitral valve prolapse (MVP). CMR revealed partial LV noncompaction from the apex to midventricular region with an NC/C ratio of 3.0 (Figure 2C and D). An implantable cardioverter-defibrillator (ICD) was implanted. After 9 years of follow-up, the LV function remained normal without ICD shocks. CK-
levels were elevated (419-1188 U/L) in the absence of neuromuscular signs. His mother (II:2) was under cardiologic surveillance because she had a VSD, MVP, CoA and a bicuspid aortic valve. The CMR showed that she complied for the diagnostic criteria for NCCM with a NC/C ratio of 2.4. She had diastolic LV dysfunction, with preserved LV systolic function and underwent multiple cardiac ablations for atrial fibrillation. At age 44 years she experienced severe bradycardia, which necessitated cardiac resuscitation, resulting from combined flecainide and metoprolol treatment. A pace-maker was implanted. Her highest CK-level was 1174 U/L. The proband’s brother (III:2) also suffered multiple episodes of syncope and was diagnosed with NCCM at age 19, with a NC/C ratio of 3.1 on CMR. His highest CK-level was 294U/L. Proband III-3 was screened cardiologically and had no signs of NCCM on echocardiography. No DNA analysis was performed.

Genetic testing

Diagnostic DNA NGS targeted testing of a panel of 54 cardiomyopathy genes, that did not include FLNC, as presented in Table 1, did not reveal a genetic cause for NCCM in the two index cases. Also single-nucleotide polymorphism-array DNA testing showed no structural DNA changes. Subsequently, whole exome sequencing was performed in the NCCM patients II:1, III:1 and III:3 from family A and II:1, III:2, and II:2 from family B. Patients II-2 from family A and III:1, III:2, and II-2 from family B were included because we suspected that a causative mutation may underlie a spectrum of cardiac phenotypes. Written informed consent was obtained from all participating family members. The investigation conforms to the principles outlined in the Declaration of Helsinki. Variants were annotated using ANNOVAR19 and filtered using an in-house developed pipeline. Only variants segregating within each family, affecting exons or splice sites, with a population frequency below 0.01 in ExAC, NFE, GnomAD. GoNL were kept. For in silico prediction of the effect of non-synonymous variants we used align GVGID,11 SIFT12 and Polyphen13 and ensemble scores LR and Radial SVM.14 We selected variants who were predicted to be damaging by 3 of 5 prediction programs. Segregating synonymous variants and variants predicted to be tolerated were excluded. In Family A, two candidate genes remained after filtering, MYH4 and FLNC, of which only the last was previously associated with cardiomyopathy. A variant in FLNC (c.6397C>T, p.(Arg2133Cys)), NM_001458.4, confirmed by sanger sequencing) segregated with the cardiac phenotype of NCCM in the three NCCM patients and in the two relatives with borderline NCCM features. A variant in the same location (p.(Arg2133His)) was previously reported in a family with HCM and classified as probably pathogenic.15 In family B, a novel FLNC variant (c.7177C>T, p.(Pro2393Ser)) was identified. The two FLNC variants were absent in the Genome Aggregation Database (http://gnomad. broadinstitute.org), affect highly conserved amino acids, and were predicted to be deleterious by multiple in silico prediction programs. No FLNC variants were found in thirteen unrelated NCCM patients without a CHD and without a pathogenic variant in 48 cardiomyopathy genes.

Histology

Right ventricular endomyocardial biopsy (RVEMB) samples from the proband of family B (III:1) were stained with hematoxylin and eosin as well as Masson’s trichrome. To visualize protein aggregation and autophagic activity in cardiomyocytes, immunohistochemistry for microtubule-associated protein 1A/1B-light chain 3 (LC3) was performed, as described previously.16 Light microscopic analysis showed nonspecific cardiomyopathic changes of myocyte hypertrophy and increased interstitial fibrosis (Figure 3A). The RV did not show an excessively thickened endocardial layer or hypertrabeculation, or intracellular aggregates or autophagic activity (Figure 3B).

Discussion and conclusions

This is the first report linking FLNC to NCCM in two families with rare FLNC variants. In these two families the cardiac phenotype included NCCM, NCCM with concurrent CoA, NCCM with concurrent VSD and MPV and also a NCCM patient with a complex CHD consisting of VSD, MVP, CoA and bicuspid aortic valve, and myocardial dysfunction. These observations suggest that missense FLNC variants may

Table 1. Diagnostic DNA NGS targeted testing of a panel of 54 cardiomyopathy genes (not including FLNC).

<table>
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<th>Diagnostic DNA panel</th>
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<tr>
<td>ABC29, ACTC1, ACTN2, BAG3, CALR3, CASQ2, DES, DSC2, DSG2, DSP, DTVN, EMD, EYA4 FH1, FKTN, GATA4J, GLA, HCN4, JPH2, JUP, LAMA4, LAMP2, LDB3, LMNA, MIB1, MYBP3, MYH6, MYH7, MYL2, MYL3, MYLK2, MYOZ1, MYOZ2, MYPN, NEB1, NECN, NKC3-5, PKP2, PLN, PRDM16, RBM20, RYR2, SCN5A, TAZ, TBX20, TCFAP2B, TMEM43, TNCC1, TNIN3, TNNT2, TPM1, TTN, TTR and VCL</td>
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cause familial NCCM with or without one or more structural heart defects. Missense
FLNC variants in the N and C terminal domains of FLNC have been associated with hypertrophic and restrictive cardiomyopathy, causing sarcomeric aggregates containing FLNC leading to sarcomere dysfunction. Other FLNC domains were associated with myofibrillar myopathy, showing similar intracellular aggregates in skeletal muscles. A large study of inherited cardiovascular disease patients showed that truncating FLNC variants were associated predominantly with an overlapping phenotype of severe dilated- and arrhythmic cardiomyopathies. The cardiomyopathy phenotypes associated with FLNC variants have not included NCCM so far, however signs of LV hypertrophy are not fulfilling NCCM diagnosis in some carriers of a truncating FLNC variants have been reported. FLNC has not been linked to CHD, so far, to the best of our knowledge.

Aortic coarctation in NCCM patients seems rare with an estimated prevalence of less than 1%. We report two families with familial NCCM and a likely pathogenic FLNC variant in which CoA occurred; in one family one patient had NCCM with concomitant CoA. In the other family a NCCM patient with the familial FLNC variant had complex congenital cardiac defects including a CoA. It remains to be elucidated how FLNC and other genes can cause a cardiomyopathy with or without a CHD, and skeletal myopathy in other patients. It may be that FLNC resembles MYH7 in that aspect. Because among all the known genes associated with cardiomyopathies as well as skeletal myopathies, MYH7 variants occurs the most frequent in NCCM, and is also linked to Ebstein anomaly with and without NCCM, HCM, DCM, or Laing distal myopathy. Suggesting that variants affecting distinct domains of sarcomeric proteins may define the spectrum of cardiac and skeletal muscle phenotypes.

Pathogenic FLNC variants are expected to disrupt the structure of the sarcomeric protein, leading to the formation of protein aggregates resulting in an impairment of the sarcomere function. One of the identified variants in this study, FLNC p.(Arg2133Cys), may have a similar effect as the reported FLNC variant p.(Arg2133His) at the same location with another amino-acid substitution, that was shown to have disrupting actin aggregates in cardiac tissue. We found elevated CK levels in two of the three NCCM patients in both affected families. Elevated CK levels were also noted in a previous study regarding FLNC variants with myopathy but also in patients with only a cardiac phenotype of HCM. For the novel FLNC variant p.(Pro2393Ser), we observed fibrosis in the RV myocardium samples, indicating a damaging effect of the variant on the cardiac muscle. Fibrosis was also observed in previous reports with FLNC mutations. The cardiac fibrosis observed in the patients with the FLNC variant suggests that similar pro-fibrotic mechanisms may be involved as observed in MYBPC3 cardiomyopathy. Similarly, to the original report no signs of intracellular aggregates or autophagic activity in the RV of the patient or in patients with a FLNC related cardiomyopathy were noted. The RV of this patient did not show evidence for hypertrophy morphologically or on imaging. However this does not exclude an effect of the FLNC variant on the left ventricle, since RV hypertrophy in NCCM is rarely reported, and the NCCM presents predominantly with a LV phenotype.

As previous studies showed, genetics plays an important role in approximately half of the NCCM cases. The genetics of NCCM are complex and affect mostly genes associated with myopathies including sarcomere or mitochondrial dysfunctioning. In this perspective FLNC fits into the genetically heterogeneous background of NCCM. Further studies are needed to assess the exact role and mechanisms of FLNC in NCCM, aortic coarctation and mitral valve abnormalities.

Figure 2. Imaging of two NCCM patients. Family A. patient III.3; A) Short axis of the left ventricle on cardiac magnetic resonance (CMR) showing the prominent trabeculae and intertrabecular recesses in NCCM; B) Echocardiogram of the LV of patient (III.3) with NCCM and excessive long chordae of the anterior mitral valve leaflet (arrow). Family B. patient III.1; C) Two-chamber long-axis of the left ventricle on CMR; D) Short axis view of the left ventricle on CMR showing the prominent trabeculae and intertrabecular recesses in NCCM.


19. Vermeer AM, van Engelen K, Postma AV, et al. Ebstein anomaly associated with left ventricular noncompaction: an autosomal dominant condition that can...


