Genetic Bases of Hypertrophic Cardiomyopathy

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Hypertrophic cardiomyopathy is characterized by myofibrillar derangement and predominant hypertrophy of the interventricular septum associated with a reduced or normal left ventricular cavity. It determines diastolic dysfunction and a tendency towards myocardial ischemia, arrhythmia, and sudden death. With a prevalence of 1 case per 500 individuals, it occurs in all age groups, from birth to the 8th decade, having a genetic origin preliminarily detected from familiar forms among carriers of the disease.

The complex character of this entity outlined in the first reports of the modern era became unequivocally evident by systematic studies developed over recent decades. The application of a methodology generated in molecular genetics by obtaining genetic maps of high resolution facilitated the redefinition of concepts related to the pathology, physiopathology, and diagnosis of hypertrophic cardiomyopathy. Genetic analysis facilitating identification of the disease in its preclinical phase and risk stratification based on molecular criteria render the future introduction of genetic therapy viable.

Hypertrophic cardiomyopathy has highly heterogeneous morphological, functional, and clinical characteristics. The phenotype is composed of ventricular hypertrophy associated with myofibrillar disorganization. Marked phenotype variation relative to the magnitude and extension of the myocardial hypertrophy is observed. Echocardiographic evaluation reveals predominant hypertrophy of the intraventricular septum of diffuse character, commonly extending to the free anterior-lateral wall of the left ventricle. Wall thickness is usually above 20mm, although measurements between 20 and 15mm have been found. Borderline thicknesses of 15mm or even 13mm have also been identified. Asymmetric forms predominate over concentric ones, representing 1 to 2% of the cases. Myofibrillar disarray extends over 5 to 30% of the myocardial tissue and has a low correlation with the degree of hypertrophy.

Hypertrophic cardiomyopathy can appear without showing its main morphological characteristics. Arrhythmias and sudden death in the absence of echocardiographic signals of left ventricular hypertrophy (LVH) in patients with the genotype, indicate incomplete phenotype expression. It is concluded that the phenotype is represented by a continuous spectrum, from macroscopically normal to severe forms showing massive myocardial hypertrophy.

The phenotype changes with age. Hypertrophic cardiomyopathy can appear at birth showing serious obstructive forms with marked hypertrophy, rapid evolution to cardiac failure, and a high level of mortality. It more frequently manifests clinically in adolescence when a detectable increase in parietal thickness of the left ventricle accompanies an increase in weight and stature. Although clinical evolution in children and adolescents is usually benign, annual mortality can reach 4.8%, higher than that shown among adults. Adolescents and young adults can manifest extreme degrees of LVH with outflow tract obstruction, cavity reduction, and evolution towards sudden death. An inverse relationship between age group and left ventricular parietal thickness may be shown. Gradual regression in hypertrophy, increased ventricle volume, and diastolic dysfunction occur in approximately 10% of individuals reaching maturity. Late forms, manifest after 65 years of age, are individualized, with a hypertrophy generally restricted to the interventricular septum.

The natural history is also heterogeneous. While some cases remain in asymptomatic form indefinitely, other patients experience sudden death or evolve to cardiac failure. Although annual mortality recorded at reference centers falls between 2 and 6%, measurements among less selected populations fall between 0.5 and 1.5% of cases. Clinical indicators like degree of hypertrophy, records of ventricular arrhythmia, and abnormal response to exercise, have a limited value for risk stratification of sudden death.
The clinical complexity characterizing hypertrophic cardiomyopathy becomes even more evident when its genetic aspects are analyzed. Transmission by autosomal dominant heredity attributes the same genetic disturbance to all affected members of a family. Although preliminary verification has estimated that only 50% of the cases show a familial character, recent more evaluations consider that truly sporadic forms are borne by less than 10% of disease carriers. Three factors may underestimate the diagnosis of genetic forms: evaluation of very small families, adoption of rigid echocardiographic criteria, and the presence of mutations of variable penetrance. Echocardiographic manifestations of LVH may be absent or have a subclinical character, especially among children and adolescents. The electrocardiogram may be useful for identifying these patients. About 50% of adult carriers of the genotype who have normal wall thickness on the echocardiogram have electrocardiographic alterations. The large clinical variability among families affected by hypertrophic cardiomyopathy suggests the involvement of more than 1 gene with phenotypic manifestations in common. In 1989, Jarcho et al, using genetic linkage analysis, mapped on chromosome 14q1, the first locus related to the disease, later described as CMH1. One year later, Geisterfer-Lowrance et al, using genetic mapping and DNA sequencing, individualized the point mutation affecting the gene that codifies the $\beta$-miosin heavy chain, considered to cause the disease.

Up to the present time, 7 further genes have been related to hypertrophic cardiomyopathy; they encode, respectively, cardiac troponin C, $\alpha$-tropomyosin, myosin-binding protein C, essential and regulatory myosin light chains, cardiac troponin I, and cardiac actin (Table I). Reports of a mutation involving the gene of titin and of an as yet unidentified gene at locus 7q3 in forms associated with the Wolff-Parkinson-White syndrome are still under evaluation. The genetic heterogeneity would partially justify the clinical variability among carriers of the disease. The finding that phenotype expression has intrafamilial variability suggests the participation of other factors, genetic or environmental.

Although from the biomolecular viewpoint, hypertrophic cardiomyopathy can be interpreted as a multiple entity, the fact that the genes affected encode sarcomere proteins gives a unifying character to the entity. About 100 mutations have been described, corresponding to 2/3 of the total. One, therefore, observes allelic heterogeneity, several mutations involving the same gene being capable of evoking the disease.

In the human genome, approximately 30,000 genes encode structural proteins and enzymes. They constitute DNA segments formed by nucleotide sequences with a particular location in the chromosome, called the chromosome locus. A nitrogen base, the deoxyribose sugar, and a phosphate group form the nucleotides. Two types of nitrogen bases exist, purines [adenine (A) and guanine (G)] and pyrimidines [cytosine (C) and thymine (T)]. Genes have a variable dimension, from a few hundred to thousands of base pairs. They have paternal and maternal copies called alleles, and their production results from the combination of both. The base-pair sequence in DNA is used to form messenger RNA (mRNA), which takes information to the ribosomes localized in the cytoplasm where protein synthesis takes place. The RNA molecule is synthesized directly from a DNA template, a process called transcription. Human genes have base sequences called exons that encode amino acids and are transcribed into mRNA and nonfunctional sequences, the introns. At the final step of the formation of mRNA, the intron sequences are removed and the exons united by a process known as splicing, taking place in ribonucleic corpuscles. The removal of the intron sequences is performed using the donor and acceptor sites located at the exon-intron borders. The base sequence of the mRNA will determine the amino acid sequence of the proteins. If an error occurs during the splicing process, a modified protein is formed. The genetic code contained in the DNA helix and in the mRNA is essential for the formation of the different types of proteins. Base triplets, the codons, form it. Sixty-four codons exist, of which 61 encode amino acids and 3 are termination codons, which have no encoding function. The genetic code describes the relationship between the base sequences and the amino acid sequence of the protein. Point mutations determine permanent changes in the base sequence of a DNA molecule by substituting a pair of bases with a reflex on the encoded protein.

### Table I – Genes related to hypertrophic cardiomyopathy and chromosome location

<table>
<thead>
<tr>
<th>Gene</th>
<th>Chromosome</th>
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<tbody>
<tr>
<td>Cardiac $\beta$ myosin heavy chain</td>
<td>14q1</td>
</tr>
<tr>
<td>Cardiac T tropinin</td>
<td>1q32</td>
</tr>
<tr>
<td>$\alpha$ tropomyosin</td>
<td>1q22</td>
</tr>
<tr>
<td>Myosin binding C protein</td>
<td>11p11.2</td>
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<tr>
<td>Essential myosin light chain</td>
<td>3p21.2-p21.3</td>
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<td>Regulatory myosin light chain</td>
<td>12q23-q24.3</td>
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<tr>
<td>Cardiac 1 tropinin</td>
<td>19p13.2-q13.2</td>
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<tr>
<td>$\alpha$ cardiac actin</td>
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**Mutations of the gene of the cardiac $\beta$-miosin heavy chain** - About 35% of the cases of hypertrophic cardiomyopathy are caused by mutations of the gene of the heavy chain of cardiac $\beta$ myosin, which encodes the predominant isoform expressed in the ventricles. The thick filaments contain 2 proteins of the cardiac $\beta$ myosin heavy chain that are connected to 4 molecules of the light myosin chains. The first contains 30% cardiac myosin, comprising 95% ventricle myosin. It is a contractile protein with enzymatic activities.
action, which in the presence of actin hydrolyzes ATP into ADP and phosphate. The molecule formed by 1,934 amino acids has an asymmetric shape and distinct segments. The globular head contains the actin binding sites and ATP. The neck region is associated with the light myosin chains, and the rod like tail, in turn, establishes the connection to myosin molecules that form the thick filaments, each made up of 400 molecules.

In their near-totality, mutations that affect the gene of the cardiac β myosin heavy chain are point mutations of the type described as “missense mutations” in which the substitution of a single base of the DNA in a given exon results in the exchange of the encoded amino acid. Deletion of the nontranslated 3’ region of the gene, inducing the loss of at least 5 amino acids in the rod of the protein has also been reported. Deletions in codons 10 and 930, which do not result in a change in the DNA reading frame, have been subsequently described.

At least 50 mutations of the gene of the cardiac β myosin heavy chain have been identified affecting unrelated families. It is estimated that the greater part has been already individualized. All cause common manifestations of hypertrophic cardiomyopathy. The mutations described cause the disease and do not constitute polymorphism connected to adjacent genes, unrelated to the disease.

Mutations with changes in the encoded amino acid determine alterations in the protein structure of the gene product. The substitution of the amino acid takes place in the protein molecule. Mutated residues are localized at the interface with actin and the binding with ATP and the myosin light chains. The mutant allele produces an anomalous protein that, when incorporated into the thick filaments, determines contractile dysfunction, even though the normal allele produces an integral protein. It is, therefore, a dominant negative mutation and may or may not determine the change in the charge of encoded amino acid, altering the biomolecular conformation of the polypeptide.

The cardiac β myosin heavy chain polypeptides are encoded in 30 KB of DNA, arranged in 40 exons; this renders the analysis of each one extremely laborious and time-consuming. The near-totality of the missense mutations is localized in the first 23 exons, which encode the globular head and the head-tail junction of the protein. This gene is highly susceptible to mutagenesis in special codons 403, 719, and 741. It is suspected that the arginine residue plays a fundamental role in the protein’s normal function. However, no mutation has a predominant character, although Arg403Gln is a frequent one. The greater part of the mutations result from the exchange of guanine or cytosine.

The mutations of the cardiac β myosin heavy chain were the first to be described, allowing greater knowledge of the genotype-phenotype relationship, degree of penetrance, and clinical evolution. Yet, the analysis of these aspects is based on the evaluation of a still limited number of cases, not rarely originating from a single family. The clinical valorization of the information presently available depends on confirmation at a higher scale, using data resulting from the identification of new affected families.

Mutations of the cardiac β myosin heavy chain are frequently associated with expressive myocardial hypertrophy with high penetrance. Although hypertrophy has a similar degree and extension in the diverse mutations of the gene, the prognosis shows variable characteristics. Consequently, risk stratification can only be determined on the basis of genetic analysis, because many mutations of the same gene have the phenotype in common.

Certain mutations in the cardiac β myosin heavy chain are considered malignant due to the high incidence of sudden death. They are associated with a change in the charge of the encoded amino acid. Benign mutation can also exhibit this characteristic. Malignant forms encode severe myocardial hypertrophy and complete penetrance. Arg403Gln, the first described mutation results from the exchange of adenine by guanine in exon 13, resulting in the substitution of arginine by glutamate in codon 403. It causes premature sudden death, which in 50% of cases occurs before the age of 30. The same mutation was described in a small Korean family with a low incidence of sudden death, despite its high penetrance. Arg719Gln and Arg453Cys are equally serious, with a high incidence of sudden death and an average survival time of 38 and 30 years, respectively. Gly716Arg, with a penetrance of 100%, precocious phenotype expression, and premature sudden death, evolves towards cardiac failure in the adult.

Mutations Glu930Lys and Arg249Gln are of intermediate risk. Leu908Val has a better prognosis: 92% of patients are alive at 60 years. Gly256Glu like Arg403Trp and Arg870His are also benign. Mutation Val606Met is described as benign but can eventually have a bad prognosis. Mutation involving the same codon, like Arg403Gln, Arg403Leu, and Arg403Trp can have a distinct prognosis and a variable degree of penetrance.

The analysis of many families with an identical mutation indicates that the mutations can have an independent origin. Mutations of the gene of the cardiac β myosin heavy chain gene are also identified in carriers of sporadic forms, demonstrating that familiar and nonfamiliar forms make up the same spectrum. It consists of spontaneous mutations transmitted to the descendants with expression on the mRNA and the carrier protein; this may not be so in the case of a very small family.

In mutations of the gene of the cardiac β myosin heavy chain, because penetrance is usually high, it is an exception for carriers of the genotype not to exteriorize the phenotype until adolescence. The high penetrance conditions the carrier for severe hypertrophy. Only a few muta-
tions, like Arg403Leu, Arg403Trp and Asn232Ser have low penetration. In these cases, hypertrophy may be mild or borderline. Variation in the degree of hypertrophy is observed between families carrying the same mutation.

The greater part of the mutated protein is expressed in the sarcomere of the myocardial fibers, directly affecting the generation of contractile force. Analysis of endomyocardial biopsies reveals the loss of sarcomere filament alignment, conditioning to miofibrillar disarray. The mutated myosin is also expressed in the skeletal musculature producing myopathy with a predominance of slow-type myofibrils and absence of mitochondria.

**Mutations of the cardiac troponin T gene** - In 1993, Watkins et al. individualized a second locus related to cardiac hypertrophy, situated in chromosome 1q32, and then named it CMH2. One year after, the same workers described 3 mutations affecting the gene Troponin T, mapped on this locus.

Cardiac T troponin mutations represent about 15% of hypertrophic cardiomyopathy cases. Various isoforms originated by recomposition are expressed in the myocardium. Thin filaments are formed by actin, tropomyosin and troponins T, I, and C. Contraction depends on actin-myosin interaction, regulated by a tropomyosin and the troponin complex. Once calcium-C troponin binding is established, the troponin-tropomyosin complex releases actin, enabling the binding of the globular heads of cardiac β myosin to the thin filaments.

The gene of cardiac T troponin is composed of 17 exons distributed over 17 KB of DNA. Missense mutation in a total of 10, localized in exons 8, 9, 11, 14, and 16 predominate. De novo mutation in exon 9, Arg92 Trp, has recently been reported. Deletion of 3 nucleotides corresponding to all of the codon of glutamic acid, ΔGlu160, was also identified. Splice site mutations, involving the donor site of intron 15, Int15G-A, generates a truncated, rapidly degraded protein, the consequence of the loss of 28 terminal amino acids. Preparations containing intact sarcomeres with a high expression of truncated cardiac troponin T show a marked reduction in their contractile function. In the presence of low calcium concentrations, relaxation is impaired by interference with the inhibitor function of cardiac I troponin.

Clinical manifestations are less heterogeneous than those evidenced by mutations of the cardiac β myosin heavy chain. For the most part, they determine similar phenotypes and show low penetrance. Hypertrophy generally of a mild or subclinical grade may be absent in 25% of the cases, according to echocardiographic findings. Maximal wall thickness is between 11 and 16mm, in contrast to that produced by mutations in the cardiac β myosin heavy chain, average 24mm. Although frequently asymptomatic, prognosis is comparable to that of malign mutations of the cardiac β myosin heavy chain. Mortality tends to be high among young men. There occurs a high incidence of death, mostly of a sudden character, before the age of 30. As a rule, delayed death due to cardiac failure is not observed. At least 6 mutations have a poor prognosis: Ile79Asn, Arg92Gln, Arg92Trp, Ala104Val, D Ghu160 and Int15G-A,56. The sole exception is that of Phe110Ile, which affects 13% of families with a mutation of the cardiac troponin T gene; in Japan it is occasionally associated with apical hypertrophy. As in the benign mutations of the cardiac β myosin heavy chain, no change occurs in the charge of the amino acid involving one of the main sites of binding with the cardiac α actin molecule.

It has been possible by reanalysis of the genome organization of the gene of cardiac troponin T to identify codon 102 as the point susceptible to mutation. Distinct characteristics were observed in one of the mutations involving this codon, Arg102Leu, in which high penetrance with a marked degree of hypertrophy was noted.

A mutation of the cardiac troponin T gene associated with extensive myofibrillar disarray without an increase in cardiac mass has also been reported in a family with sudden death. The mutation involves exon 9 with a substitution of arginine by leucine in codon 94, producing an exchange of amino acid charge and protein conformation change. It can be deduced that hypertrophy may not be the factor determining sudden death in hypertrophic cardiomyopathy but rather, the myofibrillar disarrangement associated with fibrosis. In a contemporaneous form, mutation Arg278Cys has been described in a 67-year-old subject who 10 years ago was normal from clinical and electrocardiogram points of view. All known cases until then had begun in adolescence. It is concluded that mutation of cardiac troponin T can become clinically manifest later in life, justifying cases of sudden death in middle-aged, previously asymptomatic individuals, with a normal or slightly hypertrophic left ventricle.

**Mutations of the α tropomyosin gene** - The evaluation of families that did not exhibit a linkage with CMH1 and CMH2 facilitated the identification by Thierfelder et al. in 1993 of a third locus, mapped on chromosome 15q22, called CMH3. The gene composed of 14 exons, which encodes a tropomyosin, is localized in the long arm of this chromosome and transcribes 1kb RNAm. It is only implicated in the development of a few, about 5%, of the cases of the disease.

α tropomyosin is a 284 amino acid polypeptide, which constitutes the thin filaments and is abundantly expressed in the skeletal musculature. It represents 5% of the myofibril protein. Localized in a major groove of the actin filament, it has a preponderant role in the composition and organization of the thin filaments by establishing the binding between the troponin complex and actin.

Mutations of this gene, numbering 4, are all missense mutations. Initially 2 mutations on exon 5 were described, Asp175Asn and Glu180Gly. Later, Ala63Val and
Lys70Trh were reported. Some of these mutations can interfere with calcium regulation in the thin filaments. Contrary to mutations of other genes, they are expressed in various structures like gametes, lymphocytes, and myocytes, but being only capable of producing myocardial disease.

Mutation Asp175Asn seems to predominate and produced a mutated protein expressed in the myocardium and skeletal musculature. The recurrent identification of the G-A transition in exon 5 indicates a high propensity toward mutagenesis in the 579 residue of α-tropomyosin.

De novo mutation involving the α-tropomyosin gene have been reported. They can result in transmission to descendants and can cause familial hypertrophic cardiomyopathy, as already referred to in relation to mutation Asp175Asn. Carriers of apparently sporadic forms of mutation of this gene should be warned about the risk of transmission. Spontaneous mutations appear precociously in gamete development or during embryonic life.

The clinical consequences of mutations of the α-tropomyosin gene are less known. The phenotype is characterized by generally mild hypertrophy, but variations in the degree and extension of the hypertrophy between carriers of the same mutation have been observed. From the histopathological point of view, no large difference relative to gene mutation encoding proteins composing the thick filaments exists. In Asp175Asn, hypertrophy is of a variable degree and extension among members of distinct families. Prognosis is favorable, with a life expectancy close to normal, comparable to that with benign mutations of the β myosin heavy chain. Glu180Gly produces less hypertrophy, but has a similar survival curve. Ala63Val and Lys70Thr are described as having a less favorable prognosis, with evolution towards cardiac failure. Therefore, in relation to the α-tropomyosin gene, the type of mutation determines the clinical expression.

Mutations of the gene of the myosin-binding protein C (CMH4) - In 1993, a new locus situated on the 11p11.2 chromosome called CMH4, was identified. Subsequently, Watkins et al. simultaneously with Bonne et al. described the first 3 mutations of the gene encoding the myosin-binding C-protein in unrelated families, considered as causing the disease.

The myosin-binding protein C is a myofibril protein that does not directly participate in contraction. Its expression is restricted to the myocardium. It is placed transversally in the A bands and binds the myosin of the thick filaments and titin of the elastic filaments. Its function is little known, but it modulates speed of contraction in response to adrenergic stimulation.

The sequence and organization of the gene have been recently described. Formed by the 21 KB DNA, it is made up of 35 exons. Analysis of the genetic linkage shows that 20% of the cases of hypertrophic cardiomyopathy are caused by this gene. About 28 distinct mutations have been described in unrelated families. Missense mutations and nonsense mutations, in which a single substitution of a base of the DNA leads to the appearance of the terminal codon, precociously closing the synthesis of protein, have been reported. Mutations by deletion, insertion, duplication, or splicing have also been documented. The consequences of the mutations on the structure and function of the protein are as yet not well known.

Myosin-binding protein C is composed of 1,274 amino acids, required for its incorporation into the sarcomere. Mutations with amino acid exchange presumably act by a dominant negative effect. Others like duplicating or splice site mutations modify the reading matrix of the DNA, generating truncated proteins not having the terminus responsible for the binding to myosin. The truncated protein can bind in a defective manner to myosin or not be incorporated into A band of the sarcomere, preventing the interaction of the contractile unit with the cytoskeleton. Another possibility is that due to the low stability of the transcript, because no production occurs of the stable protein, the mutation acts as an null allele. In the recently described splice site mutation, the isolated insertion of guanine in exon 25 leads to the loss of 40 bases on the mRNA, which is then processed in an abnormal manner. The truncated protein generated could not be isolated by endomyocardial biopsy, presumably because it is expressed at very low concentrations or by not being present. Insufficient synthesis or rapid degradation may justify lack of detection of the mutated protein.

Mutations of the gene of the myosin-binding protein C have incomplete penetrance with mild hypertrophy and a generally favorable clinical evolution. Many carriers of the genotype do not exteriorize the phenotype, revealing normal thickness on echocardiography and, frequently, absence of electrocardiographic alterations. Only 58% of the affected members express some degree of hypertrophy prior to reaching 50 years of age. A tendency exists for penetration to remain incomplete until the 5th or 6th decades of life. Phenotype expression although delayed is usually uniform.

Mutations of this gene go against the conception that carriers of the disease should develop clinical manifestations of it prior to the beginning of adulthood. Late manifestation may render diagnosis of familial forms and identification of a autosomal dominant inheritance difficult. Despite the tendency towards benign evolution with a more favorable survival curve in comparison to mutations of other genes, increased mortality occurs conditioned by age and the development of hypertrophy. The greatest portion of deaths related to cardiopathy have a sudden character. From the moment when it becomes clinically manifest, it does not have a truly benign character. In a recent evaluation, prognosis was more favorable than in mutations of...
Mutations of the genes of the essential and regulatory myosin light chains - Mutations of the genes that encode, respectively, the essential and regulatory myosin light chains were recognized in 1996 as being associated with hypertrophic cardiomyopathy. Myosin is a hexameric protein, formed by 2 heavy chains and 2 pairs of light chains, designated as essential and regulatory. The essential myosin light chain is a polypeptide with 195 amino acids expressed in the myocardium and skeletal muscle. It has inotropic properties, probably related to the activation of the thin filaments. When removed from skeletal muscle in vitro, it reduces the speed of movement of actin and the generation of isometric force. The regulatory myosin light chain, composed of 166 amino acids, is expressed in the same structures. Rupture of the gene results in sarcomere disorganization and gives rise to the embryonic form of dilated cardiomyopathy. The light chains could have the function of stabilizing the structure that bears the globular head of the protein in cardiac β myosin heavy chain molecule.

Considering that mutations of the cardiac β myosin heavy chain affect the interface with the light chains, Poetter et al proposed the hypothesis that this protein when mutated could cause the disease. To this effect, the DNA of 383 unrelated families affected by hypertrophic cardiomyopathy was screened. The amplification of 7 exons that make up the gene of the light chain of essential myosin located in chromosome 3p revealed 2 point mutations with amino acid substitution. Six of 13 affected members with mutation Met149Val had the same phenotype, characterized by hypertrophy of papillary muscles and adjacent myocardium, generating medial ventricular obstruction. This less frequent form had not been previously demonstrated to have familial character. This mutation demonstrated high penetrance and a poor prognosis. Screening of another 16 unrelated individuals identified mutation Arg154His associated with the same phenotype.

Distinct mutations involving the gene of the regulatory myosin light chain have been simultaneously described by the same research group. The gene mapped in chromosome 12q is distributed in 12 KB of DNA and covers 7 exons. Three mutations with amino acid exchange have been documented: Ala13Thr, Glu22 Lys, and Pro94Arg, equally associated with mid-ventricular obstructive forms. The expression of these mutations was shown to be variable and of decreasing penetrance. Two other mutations associated with typical familial forms have subsequently been described.

Mutations of the light chains of respectively essential and regulatory myosin, determine variable phenotypes between families and between members of the same family. Mutations of both genes are associated with myopathic alterations, documented on skeletal muscle biopsy.

Mutations of the cardiac troponin I gene - In 1997, Kimura et al described the 7th gene responsible for hypertrophic cardiomyopathy, encoding cardiac troponin I. Localized on chromosome 19p, the gene is constituted of 6.2 KB DNA arranged in 8 exons. Cardiac troponin I is a polypeptide formed by 210 amino acids, integrating the troponin complex of the thin filaments. It exerts a modulator influence on the calcium-dependent actin-myosin interaction. The region of the molecule containing the site binding to actin is essential for the protein to exert inhibitory action on contraction. The C terminal, in turn, effects the binding to troponin C.

Six mutations have been originally described in this gene, 5 missense mutations and one deletion without disruption of the reading frame. Recently, deletion on exon 8 producing loss of 8 amino acids and of the stop codon was described for the first time in a Western family. Because the protein has its expression restricted to the myocardium, the effect of the mutations is limited to the heart. The mechanisms of action of these mutations are still hypothetical. They reach exons 7 and 8, corresponding to terminal C of the molecule. In this way, they become capable of interfering with the inhibitory function exerted by the protein and condition the loss of regulation of the contractile function, leading to hypercontractibility. Therefore cardiac troponin I mutations may act by mechanisms distinct from those of the other mutations.

Mutations of troponin I genes have a heterogeneous character. Although some are associated with typical forms of hypertrophic cardiomyopathy, others like Arg162Trp and Gly203Ser determine apical hypertrophy. Occasional association with Wolff-Parkinson-White syndrome has been noted. It is possible to individualize cases of typical forms and also of apical hypertrophy in the same family among carriers of the same mutation. Apical hypertrophic cardiomyopathy, predominating in the Orient, was preliminarily considered as sporadic in nature. Its identification...
among carriers of mutations of cardiac troponin I and myosin binding protein C indicated that the apical form is part of the wide clinical spectrum of hypertrophic cardiomyopathy as a sarcomere disease.

A report exists of spontaneous mutation with substitution of lysine for glutamine in codon 206 \(^{37}\). In the typical forms described in the West, hypertrophy has a mild or moderate degree with septal thickness of 15 or 16mm, or evidence of a subclinical character with expression restricted to the electrocardiogram \(^{35}\).

**Mutation of the cardiac \(\alpha\) actin gene** - The gene encoding \(\alpha\) cardiac actin was identified in 1999 by Mogensen et al \(^{38}\) as related to the disease in a family with hypertrophic cardiomyopathy. The major protein of the thin filaments, actin is directly involved in the generation of the contractile force of the sarcomere as well as in its transmission to the cell medium. The point mutation of the gene, mapped on chromosome 15, consists of the substitution of guanine for thymine in exon 5 with exchange of alanine by serine in codon 295. The mutation is located on the surface of the protein molecule close to the site of binding to \(\beta\) myosin. It is possible that it causes distortion of the topology of the polypeptide, harming the interaction with the thick filaments and the generation of contractile force in the sarcomere. The mutation related to hypertrophic cardiomyopathy is located close to 2 other point mutations recently associated with familial forms of dilated cardiomyopathy. The cardiac active gene is the first one to be simultaneously related to 2 distinct forms of cardiomyopathy \(^{38}\).

Cardiomyopathy produced by this gene has high penetrance with a predominance of asymptomatic forms and beginning in variable age groups. The phenotype is heterogeneous. Records exist of massive septal hypertrophy, late evolution towards serious systolic dysfunction, and an association with the Wolff-Parkinson-White syndrome in the presence of borderline left ventricular hypertrophy. Many carriers of the genotype exhibit only prominent electrocardiographic alterations at the basal parts of the septum \(^{38}\).

**The development of myocardial hypertrophy** – The mechanisms by which sarcomere protein mutations determine myocardial hypertrophy are still of a speculative character. The major part of information about the physiopathology of mutations is based on transgenic experimental models and in vitro analyses. At the molecular level, mutations determine structural and functional alterations in the sarcomere, which impair the generation of contractile force \(^{28,42,56,59}\). Reduction in contractility, however, does not have a reflex on the global performance of the left ventricle, which characteristically is hyperdynamic or normal. Myocardial hypertrophy, of adaptive and compensatory character, would develop by a series of as yet unknown processes, presumably common to all mutations \(^{42,70}\).

It is acknowledged that hypertrophy distribution depends more on hemodynamic factors than on the regio-

**Genetic diagnosis** – Genotyping becomes the rational basis of genetic counseling and the identification of patients with a high risk of sudden death, requiring precocious therapeutic intervention. Genetic diagnosis among children and adolescents is decisive in cases of reduced or absent penetrance, or ambiguous situations like those of athletes or persons with systemic arterial hypertension and left ventricular hypertrophy. In affected families, it enables the release of normal members and the precocious identification of malignant mutations. However, genetic analysis is restricted to a few research centers having as their principal objective the identification of new genes and mutations. Large-scale application of information arising from genetic diagnosis in clinical and ethical contexts is still a complex process in view of the risk of doubtful interpretations and unjustified discrimination.
Genetic studies have made a decisive contribution by enabling the characterization of hypertrophic cardiomyopathy as a sarcomere disease and identification of the predominance of familial forms. The spectral character of the disease becomes evident from the corroboration that in affected families 20% to 30% of the carriers of the genotype do not meet classical echocardiographic criteria of left ventricular hypertrophy, even though being susceptible to sudden death. In this group, the record of minimal alterations on the electrocardiogram and echocardiogram have greater significance than in the population in general. One verifies that prognosis is determined by the genetic substrate and that phenotype and clinical evolution vary according to the gene and type of mutation (Table I).

The expectation for the coming years is that remaining genes and their respective mutations will be identified. The evaluation of the effect of different mutations on the structure and function of the sarcomere is equally decisive, depending above all on the expansion of experimental studies based on transgenic animals. It is of fundamental importance to expand the analysis of genotype-phenotype relationships, still limited by the small number of families evaluated and by allele heterogeneity. The identification of modifying genes and of the mechanisms implicated in the establishment of myocardial hypertrophy should be effected, probably over the long range. Elucidation of these processes is essential for the introduction in the future of genetic therapy as a decisive form of intervention in the development of hypertrophic cardiomyopathy.

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<th>Chart I – Clinical characteristics of hypertrophic cardiomyopathy in its respective genetic forms</th>
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<tr>
<td><strong>Gene</strong></td>
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<td>β cardiac myosin heavy chain</td>
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<tr>
<td>Myosin-binding protein C</td>
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<td>Essential and regulatory myosin light chain</td>
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<td>Cardiac I troponin</td>
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<td>α cardiac actin</td>
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