

ARTICLE

Combined approach for finding susceptibility genes in DISH/
chondrocalcinosis families: whole-genome-wide linkage and
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Twelve families with exuberant and early-onset calcium pyrophosphate dehydrate chondrocalcinosis (CC) and diffuse idiopathic skeletal hyperostosis (DISH), hereafter designated DISH/CC, were identified in Terceira Island, the Azores, Portugal. Ninety-two (92) individuals from these families were selected for whole-genome-wide linkage analysis. An identity-by-descent (IBD) analysis was performed in 10 individuals from 5 of the investigated pedigrees. The chromosome area with the maximal logarithm of the odds score (1.32; $P=0.007$) was not identified using the IBD/identity-by-state (IBS) analysis; therefore, it was not investigated further. From the IBD/IBS analysis, two candidate genes, *LEMD3* and *RSPO4*, were identified and sequenced. Nine genetic variants were identified in the *RSPO4* gene; one regulatory variant (rs146447064) was significantly more frequent in control individuals than in DISH/CC patients ($P=0.03$). Four variants were identified in *LEMD3*, and the rs201930700 variant was further investigated using segregation analysis. None of the genetic variants in *RSPO4* or *LEMD3* segregated within the studied families. Therefore, although a major genetic effect was shown to determine DISH/CC occurrence within these families, the specific genetic variants involved were not identified.

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INTRODUCTION

Diffuse idiopathic skeletal hyperostosis (DISH, MIM 106400) and chondrocalcinosis (CC, MIM 118600) are diseases characterized by ectopic calcification. DISH is characterized by the ossification of entheses in the axial and peripheral skeleton that affect the anterior spinous longitudinal ligament, in particular the right side of the spine, with preservation of the intervertebral disc space.¹ CC is characterized by the deposition of calcium-containing crystals in articular cartilage, synovial membranes and, less often, in periarticular soft tissues.^{2,3} *ANKH* mutations are the only known cause of a very small number of cases of monogenic CC (MIM 118600)^{4–7} as well as craniometaphyseal dysplasia (MIM 123000).^{8–10} The *ANKH* gene maps to chromosome 5p15.1 and encodes the multipass transmembrane protein ANK that transports intracellular inorganic pyrophosphate to the extracellular milieu,¹¹ where it acts as a potent inhibitor of mineralization.¹² The etiology of DISH is still unknown, but several lines of evidence suggest that genetic factors might be involved in its etiology.^{13–15} Very few genetic studies on DISH have been published, and until now, only *COL6A1*^{16,17} and *FGF2*¹⁸ have been shown to have a positive association with DISH susceptibility. Moreover, all the variants in both genes that showed significant associations were located in non-coding regions and are very common variants within the general population, suggesting that these variants have a minor effect on DISH susceptibility. DISH can coexist with ossification of the posterior longitudinal ligament (MIM 602475), a disease in which

the genetic background is considered relevant to its etiology. Unlike DISH, ossification of the posterior longitudinal ligament has been extensively investigated, and despite some conflicting studies, a great number of susceptibility genes have been reported. These genes include collagen 6A1 and 11A2 (*COL6A1* and *COL11A2*, respectively),^{17,19} bone morphogenetic protein 2 and 4 (*BMP-2* and *BMP-4*, respectively),^{20,21} ectonucleotide pyrophosphatase/phosphodiesterase 1 (*ENPP1*),²² transforming growth factor 1 and 3 receptor (*TGFβ1* and *TGFβ3*, respectively),^{23,24} estrogen receptor 1 (*ESR1*)²⁴ and R-spondin 2 (*RSPO2*)^{25,26} among others. However, as occurs in DISH, the genetic variants with a positive association with ossification of the posterior longitudinal ligament appear to have a minor effect on disease susceptibility, suggesting that the heritable component of its etiology is likely to be polygenic in most cases.

Genetic and clinical links exist between CC and DISH, and there is evidence that disordered pyrophosphate metabolism may have an important role; in general, conditions that favor increases in inorganic pyrophosphate promote calcium pyrophosphate dehydrate crystal formation.²⁷ There is evidence of genetic involvement in CC and forms of spinal ossification in both human⁵ and animal models; the *ank* mouse develops severe hydroxyapatite CC and spinal ossification,¹¹ and the tip-toe walking (*ttw*) mouse is the main model for human ossification of the posterior longitudinal ligament and displays cartilage calcification.²⁸

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The coexistence of DISH with CC is very common in Terceira Island—Azores, suggesting that both diseases, hereafter designated the DISH/CC phenotype, share the same pathogenic mechanism.²⁹ A similar phenotype has also been reported in several previous studies.^{30,31} Our main objective was to elucidate the genetic factors underlying this condition and to establish whether this disease is a novel form of pyrophosphate arthropathy or a simple association between two different disorders with extremely high prevalence on this island in the Azores. A formal segregation study in these families suggested that the condition was caused by a Mendelian inherited disorder with an autosomal-dominant model of transmission.³² Therefore, we performed a whole-genome linkage analysis followed by ‘identity-by-state (IBS)/identity-by-descent (IBD)’ mapping to determine the genetic cause of this phenotype. IBD mapping is a statistical method for the detection of genetic loci that share an ancestral segment among ‘unrelated’ affected pairs of individuals. IBD mapping is a more robust approach than allelic heterogeneity and can be used as a complementary method to genome-wide linkage studies to identify rare inherited variants when combined with sequencing data.³³

MATERIALS AND METHODS

Collections used

Twelve pedigrees with 92 individuals were used for the linkage analysis.

Ten affected individuals from five pedigrees previously used for linkage analysis were selected for IBD/IBS analysis. The radiological characterization of these individuals is presented in Table 1.

This study was approved by the HSEIT Ethics Committee and all participants provided informed consent. Peripheral blood was collected for biochemical analysis and DNA isolation.

Microsatellite genotyping

Highly informative microsatellite markers from the Linkage Mapping Set V2 (Applied Biosystems, Foster City, CA, USA) were used for the linkage analysis. Microsatellites were amplified in 20 µl PCR reactions using optimized reaction conditions. To simplify the genotyping process, PCR products were pooled based on the fluorochrome used, the allele-size range and avoiding fluorochromes with overlapping spectra or overlapping alleles. The 96-well plates were prepared by adding a mixture of 10 µl of Hi-Di formamide (Applied Biosystems) and 0.3 µl of the internal line standard GeneScan 400HD-ROX (Applied Biosystems) to each well. Samples were denatured and immediately run on an ABI automated DNA genetic analyzer 3700 and 3130XL. Microsatellite genotype calling was performed semiautomatically using the GENOTYPER software (Applied Biosystems) and the GeneMapper Software Version 4.0 (Applied Biosystems).

Quality control and linkage analysis

To minimize data errors, extensive checking procedures were undertaken. Genetic Analysis System (GAS version 2.0; Alan Young, Oxford University, Oxford, UK) was used to assign discrete allele numbers to the alleles. Mendelian inheritance was checked with GENOTYPER and GeneMapper (Applied Biosystems). PEDCHECK³⁴ was used to screen all the data for previously undetected Mendelian inconsistencies. Statistical analysis and parametric and non-parametric analyses were performed using VITESSE,³⁵ GENEHUNTER and MERLIN, respectively.³⁶ Marker positions were obtained from the National Center for Biotechnology Information database (NCBI). Parametric linkage analysis was performed assuming autosomal-dominant transmission, a penetrance of 90–95%, phenocopy rates of 0.05–0.1% and minor allele frequencies of 0.1–1%.

IBD/IBS analysis

Genotyping was performed with Illumina (San Diego, CA, USA) 370CNV BeadChips. To perform the analysis, 50,000 markers with the following characteristics were chosen: (a) not in a linkage disequilibrium with each other ($r^2 < 0.1$) and (b) had a minor allele frequency > 0.3 . These were chosen using the TASCOC control set (approximately 400 individuals with predominantly UK ancestries from Tasmania, Australia).³⁷ IBD segments were then inferred for all pairs of cases across the genome using PLINK³⁸ with the ‘Spairs’ statistic, which represents the number of pairs of cases that appear to have IBD inherited chromosomal segments at each point on the genome.

Using the fastIBD option in the software package BEAGLE (2),³⁹ it was possible to identify IBD regions across families while also taking into account the linkage disequilibrium. The analysis was set for a minimum IBD segment length of 1 cM and an IBD detection threshold of 10^{-10} . The genome-wide P -value was estimated knowing that there were 366 segments with an average of 3.1 case pairs shared across the genome. The number of case pairs shared across the genome approximately followed a Poisson distribution. As the 366 segments were not independent across the genome, principal component analysis was used to show that the 366 multiple tests performed were approximately equivalent to 202 independent tests.⁴⁰ The genome wide P -value was obtained by estimating the probability of obtaining a maximum number of case pairs shared out of 202 draws from the Poisson distribution ($\lambda = 3.1$) by chance. As a maximum of 10 cases were observed to share IBD, this probability was estimated to be 0.077 by assuming that the number of case pairs shared had a Poisson distribution with a mean of 3.1.

Sanger sequencing

RSPO4 gene primer pairs were designed using the Primer3 software (Open source developed and maintained by Whitehead Institute for Biomedical Research, Cambridge, MA, USA). Regulatory coding regions, including intron–exon boundaries, were amplified and sequenced in 55 patients with DISH/CC disease and 36 unaffected control subjects. The *LEMD3* gene was sequenced using the primers previously described by Hellemans et al.⁴¹ The amplification conditions for all the primers are available upon request. PCR fragments were purified with ExoSAP-IT (Exonuclease I and Shrimp Alkaline Phosphatase) and sequenced using ABI Big Dye chemistry

Table 1. Radiological characterization of the patients used for the IBD/IBS analysis

Code	Sex	Family bond	Axial	Elbow	Knee
1-1.1	M	Father and daughter	DISH	Periarticular calcification	Calcification
1-1.3	F		OALL	Enthesopathy	Normal
2a-2.32	M	Brothers	Normal	Enthesopathy	Normal
2a-2.34	M		Normal	Osteophytosis, enthesopathy	Chondrocalcinosis
2b-2.35	F	First-degree cousins	DISH	Capsular Calcification	Osteophytosis; arthrosis
2b-2.39	M		DISH	Capsular calcification; osteophytosis	Chondrocalcinosis
7-7.3	M	Brothers	OALL	Enthesopathy; capsular calcification	NA
7-7.4	F		OALL	Enthesopathy; osteophytosis; capsular calcification	Enthesopathy; osteophytosis; arthrosis
11-11.4	M	Brothers	DISH	Periarticular calcification	Enthesopathy; arthrosis
11-11.5	M		DISH	Periarticular calcification	Normal (with Pseudogout episodes)

Abbreviations: DISH, diffuse Idiopathic skeletal hyperostosis; enthesopathy, new bone formation at enthuses; NA, not available; OALL, ossification of the anterior longitudinal ligament; osteophytosis, outgrowths of bone tissue.

(unidirectional or, when necessary, bidirectional) followed by purification with EDTA/sodium acetate and ethanol precipitation. Sequencing products were run on an automated DNA sequencer (ABI 3130XL, Applied Biosystems), and genetic variants were screened by sequencing analysis with SeqScape (Applied Biosystems). Base calling for heterozygous positions occurred when the smaller peak of two co-incident peaks was >25% of the maximum peak.

Whenever possible, the genetic variants found in the *RSPO4* gene were screened across all family members when an affected individual was identified. The *LEMD3* rs201930700 variant was typed in a representative randomized group of 124 individuals from Terceira Island.

Statistical analysis

The Hardy–Weinberg equilibrium and all other statistical analyses were performed using the PLINK software.³⁸ Association was tested between the DISH/CC disease and allelic variants with the Cochran–Armitage trending test, dominant and recessive gene action tests with 1 degree of freedom and a genotypic test with two degrees of freedom. The Fisher's exact test was used to assess the differences in the allele frequencies between the 55 patients with DISH/CC and the 36 control individuals. For all statistical tests, a *P*-value ≤ 0.05 was considered statistically significant.

RESULTS

Linkage analysis

A whole-genome linkage scan, including 92 individuals from 12 pedigrees, was performed on all autosomes, with 189 microsatellite markers and an average distance between markers of 17 cM. Parametric and non-parametric analyses were also performed. The best results were obtained with a non-parametric linkage analysis for chromosome 16 (single point 2.38 and multipoint 1.16). Because the average distance between markers was >10 cM, one extra set of 137 markers, which was distributed on all autosomes, was further amplified and reanalyzed to enlarge the power of the study. A total of 327 microsatellites were included in the analysis. Owing to its complexity, pedigree 2 was excluded from the analysis because of computation time and memory constraints. When assuming an autosomal-dominant mode of inheritance with a minor allele frequency of 0.1%, phenocopy rate of 0.1% and penetrance of 90%, the parametric analysis did not give significant results.

Because there was only a nominally suggestive linkage to an area on chromosome 16, a new set of 13 microsatellite markers that covered this entire chromosome with an average spacing of 8.99 cM was further amplified and gave more significant results (Table 2).

The most significant result on chromosome 16 was obtained for the D16S3100 marker; it had a logarithm of the odds score of 1.32 and a *P*-value of 0.007. The chromosomal area (16p12.1) directly surrounding marker D16S3100 in the 26.020–27.120 K region had only 12 annotated genes, among which 6 were uncharacterized, 3 were pseudogenes and 1 microRNA and 1 open reading frame were also identified. According to the present knowledge, the only gene of putative interest in this region was *HS3ST4*, which encodes the enzyme heparin sulfate (glucosamine) 3-O-sulfotransferase. This region was not further investigated owing to a lack of replication using IBD analysis.

IBD sharing

There were a total of 10 cases corresponding to 45 pairs of cases in the IBD analysis. The greatest number of case pairs shared was 10, which was on chromosome 12. The 10 pairs were shared by 3 families. The significance of the association with chromosome 12 included a single-point *P*-value = 0.0014 and a genome-wide *P*-value = 0.077.

When using BEAGLE on the CC data set with a minimum IBD segment length of 1 cM and an IBD detection threshold of 10^{-10} , regions on chromosomes 12 and 20 revealed the sharing of 10 pairs (Table 3). The IBD analysis did not replicate the weak suggestive linkage to an area surrounding the D16S3100 marker on chromosome 16.

The chromosome regions considered to be of interest were selected considering the maximum number of pairs shared. Two zones, one in chromosome 12 and one in chromosome 20, had a maximum of 10 pairs shared and were therefore further

Table 3. Chromosomal regions shared between pairs identified using fastIBD and PLINK

Chromosome	Start position (hg18)	End position	Pairs sharing fastIBD	Pairs sharing PLINK
4	7403378	8100956	9	3
5	138687057	141055614	9	3
12	65667554	66378203	9	9
12	66378203	68670915	10	10
13	25442369	25501361	9	6
20	821749	1266214	10	7
20	5157217	6074302	9	8

Table 2. Results from a single-point analysis (MERLIN) for 11 pedigrees on chromosome 16

Chromosome 16	Location	Z-mean	P-value	Delta	LOD	P-value
Markers	cM	Max = -3.39	Max = 0.00	Max = 0.696	Max = 5.62	Max = 0.00
D16S521	1.08	0.33	0.4	0.117	0.03	0.4
D16S3027	8.71	0.43	0.3	0.150	0.05	0.3
D16S423	14.05	0.42	0.3	0.158	0.05	0.3
D16S418	14.77	0.49	0.3	0.184	0.06	0.3
D16S3075	23.28	0.48	0.3	0.286	0.10	0.2
D16S3046	40.65	1.09	0.14	0.539	0.45	0.08
D16S3100	52.26	1.83	0.03	0.696	1.32	0.007
D16S415	67.62	1.38	0.08	0.696	0.86	0.02
D16S3057	77.13	0.33	0.4	0.271	0.06	0.3
D16S514	81.15	0.10	0.5	0.072	0.00	0.4
D16S515	92.06	0.80	0.2	0.471	0.26	0.14
D16S516	95.45	0.36	0.4	0.229	0.06	0.3
D16S505	108.96	0.52	0.3	0.557	0.20	0.2

Abbreviation: LOD, logarithm of the odds.

investigated. In contrast with previous reports that identified *ANKH* as a CC-associated gene, it was not identified on chromosome 5 in our study.

A total of 167 previously characterized genes were identified in the regions of interest on chromosomes 12 and 20. There were 52 genes in the region on chromosome 12 (65667554–68670915), of which only 35 were annotated. Just next to this region in chromosome 12 (65169571–652483279), one gene that had already been associated with disordered calcification, *LEMD3*, was identified. This gene encodes a LEM domain-containing protein. The encoded protein functions to antagonize TGF- β and BMP signaling at the inner nuclear membrane. Mutations in this gene have been associated with osteopoikilosis (MIM 166 700), Buschke–Ollendorff syndrome (MIM 166700) and melorheostosis (MIM 155950).⁴¹

There were 148 genes in chromosome 20 (chromosome region from 821749 to 6074302), of which 115 were annotated. The best candidate gene in chromosome 20 was *RSPO4* (chromosome 20:958452–1002284). The *RSPO4* gene encodes a member of the R-spondin family of proteins that share a common domain organization consisting of a signal peptide, cysteine-rich/furin-like domain, thrombospondin domain and a C-terminal basic region. The encoded protein may be involved in the activation of the Wnt/ β -catenin signaling pathways.⁴² WNT signaling is critical for nail development and mutations in different WNT-associated genes have been identified in disorders involving the nails. *RSPO4* mutations have been particularly associated with autosomal-recessive congenital anonychia (NDNC4; MIM 206800).^{43,44}

Sequencing results

RSPO4 sequencing. Nine genetic variants were identified in the *RSPO4* gene, including three missense variants (rs6140807 and rs61740632, rs201485021), one splice site variant (rs775644973), two synonymous (rs150446609 and rs41275604) and three regulatory region variants (rs146447064, rs149154047 and rs6056520) (Table 4). All of them were in Hardy–Weinberg equilibrium.

Two variants in the regulatory region of the *RSPO4* gene (rs146447064 and rs149154047) were located in a fully conserved region and are rare (Table 4). The same was observed for the synonymous variant rs150446609, which was very rare (< 0.01) and was located in a fully conserved chromosome region. The missense variant rs201485021 located in exon 3 has a SIFT score of 0 and a PolyPhen value of 1, indicating that it has deleterious and damaging effects on the protein, respectively. Additionally, this variant was located in a fully conserved region and was extremely rare (minor allele frequency < 0.01); in the ‘1000 genomes’ (genomes from 26 different populations), the variant was identified in only 2 males from an Iberian population in Spain. In our study, this variant was found in one female in our control group ($n=36$). The other two missense variants (rs6140807 and rs61740632) had SIFT and PolyPhen values indicative of minor effects on the protein (tolerated and benign, respectively) and were both relatively rare. We found an extremely rare HGMD mutation (rs775644973 or CS065613) that has previously been associated with congenital anonychia.⁴⁴ The variant in heterozygosity was found in 1 asymptomatic female in our group of 55 DISH/CC patients.

Table 4. Genetic variants identified in the *RSPO4* gene and functional significance information

Exon/intron	Variant	SNP_ID	Type of variant	MAF	SIFT	PolyPhen	Nucleotide conservation
Upstream gene	c.-131C>T	rs146447064	Regulatory reg.	0.01	NA	NA	Fully conserved
Upstream gene	c.-115G>A	rs149154047	Regulatory reg.	0.01	NA	NA	Fully conserved
5'	c.-85G>A	rs6056520	Regulatory reg.	0.35	NA	NA	1 non-conserved species (mouse lemur)
Exon 1	c.12A>C	rs150446609	Synonymous p.Pro4=	< 0.01	NA	NA	Fully conserved
Intron 1	c.79+1G>A	rs775644973	Splice donor (HGMD mutation)	< 0.01	NA	NA	Fully conserved
Exon 3	c.317G>A	rs6140807	Missense p.Arg106Gln	0.03	0.21	0.072	2 non-conserved species (pig and hedgehog)
	c.367C>G	rs201485021	Missense p.Pro123Ala	< 0.01	0	1	Fully conserved
Exon 4	c.471C>T	rs41275604	Synonymous p. Cys157=	0.01	NA	NA	1 non-conserved species (hedgehog)
Exon 4	c.524A>T	rs61740632	Missense p.His175Pro	0.01	0.3	0	4 non-conserved species (mouse lemur, Pig, hedgehog and elephant)

Abbreviations: MAF, minor allele frequency; NA, not available; SNP, single-nucleotide polymorphism. Nucleotide conservation was obtained using alignments of the human, chimpanzee, mouse lemur, pig, hedgehog and elephant genes available in the Ensembl database (<http://www.ensembl.org/index.html>) (accessed on January 2017).

Table 5. Genetic variants identified in the *LEMD3* gene in four probands that have been previously investigated and information about their functional significance

Exon/intron	Variant	SNP_ID	Type of variant	MAF	SIFT	PolyPhen	Nucleotide conservation
Intron 1	c.1523-12C>T	rs11175678	Intronic	0.04	NA	NA	Non-conserved
Intron 5	c.1695+100G>A	rs11610822	Intronic	0.25	NA	NA	1 non-conserved species (mouse lemur)
Intron 7	c.2024-34_2024-31delGATT	rs10534559	Intronic	?	NA	NA	Non-conserved
Exon 13	c.2701C>T	rs201930700	Missense	< 0.01	0	0.995	Fully conserved

Abbreviations: MAF, minor allele frequency; NA, not available; SNP, single-nucleotide polymorphism.

LEMD3 sequencing. *LEMD3* was sequenced in the four probands from the families who shared the maximum number of pairs in the IBD/IBS analysis. The identified genetic variants can be observed in Table 5.

Variant **rs201930700** was located in exon 13 of the *LEMD3* gene and had a SIFT score of 0 and a PolyPhen value of 0.995, which indicated that this variant has deleterious and damaging effects on the protein, respectively. The variant was extremely rare. The minor allele frequency was unknown because there was insufficient data to establish population frequency; it was identified in only 5 of the 121,412 alleles (ExAc_Aggregated_Populations), indicating that it was unquestionably rare (minor allele frequency 0.00004). A cohort of 124 individuals, which is representative of the Terceira Island population, was typed for this mutation, and 1 other individual carrying the variant was identified. This individual and their relatives were examined (interviewed for clinical purposes, X-rayed and typed).

The rs201930700 variant was in Hardy–Weinberg equilibrium. Of the five individuals studied in the AZ3 family, four individuals (two males and two females) were affected by DISH/CC and were carriers of the rs201930700 variant and one individual (male) was affected by DISH/CC but was a non-carrier of the variant. In the AZ3 family, it was impossible to verify segregation as all the individuals studied were affected by DISH/CC affected; thus an association test was not performed.

Segregation analysis. Segregation analysis involved typing eight genetic variants in the seven DISH/CC pedigrees where the variants were found, including the seven variants in *RSPO4* (rs146447064, rs149154047, rs6056520, rs150446609, rs6140807, rs41275604 and rs61740632) and variant rs201930700 in *LEMD3*.

Most of the analyzed families were uninformative because of the large number of affected versus unaffected individuals. Variants did not segregate within the investigated informative families (see Supplementary Figure 1 and Supplementary Table 1 for more information).

Association study. The *RSPO4* regulatory region variant **rs146447064** was significantly ($P=0.03$) more frequent in the controls (14%) than in the DISH/CC patients (5%), and when adjusted for gender, it was significant in females ($P=0.0333$) but not in males (Table 6).

Similar results were obtained using the Cochran–Armitage trend and allelic tests (see Supplementary Table 2). The other *RSPO4* regulatory region variant, rs149154047, was also significantly different in frequency between the DISH/CC males (3%) and control males (14%) ($P=0.0502$) but not in females. Similar results were obtained using the Cochran–Armitage trend test ($P=0.02453$, $\text{CHISQ}=5.057$, $\text{Df}=1$) and the allelic test ($P=0.02958$, $\text{CHISQ}=4.734$, $\text{Df}=1$) (see Supplementary Table 2).

DISCUSSION

DISH/CC is a poorly understood phenotype characterized by peripheral and axial enthesopathic calcifications, which fulfill the radiological criteria for DISH and are associated with calcium pyrophosphate dehydrate CC in some cases.^{31,45,46} The variable clinical and radiological presentation of patients with DISH/CC in the families investigated raises the possibility of genetic heterogeneity. The whole-genome linkage analysis found no chromosomal region with clear linkage in the affected families. The following reasons can be postulated to explain these results: (1) the phenotypic heterogeneity observed within and among pedigrees may result from genetic heterogeneity; (2) unknown environmental factors may be affecting the phenotypes and causing difficulties in the analysis; and (3) the condition may not be a simple Mendelian disorder caused by a large effect rare variant but instead may have a more complex and polygenic genetic architecture. The region on chromosome 16 that showed weak suggestive linkage was not replicated by the IBD/IBS study and was not further investigated.

The concurrence of DISH and CC suggests a shared pathogenic mechanism.³² The etiology of DISH/CC is unknown. However, because it is a bone-forming disease, it is expected that genes related to the calcification and ossification process are implicated in its etiology. In the second strategy utilized, the IBS/IBD analysis, two chromosomal areas were identified and further investigations performed. Two genes, *RSPO4* and *LEMD3*, were selected as good candidates based on their functions. Two variants in the *RSPO4* gene (rs146447064 and rs14915407), which were significantly more frequent in the controls than in DISH/CC patients, may have a possible protective role in the DISH/CC phenotype, though the level of significance observed was not definitive.

As already mentioned, the *RSPO4* protein is known to have a major role in activating the Wnt/ β -catenin signaling pathways.⁴² It

Table 6. Fisher's exact test for genetic variants found in the *RSPO4* gene in Azorean patients with DISH/CC and controls without DISH/CC disease

SNP	Alleles	MAF						Fisher's exact test									
		DISH/CC			Controls			DISH/CC vs controls									
		All, N = 55	Female, N = 19	Male, N = 36	All, N = 36	Female, N = 22	Male, N = 14	All	Females	Males	OR	P	OR	P	OR	P	
rs146447064	C/T	0.045	0.026	0.056	0.139	0.182	0.071	0.30	0.03	0.12	0.03	0.76	0.67				
rs149154047	G/A	0.045	0.079	0.028	0.097	0.068	0.143	0.44	0.22	1.17	1	0.17	0.05				
rs6056520	C/T	0.282	0.184	0.333	0.194	0.182	0.214	1.63	0.22	1.02	1	1.83	0.33				
rs150446609	A/C	0.045	0.053	0.042	0.014	0.023	0	3.38	0.41	2.39	0.59	NA	0.56				
rs775644973	G/A	0.009	0.026	0	0	0	0	NA	1	NA	0.46	NA	1				
rs6140807	G/A	0.182	0	0.028	0	0	0	NA	0.52	NA	1	NA	1				
rs201485021	C/G	0	0	0	0.014	0.023	0	0	0.40	0	1	NA	1				
rs41275604	C/T	0.036	0.053	0.028	0.028	0.045	0	1.32	1	1.17	1	NA	1				
rs61740632	A/C	0.036	0	0.056	0.014	0.023	0	2.68	0.65	0	1	NA	0.57				

Abbreviations: CC, chondrocalcinosis; DISH, diffuse idiopathic skeletal hyperostosis; MAF, minor allele frequency; NA, not applicable; OR, odds ratio; SNP, single-nucleotide polymorphism. The minor allele is represented in bold.

is known that induction of the Wnt signaling pathway promotes bone formation, while inactivation of this pathway leads to an osteopenia state.⁴⁷ Based on this knowledge, we hypothesized that rs146447064 and rs14915407 were variants associated with a reduction in *RSPO4* gene expression, thus reducing Wnt activation and consequently protecting against new bone formation. Obviously, further studies are needed to test this theory.

The *LEMD3* gene missense variant (rs201930700) was found in two apparently unrelated families. This variant has been identified very few times previously and both families in which it was typed presented the DISH/CC phenotype. The variant identified caused the substitution of amino acid 901 from a large and basic arginine to a large and aromatic tryptophan. Phylogenetically, according to the Ensembl database, the modified nucleotide was highly conserved in all vertebrates and was located in the carboxy-terminal nucleoplasmic region of the MAN1 protein. This region (amino acids 782–911) is predicted to be an RNA recognition motif-like (RRM-like) protein interaction domain called the U2AF homology motif.^{48,49} The conserved region that contains the UHM domain (U2AF homology motif kinase 1) is exclusive to MAN1 proteins and is essential for smad2 and smad3 binding.⁵⁰ It is unknown whether interactions between MAN1 and Smad1 or Smad2 and Smad3 inhibit BMP and TGF- β signaling, respectively.^{51,52} It has been reported that heterozygous loss-of-function mutations in *LEMD3* enhance TGF- β signaling and lead to sclerosing bone dysplasia, osteopoikilosis and Buschke–Ollendorff syndrome.⁴¹ As the identified variant (rs201930700) was deleterious but was not a loss-of-function mutation, the carriers of the variant did not present any signs of osteopoikilosis or Buschke–Ollendorff Syndrome. At this point, it was difficult to ascertain the effect produced on the DISH/CC phenotype by the rs201930700 variant. We postulate that the rs201930700 variant may promote enhanced TGF- β signaling, leading to increased bone formation. Our hypothesis was not confirmed by the segregation analysis, which might be due to the characteristics of the pedigrees used, as almost all the individuals were DISH/CC affected, making it very difficult to verify segregation. Other studies will be necessary to verify the impact of this rare variant on the phenotype to establish a possible association.

In conclusion, a combined strategy led us to investigate two candidate genes, *RSPO4* and *LEMD3*, and the results obtained suggested a small and protective role for the two *RSPO4* gene regulatory variants, probably by altering *RSPO4* gene expression. At this point, it is impossible to ascertain the relevance of the extremely rare variant in *LEMD3* (rs201930700) to the DISH/CC phenotype. Although the results are interesting, future studies are required to confirm and assess their relevance for DISH/CC pathologies.

COMPETING INTERESTS

The authors declare no conflict of interest.

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