

Generalised Osteoarthritis: from Mendelian Disorder to Complex Disease $\operatorname{Min,\ J.L.}$

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CHAPTER

5.2

Mutation analysis of candidate genes within the 2q33.3 linkage area for familial early onset generalised osteoarthritis

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Abstract

In a genome-wide linkage scan of seven families with familial early onset osteoarthritis (FOA), we mapped a FOA locus to a 5 cM region on chromosome 2q33.3-2q34 with a maximum LOD score of 6.05. To identify causal variants, 20 positional candidate genes were sequenced for coding, splicing, and 5' and 3' untranslated regions. The pathogenicity of possible disease causing variants was evaluated using predicted effects on protein structure and function, splicing enhancers, degree of conservation and frequency in 790 unrelated subjects from the population-based Rotterdam study scored for the presence of radiographic signs of OA (ROA). Nine novel variants, identified in NRP2, KIAA1571, ADAM23, IDH1, PIP5K3 and PTHR2, cosegregated with FOA, of which two were promising. IDH1 Y183C cosegregated in one family, involved a conserved amino acid change and showed a damaging effect predicted by PolyPhen and SIFT. In the Rotterdam sample, carriers of IDH1 Y183C (0.02) had an increased but insignificant risk for generalised ROA. The second variant, NRP2 c.1938-21T>C cosegregated in three families. In the Rotterdam sample, carriers conferred a significant increased risk of 2.1 (95% CI, 1.1-4.1, P = 0.032), to have generalised ROA. Furthermore, two variants (NRP2 c.1938-21T>C and IDH1 c.933-28C>T) occurred together on the haplotypes segregating with FOA in two out families. This haplotype was rare in the Rotterdam sample (0.0013). Two promising variants in or near NRP2 and IDH1, may be not sufficient for the onset of FOA alone but might have a modulating role with FOA. Confirmation in other OA populations is required.

Introduction

Osteoarthritis (OA) is the most common joint disease and is the leading cause of disability among the elderly. It is a degenerative disease of the joints characterised by cartilage degeneration and subchondral bone remodeling. Clinically, OA manifests itself with pain, stiffness, disability and loss of joint function. Twin, sibling and segregation studies have highlighted substantial heritability for primary OA ranging between 39%-80% depending on the sex and joint location^{1,2}. OA often arises as a complex trait, but rare familial forms with autosomal dominant transmission have been reported³⁻⁵. In familial forms of osteochondrodysplasia often displayed with secondary early onset OA, mutations in several structural genes of the extracellular cartilage matrix (ECM) including COL2A1, COL9A1, COMP have been identified^{6;7}. There is no convincing evidence yet that these genes are implicated in the susceptibility for primary early onset OA occurring without dysplasia⁷⁻⁹. Linkage and association studies on the basis of joint- or sex-specific OA definitions have yielded several loci associated with OA susceptibility including the interleukin-1 cluster (IL1A, IL1B, IL1RN)¹⁰⁻¹², ESR1^{13;14}, VDR^{15;16}, MATN3^{17;18}, FRZB^{19;20}, IL4R²¹, ASPN²² and CALM1²³.

Previously, we reported on a genome-wide linkage scan which revealed evidence for linkage at 2q33.3-2q34 with a two point LOD score of 6.05 (theta = 0.00) and a multipoint NPL score of 4.70 (P = 0.0013) in seven extended families with early onset generalised OA (FOA) without dysplasia²⁴. This region of 5 cM highlighted two strong candidate genes, PTHR2 and FZD5, which were unlikely to explain the observed linkage signal²⁴. None of the OA susceptibility genes mentioned are located in this linkage region. However, this interval is nearby the locus 2g32 identified in the UK study in which female siblings with hip replacement were investigated. In this study, a functional polymorphism in FRZB, a gene that antagonises the Wingless signalling, conferred susceptibility for hip OA in females²⁵. Our subsequent investigation of FRZB demonstrated a significant association with radiographic generalised OA in a random sample from the population-based Rotterdam study and with clinical generalised OA in affected sibling pairs from the GARP study²⁶. The high LOD score in the early onset OA families and absence of variants in the two most promising genes prompted us to perform an extended systemic mutation analysis of candidate genes in the 2q33.3-2q34 linkage area. In the present study, we report results of mutation screening of the entire coding region, splice sites, and 5' and 3' untranslated regions (UTR) of 19 positional candidate genes and the FRZB gene in affected family members with early onset generalised OA.

Materials and methods

Families

Families containing patients that express primary generalised OA in each generation were collected from different parts of the Netherlands. Informed consent was obtained from all patients and the medical ethics committee of the Leiden University Medical Centre approved the study. Probands were recognised through Rheumatology outpatient clinics. Family members were recruited via probands. Initially, we used questionnaires to select eligible families. For eligible families, complete medical history and available radiographs were obtained from Rheumatologists of almost all affected family members (81%). Radiographs were reevaluated for signs of chondrodysplasia, spinal dysplasia or abnormal development of the epiphyses of the peripheral joints. These features were absent in all families. The presence of radiographic OA (ROA) was assessed according to Kellgren/Lawrence criteria²⁷ by an experienced reader. Some individuals had marked Heberden's nodes and ankle OA. A selection of affected and unaffected individuals of family 1, 2, 4, and 7 were additionally visited for physical examination in order to prevent misclassification. The mean age of onset of OA in these patients was 33 years ranging between 20 and 50 years. The phenotype within these families is characterised by distinct progressive OA in the absence of mild or severe chondrodysplasia, however, with symptoms and ROA at multiple joint sites simultaneously including involvement of the hands with noduli, knees, hips, ankle and spine. Individuals with clinical and radiographic evidence of OA in two or more joint sites before the age of 50 years were considered affected. Extensive description of the phenotype in family 1, which is representative for the phenotypes also of the other extended families included, is described elsewhere ²⁸. All clinical diagnostic decisions were made independent to genetic linkage analysis and homogeneity of the phenotype between different families was checked.

The Rotterdam sample

The Rotterdam study, which comprises 7,983 Caucasian participants is a prospective, population-based cohort study of the determinants and prognosis of chronic diseases in the elderly²⁹. The medical ethics committee of the Erasmus University Medical Centre approved the study, and informed consent was obtained from all subjects. In a random sample of 809 unrelated subjects (the Rotterdam sample), ages 55-65 years, radiographs were scored for the presence of radiographic OA (ROA) in two knees, two hips³⁰, 36 hand joints and three levels of the thoracocolumbar spine^{30,31}. All radiographs were scored according to the Kellgren/Lawrence grading system (grades 0-4)²⁷ by two independent readers, blinded to all other data of the

participant. Definite ROA at a particular joint site was defined as a Kellgren/Lawrence score of two or more²⁷. In the hands, 36 separate joints were scored comprising eight joint groups: distal interphalangeal joints, the interphalangeal joint of the thumb, the proximal interphalangeal joints, the metacarpalphalangeal joints, the first carpometacarpal joints, the trapezoscaphoideal joints, the radionavicular joints and the distal radioulnar joints. By definition, ROA of the spine is confined to the apophyseal joints, but these joints could not be assessed on the lateral radiographs of the spine that were available. Instead, we assessed disc degeneration (DD) of the spine, at three levels i.e. thoracic (Th4 to Th12), lumbar (L1 to L4 or L5) and lumbosacral (L5-S1 or L5-L6).

We analysed and evaluated the occurrence and the generalised OA status of carriers of novel variants in the population-based sample as a qualitative trait using previously described definitions³². In brief, subjects with two or more of the following four criteria were considered as affected with generalised ROA: hand ROA in three or more hand joint groups (the right and left hands were considered separately), spinal DD in two or more disc levels, knee ROA in one or two knees and hip ROA in one or two hips. Subjects affected with generalised ROA were compared to the complete Rotterdam sample from which subjects with generalised ROA were excluded. Phenotypic data for assessment of generalised ROA was available for 790 subjects.

Mutation analysis

Genomic DNA was isolated from EDTA blood of affected and unaffected family members. Initially, three affected family members (individual 14 from family 1, individual 10 from family 2 and individual 9 from family 4) were screened for possible mutations by direct forward and reverse sequencing from both ends (Figure 1). If a novel variant was identified, unaffected family members of these families and family members of remaining families were sequenced. Reference sequences corresponding to all coding and 5' and 3' UTR regions of the genes were obtained from the UCSC genome browser assembly May 2004 (http://genome.ucsc.edu/) or the Ensembl Genome database v35 (www.ensembl.org), NCBI build 35. Table 1 shows the Genbank numbers. To amplify exons, forward and reverse primer sets (primer sequences upon request) were designed with at least 25 bp flanking intronic sequences using Primer3 (http://www.broad.mit.edu/cgibin/primer/primer3_www.cgi) with the conditions described by³³. 3'UTR of NRP2 and exons 1-27 of PIP5K3 have not been sequenced due to current genome browser updates. PCR amplifications were carried out in a volume of 15 µl that contained 15 ng genomic DNA, 4.1 pmol of the PCR primers, 1.5 mM MgCl₂, 0.2 mM and 0.6 units of rTaq polymerase (Amersham Biosciences) or 0.6 units of HotfirePol® DNA

polymerase and solution S (Solis Biodyne) for GC-rich regions or standard conditions of the GC-rich PCR system (Roche). Reactions were cycled at 94°C for respectively 1 min or 15 min for GC-rich regions and then cycled for 35 cycles of 94°C for 30 sec, 57°C for 1 min 15 sec, 72°C for 30 sec, and finally incubated for 6 min at 72°C on B&L primus HT cyclers. PCR products were purified using Multiscreen 96 well plates (Millipore) filled with Sephadex (Amersham Biosciences) and quantified on 1.5% agarose gels. PCR products were sequenced for possible mutations using an ABI3730 capillary sequencer with Big Dye chemistry (Applied Biosystems).

Validation of possible mutations

To investigate whether variants were novel or common SNPs, all variants observed in affected family members were blasted for existence using National Centre of Biotechnology Information (NCBI) SNP blast, build 124 (http://www.ncbi.nlm.nih.gov/SNP/snpblastByChr.html). We applied the nomenclature of den Dunnen et al. (2001) to describe these variants³⁴. When a novel variant was found, segregation analysis of the variant with OA within all family members of the seven families was performed. The impact of a novel variant involving an amino acid change was examined using PolyPhen (http://tux.emblheidelberg.de/ramensky/index.shtml) or SIFT (http://blocks.fhcrc.org/sift/SIFT.html)³⁵. PolyPhen is a computational tool that predicts possible impact of an amino acid substitution on the structure and function of a human protein using straightforward physical and comparative considerations³⁶. SIFT (sorting intolerant from tolerant) estimates tolerance indices that predict tolerated and deleterious substitutions for every position of the query protein sequence using multiple alignment information³⁷. To test the effect on the splicing process, exonic variants were screened for exonic splicing enhancers sequences using http://exon.cshl.edu/ESE38. Conservation was determined using the Multiz Alignments and Conservation track of the UCSC genome browser version 140 (http://genome.ucsc.edu/) which shows a measure of evolutionary conservation in 17 vertebrates, including mammalian, amphibian, bird, and fish species.

Genotyping

Nine novel variants were genotyped in 790 random subjects from the population-based Rotterdam study³⁹. All variants were in Hardy-Weinberg equilibrium (HWE). Variants were genotyped using Sequenom homogenous Mass Extend MassARRAY System (Sequenom Inc, San Diego, CA) using standard conditions. Genotypes were analysed using Genotyper version 3.0 software (Sequenom Inc.).

Statistical analysis

HWE was calculated with an exact HWE test for rare alleles implemented in R version 2.3.1, (http://www.r-project.org/). A logistic regression model was fitted to measure the strength of association, which is expressed as odds ratios (OR) with 95% confidence intervals (95% CI) adjusted for age (years), body mass index (BMI in kg/m²), and sex. In these analyses, homo- and heterozygous carriers of the risk allele were pooled. All analyses were performed with SPSS version 11 software (SPSS, Chicago, IL).

Results

Families

Previously, we performed a genome-wide linkage scan of seven extended families with early onset OA without any dysplasia. The scan revealed significant evidence for linkage to chromosome 2q33.3-2q34, implicating a 5 cM interval (4.6 Mb) between markers D2S1384 and D2S217824. In family 2 and 4, most affected individuals carry identical heterozygous haplotypes resulting in two possible haplotypes explaining the linkage (Figure 1). In family 2, the most likely haplotype 3678 (A1) allows one phenocopy (individual 9) and the second likely haplotype 7658 (A2) allows two phenocopies (subject 6 and 17). In family 4, the most likely haplotype 5515232 (A1) allows no phenocopies and the second likely haplotype 5144984 (A2) allows one phenocopy (individual 7). As illustrated in Figure 1, recombinations in individuals conferring either haplotype A1 or A2 confined this linkage interval. To detect sequence changes, we initially screened three affected family members from three different families contributing most to the linkage: individual 14 in family 1, individual 10 in family 2 and individual 9 in family 4. When a novel variant was found, segregation analysis of the variant with OA within all family members of the seven families was performed.

Mutation screening of positional candidate genes

On the basis of these result, we performed extended mutation analysis of the chromosome 2 locus. Using the human genome resources, 18 known RefSeq genes, nine predicted RefSeq genes (four genes with model status and five genes with predicted status in NCBI build 35) have been identified within the linkage area of chromosome 2 as shown in Table 1. Nearby the linkage region, at marker D2S72, the

ICOS-CTLA4-CD28 cluster is located which is implicated in cytokine secretion and T-cell immunity⁴⁰⁻⁴². *FRZB*, previously associated with hip OA and generalised OA is located more upstream at 2q32^{43;44}. The entire coding region, splice sites, and 5′ and 3′ UTRs of *FRZB*, *CTLA4*, *CD28*, *ICOS*, *NRP2*, *NM*_017759, *NDUFS1*, *EEF1B2*, *GPR1*, *XM*_371590, *ADAM23*, *MDH1B*, *CPO*, *KLF7*, *CREB1*, *NM*_030804, *FZD5*, *IDH1*, *PIP5K3* and *PTHR2* were sequenced comprising 17 RefSeq genes and three predicted genes. Results of the screening of *PTHR2* and *FZD5* have previously been reported²⁴. If the considered variant appeared novel in the current dbSNP database (build 124) and cosegregated with OA in the affected family, it was evaluated according to the criteria previously mentioned.

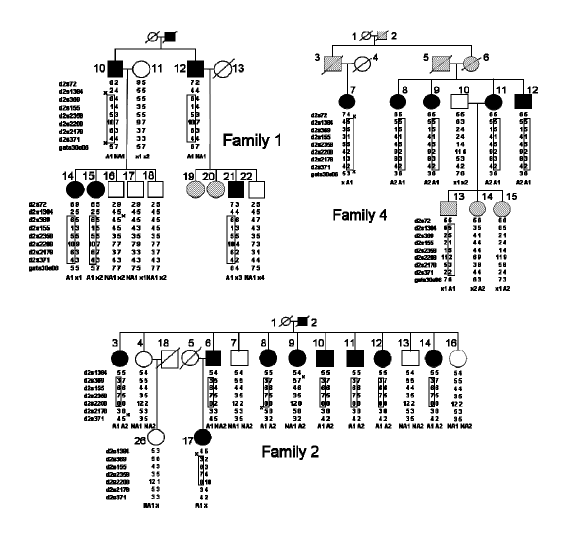


Figure 1 Family pedigrees segregating for generalised early onset OA and chromosome 2q33.3-2q34 loci. Blackened circles and squares indicate affected females and males, respectively. White individuals represent unaffected family members. Crosses indicate recombinations. Shaded individuals indicate individuals with unknown disease status. The black lined box shows the most likely haplotype (A1), allowing one phenocopy (individual 9) in family 2.

Table 1 Known genes between the markers D2S72 and D2S2178 based on RefSeq, mRNA, TrEMBL and Swiss-Prot prediction using Ensembl genome database v35

Marker	Genbank ID	HUGO ID	Description	No. Exons
	NM_001037631, NM_005214	CTLA4	Cytotoxic T-lymphocyte protein 4 precursor	3,4
	NM_006139	CD28	T-cell specific surface glycoprotein CD28 precursor	4,5
D2S72	NM_012092	ICOS	Inducible T-cell co-stimulator precursor	5
D2S1384	NM_057177, NM_205863, NM_152526	ALS2CR19	Amyotrophic lateral sclerosis 2 chromosomal region candidate gene protein 19	22,23,23
	BC009222, NM_201264, NM_018534, NM_201279, NM_003872, NM_201266	NRP2	Neurophilin-2 precursor	5,16,16, 16,17,17
	NM_017759	NM_017759	Unknown (NP_060229)	12
D2S155	NM_005006	NDUFS1	NADH-ubiquinone oxidoreductase 75 kDA subunit	19
	NM_001959, NM_021121	EEF1B2	Elongation factor 1-beta	6,7
			Probable G protein-coupled receptor	3
	XM_371590	XM_371590	Unknown (Q9HCK1)	1
	XM_496742	XM_496742	Unknown (Q9BZ60)	1
D2S369	NM_003812	ADAM23	A disintegrin and metalloproteinase 23 preproprotein	26
D2S2358	NM_206892	MDH1B	malate dehydrogenase 1B (NP_996775)	10
	NM_014929	NM_014929	Unknown (KIAA0971)	12
	NM_173077	CPO	Carboxypeptidase O	9
	NM_003709	KLF7	Krueppel-like factor 7	4
	NM_004379, NM134442	CREB1	cAMP-response element binding protein	8,9
	NM_145280	NM_145280	hepatocellular carcinoma-associated antigen (NP_660323)	4
	NM_152523	NM_152523	hypothetical protein FLJ40432 (NP_689736)	11
	NM_030804	NM_030804	Unknown (CB031)	1
	NM_003468	FZD5	Frizzled 5 precursor	2
D2S2208	XM_371591	XM_371591	Unknown (Q8WW68)	1
	NM_006891	CRYGD	Gamma crystallin D	3
	NM_020989	CRYGC	Gamma crystallin C	3
	NM_005210	CRYGB	Gamma crystallin B	3
	NM_014617	CRYGA	Gamma crystallin A	3
	XM_371592	XM_371592	similar to RIKEN cDNA D630023F18	9
	NM_005896	IDH1	Isocitrate dehydrogenase 1	10
	NM_152671, NM_015040	PIP5K3	FYVE finger containing phosphonositide	10,42
D2S2178	NM_005048	PTHR2	Parathyroid hormone receptor precursor	13

Table 2 Possible mutations segregating in FOA families

Gene	Gene	DNA nt	aa change	Sequence	Splicing ³	PolyPhen ⁴	SIFT ⁵	Segregation family ⁶
	Location	change ¹	_	conservation ²		-		
NRP2	3'UTR ⁷	c.941A>C		no				F4:A1
NRP2	intron 7^8	c.1938-21T>C		no	possible			F2:A1, F4:A2, F1:A1
KIAA1571	exon 1	c.6368G>T	R2133S	no	yes	benign (1.35)9	tolerated (0.00)9	F4:A1, F7:A1
ADAM23	intron 20	c.2065+24C>T		no	possible			F2:A2
IDH1	exon 6	c.782A>G	Y183C	yes	no	damaging (2.89)	not tolerated (0.00)	F2:A2
IDH1	intron 6	c.933-28C>T		no	possible			F2:A1, F4:A2
PIP5K3	3'UTR	c.8429T>A		yes	•			F4:A2
PIP5K3	3'UTR	c.8434insC		no				F4:A2
PTHR2	exon 6	c.786G>T	A225S	no	no	benign (0.12)	tolerated (0.77)	F4:A2

 $^{^1}$ Numbers are based on positions in BC009222 (gi:33874364), NM_201267 (gi:41872566), XM_371590 (gi:42656560), NM_003812 (gi:73765550), NM_005896 (gi:28178824), NM_015040 (gi:50881947), NM_005048 (gi:39995097).

² Sequence conservation based on vertebrate Multiz Alignment and Conservation of 17 species (UCSC Genome browser).

³ Splicing effects in exons have been predicted by ESE finder.

⁴ PolyPhen prediction PSIC score difference.

⁵ SIFT prediction with normalised probability.

⁶ A1 represents the most likely haplotype and A2 the second likely affected haplotype.

⁷ This variant is located in one of the six transcripts (BC009222, gi:33874364).

⁸ This variant is located in six of the seven transcripts (NM_201267 (gi:41872566), NM_003872 (gi:41872532), NM_201279 (gi:41872571), NM_018534 (gi:41872543), NM_201266 (gi:41872561), NM_201264 (gi:41872556)).

⁹ Poor predictions due to little information.

Evaluation of novel variants

The initial screening indicated 26 novel variants (17 SNPs and 9 insertion/deletion polymorphisms). From these 26 variants, only nine promising variants cosegregated with OA within one or more families as illustrated in Table 2. Three of these variants were found in coding regions and involved an amino acid change: (*KIAA1571* R2133S, *IDH1* Y183C and *PTHR2* A225S). Six variants were located in UTR regions (*PIP5K3* c.8429T>A, *PIP5K3* c.8434insC and *NRP2* c.941A>C) or in vicinity of exon/intron boundaries (*NRP2* c.1938-21T>C, *ADAM23* c.2065+24C>T and *IDH1* c.933-28C>T). Using PolyPhen, SIFT and ESE finder analysis to predict possible functional effects of these variants, two variants emerged as potential mutations: *KIAA1571* R2133S and *IDH1* Y183C (Table 2). In addition, three variants (*NRP2* c.941A>C, *PIP5K3* c.8429T>A and *PIP5K3* c.8434insC) were conserved across other species and might be of functional importance.

The predicted *KIAA1571* gene probably belongs, as predicted in Unigene, to the fibronectin type III and M protein repeat family in C. elegans. Fibronectin is a component of the ECM and *KIAA1571* may therefore be an excellent candidate gene. The G/T nucleotide change in the third exon of *KIAA1571* (Q9HCK1) results in disruption of exonic splicer enhancer motifs which serves as binding site for Serine/Arginine protein 40 and 55 and might be therefore a functional variant. However, because this gene is a predicted gene, little is known about other possible predicted functional effects on the protein. The novel variant *KIAA1571* R2133S cosegregated in family 4 and 7 with OA and showed a rare population frequency of 0.01 corresponding to nine carriers of 763 genotyped (Table 3). In the Rotterdam sample, we did not observe a significant association of this variant with generalised ROA. It is unlikely that this variant is a causal mutation.

Isocitrate dehydrogenase 1 (*IDH1*) encodes a cytoplasmic enzyme which catalyzes the oxidative decarboxylation of isocitrate to 2-oxoglutarate and has a significant role in cytoplasmic NADPH production⁴⁵. In *IDH1*, two variants (Y183C and c.933-28C>T) cosegregated with the OA phenotype. *IDH1* Y183C cosegregated in affected family members in family 2 (A2, Figure 1). This variant was located in exon 6 encoding the isocitrate/isopropylmalate dehydrogenase domain (PF00180) of IDH1, predicted to be probably damaging for the protein structure/function by SIFT and PolyPhen, and highly conserved across all species investigated. Based on these results, this variant could be functional for the onset of generalised OA. In the Rotterdam sample, we observed 14 carriers out of 767 genotyped corresponding to a frequency of 0.02. In addition, carriers of this variant conferred an odds ratio, adjusted for age, BMI and sex of 2.8 (95% CI, 0.82-9.7, P = 0.10) to have generalised ROA, as shown in Table 3.

Another variant in this gene, IDH1 c.933-28C>T, was identified in family 2 and 4, near the intron/exon boundary of exon 7. This variant was not conserved across other species, and was not associated with generalised ROA in the Rotterdam sample (frequency 0.04).

Neuropilin 2 (*NRP2*), is an interesting gene because it encodes for the co-receptor of vascular endothelial growth factor₁₆₅ (VEGF ₁₆₅) which is an essential factor for endochondral ossification^{46;47}. Furthermore, VEGF and its receptors are expressed in OA cartilage and VEGF stimulates production of ECM degrading matrix metalloproteinases (MMPs)^{48;49}. In the *NRP2* gene, two novel variants were found: c.941A>C and c.1938-21T>C. In family 4, *NRP2* c.941A>C was identified in a residue with a low conservation score. This variant showed a frequency of 0.03 in the random population and no significant association with generalised ROA was observed. The second *NRP2* variant, c.1938-21T>C, was not conserved and cosegregated in three families (1, 2, and 4) and was more frequent in the population (0.07). Carriers of at least one risk allele of the *NRP2* c.1938-21T>C variant, conferred a significant increased risk of 2.1 (95% CI, 1.1-4.1, P = 0.032), adjusted for age, sex, BMI, to have generalised ROA (Table 3).

Phosphatidylinositol-3-phosphate/phosphatidylinositol 5-kinase, type III (PIP5K3) catalyzes the phosphorylation of phosphatidylinositol-4-phosphate and has a role in endosome-related membrane trafficking⁵⁰. We found two novel variants (PIP5K3 c.8429T>A and PIP5K3 c.8434insC) in the 3'UTR region of PIP5K3 in family 4. PIP5K3 c.8429T>A involved a highly conserved residue. PIP5K3 c.8434insC was not conserved. In the Rotterdam sample, we observed that PIP5K3 c.8429T>A and PIP5K3 c.8434insC showed a population frequency of 0.04 and were in complete LD (D'= 1, r^2 = 1). In the Rotterdam sample, no significant associations of these variants with generalised ROA were observed excluding a possible pathogenic role in relation to the onset of FOA. Even though PIP5K3 c.8429T>A occurred in a conserved residue, it is likely that these variants are neutral polymorphisms.

Table 3 Frequencies of novel segregating variants in population-based sample scored for generalised ROA (GOA)

Variant	Total	No GOA ¹	GOA ¹	P^2
	(n = 790)	(n = 659)	(n = 131)	
NRP2 c.941A>C	0.03(743)	0.03(601/17)	0.02(123/2)	0.44
NRP2 c.1938-21T>C3	0.07(741)	0.07(574/40)	0.12(112/15)	0.03
KIAA1571 c.6368G>T	0.01(763)	0.01(630/7)	0.02(124/2)	0.47
<i>ADAM</i> 23 c.2065+24C>T	0.04(741)	0.04(591/22)	0.03(124/4)	0.79
IDH1 c.782A>G	0.02(767)	0.01(631/9)	0.04(122/5)	0.10
<i>IDH1</i> c.933-28C>T	0.04(773)	0.04(622/23)	0.04(123/5)	0.94
PIP5K3 c.8429T>A3	0.04(756)	0.04(603/25)	0.04(123/5)	0.83
<i>PIP5K3</i> c.8434insC ³	0.04(734)	0.04(592/24)	0.04(113/5)	0.96

¹ Frequency (number of carriers of two common alleles/ number of carriers of one or two rare alleles).

 $^{^{2}\}textit{P}$ values adjusted for age, BMI and sex.

³One, one and two homozygous carriers for NRP2 c.1938-21T>C, PIP5K3 c.8429T>A and PIP5K3 c.8434insC respectively.

Finally, we also examined whether some novel variants were inherited together in different families to identify a possible LD pattern or genetic interaction resulting in a high LOD score linked to OA. In the seven families, only two variants, *NRP2* c.1938-21T>C and *IDH1* c.933-28C>T occurred together on haplotype A1 in family 2 and on haplotype A2 in family 4. In the random population, this inheritance pattern was observed only once in 754 genotyped subjects (0.0013). This individual had spinal DD at three disc levels which has a prevalence of 0.04 in the random population.

Discussion

Upon screening 20 genes localised within or near the area of linkage in three families that contributed most to the linkage, we identified nine novel variants cosegregating with OA. We evaluated the significance of the novel variants by prediction of pathogenicity using *in silico* functional analysis and by establishing the frequency in the random population. The *IDH1* Y183C variant was the most promising variant since it cosegregated on haplotype A2 with the OA phenotype in family 2, was predicted to be probably damaging for the protein structure/function and concerned a highly conserved residue. Among carriers of this variant in the general population (frequency 0.02), the risk of generalised ROA was 2.8 (95% CI, 0.82-9.7, P = 0.10). Given these results and the finding that family 2 already showed a significant linkage on its own²⁴, this variant may contribute to the FOA susceptibility in family 2.

The risk allele showed reduced penetrance among carriers in the Rotterdam study which might not be expected for a mutation causing a severe disorder such as FOA. It is expected that causal variants for severe OA are very rare and strongly correlated with generalised ROA. The reduced penetrance of this variant might reflect genetic interaction with other causal variants at this or other loci. Therefore, the pathogenic potential of this variant should be confirmed further in other populations with advanced OA or supported by functional assays. *IDH1* supplies NADPH for antioxidant systems suggesting a regulatory role in cellular defense against oxidative stress and in senescence⁵¹. Little is known about a possible role of *IDH1* in cartilage but we speculate that increased oxidative stress could make chondrocytes more susceptible to cell death which might contribute to the onset of OA.

A second promising variant, NRP2 c.1938-21T>C, emerged from our mutation analysis which cosegregated in three families (1, 2, and 4) contributing most to the linkage. Carriers of this variant conferred a significant increased risk of 2.1 (95% CI, 1.1-4.1, P = 0.032), to have generalised ROA. Given the low effect size and the high frequency (0.07) in the random population, this variant may be not sufficient alone for the onset of FOA but it might be that this variant modulates OA susceptibility

and interacts genetically with another variant. *NRP2* acts as a co-receptor of VEGF₁₆₅, which is produced from hypertrophic chondrocytes and is also expressed in OA cartilage^{52;53}. VEGF is an essential coordinator of growth plate morphogenesis and triggers cartilage remodeling⁵⁴. VEGF may contribute to OA cartilage destruction through stimulation of MMPs⁵⁵⁻⁵⁷.

The *NRP2* c.1938-21T>C occurred together with *IDH1* c.933-28C>T on haplotype A1 in family 2 and on haplotype A2 in family 4. This haplotype is extremely rare in the random population (0.0013). These results may suggest that family 2 and 4 might have a common ancestor with a FOA susceptibility haplotype. At this point, we conclude that possibly a causal variant in the LD pattern driven by both variants may have contribution to OA in family 2 and 4. Alternatively, both variants could be in LD with a causal variant each indicating an oligogenic disorder or a causal variant with a genetic modifier. More research is necessary to fully investigate the role of this haplotype in the OA phenotype.

How can we reconcile these results; what do they tell us about the proposed role of these genes in OA and why is there no compelling evidence for a genetic variant responsible for the observed LOD score of 6.0 in the families with early onset OA? Although the frequencies of carriers of novel variants identified in the OA families were rare (0.01-0.07%) in the random population-based sample as expected for such a severe phenotype as FOA, detected associations with generalised ROA did not achieve a high level of significance. This may be explained by the low frequency of the carriers, the absence of a clinical OA assessment and the relatively high frequency of generalised ROA above 55 years (0.17) which can result in reduced power of the statistical test, misclassification and spurious associations. We also analysed the presence of ROA among carriers in the random population as quantitative trait using a sumscore of the number of affected joints²⁴ which revealed similar findings (data not shown).

There is a possibility that the family members could be affected with the generalised ROA by chance alone although the phenotype is more severe (clinical OA before age 50). In family 2 and family 4, we observed two possible haplotypes segregating with the disease allowing one or two phenocopies suggesting that these individuals could be sporadic patients. Both haplotypes (A1 and A2) confined the linkage area. Consistent with an age-related disorder, we were not able to perform further segregation analysis because first degree relatives died or are too young to reveal symptoms of the disease.

We prioritised genes for sequencing based on the function known in the literature and human genome resources. Consequently, we have not selected genes which have clearly a function completely unrelated to OA like the gamma crystallins which is associated with cataract and lens morphology⁵⁸ or ALS2CR19 which plays a role in

asymmetrical cell division and cell polarization processes⁵⁹ or a few predicted genes like NM_014929 with unknown function. Obviously, these genes could carry an OA causing allele and will be screened for mutations in the future. In addition, we selected families that contributing most to the linkage for the initial mutation survey. Apparently, remaining four families could harbor causal variants in one of the screened genes. Alternatively, it is possible that affected family members may have a noncoding regulatory mutation in promoter or intron or a heterozygous deletion of one or more of the exons in one of the screened genes that has not yet been detected in our study. In addition, predictions of the on the basis of computational algorithms as PolyPhen, SIFT and ESE finder are difficult to interpret. Furthermore, even variants that are associated with disease may be in linkage disequilibrium with the true causal variant or are rare polymorphisms.

Although we were able to identify nine novel variants cosegregating with the FOA phenotype by our extended mutation analysis, we may found no robust evidence for a major disease gene responsible for the observed linkage to the FOA phenotype. Our results, however, might indicate a possible modulating role for variants in or near *NRP2* and *IDH1*. Further mutation analysis of the linkage area on chromosome 2q33.3-2q34 and confirmation of the most promising variants in other populations with advanced OA is needed.

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