

Contribution of *SNRNP200* sequence variations to retinitis pigmentosa

X Zhang¹, TYY Lai¹, SWY Chiang¹, POS Tam¹, DTL Liu^{1,2}, CKM Chan^{1,3}, CP Pang^{1,2}, C Zhao⁴ and LJ Chen^{1,2}

Abstract

Purpose Mutations in the *SNRNP200* gene have been reported to cause autosomal dominant retinitis pigmentosa (adRP). In this study, we evaluate the mutation profile of *SNRNP200* in a cohort of southern Chinese RP patients.

Methods Twenty adRP patients from 11 families and 165 index patients with non-syndromic RP with mixed inheritance patterns were screened for mutations in the mutation hotspots of *SNRNP200*. These included exons 12–16, 22–32, and 38–45, which covered the two helicase ATP-binding domains in DEAD-box and two sec-63 domains. The targeted regions were amplified by polymerase chain reaction and analyzed by direct DNA sequencing, followed by *in silico* analyses.

Results Totally 26 variants were identified, 18 of which were novel. Three non-synonymous variants (p.C502R, p.R1779H and p.I698V) were found exclusively in patients. Two of them, p.C502R and p.R1779H, were each identified in one simplex RP patient, whereas p.I698V occurred in one patient with unknown inheritance pattern. All three residues are highly conserved in *SNRNP200* orthologs. Nevertheless, only p.C502R and p.R1779H were predicted to affect protein function by *in silico* analyses, suggesting these two variants are likely to be disease-causing mutations. Notably, all mutations previously identified in other study populations were not detected in this study.

Conclusions Our results reveal a distinct mutation profile of the *SNRNP200* gene in a southern Chinese cohort of RP patients. The identification of two novel candidate

mutations in two respective patients affirmed that *SNRNP200* contributes to a proportion of overall RP.

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Introduction

Retinitis pigmentosa (RP; MIM268000; Mendelian Inheritance in Man; National Center for Biotechnology Information, Bethesda, MD, USA) refers to a clinically and genetically diverse group of diffuse retinal dystrophies. The prevalence of RP is about 1/4000 to 1/6000 in certain populations^{1–3} with no significant ethnic difference and affects more than 1 million individuals worldwide.⁴ Presenting symptoms of RP are various, but the typical ones include the following: (1) nyctalopia (night blindness) as a hallmark, most commonly the earliest symptom in RP; (2) constricted visual fields, usually peripheral, and even central visual loss in advanced cases; and (3) photopsia (seeing flashes of light).⁵ These symptoms usually occur during the third decade,⁶ although they can occur earlier depending on the inheritance pattern.⁷ The fundus hallmarks include arteriolar attenuation, mid-peripheral perivascular bone-spicule pigmentation, tessellated fundus appearance, and waxy disc pallor.⁸ RP can occur as a simplex disorder without known family history (35–50%) or inherit in autosomal dominant (ad) (10–30%), autosomal recessive (ar) (10–45%), and X-linked (xl) (0–15%) patterns.^{4,9} Also it may present as part of systemic disorders, which are usually autosomal recessive.¹⁰

¹Department of Ophthalmology and Visual Sciences, The Chinese University of Hong Kong, Hong Kong, China

²Department of Ophthalmology and Visual Sciences, Prince of Wales Hospital, The Chinese University of Hong Kong, Hong Kong, China

³Hong Kong Eye Hospital, Hong Kong, China

⁴Department of Ophthalmology, Nanjing Medical University First Affiliated Hospital, Nanjing, China

Correspondence: LJ Chen, Department of Ophthalmology and Visual Sciences, The Chinese University of Hong Kong, Hong Kong Eye Hospital, 147K Argyle Street, Kowloon, Hong Kong 852, China.
 Tel: +852 39435810;
 Fax: +852 27159490.
 E-mail: lijia_chen@cuhk.edu.hk

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Genetically RP is a highly heterogeneous condition. More than 60 RP genes have been identified, while 23, 36, and 3 genes were responsible for adRP, arRP, and xlRP, respectively (RetNet, the Retinal Information Network, University of Texas Houston Health Science Center, Houston, TX, accessed on 22 November 2012). Despite this large number of genes, the vast majority was mapped in populations other than Chinese. Up to date no single gene or mutation contributes to a large proportion of RP in Chinese has been reported,^{11–19} as does p.P23H of *rhodopsin* (*RHO*, MIM 180380) in Caucasians. The *RP33* locus, initially mapped to chromosomal region 2cen-q12.1, was identified in a large Chinese family with adRP.²⁰ Subsequent studies had identified two mutations, p.S1087L²¹ and p.R1090L,²² in the *small nuclear ribonucleoprotein 200 kDa* (*U5*) (*SNRNP200*, MIM 601664) gene to be segregated with RP in two respective large Chinese adRP families. *SNRNP200* is a member of the RNA intron-splicing factor protein family and is widely expressed in human tissues. It encodes a U5 small nuclear ribonucleoprotein 200 kDa helicase, which is a 2136-amino acid splice-complex enzyme containing two pairs of DEAD-box, each followed by a Secretary(sec)-63 domain. DEAD-box proteins are ubiquitous enzymes that use ATP to rearrange RNA and RNA-protein structures. They consist of a helicase core containing 12 conserved motifs involved in ATP binding, RNA binding, and ATP hydrolysis.²³ They all possess an ATPase activity that is stimulated by RNA, giving rise to a broad range of biochemical effects *in vitro*. Thus, they can often unwind short RNA duplexes. This property, together with their structural similarity to the well-characterized DNA helicases, has led to their common designation as 'RNA helicases.' However, they also can displace RNA-bound proteins, accelerate RNA strand annealing or RNA folding, and act as RNA clamps to form stable ribonucleoprotein (RNP) complexes.²³ In yeast, the sec-63 domain is required to reconstitute post-translational translocation in protoliposomes.²⁴ Yeast BiP/Kar2p binds to a DnaJ domain of sec-63 and mediates the import of polypeptides into the endoplasmic reticulum (ER) lumen²⁵ and the export of misfolded proteins into the cytoplasm.²⁶ The DnaJ domain of sec-63 is known to face the ER lumen.²⁷ The large cytoplasmic portion of sec-63 has no known homolog in other proteins. Sequence analysis demonstrated significant similarity between this region and two regions of the helicases family typified by yeast Brr2p and human U5-200 kD.^{28,29} These RNA helicases contain two sec63-like domains, each lying C-terminal to an ATPase domain. The Brr2p N-terminal ATPase domain is essential for *in vitro* RNA unwinding in U4/U6 small ribonucleoprotein particles (snRNPs).^{30,31} Although the functions of the helicase

sec-63-like domains remain unknown, the associations of the translocon and Brr2p with RNA-protein complexes (the ribosome and spliceosome, respectively) indicate that sec-63 may act as a ribosome receptor. The two initially reported *SNRNP200* mutations are both located in the first sec-63 domain, suggesting this domain could be a mutation hotspot in individuals of Chinese descent. Further, the contribution of *SNRNP200* to RP has also been investigated in a group of Americans with mixed descent.³² The p.S1087L was also identified in this study, along with another three disease-causing mutations (DCM), p.R681C, p.R681H, and p.Y689C. Besides, a new mutation p.Q885E was recently identified in another four generations of Chinese family.³³ All four mutations (p.R681C, p.R681H, p.Y689C, and p.Q885E) are located in the first DEAD-box of *SNRNP200*.

In this study, we screened the regions containing all the DEAD-box and sec-63 domains in *SNRNP200* that may cover the mutation hotspots in a southern Chinese cohort of 185 RP patients and 178 control subjects in order to further evaluate the contributions of *SNRNP200* in Chinese RP.

Materials and methods

Study subjects

Twenty adRP patients from 11 families and 165 unrelated non-syndromic RP patients were recruited from the Hong Kong Eye Hospital and the Prince of Wales Hospital, Hong Kong. All were Han Chinese. The diagnosis was based upon typical RP presentations according to ocular examinations that included slit-lamp biomicroscopy, full-field electroretinography, visual fields, and fundus photography. Besides, best-corrected visual acuity (BCVA) and intraocular pressure as assessed by applanation tonometry were recorded. Patients with syndromic RP, such as Bardet-Biedl syndrome and Usher syndrome, or end-stage syphilitic neuroretinitis, and cancer-associated retinopathy were excluded. A group of 178 unrelated healthy control subjects, aged above 60 years, were also recruited. They were given complete ophthalmic examinations and confirmed free of RP or other major eye diseases except for mild senile cataract. This study was approved by the Ethics Committee on Human Research, the Chinese University of Hong Kong. Informed consents were obtained from all study subjects. All procedures were conducted in accordance with the tenets of the Declaration of Helsinki.

Mutational screening of the *SNRNP200* gene

In order to evaluate the contribution of the potentially mutation hotspots in *SNRNP200*, we screened 24 exons

of the gene, including exons 12–16, 22–32, and 38–45, in all the study subjects. Exons 12–15 and exons 30–32 cover the two helicase ATP-binding domains, whereas exon 16 covers the first connection of the helicase ATP-binding domain and the helicase CTER, which could be the hotspot in Americans.³² Exons 22–29 and exons 38–45 cover the two sec-63 domains.

Genomic DNA from whole blood was extracted using the QIAamp DNA Blood kit (Qiagen, Valencia, CA, USA) according to the manufacturer's instructions. Polymerase chain reaction (PCR) primers were designed according to the sequence of human *SNRNP200* (ENSG00000144028) from the Ensembl database (primer information provided upon request). A total of 14 amplicons covering the regions, including exons 12–16, 22–32, and 38–45, and the corresponding exon–intron boundaries were analyzed by PCR and direct DNA sequencing with the Big-Dye Terminator Cycle Sequencing Reaction Kit (version 3.1, Applied Biosystems, Inc. (ABI), Foster City, CA, USA) on an automated DNA sequencer (3130XL; ABI) according to the manufacturer's protocol.

Analyses of variants

The detected variants were defined as 'novel' if it has neither been reported in the literature nor registered in the single nucleotide polymorphism database (dbSNP). Several criteria were observed to select and prioritize potentially disease-causing variants: (1) the variant was predicted to alter the amino-acid sequence of the protein; (2) it occurred exclusively in patients and was absent in controls; and (3) it was predicted to alter the protein structure or function by *in silico* analyses. In this study, we applied five *in silico* programs. Three of them, PolyPhen, SIFT, and PMUT, had been described in our previous study.¹¹ Here, two new programs were used to predict the effect of each missense mutation: PolyPhen-2 (Polymorphism Phenotyping version 2, <http://genetics.bwh.harvard.edu/pph2/>; accessed on 10 November 2012)³⁴ and SIFT BLink (Sorts Intolerant From Tolerant Blink, http://sift.jcvi.org/www/SIFT_BLink_submit.html; accessed on 10 November, 2012).³⁵ In PolyPhen-2, a position-specific independent counts (PSIC) score, based on multiple sequences alignment of observation and three-dimensional protein structural information, was given to each variant in the context of two data sets: HumDiv and HumVar. We chose HumVar because it is the preferred model for diagnosis of Mendelian diseases. A mutation with a PSIC score of >2.0 was classified as 'probably damaging,' a score of 1.5 to 2.0 was classified as 'possibly damaging,' and a score of <1.5 was classified as 'benign'.³⁴ In SIFT BLink, the protein sequence ID NP_054733.2

(GI: 40217847) was used with default parameter setting. The specific amino-acid location in the protein was compared among various organisms from pre-computed NCBI BLAST searches. The output score represented a normalized probability for how the amino-acid substitution was tolerated. A score of <0.05 gave a prediction of 'affect protein function' and a score of >0.05 gave a prediction of 'tolerated'.³⁵

The influence of a substitution on any functional motifs was studied by the PROSITE database (ScanProsite, <http://prosite.expasy.org> Swiss Institute of Bioinformatics, Lausanne, Switzerland; accessed on 10 November 2012). The helicase ATP-binding domain and the helicase CTER domain were revealed by the program. Further, the SMART software (Simple Modular Architecture Research Tool, European Molecular Biology Laboratory, Heidelberg, Germany; accessed on 10 November 2012 <http://smart.embl-heidelberg.de/>) was used to locate the sec-63 domain in the *SNRNP200* protein. To predict coding exons and splice sites in *SNRNP200*, we used NetGene2 Server (Technical University of Denmark, Lyngby, Copenhagen, Denmark; accessed on 10 November 2012 <http://www.cbs.dtu.dk/services/NetGene2/>).³⁶ We also investigated evolutionary conservation of amino acids in *SNRNP200* using the UCSC Genome Browser (<http://genome.ucsc.edu/cgi-bin/hgGateway>, accessed on 9 January 2012).³⁷ An amino acid that is shown to be highly conserved among different species indicated its importance in maintaining protein function.³⁸

Statistical analysis

Genotype and allele frequencies of any common variants were compared between patients and controls using the χ^2 -test in SPSS (ver.16.0, SPSS Inc., Chicago, IL, USA). The Bonferroni method was used to correct the *P*-values in multiple comparisons. A corrected *P*-value (*P*_{corr}) of less than 0.05 was defined as statistically significant.

Results

Characteristics of the RP patients

Among the 11 recruited adRP families, there were 10 male and 10 female affected members. Whereas the 165 non-syndromic RP patients consisted of 87 males and 78 females, with age ranging from 6 to 84 years. On the basis of the family history, 11 (6.7%) of them were classified as adRP, 26 (15.8%) as arRP, 2 (1.2%) as xLRP, and 97 (58.8%) as simplex RP (sRP). Besides, 29 (17.6%) patients could not be classified because of the absence of family information and were denoted as 'unknown' inheritance pattern. Therefore, these 165 patients represented a

group of southern Chinese non-syndromic RP patients with mixed inheritance.

Sequence variants detected in the SNRNP200 gene

A total of 26 variants were identified in the sequenced regions that cover the entire two helicase ATP-binding domains and the two Sec-63 domains (Table 1). Eight of them (p.L1184L, p.S1218S, c.5134-6C>G, p.L1773L, c.5324-31G>C, p.A1819A, c.5488 + 82C>T, and c.5755-20A>G) were common SNPs registered in the dbSNP database. They were found in both patients and control subjects. Association analyses showed that all these eight SNPs were not associated with RP ($P_{\text{corr}} > 0.05$). Further, none of these variants leads to a missense change, implying that they are unlikely to be disease causing.

Apart from the eight SNPs, we identified 18 novel variants. Nine were intronic, of which five (c.1515 + 67T>C, c.3365 + 145G>A, c.3366-25A>G, c.5134-17C>T, and c.5324-41G>A) were each found in one patient but not in controls. Although we cannot rule out the pathogenicity of these intronic changes as family members were not available for segregation analysis, none of these variants is located in splicing sites according to the NetGene2 Server prediction (data not shown), making them less likely to be disease causing. Two intronic variants (c.2036 + 74A>G and c.2036 + 93G>T) were each found in two control subjects, whereas the remaining two (c.2036 + 107delT and c.3639 + 53_c.3639 + 93del) were found in one control each. They are less likely to be pathogenic. Of the nine coding variants, four synonymous ones, p.G477G, p.E494E, p.S1230S, and p.I2029I, were found in one patient each, whereas p.L2019L was found in one control subject. By using SIFT and SIFT BLINK, the only *in silico* analysis available for synonymous variants showed these five variants were benign with low potential of disease causing (data not shown). Among the four non-synonymous variants, p.A1995T was found in both patients and controls, whereas p.C502R, p.I698V, and p.R1779H were each found in one patient but not in controls (Figure 1).

All the four missense variants were tested by using the *in silico* programs and multiple protein sequences alignment. Three residues (p.C502, p.I698, and p.R1779) were absolutely conserved, whereas the p.A1995 was highly conserved across different species, indicating possible impact on the protein function (Figure 2a). According to the PROSITE and SMART, p.C502 is located in the first helicase ATP-binding domain, p.R1779 is found between the second HELICASE CTR domain and the second sec-63 domain (Figure 2b), p.I698 is in the first helicase CTR domain, whereas p.A1995 is in the second

sec-63 domain. The p.A1995T was predicted to be benign by all of the *in silico* programs and was found in one control subject, suggesting a low chance of being a DCM. The p.C502R, which was found in one sRP patient, was predicted to be pathological, whereas p.I698V was shown to be benign by all five programs (Table 2). For p.R1779H, which was found in another sRP patient, the predictions were inconsistent. As it has been suggested that the Polyphen 2 program could be more reliable,³⁴ p.R1779H is considered pathological. With respect to the carriers of the two potential DCMs, the patient carrying p.C502R is an 82-year-old lady who was diagnosed with RP at the age of 64 years with typical RP fundus changes (Figure 3). Her BCVA was 20/200 and hand movement in her right and left eyes, respectively, with severe visual field damage in both eyes (Figure 3) during her last visit. The patient with p.R1779H had disease onset at the age of 53 years and had typical manifestations of RP (data not shown). These two patients had no other affected family members by the time of recruitment, suggestive of simplex RP. Mutation in some other RP genes, including *RHO*, *RPL*, *NR2E3*, *NRL*, and *BEST1*, had all been excluded in these patients.^{11,12,18}

Discussion

In this study, we have evaluated the mutation profile in 24 exons containing the hotspots in *SNRNP200* among a cohort of southern Han Chinese RP patients and controls. A total of 18 novel variants were detected, among which three non-synonymous changes, p.C502R, p.I698V, and p.R1779H, were found exclusively in patients. Moreover, p.C502R and p.R1779H were predicted to be pathological by *in silico* programs. They are likely to be disease-causing mutations of RP. If they are genuinely pathogenic, the *SNRNP200* gene might have accounted for ~1.1% (2/176, 11 of them from families and 165 were simplex) of overall RP, albeit we only screened the potential mutation hotspots of the gene. In contrast, another non-synonymous variant p.A1995T, which occurred in both patients and controls, was predicted to be benign, suggesting it is unlikely to be pathogenic. Notably, the three DCMs p.Q885E, p.S1087L, and p.R1090L, which were previously identified in three respective adRP pedigrees from northern China,^{21,22,33} were not found in our study. In the first sec-63 domain region, which harbors two of these three mutations (p.S1087L and p.R1090L), only three common SNPs (p.L1184L, p.S1218S, and p.S1230s) and three intronic changes (c.3365 + 145 G>A, c.3366-25 A>G, and c.3639 + 53_c.3639 + 93del) were detected in our study subjects. Likewise, the mutations that were detected in the Caucasian cohort³² were also not identified in our present study. Therefore, results of our study enrich

Table 1 Sequence variants detected in exons 12–16, 22–32, and 38–45 of the SNRNP200 gene among 165 Chinese RP patients and 20 affected members from 11 adRP families

Number	Location	Nucleotide change	Residual change	Description	MAF in HapMap (CHB)	Variation frequency ^a		
						Mixed RP Patients	adRP family Affected members	Controls
1	Exon 12	c.1431C>T	p.G477G	Novel	/	0/1/165	0/0/20	0/0/178
2	Exon 12	c.1482G>A	p.E494E	Novel	/	0/0/165	0/1/20	0/0/178
3	Exon 12	c.1504T>C	p.C502R	Novel	/	0/1/165	0/0/20	0/0/178
4	Intron 12	c.1515+67T>C	/	Novel	/	0/1/165	0/0/20	0/0/178
5	Intron 15	c.2036+74A>G	/	Novel	/	0/0/165	0/0/20	0/2/178
6	Intron 15	c.2036+93G>T	/	Novel	/	0/0/165	0/0/20	0/2/178
7	Intron 15	c.2036+107delT	/	Novel	/	0/0/165	0/0/20	0/1/178
8	Exon 16	c.2092A>G	p.I698V	Novel	/	0/1/165	0/0/20	0/0/178
9	Intron 25	c.3365+145 G>A	/	Novel	/	0/1/165	0/0/20	0/0/178
10	Intron 25	c.3366-25 A>G	/	Novel	/	0/1/165	0/0/20	0/0/178
11	Exon 27	c.3550T>C	p.L1184L	rs3171927	0.155	5/50/165	0/3/20	1/51/178
12	Intron 27	c.3639+53_c.3639+93del	/	Novel	/	0/0/165	0/0/20	0/1/178
13	Exon 28	c.3654T>C	p.S1218S	rs3214060	0.088	0/25/165	0/5/20	1/14/178
14	Exon 28	c.3690C>T	p.S1230S	Novel	/	0/1/165	0/0/20	0/0/178
15	Intron 36	c.5134-17C>T	/	Novel	/	0/1/165	0/0/20	0/0/178
16	Intron 36	c.5134-6C>G	/	rs3214062	0.005	0/2/165	0/0/20	0/2/179
17	Exon 37	c.5317C>T	p.L1773L	rs772175	0.474	33/81/165	2/10/20	25/78/178
18	Exon 37	c.5336G>A	p.R1779H	Novel	/	0/1/165	0/0/20	0/0/178
19	Intron 37	c.5324-41G>A	/	Novel	/	0/1/165	0/0/20	0/0/178
20	Intron 37	c.5324-31G>C	/	rs78519182	0.026	0/4/165	0/0/20	0/2/178
21	Exon 38	c.5457C>T	p.A1819A	rs148995951	N/A	0/0/165	0/0/20	0/1/178
22	Intron 38	c.5488+82C>T	/	rs72937668	0	0/2/165	0/0/20	0/0/178
23	Intron 40	c.5755-20A>G	/	rs3214063	0.216	10/57/165	2/5/20	5/61/178
24	Exon 42	c.5983G>A	p.A1995T	Novel	/	0/1/165	0/0/20	0/1/178
25	Exon 42	c.6057A>G	p.L2019L	Novel	/	0/0/165	0/0/20	0/1/178
26	Exon 42	c.6087C>A	p.I2029I	Novel	/	0/1/165	0/0/20	0/0/178

Abbreviations: CHB, Chinese Han Beijing in HapMap populations; MAF, minor allele frequency; N/A, not available.

^aThe variation frequency was presented in ‘Homozygote variant genotypes/Heterozygote variant genotypes/Total genotypic counts’ fashion.

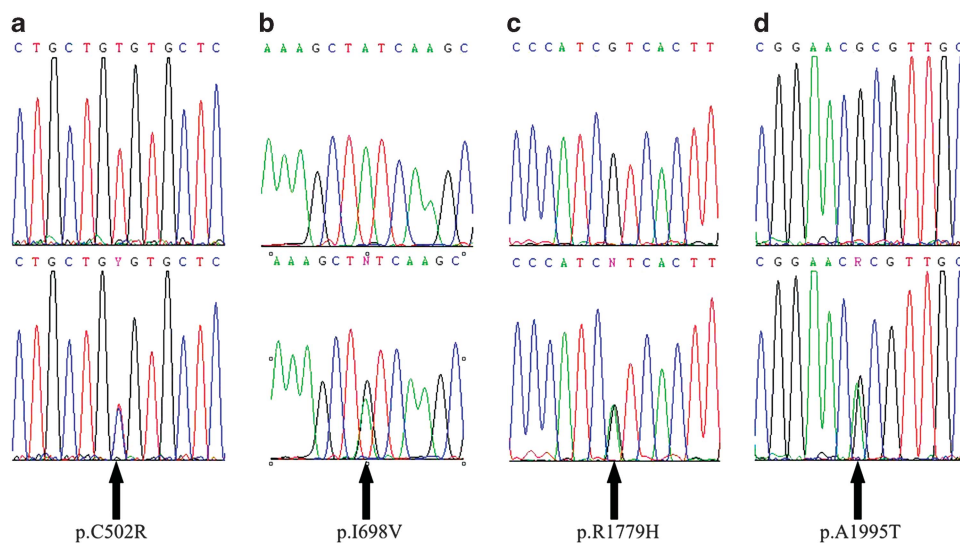


Figure 1 The forward sequence chromatograms of (a) p.C502R; (b) p.I698V; (c) p.R1779H; and (d) p.A1995T. Chromatograms for the normal allele are shown above those for the mutant allele. Arrows indicate positions of mutations.

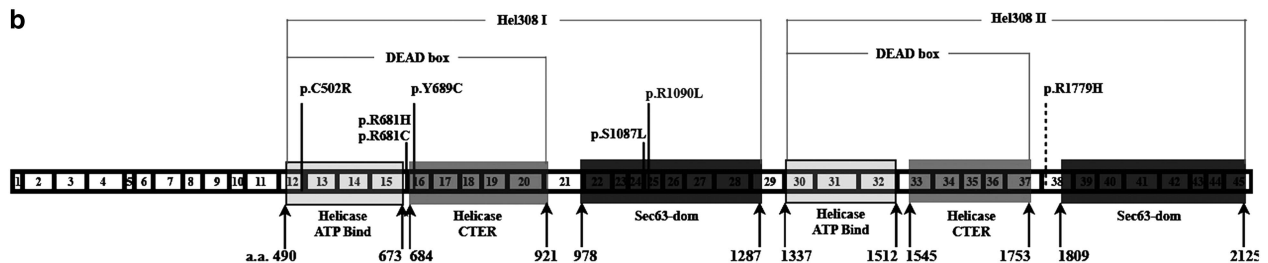
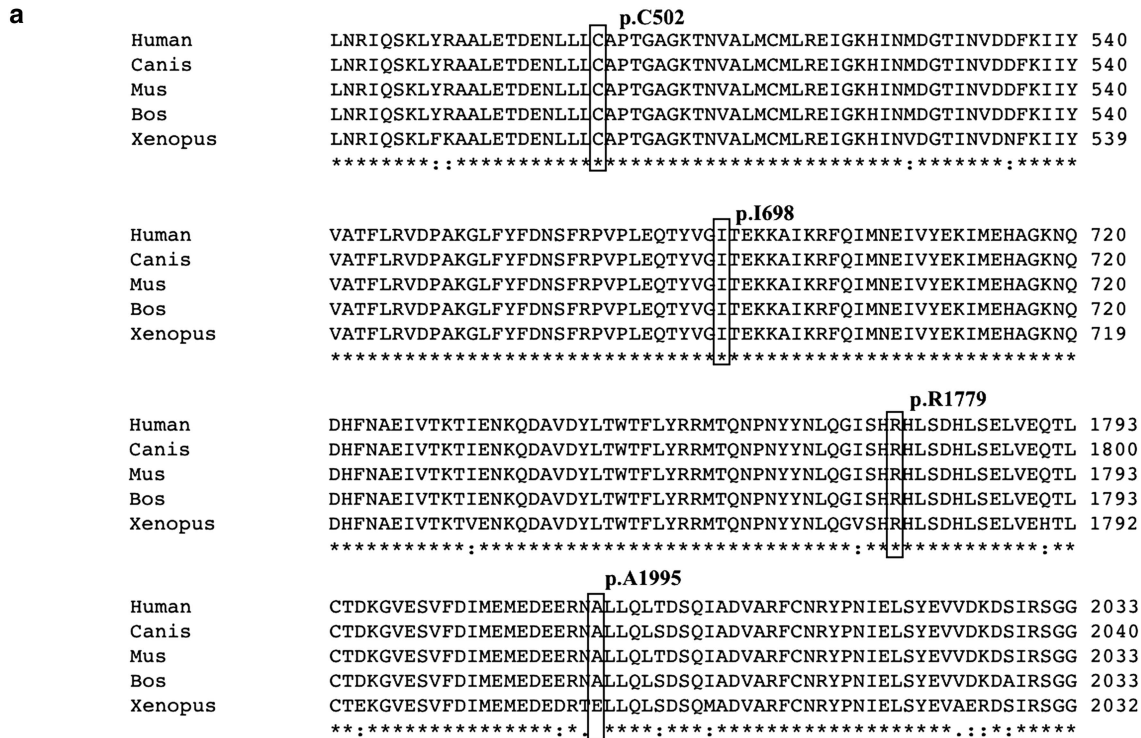


Figure 2 (a) Multiple protein sequence alignment of SNRNP200 (partially), showing the location of p.C502, p.I698, p.R1779, and p.A1995, while the first three residues are absolutely conserved and the last residue are well conserved across different species. The accession numbers of SNRNP200 protein sequences of different species are as follows: Human NP_054733.2, Canis (*Canis lupus familiaris*) XP_532949.2, Mus (*Mus musculus*) NP_796188.2, Bos (*Bos taurus*) NP_001193092.1, Xenopus (*Xenopus (Silurana) tropicalis*) XP_002932581.1. (b) Protein structure and all reported disease-causing mutations in SNRNP200 with transcript reference No. ENST00000323853. 1–45 demonstrated exon 1–45 with corresponding length. Domain location reference software: PROSITE for helicase ATP-binding and helicase CTER domains; SMART for Sec-63 dom; a.a.: amino acid sequence number of SNRNP200; mutations reference: p.C502R (present study); p.R681H;³² p.R681C;³² p.Y689C;³² p.Q885E;³³ p.S1087L;^{21,32} p.R1090L;²¹ p.R1779H (present study, dash line indicated probably be disease causing mutation).

the SNRNP200 mutation spectrum and thus substantiate its contribution to RP.

The spliceosome for pre-mRNA splicing is a specialized RNA and dynamic protein complex. The exons ligation relies on spliceosome splicing out the introns from a transcribed pre-mRNA segment.³⁹ The major spliceosome composes of four small nuclear RNA proteins, called snRNPs, U1, U2, U4/U6, and U5, and a range of non-snRNP splicing factors. The individual snRNP particles participate in the splicing cycle in a highly dynamic manner. Lauber *et al.*²⁹ demonstrated that human SNRNP200 encoded a U5-specific 220 kDa protein that is homologous to yeast SNU246 encoding

a U5-specific protein Brr2, while both of these two proteins contain two conserved domains being characteristic of the DEXH-box protein family of the putative RNA helicases and RNA-stimulated ATPases. They also showed disruption of SNU246 in yeast is lethal and leads to a splicing defect *in vivo*.²⁹ Brr2 is a DExD/H-box helicase responsible for U4/U6 unwinding during spliceosomal activation. The two reported mutations, p.R1090L and p.S1087L, are well conserved in many species. The yeast analog of these two mutations (p.R1107L and p.N1104L) were shown to compromise RNA unwinding in budding yeast.²¹ These indicate that the two mutations in the U5-specific 220 kDa

Table 2 Carriers and pathogenic potentials of the four novel *SNRNP200* non-synonymous variants

Variants	Exon	Case		Control	Descriptions	PolyPhen		PolyPhen 2		SIFT		SIFT BLink		PMUT	
		Type	Sex/Age			Prediction	PSIC score	Prediction	HumanVar score	Prediction	Score	Prediction	Score	Prediction	RI
c.1504T>C (p.C502R)	12	sRP	F/71	0/0/178	Novel	Probably damaging	4.14	Probably damaging	1.00	Damaging	0	Affected Protein Function	0	Pathologic	4
c.2092A>G (p.I698V)	16	—	NA	0/0/178	Novel	Benign	0.74	Benign	0.00	Tolerated	0.12	Tolerated	0.23	Neutral	9
c.5356G>A (p.R1779H)	38	sRP	F/53	0/0/178	Novel	Benign	0.27	Probably damaging	0.95	Damaging	0.04	Affected Protein Function	0.01	Neutral	0
c.5983G>A (p.A1995T)	42	—	M/62	0/1/178	Novel	Benign	0.20	Benign	0.00	Tolerated	0.14	Tolerated	0.31	Neutral	5

Abbreviations: Age, age (years) at recruitment; PSIC, position-specific independent counts, a scoring system used in PolyPhen; RI, reliability index; sRP, simplex retinitis pigmentosa; Type, type of RP.

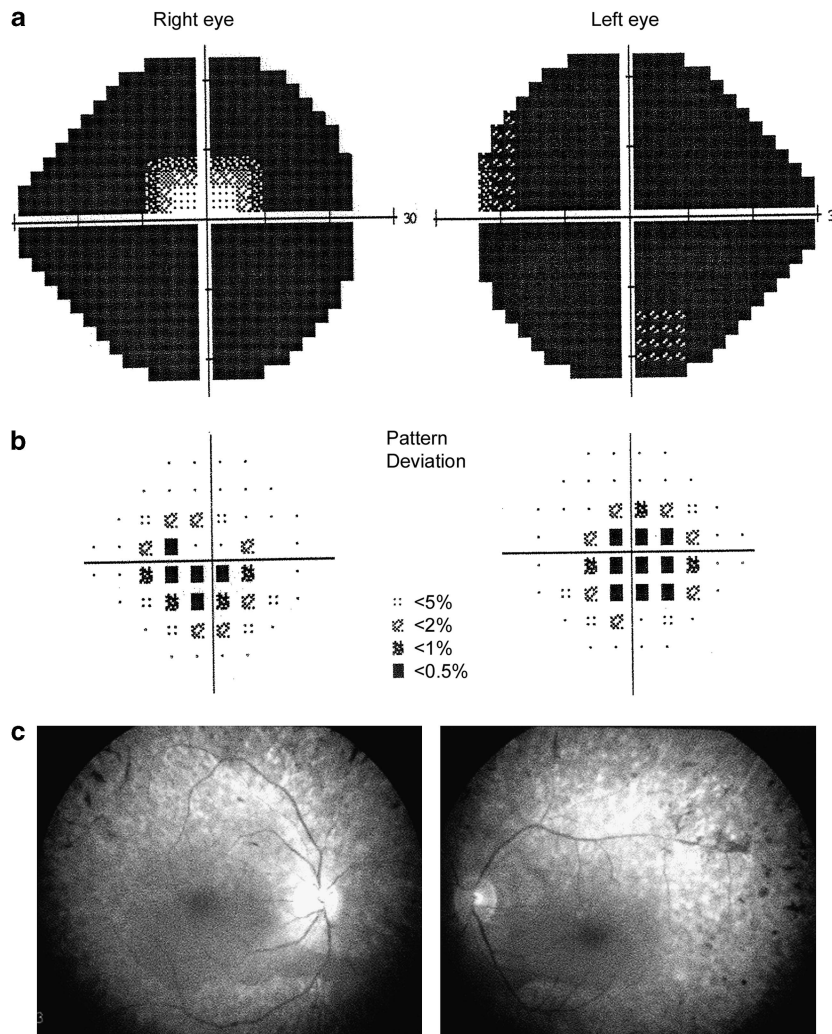


Figure 3 The visual field tests (a and b) and fundus photos (c) of the p.C502R carrier during the last visit with the typical presentations of retinitis pigmentosa.

protein may lead to pathogenesis through impaired RNA unwinding and thus splicing. Notably all five reported mutations are located within hel308-I. The mutations p.R681H, p.R681C, and p.Y689C are located in the boundaries between the helicase ATP-binding and helicase CTR domains, whereas p.S1087L and p.R1089L are located within the sec-63 domain. The mutation p.C502R detected in this study is located in the helicase ATP-binding domain. Hel308-I has the highest sequence conservation among different species, suggesting a critical role in the helicase activity. It has a direct role in the unwinding of the U4/U6 helix.⁴⁰ Structural defect of Hel308-I may impair helicase/ATPase activity, leading to defects in spliceosome catalysis. In contrast to the unwinding function of Hel308-I, the Hel308-II module of Brr2 does not have ATPase activity and is also unlikely to have helicase activity.⁴⁰ This may explain the lack of mutation reported on Hel308-II in RP, as the structural defect may not directly affect the spliceosome. Hel308-II has been demonstrated to interact with Prp8 and Snu114 both *in vitro* and *in vivo*. Further, the Prp8-CTR facilitates the binding of the Brr2/Prp8-CTR complex to U4/U6 in yeast.⁴¹ In this current study, we have identified a potentially pathogenic mutation, p.R1779H, which is located between the second helicase CTR and sec-63 domains. It is the first candidate mutation found in the hel308-II module for RP. Although it does not disrupt the helicase directly, it may have a role in mediating the regulation of Brr2 activity or the interaction with other spliceosomal proteins.

To date, the RNA intron-splicing factors have only been implicated in adRP.^{21,32,42–44} Although it is clear that the retina requires a relatively high level of RNA splicing activity for optimal tissue-specific physiological function,⁴⁵ how the defects in this essential macromolecular complex transform into a photoreceptor-specific phenotype is unknown. The underlying reason for the RNA intron-splicing factors to be related to adRP is also unclear. In our current study, two possible mutations were found in sRP patients, suggesting that *SNRNP200* may also have a role in other forms of RP. However, as the family members of the two patients were not available for clinical and genetic investigations and these two mutations were heterozygous, it is unclear whether they represent *de novo* mutations or dominant mutations with reduced penetrance.

In our Chinese study subjects, we did not detect the *SNRNP200* mutations that were previously found for adRP,^{21,22,32} suggesting that the mutation spectra of *SNRNP200* vary across different populations or even within the same ethnic group. Apart from showing the contribution of *SNRNP200* to a proportion of overall RP, our results revealed a highly mutable *SNRNP200* in RP. Even in a relatively small number of adRP patients in our

cohort from southern China, we detected a large number of *SNRNP200* variants that are either potentially causal or associated with RP. These were found within the 24 exons containing known mutable regions. Therefore, screening all the 45 coding exons of *SNRNP200* is worthwhile in order to reveal the significance of *SNRNP200* in RP. In addition, RP families should also be screened to ascertain segregation of *SNRNP200* mutations with RP. As our data indicate possible direct causation of specific *SNRNP200* variants on RP, functional analysis of potential disease-causing mutants should be warranted.

Summary

What was known before

- One of the RNA intron-splicing factors—the *SNRNP200* gene—has been reported to cause retinitis pigmentosa.
- All six reported mutations (p.R681C, p.R681H, p.Y689C, p.Q885E, p.S1087L, and p.R1090L) are all located in the Hel308 I molecule.
- All reported mutations were identified only in adRP patients.

What this study adds

- This is the first evaluation and relative prevalence of *SNRNP200* in a large series of southern Chinese patients rather than one RP family.
- We found that the profile differed from other reported populations on the hotspots of this gene and identified two novel candidate mutations (p.C502R and p.R1779H) in two RP patients.

Conflict of interest

The authors declare no conflict of interest.

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