Carrier frequency and incidence estimation of RPE65-associated inherited retinal diseases in East Asian population by population database-based analysis

Eun Hye Cho¹, Jong Eun Park²*, Taeheon Lee³, Kyeongsu Ha³ and Chang-Seok Ki³

Abstract

Background Inherited retinal diseases (IRDs) are clinically and genetically heterogeneous disorders leading to visual impairment and blindness. Because gene therapy for RPE65-associated IRDs was recently approved, it is necessary to predict the carrier frequency and prevalence for RPE65-associated IRDs. This study aimed to analyze the carrier frequency and expected incidence of RPE65-associated IRDs in East Asians and Koreans using exome data from the Genome Aggregation Database (gnomAD) and the Korean Reference Genome Database (KRGDB).

Methods We analyzed 9,197 exomes for East Asian populations from gnomAD comprising 1,909 Korean and 1,722 Korean genomes from KRGDB. All identified RPE65 variants were classified according to the 2015 American College of Medical Genetics and Genomics and the Association for Molecular Pathology guidelines.

Results The total carrier frequencies of East Asians and Koreans from both gnomAD and KRGDB were 0.10% (11/10,919) and 0.06% (2/3,631), respectively. The estimated incidence of RPE65-associated IRDs was 1/3,941,308 in East Asians and 1/13,184,161 in Koreans.

Conclusion This study identified carrier frequencies of RPE65-associated IRDs in East Asians and Koreans using gnomAD and KRGDB. We confirmed that the carrier frequency of RPE65-associated IRDs patients was low in Koreans among all East Asian populations, and the incidence was also predicted to be lower than in other East Asian populations. The variant spectrum of RPE65 gene in East Asian and Korean populations differed greatly from those of other ethnic groups.

Keywords RPE65-associated IRD, RPE65, gnomAD, KRGDB, Carrier frequency, East Asian
Background
Inherited retinal diseases (IRDs) are clinically and genetically heterogeneous disorders leading to visual impairment and blindness and affect approximately one in 2,000 individuals worldwide [1]. IRDs include several phenotypes, of which retinitis pigmentosa (RP) is the most common form and Leber congenital amaurosis (LCA) is the most severe form. To date, 280 genes have been discovered that are associated with IRDs [2].

*RPE65* encodes retinoid isomerohydrolase, which is responsible for converting all-trans-retinyl ester to 11-cis-retinol [3]. *RPE65* is associated with three IRD phenotypes, autosomal recessive LCA 2 (MIM 204,100), autosomal dominant RP 20 (MIM 613,794), and autosomal dominant RP 87 with choroidal involvement (MIM 618,697) and accounts for 2–16% of LCA and 1.02–2.7% of autosomal recessive RP [4].

On December 19, 2017, Voretigene neparvovec-rzyl (VN, LUXUTRNA, Spark Therapeutics, Philadelphia, PA, USA) was approved by the Food and Drug Administration (FDA) for the treatment of *RPE65*-associated IRDs, becoming the first FDA-approved gene therapy for a genetic disease [5]. In Korea, it was also approved by the Ministry of Food and Drug Safety of Korea in September 2021. Therefore, it is important to know the carrier frequency of *RPE65* to predict the incidence of *RPE65*-associated IRDs, which is useful information for evaluating the potential number of patients for VN therapy.

The Genome Aggregation Database (gnomAD) is a widely used genomic database worldwide, and gnomAD is consists of 125,748 exomes and 4,359 genomes [6]. The gnomAD contains exome data collected from 9,197 East Asians, including 1,909 Koreans. It is suitable for East Asian studies as it has the largest amount of data from East Asians among the public genomic databases. The Korean Reference Genome Database (KRGDB) is a large-scale variant database of whole genome sequencing data of 1,722 Koreans and contains single nucleotide and short insertion/deletion variants [7]. The *RPE65* gene variant was interpreted according to the 2015 American College of Medical Genetics and Genomics and Association for Molecular Pathology guidelines (2015 ACMG/AMP guidelines), which has been widely adopted in clinical practice [8]. In this study, we analyze the carrier frequency of *RPE65* and expected incidence of *RPE65*-associated IRDs in East Asian populations using the population database from the gnomAD and KRGDB through 2015 ACMG/AMP guidelines.

Methods
East asian population data
Three representative population databases were used to analyze the carrier frequency of the *RPE65* gene in East Asians. The gnomAD data (v2.1.1) for the *RPE65* gene was obtained from gnomAD (https://gnomad.broadinstitute.org/, accessed on 13 September 2021). We analyzed 9,197 East Asian exomes of which 1,909 were from Koreans, 76 were from Japanese, and 7,212 were from other East Asian populations. The filtered variants that were flagged in gnomAD as failing ‘InbreedingCoeff’, ‘AC0’, or ‘RF’ QC filters were excluded from the analysis. The KRGDB was used as the Korean database, and the database consists of 1,722 Koreans (http://coda.nih.go.kr/coda/KRGDB/index.jsp, accessed on 25 September 2021).

Variant classification and statistical analysis
All variants were interpreted using the 2015 ACMG/AMP guidelines and Sequence Variant Interpretation (SVI) general recommendations for ACMG/AMP criteria by ClinGen (https://clinicalgenome.org/working-groups/sequence-variant-interpretation/, accessed on 1 November 2021). The 2015 ACMG/AMP guidelines recommend the classification of variants into five categories: pathogenic variants (PV), likely pathogenic variants (LPV), variants of uncertain significance, likely benign variants, and benign variants. REVEL [9] and SpliceAI [10] were used to predict variant pathogenicity. All variants identified in gnomAD were additionally classified according to the Human Gene Mutation Database (HGMD) and ClinVar. The HGMD professional database (http://www.hgmd.org/, release 2021.04) is a comprehensive collection of germline variants categorized into six categories. ClinVar (https://www.ncbi.nlm.nih.gov/clinvar/, accessed on 5 February 2022) is a freely available archive that provides the classification of variants interpreted by clinical laboratories.

*RPE65* carrier frequency and incidence estimation
East Asian and Korean carrier frequencies were calculated for *RPE65* gene using gnomAD and KRGDB. We used those classified as PV and LPV according to the 2015 ACMG/AMP guidelines, the disease-causing variant (DM) in HGMD and those classified as PV and LPV in ClinVar for carrier frequency analysis. Thereafter, we estimated the incidence of *RPE65*-associated IRDs based on frequency and the Hardy–Weinberg equilibrium principle \(1 = p^2 + 2pq + q^2\). The Hardy-Weinberg equilibrium states that allele frequencies in a population remain constant from generation to generation in the absence of disturbance including mutation, natural selection, nonrandom mating, genetic drift, and gene flow [11]. The detailed prediction method was described in a previous study [12]. MedCalc ver. 11.5.1.0 (MedCalc Software, Maakerke, Belgium) was used for statistical analysis, and 95% confidence intervals (CIs) were calculated for each value.
Results
In total, 9,197 East Asian exomes including 1,909 Korean exomes from gnomAD and 1,722 Korean genomes from KRGDB were analyzed for \textit{RPE65} gene variants. The classification of these variants according to the 2015 ACMG/AMP guidelines, HGMD, ClinVar is summarized in Table 1. A total of nine variants were classified as PV/LPV according to the 2015 ACMG/AMP guidelines. Total carrier frequencies in East Asians and Koreans from gnomAD and KRGDB were 0.10% (11/10,919) and 0.06% (2/3,631), respectively. The estimated incidence of \textit{RPE65}-associated IRDs was 1/3,941,308 in East Asians and 1/13,184,161 in Koreans. According to the ClinVar, carrier frequency of East Asian and Korean was 0.08% and 0.03%, respectively. The estimated incidence of \textit{RPE65}-associated IRDs was 1/5,887,633 in East Asians and 1/18,510,805 in Koreans. According to the HGMD, carrier frequencies of East Asians and Koreans were 1.24% and 0.19%, respectively. The estimated incidence of \textit{RPE65}-associated IRDs was 1/26,167 in East Asians and 1/1,076,258 in Koreans.

\textit{RPE65} PVs/LPVs found in East Asians and Koreans are summarized in Table 2. Of the nine variants classified as PV/LPV identified in East Asians, only two variants were identified in Koreans: c.858+1G>T and c.335G>A (p.Cys112Tyr). When comparing the PVs/LPVs found in East Asians and Koreans with other ethnicities, PVs/LPVs identified in East Asians and Koreans were not found in Ashkenazi Jewish, European (Finnish), and other populations.

Discussion
In this study, the carrier frequency and estimated incidence of \textit{RPE65}-associated IRDs were analyzed for East Asians and Koreans using gnomAD and KRGDB. The carrier frequency of East Asians was 0.10% and of Koreans was 0.06%. Based on disease classification databases, HGMD and ClinVar, carrier frequencies were 0.08–1.24% in East Asians and 0.03–0.19% in Koreans. According to the previous study of Hanany et al., carrier frequency of IRDs in East Asians was about 40%, the second highest after Europeans, and the estimated genetic prevalence of IRDs was the highest in East Asians at 1/1,003 [13]. \textit{EYS} and \textit{USH2A} were the most prevalent genes in East Asians with carrier frequencies of 2.52% and 2.48%, respectively. On the other hand, \textit{RPE65}
| Nucleotide change | Amino acid change | KRGDB allele frequency (n = 1,722) | gnomAD | REVEL 2015 | 2015 ACMG/AMP (criteria) | Reference |
|------------------|------------------|-----------------------------------|--------|------------|--------------------------|-----------|
| c.1451G > A     | p. Gly484Asp     | 0                                 | 0      | 5.45E-05   | 0                        | 0.984     |
| c.1399 C > G    | p. Pro467Ala     | 0                                 | 0      | 1.09E-04   | 0                        | 0.8909    |
| c.858 + 1del    | p.?              | 0                                 | 0      | 5.44E-05   | 0                        | -         |
| c.858 + 1G > T  | p.?              | 4.55E-04                          | 0      | 0          | 0                        | -         |
| c.545 A > G     | p. His182Arg     | 0                                 | 0      | 5.44E-05   | 0                        | 0.8999    |
| c.335G > A      | p. Cys112Tyr     | 2.62E-04                          | 0      | 5.44E-05   | 0                        | 0.9169    |
| c.272G > A      | p.Arg91Gln       | 0                                 | 0      | 5.44E-05   | 3.70E-04                 | 0.6589    |
| c.271 G > T     | p.Arg91Trp       | 0                                 | 0      | 1.09E-04   | 3.70E-04                 | 0.8519    |
| c.200T > G      | p.Leu67Arg       | 0                                 | 0      | 5.44E-05   | 0                        | 0.98      |

gnomAD: Genome Aggregation Database, KRGDB: Korean Reference Genome Database, 2015 ACMG/AMP: 2015 American College of Medical Genetics and Genomics and Association for Molecular Pathology guidelines, LPV: likely pathogenic variant
is rare among East Asians with a carrier frequency of 0.20% [13]. The carrier frequency of RPE65 was different in each continental population, which was highest in Africans (0.55%), followed by non-Finnish Europeans (0.28%), Latinos (0.27%), South Asians (0.20%) and Finnish (0.12%) [13]. In the present study, carrier frequency in East Asians was even lower at 0.10%. In addition, there might be difference in sub-continental population. In Koreans, the carrier frequency of RPE65 was 0.05% in gnomAD Korean and 0.06% in KRGDB, which were lower than those of East Asians.

The carrier frequency was different according to the variant interpretation methods. The carrier frequency of 0.20% in the previous study by Hanany et al. was based on variant classification that used parameters including allele frequency, segregation analysis, biochemical analyses, presence of the variant in patients vs. controls, and biallelically vs. monoallelically [13]. In the present study, the carrier frequency based on HGMD was highest, and the carrier frequency based on 2015 ACMG/AMP guidelines and ClinVar was similar. According to the previous study of Park et al., the variants classified as DM in HGMD showed a 91.62% concordance rate compared to PV/LPV by 2015 ACMG/AMP guidelines [14]. The accuracy of variant classification in ClinVar gradually improved gradually and has been influenced by review status, represented as the number of stars [15]. Because the higher number of stars indicates a higher the concordance rate, the reliability of the classification in ClinVar can be evaluated as the number of stars. To prevent overestimation of carrier frequency, variant classification in HGMD and ClinVar should be accepted with caution. Although the 2015 ACMG/AMP guidelines were not perfect, it is considered as the standard guideline. Therefore, the carrier frequency in the present study was considered accurate to estimate the incidence of RPE65-associated IRDs.

According to data from the Korean Statistical Information Service (http://kosis.kr/; accessed on 10 June 2022) in 2020, the total population of Korea was 51.8 million with 272,337 births. Based on the Korean carrier frequency identified in this study, the number of carriers is estimated to be about 31,000 in total and 163 in newborns per year. The estimated incidence of RPE65-associated IRDs in Korea based on the Hardy-Weinberg equilibrium is approximately 0.02 cases per year. Therefore, the number of patients eligible for VN was expected to be very low in Korea.

According to the Leiden open variation database (LOVD), the most common PVs/LPVs in RPE65 were c.271 C>T, c.1102T>G, and c.11+5G>A, which accounts for 26.7% of PVs/LPVs identified in RPE65 [16]. Among them, only c.271 C>T was present in the East Asian populations. c.1102T>C is known as founder mutation in the Dutch populations but is absent in the East Asian populations [16, 17]. c.271 C>T was the most common PVs/LPVs in gnomAD East Asian, which was consistent with the findings of LOVD, but it was absent in gnomAD Korean and KRGDB. c.271 C>T was frequently identified in Saudi Arabia and Tunisia [16] and was identified in one Korean patient with LCA [18]. Only one PV/LPV was present in gnomAD Korean and in KRGDB: c.335G>A in gnomAD Korean and c.858+1G>T in KRGDB. c.335G>A was identified as homozygous in a Chinese patient with autosomal recessive RP [19]. c.858+1G>T was identified as homozygous in two Indian families with LCA [20, 21] and in one Korean LCA patient as compound heterozygous with c.271 C>T [18]. Based on these findings, it was found that the variant spectrum of RPE65 varies in different ethnicities, and Koreans showed a variant spectrum similar to that of the overall Asian populations.

Genotype-phenotype correlation analysis identified that truncating variants were associated with severe phenotype and early disease onset [22, 23]. Most of the PVs/LPVs identified in the East Asian database consisted of missense variant, and splicing variants accounted for only a small portion. In addition, the most common variants were missense variants: c.271 C>T and c.1399 C>G. Therefore, the variant spectrum in the patients of RPE65-associated IRDs was expected to consist mostly of missense variants. This was consistent with LOVD, in which the most common variant is the missense variant, c.271 C>T. Indeed, previous studies showed that missense variants were most frequent in patients of RPE65-associated IRDs regardless of ethnicity [22, 23].

Since a treatment has been approved for RPE65-associated IRDS, identification of RPE65 patients has become important. Recently, the RPE65 gene was added in ACMG secondary findings v3.0 because gene therapy treatment that may be more efficacious earlier in disease progression of patients is helpful in early diagnosis [24]. Therefore, it is meaningful to predict the carrier and prevalence of RPE65-associated IRDs in East Asians and Koreans through this study.

This study has some limitations. First, we did not analyze structural variations such as large deletion/insertion of the RPE65 gene. Currently, only two cases of large deletion of RPE65 have been reported [25, 26]. Second, we analyzed two separate population databases, gnomAD and KRGDB, of which the strategies for variants calling and filtering can differ. It might affect the carrier frequency calculated using each database. Third, although rare, RPE65 was associated with autosomal dominant RP 87 with choroidal involvement (RP87), which was characterized with incomplete penetrance and variable expressivity [27, 28]. Therefore, the carrier frequency estimated in this study might include patients with RP87 as well as
carriers of RPE65-associated IRD. To date, only one causative variant, c.1430 A>G, was identified to cause RP87 [27, 29], which was not identified in this study. Nonetheless, this study has several advantages. This is the largest study among those performed in East Asia that analyzed the entire RPE65 gene. To the best of our knowledge, there are no large-scale population studies of carrier frequency and estimated RPE65-associated IRDs incidence in Koreans. We believe that this study more accurately predicts the carrier frequency of RPE65-associated IRDs in East Asia and Korea.

**Conclusion**
This study identified the carrier frequencies in East Asians and Koreans using gnomAD and KRGDB. We confirmed that the carrier frequency of RPE65-associated IRDs patients was low in Koreans among East Asians, and the incidence was also predicted to be lower than in other East Asian populations. The variant spectrum of RPE65 genes in East Asian and Korean populations differed greatly from those of other ethnic groups. Our data are expected to serve as a reference for further investigations of RPE65-associated IRDs in East Asian and Korean populations.

**Abbreviations**

2015 ACMG/AMP guidelines, 2015 American College of Medical Genetics and Genomics and the Association for Molecular Pathology guideline

ACMG American College of Medical Genetics

CIs Confidence intervals

DM Disease-causing variant

gnomAD Genome Aggregation Database

HGMD Human Gene Mutation Database

IRD Inherited retinal diseases

KRGDB Korean Reference Genome Database

LCA Leber congenital amaurosis

LOVD Leiden open variation database

LPV Likely pathogenic variant

PV Pathogenic variant

PR Retinitis pigmentosa

VN Voretigene neaparvovec-czyl

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**Authors’ contributions**

EHC, JEP participated in analysis and interpretation of the data and the drafting of the manuscript. TL and KH participated in acquisition and analysis of data. CSK and JEP participated in the study concept and design, and drafting of the manuscript and for important intellectual content.

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**Data availability**

All data are available by corresponding author upon reasonable request.

**Declarations**

**Ethics approval and consent to participate**
No ethical approval was required.

**Consent for publication**
Not applicable.

**Competing interests**
The authors have no competing interests to declare.

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