Statistical driver genes as a means to uncover missing heritability for age-related macular degeneration

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Abstract

Background: Age-related macular degeneration (AMD) is a progressive retinal disease contributing to blindness worldwide. Multiple estimates for AMD heritability ($h^2$) exist; however, a substantial proportion of $h^2$ is not attributable to known genomic loci. The International AMD Genomics Consortium (IAMDGC) gathered the largest dataset of advanced AMD (ADV) cases and controls available and identified 34 loci containing 52 independent risk variants defining known AMD $h^2$. To better define AMD heterogeneity, we used Pathway Analysis by Randomization Incorporating Structure (PARIS) on the IAMDGC data and identified 8 statistical driver genes (SDGs), including 2 novel SDGs not discovered by the IAMDGC. We chose to further investigate these pathway-based risk genes and determine their contribution to ADV $h^2$, as well as the differential ADV subtype $h^2$.

Methods: We performed genomic-relatedness-based restricted maximum-likelihood (GREML) analyses on ADV, geographic atrophy (GA), and choroidal neovascularization (CNV) subtypes to investigate the $h^2$ of genotyped variants on the full DNA array chip, 34 risk loci ($n = 2758$ common variants), 52 variants from the IAMDGC 2016 GWAS, and the 8 SDGs, specifically the novel 2 SDGs, PPARD and PLCG2.

Results: Via GREML, full chip $h^2$ was 44.05% for ADV, 46.37% for GA, and 62.03% for CNV. The lead 52 variants' $h^2$ (ADV: 14.52%, GA: 8.02%, CNV: 13.62%) and 34 loci $h^2$ (ADV: 13.73%, GA: 8.81%, CNV: 12.89%) indicate that known variants contribute ~14% to ADV $h^2$, as well as the differential ADV subtype $h^2$.

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Background

Genome-wide association studies (GWAS) have been instrumental in identifying genomic variants associated with complex traits for over 10 years [1]. GWAS detect such associations by comparing allele frequencies in individuals with and without a trait of interest in a specific population [2]. These methods have been successfully applied to find large numbers of disease-associated variants that contribute to the trait’s heritability [3]. Heritability is defined as the fraction of phenotypic variance explained by genetic variation in the context of a specific range of environmental variation [3]. Broad-sense heritability ($H^2$) is the proportion of phenotypic variance that includes dominance and epistasis; whereas, narrow-sense heritability ($h^2$) is the proportion of phenotypic variation of additive genetic effects. Common variants may capture up to about two-thirds of narrow-sense heritability for age-related macular degeneration (AMD), but despite this, much of AMD heritability is still unexplained by known variants [4]. The topic of missing heritability has been discussed, especially regarding complex diseases, and may be attributable to non-additive effects of genomic variants that are not discernible from traditional GWAS [3].

GWAS have been remarkably successful for identifying genomic loci contributing substantially to AMD risk. This progressive, adult-onset condition is among the leading causes of blindness in the world in individuals over 60 and is expected to become a significant health burden as the aging population increases in size [5]. AMD leads to the decline of central vision in patients as a result of lipid deposits (drusen), photoreceptor loss, and inflammation in the macula [6, 7]. It is clinically characterized into multiple subtypes: early, intermediate, or advanced AMD stages based on disease severity. Advanced AMD (ADV) is further sub-categorized into geographic atrophy (GA) or choroidal neovascularization (CNV).

The International AMD Genomics Consortium (IAMDGC) performed the largest case-control GWAS to date for ADV and identified 52 independent common and rare variants from 34 susceptibility loci, the highest risk loci being $CFH$ and $ARMS2/HTRA1$ genes [8]. These 52 genomic variants contribute to about half of the genomic heritability for ADV, which leaves nearly half of ADV heritability unexplained [8].

In contrast to traditional case-control GWAS, in silico pathway analyses of GWAS summary statistics identify biological pathways, which are defined by interactions of genes for a common biological function, harboring excesses of genomic variants that may be associated with a trait [9, 10]. They accomplish this by grouping variants into features that are then merged into pathways based on curated information in publicly available pathway databases [9, 10]. Because pathway analyses focus broadly on the collection of nominal genetic variants in biological pathways, they are not limited to assessing additive effects of individual variants on the trait and may be leveraged to identify genetic variance with non-additive effects. Ultimately, these analyses provide insights into trait-associated biological processes and suggest which genes are most pertinent for these pathway-level associations [11, 12]. However, they do not estimate the contribution of genetic variants in these genes and pathways to the trait’s heritability [13].

To uncover genomic loci undetectable by GWAS, we performed in silico, knowledge-driven pathway analyses of the summary statistics from the IAMDGC 2016 GWAS [14] using Pathway Analysis by Randomization Incorporating Structure (PARIS) [11, 12]. In our comprehensive approach, we utilized multiple pathway databases to determine which genes were consistently contributing to significant pathway signals for ADV [14]. We identified eight statistical driver genes (SDGs) that were significantly contributing to the significant AMD-associated pathways from PARIS: $C2$, $C3$, $LIPC$, $MICA$, $NOTCH4$, $PLCG2$, $PPARA$, and $RADS1B$. Of these eight SDGs, two genes ($PLCG2$ and $PPARA$) fell outside of the 34 AMD susceptibility loci identified by the IAMDGC GWAS [8]; we showed that these loci may be associated with ADV risk [14].

While the 2016 IAMDGC GWAS uncovered several AMD loci that explain a large portion of AMD heritability [8], their study did not investigate potential non-additive effects of AMD risk genes. Pathway analyses of GWAS data consider known biological relationships among genes in a pathway; therefore, we were able to identify two novel AMD genes ($PLCG2$ and $PPARA$) that were not found in the IAMDGC GWAS. To further examine the potential...
role of the 2 novel SDGs, we calculated the proportion of ADV variance explained by (i) common variants in PPARA and PLCG2, (ii) the 8 SDGs identified by pathway analysis, and (iii) known and novel AMD loci identified by the IAMDGC. We also applied this approach to the subtypes of ADV (GA and CNV) to elucidate whether these variants contribute more to the heritability of AMD in a particular subtype of ADV. We further interrogated the possible epistatic interactions among lead variants in the known AMD genes as well as our novel SDGs to elucidate if their contributions to AMD heritability could be attributable to non-additive effects.

**Methods**

**Statistical driver genes for advanced AMD**

We performed in silico pathway analyses using Pathway Analysis by Randomization Incorporating Structure (PARIS, v.2.4) [12] on the largest available ADV case-control GWAS results from the IAMDGC [8]. This included summary statistics for 445,115 directly genotyped variants on 16,144 advanced AMD cases and 17,832 controls [8]. The ADV cases include GA-specific cases (n = 3235), CNV-specific cases (n = 10,749), and individuals with both GA and CNV (n = 2160) [8]. Samples were genotyped with the Illumina HumanCoreExome Array as previously described [8] and are accessible through the database of Genotypes and Phenotypes (dbGAP; Accession: phs001039.v1.p1). Our knowledge-driven pathway analyses utilized three pathway databases (KEGG, Reactome, and GO) and led to the discovery of eight statistical driver genes for ADV (Table 1) [14]. Statistical driver genes (SDGs) were defined as genes that were strongly contributing (gene-level p < 0.0001) to the statistical signal of the significant pathways (pathway-level p < 0.0001) identified by PARIS. Two of these SDGs (PPARA and PLCG2) remained significant following the exclusion of the 34 known AMD loci identified by the IAMDGC from the pathway analysis because they fall outside of the known loci boundaries.

**Variant selection and genotype extraction**

For our heritability estimates, we extracted genotypes for variants in one of these seven variant criteria subsets (Table 3):

- IAMDGC Chip: Variants that were directly genotyped by the IAMDGC on the Illumina HumanCoreExome chip with custom content as previously described [8]
- 8 SDGs ±50 kb: Variants in or within 50 kilobasepairs (kb) of the eight SDGs (C2, C3, LIPC, MICA, NOTCH4, PLCG2, PPARA, and RAD51B) [14]
- 2 Novel SDGs ±50 kb: Variants in or within 50 kb of the PLCG2 and PPARA genes [14]
- Lead IAMDGC Variants: Variants that were identified as one of the 52 lead variants from the IAMDGC 2016 GWAS [8]
- 34 AMD Loci: Variants in the 34 susceptibility loci identified by the IAMDGC 2016 GWAS (Supplementary Table 5 in [8])
- 34 AMD Loci and 2 Novel SDGs ±50 kb: Variants that occur in the 34 susceptibility loci identified by the IAMDGC 2016 GWAS (Supplementary Table 5 in [8]) and variants that fall in or within 50 kb of the PLCG2 and PPARA genes [14]
- 34 AMD Loci and 8 SDGs ±50 kb: Variants in the 34 susceptibility loci identified by the IAMDGC 2016 GWAS (Supplementary Table 5 in [8]) and variants that fall in or within 50 kb of the eight SDGs [14]

Gene and loci boundaries were based on build GRCh37 of the human genome. Using PLINK v1.90 beta [15, 16], we filtered the variants from the 34 loci and SDGs based on minor allele frequency (MAF) and genotyping call rate to exclude variants that had MAF < 0.01 and missing genotype rate > 0.01. The variants from the IAMDGC chip and the 52 lead variants were not filtered for MAF or call rate; therefore, these two variant sets include common and rare variants. Penetrance was not taken into account for the SDGs [14] because their association was based on summary statistics from the IAMDGC 2016 GWAS. All variant sets were extracted from the ADV case-control data, GA-specific case-control data (GA), and CNV-specific case-control data (CNV) (Table 2).

**Estimation of AMD heritability with GCTA GREML**

Genetic relationship matrices (GRMs) were constructed using Genome-wide Complex Trait Analysis

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**Table 1**

| Gene   | Chromosome | Full Gene Name               |
|--------|------------|------------------------------|
| C2     | 6          | complement C2               |
| MICA   | 6          | MHC class I polypeptide-related sequence A |
| NOTCH4 | 6          | notch receptor 4            |
| RAD51B | 14         | RAD51 paralog B             |
| LIPC   | 15         | lipase C, hepatic type      |
| PLCG2  | 16         | phospholipase C gamma 2     |
| C3     | 19         | complement C3               |
| PPARA  | 22         | peroxisome proliferator activated receptor alpha |

The eight statistical driver genes were identified with pathway analysis using PARIS and multiple biological pathway databases [14]. Two of these genes (PLCG2 and PPARA) were not previously identified as a part of the IAMDGC susceptibility loci and are noted in bold. Full gene names are based on the HUGO Gene Nomenclature Committee (HGNC).
heritability \((h^2)\) for each category of late AMD disease states (ADV, GA, and CNV), and for each subset of variants we selected (Tables 2 and 3, respectively). This included variants within PPARA and PLCG2, variants within 8 previously identified SDGs, the 34 loci, the 52 lead IAMDGC variants, and the full directly genotyped IAMDGC chip. To obtain invertible and reliable GRMs, variants were filtered by minor allele frequency (MAF < 0.01) and missingness (missing genotype rate > 0.01) when constructing the GRMs for the 34 AMD loci and the SDGs. Without this filtering step, GCTA is unable to create functional (invertible) GRMs for subsequent restricted maximum-likelihood (REML) analyses, which estimate the proportion of phenotypic variance attributable to additive genetic variance [17].

We used the genomic-relatedness-based restricted maximum-likelihood (GREML) approach in the GCTA v1.91.3beta software to estimate narrow-sense heritability \((h^2)\) for each of the three advanced AMD datasets (ADV, GA, and CNV) and for each of the 7 variant sets. The GREML approach does not measure dominance variance or epistatic interactions. We also estimated heritability adjusting for age information available from the IAMDGC (age at diagnosis for cases and age at exam for controls) and ten principal components (PC) in the GREML analysis of the full chip. The ten PCs were calculated from genome-wide chip data using PLINK v1.90 beta [15, 16], which is a computational tool for analyzing large-scale association studies.

We aimed to determine the proportion of ADV, GA- and CNV-specific heritability explained by variants in and within 50 kilobasepairs (kb) of the SDGs. In addition, we performed pairwise linkage disequilibrium (LD) analyses using two computational tools: LDMatrix [18] and SNiPA [19]. Variant pairs with \(r^2 > 0.7\) were determined to be the same signal and not considered in downstream assessments. We further assessed whether or not each variant in the pair was considered part of an SDG, one of the AMD loci, or overlapped between the two groups of variants.

**Epistatic interaction analyses**

To investigate heritability attributable to non-additive interactions among the lead AMD variants from the IAMDGC GWAS [8] and the novel SDGs [14], we performed pairwise logistic regression-based epistasis analyses using PLINK v1.90 beta [15, 16]. Analyses were performed over the full set of variants in or within 50 kb of the 2 SDGs and the 52 lead variants from the 2016 IAMDGC GWAS [8]; therefore, the threshold for significance was set at \(2.31 \times 10^{-6}\) for multiple testing correction (Bonferroni correction for 21,675 tests). If both variants in an epistatic interaction were from the same gene/locus, we found their LD in the European population (Bonferroni correction for 21,675 tests). We also estimated \(h^2\) based on a population prevalence of ADV in individuals of European descent (0.5%, [5]), as the IAMDGC samples in this study were filtered to unrelated individuals of European ancestry [8]. We validated that the estimates we observed were not likely artifacts by re-performing our analyses of the ADV data and a random set of 1112 variants (equal to the number of common variants in the 8 SDGs) that met the same filtering criteria for common variants in the 8 SDGs. In addition, we performed the same analysis using random sets of 79 variants (equal to the number of variants in the 2 SDGs alone) that met the same filtering criteria for the common variants in the 2 SDGs to validate that the estimates we observed were not likely to be artifacts.

**Pairwise LD analysis of SDGs and 34 AMD loci**

To evaluate the linkage and independence of the variants in the 8 SDGs and the 34 AMD loci (Table 3), we examined pairwise linkage disequilibrium (LD) using two computational tools: LDMatrix [18] and SNiPA [19]. Variant pairs with \(r^2 > 0.7\) were determined to be the same signal and not considered in downstream assessments. We further assessed whether or not each variant in the pair was considered part of an SDG, one of the AMD loci, or overlapped between the two groups of variants.

**Results**

**Study data for ADV, GA, and CNV analyses**

We aimed to determine the proportion of ADV, GA-specific, and CNV-specific heritability explained by variants in and within 50 kilobasepairs (kb) of the SDGs identified by PARIS (Tables 2 and 3). We extracted 2173 variants from all 8 previously identified SDGs and 234

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**Table 2** Demographics of participants in the study data

| Data               | ADV | GA | CNV |
|--------------------|-----|----|-----|
| Total Samples      | 33,976 | 21,067 | 28,581 |
| Cases              | 16,144 | 3235 | 10,749 |
| Controls           | 17,832 | 17,832 | 17,832 |

Values represent counts of samples in the IAMDGC data

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**Table 3** Characteristics of the marker data extracted from the IAMDGC exome chip

| Data               | ADV | GA | CNV |
|--------------------|-----|----|-----|
| IAMDGC Chip        | 553,261 | 553,261 | 553,261 |
| 8 SDGs ±50 kb      | 1122 | 1122 | 1122 |
| 2 Novel SDGs ±50 kb| 79 | 79 | 79 |
| Lead IAMDGC Variants| 52 | 52 | 52 |
| 34 AMD Loci        | 2758 | 2758 | 2758 |
| 34 AMD Loci and 2 Novel SDGs ±50 kb| 2835 | 2992 | 2992 |
| 34 AMD Loci and 8 SDGs ±50 kb| 3351 | 4411 | 3351 |

Values represent counts of pruned variants based on MAF and call rate. The 8 SDGs are C2, C3, LIPC, MICA, NOTCH4, PLCG2, PPARA, and RAD51B. The 2 novel SDGs are PLCG2 and PPARA. The different numbers of variants for the expanded loci analyses were attributable to the different MAFs observed for each of the advanced AMD subtypes: ADV, GA, and CNV.
variants from the 2 novel SDGs (PPARA and PLCG2) based on their gene boundaries (Methods). However, we found that several of the variants in the SDGs either had very low MAF or had a low genotype call rate in the samples we analyzed and, therefore, they were removed prior to GRM creation (Table 3).

**Narrow-sense heritability explained by variants in SDGs for ADV, GA, and CNV**

To determine whether the SDGs contribute to the missing heritability of ADV or its subtypes (GA and CNV), we performed GREML analyses of variants from the SDGs. We found that the common, high call rate variants in the 8 SDGs was 3.76% (S.E. = 0.39) (Table 4). This was higher than the estimate observed for GA and comparable to CNV $h^2$ estimate (2.53 and 3.71%, respectively) (Table 4). The 2 SDGs contribute to 0.097, 0.12, and 0.18% ADV, GA, and CNV risk, respectively (Table 4).

To compare the SDG heritability estimates for ADV, GA, and CNV to those observed for known AMD loci, we performed GREML analyses of the common variants with high call rates within the 34 loci identified by the IAMDGC [8] for each disease subtype. We found that common variants from the known loci contribute to 13.73% (S.E. = 0.83) ADV risk, 8.81% (S.E. = 0.75) GA risk, and 12.89% (SE = 0.82) CNV risk (Table 4). The 52 lead variants identified by the IAMDGC alone explain 14.52% (S.E. = 2.48) of ADV risk. By comparison, the $h^2$ estimates for these variants were lower (8.02 and 13.62%) for GA and CNV, respectively.

Given the individual estimates for the SDGs and 34 loci, we performed GREML analyses on combinations of the SDGs and 34 loci for ADV, GA, and CNV. In all our analyses, we found that the $h^2$ estimates were very similar (Table 4). Together, the 34 loci and 8 SDGs contribute to 13.06, 8.78, and 12.59% $h^2$ for ADV, GA, and CNV, respectively (Table 4). The $h^2$ estimates derived from the 34 loci and 2 SDGs are comparable to these values (ADV: 13.51%, GA: 8.60%, and CNV: 12.20%) (Table 4). To interrogate existing linkage among the SDG variants and those in the 34 AMD loci, we performed pairwise linkage disequilibrium (LD) analyses of these variants. No pairs of variants in high LD ($r^2 > 0.7$) were found between any of the variants in the 8 SDGs and the expanded 34 loci. This indicates that the variants we compared did not have pre-existing LD outside of the known AMD loci and were independent of each other.

To replicate chip heritability calculations from the 2016 GWAS published by the IAMDGC, we performed GREML analyses of the full datasets (i.e. chip heritability) for ADV, GA, and CNV. The chip heritability for ADV was 44.05% (S.E. = 1.29) (Table 5). We achieved similar values (44.16% (S.E. = 1.29)) when we reperformed our analyses with the first 10 principal components (PCs) calculated for the full chip and ADV cases and controls. Chip heritability estimates were higher for GA and CNV (46.37 and 62.03%, respectively) than the heritability estimate for the combined ADV dataset (Table 5). These values were similar to the estimates calculated for GA and CNV chip heritability including PCs for those respective datasets (46.50 and 62.18%, respectively). Values decreased after incorporating age data available from the IAMDGC (age at diagnosis for cases and age at exam for controls) and the first 10 PCs into the GREML analyses for chip heritability of the ADV and its subtypes (Table 5).

To further verify that contributions to ADV heritability from the common variants in the 8 SDGs were unlikely due to chance, we selected randomized variants from the autosomal genome that met the same MAF criteria we had used before. We recreated the GRM for the ADV data and these variants and performed the same GREML analyses we had on the 8 SDGs. The 1122 randomized variants explain 1.76% (S.E. = 0.21), indicating that the ADV heritability estimate for the 8 SDG (3.76%) was not likely due to chance. In addition, we ran several

| Data                        | ADV Estimate (%) | S.E. (%) | GA Estimate (%) | S.E. (%) | CNV Estimate (%) | S.E. (%) |
|-----------------------------|------------------|----------|-----------------|----------|------------------|----------|
| IAMDGC Chip                 | 44.05            | 1.29     | 46.37           | 2.13     | 62.03            | 1.47     |
| 8 SDGs ± 50 kb              | 3.76             | 0.39     | 2.53            | 0.36     | 3.71             | 0.41     |
| 2 Novel SDGs ± 50 kb        | 0.097            | 0.048    | 0.12            | 0.076    | 0.18             | 0.076    |
| Lead IAMDGC Variants        | 14.52            | 2.48     | 8.02            | 1.50     | 13.62            | 2.36     |
| 34 AMD Loci                 | 13.73            | 0.83     | 8.81            | 0.75     | 12.89            | 0.82     |
| 34 AMD Loci and 2 Novel SDGs ± 50 kb | 13.51 | 0.81 | 8.60 | 0.70 | 12.20 | 0.74 |
| 34 AMD Loci and 8 SDGs ± 50 kb | 13.06 | 0.73 | 8.78 | 0.63 | 12.59 | 0.74 |

For the analyses including the SDGs and/or 34 loci, we pruned the variants based on minor allele frequency and genotype call rate in the samples. For the estimates calculated for the full IAMDGC chip and lead IAMDGC variants, we did not apply variant filtering. kb: kilobasepairs
Table 5 Chip heritability estimates calculated by GREML for advanced AMD (ADV), GA, and CNV using quantitative covariates

| Analyses                           | ADV Estimate (%) | ADV S.E. (%) | GA Estimate (%) | GA S.E. (%) | CNV Estimate (%) | CNV S.E. (%) |
|------------------------------------|------------------|-------------|-----------------|-------------|------------------|-------------|
| Full Chip (no covariates)          | 44.05            | 1.29        | 46.37           | 2.13        | 62.03            | 1.47        |
| With Age                           | 57.96            | 1.31        | 45.41           | 2.16        | 59.71            | 1.51        |
| With 10 PCs                        | 44.16            | 1.29        | 46.50           | 2.13        | 62.18            | 1.48        |
| With Age + 10 PCs                  | 42.06            | 1.31        | 41.24           | 1.97        | 60.21            | 1.51        |

These estimates were based on 15,656 advanced AMD (ADV) cases (2964 GA-specific cases and 10,340 CNV-specific cases) and 17,832 controls with age information in the IAMDGC data or with an age over 50. Covariates included age information available from the IAMDGC (age at diagnosis for cases and age at exam for controls) and 10 PCs calculated for each of the datasets

Epistasis analyses

To interrogate possible interactions among the 52 AMD variants identified by the IAMDGC and the variants in the novel SDGs (PPARA and PLCG2), we performed logistic regression-based epistasis analyses using PLINK [15, 16]. Although we did not identify any significant interactions, several modest epistatic interactions \((p < 0.001)\) were uncovered between known AMD variants and variants in PPARA and PLCG2 (Table 6). Interactions between PPARA and PLCG2 variants were also identified but did not reach the significance threshold correcting for multiple testing (Table 7).

Discussion

In this study, we estimated the proportion of ADV heritability attributable to the SDGs we previously identified by pathway analysis of the summary statistics from the IAMDGC 2016 GWAS [14]. This included common variants from the 8 SDGs that exhibited significant signals across significant pathways \((p < 0.0001)\) from KEGG, Reactome, and GO pathway databases and the 2 SDGs (PPARA and PLCG2) that fell outside of the known AMD loci identified by the IAMDGC in their recent GWAS. To compare our results with those obtained by the IAMDGC [8], we calculated heritability estimates for the whole DNA array chip, 34 AMD loci, 52 lead variants from the 34 loci, and combinations of the SDGs and the 34 loci. The estimates and 95% confidence intervals we calculated for ADV (41.52–46.58%) and GA (42.20–50.54%) chip heritability overlap with the 95% confidence intervals for chip heritability determined by the IAMDGC for ADV (44.5–48.8%, [8]) and GA (47.2–57.4%, [8]). By contrast, our estimate for CNV chip heritability (59.15–64.91%) was much higher than what was calculated by the IAMDGC (42.2–46.5%), [8]).

Our estimate for ADV heritability based on the 52 lead variants is lower than that observed for these variants by the IAMDGC (14.52% vs. 27.2%, respectively) [8]. This difference is likely due to the different methods used to estimate these values. The IAMDGC [8] calculated their estimates by a theoretical, population-based formula based on the log odds ratios, allele frequencies of the 52 variants and the assumed trait prevalence. This formula assumes that all markers are independent, and therefore that all contributions to genetic variance are additive [20]. The IAMDGC also assumed disease prevalence of 1, 5%, or 10% in their analyses [8]. The addition of the two lead variants of the PLCG2 and PPARA genes does

Table 6 Epistatic interactions from pairwise logistic regression-based epistasis testing between variants in or within 50 kb of the 2 novel SDGs (PPARA and PLCG2) and the 52 AMD-associated index variants from the 2016 IAMDGC GWAS

| AMD Locus from IAMDGC GWAS | AMD SDG Locus | Epistatic Interaction | OR | P |
|---------------------------|--------------|----------------------|----|---|
| Chr Location (bp) | rsID | Locus Name | Chr Location (bp) | rsID | Locus Name | P-values |
| 1 | 196,704,632 | rs10922109 | CFH | 22 | 46,681,141 | rs35883013 | PPARA | 1.55 | 0.00015 |
| 9 | 76,617,720 | rs10781182 | MIR6130/RORB | 16 | 81,932,165 | rs4889432 | PLCG2 | 1.09 | 0.00048 |
| 5 | 39,327,888 | rs62558361 | C9 | 16 | 81,821,531 | rs8043845 | PLCG2 | 0.66 | 0.00073 |
| 14 | 68,769,199 | rs777567 | RADS1B | 16 | 81,900,628 | rs4243218 | PLCG2 | 1.08 | 0.00079 |
| 6 | 3,255,581 | rs204993 | C2/C3/CFB | 22 | 46,685,754 | rs182313981 | PPARA | 0.63 | 0.00093 |
| 19 | 45,411,941 | rs429358 | APOE | 22 | 46,685,754 | rs182313981 | PPARA | 0.53 | 0.00099 |

Locations are given in base pairs (bp) based on build 37 of the human genome. None of these interactions are significant after correcting for multiple testing \((p < 2.31 \times 10^{-5})\). Chr: Chromosome, \(P\): p-values from the logistic regression test, distributed as \(\chi^2\) with 1 d.f
not increase our heritability estimates for ADV (14.54%). These results reinforce the notion that the variants in these SDGs in isolate are not significant but in aggregate contribute strongly to the statistical signals we previously observed for AMD-associated pathways. Their association with AMD is likely not additive but rather as a consequence of their interactions within AMD-associated pathways, demonstrating the benefit of using pathway analysis to identify genetic variance with non-additive effects.

Heritability estimates based on each of our variant sets varied based on the advanced AMD subtype analyzed (ADV, GA, or CNV). With the exception of the chip genotypes, the values estimated for ADV and CNV were much higher than those calculated for GA. This may be due to the lower sample size of GA cases in the known loci. Only a few variants were in LD ($r^2 > 0.7$), but these variants were only connected to one variant in an SDG locus. By contrast, in the analysis of the common variants from the 34 loci and 2 SDGs, the 34 loci themselves were not expanded despite the addition of the 2 SDGs.

The variance explained by genetic variants in genes from AMD-related pathway defined in the literature has been previously explored using GCTA [13]. The 19 then-known AMD associated variants explained 13.3% of AMD risk in general, and significant additional heritability was attributable to variants in inflammatory and complement pathways when accounting for the known risk variants [13]. Other pathways, including angiogenesis and apoptosis, did not significantly contribute to AMD heritability estimates [13]. By contrast, in our approach, pathways were identified via in silico pathway analysis of large-scale GWAS data with PARIS and multiple curated pathway databases. We then focused specifically on the SDGs that significantly contribute to AMD-associated pathways in our analysis. Gene expression profiles for the two novel SDGs (PLCG2 and PPARA) have been observed in retinas from AMD cases and controls [27]. In retinal tissue from unaffected individuals, PPARA is highly expressed; whereas, PLCG2 is weakly expressed [28, 29]. Neither of these genes were significantly expressed in an age-adjusted analysis of CNV retinas [27].

While we determined that common variants from the SDGs contribute to the ADV, GA, and CNV heritability, this study had several limitations. Even with our estimates, there is a substantial portion of ADV, GA, and CNV variance left unexplained by the loci interrogated in this study. Additional sources of heritability not examined in this study include rare variants, structural variants, further investigations into epistasis, and epigenetic effects. Seven rare variants were among the 52 independent, genome-wide significant markers identified by the IAMDGC in their recent GWAS [8]. In this study, we excluded rare variants (MAF < 1%) from our GREML
analyses for the known loci and the SDGs, removing about half of variants from the 8 SDG variants and about two-thirds of the variants from PPARA and PLCG2 alone. Rare variants were not excluded in our previous pathway analyses because PARIS does not take MAF into account when identifying associated pathways [11, 12]. Therefore, we were unable to consider the complete contributions of these variants to disease heritability. Additionally, the IAMDGC data repository we utilized in this study does not currently have further information on environmental exposures or behaviors of the study participants that may contribute to AMD risk, such as smoking status and diet. Therefore, these non-genetic factors could not be included in our study.

Given the non-additive nature of our heritability estimates for the combinations of known AMD loci and the SDGs for ADV and its subtypes, we hypothesize that the variants in these loci may be interacting with one another. Our epistasis analyses did not reveal any significant epistatic interactions among the known AMD variants and the variants in the 2 novel SDGs. However, this is only an initial look at the possible representation of epistatic interactions between these genes. We currently only examined epistatic effects directly between the common, high call rate variants in the 2 novel SDG and the 52 lead variants from the IAMDGC; therefore there could be additional epistatic interactions among other variants in PPARA and PLCG2 with the 34 AMD loci. In addition, large-scale genome-wide epistatic effects have not been explored. Our identification of modest interactions among these loci, including between the novel SDGs, suggests that there may be region-wide interactions that are individually too weak to discover using these analyses. Further studies should be performed to confirm potential epistatic interactions between variants in the known loci and the SDGs. Additionally, because we performed our analyses on the largest dataset of ADV cases and controls currently available, we are unable to replicate our findings with a comparable, independent dataset. As with many genetic studies, our study only included individuals of European descent. Additional work should be done to elucidate the contributions of the SDGs to AMD heritability in diverse populations because different populations may have different heritability estimates for ADV and its subtypes [30, 31].

Conclusions

Our study elucidated the contribution of pathway SDGs and known AMD loci to the heritability of ADV and its subtypes. Heritability estimates for particular ADV subtypes were previously uncharacterized. The SDGs we analyzed in this study were previously identified from pathway analyses utilizing multiple pathway databases. This more comprehensive approach uncovered an appreciable portion of ADV heritability that had not been previously characterized. While they do not demonstrate an additive amount of heritability to that estimated for the 34 AMD susceptibility loci identified by the IAMDGC, we suspect that this is due to interaction effects or the exclusion of rare variants from our analyses. It has been previously shown that additional AMD loci (RLBP1 and CLL11) can be identified by accounting for gene x age interaction effects [32]. We propose that identifying statistical driver genes from in silico pathway analyses of GWAS data may be a valid approach to recognizing patterns of heritability (including non-additive contributions) from large-scale genomic data that are undetectable by GWAS. We applied this approach to ADV and its subtypes, but it could be applied to uncover novel loci associated to other complex traits for which GWAS have been performed. Additionally, pathway analysis provides biological context for the loci in GWAS, which could aid in understanding the underlying mechanisms of traits and developing targeted treatments for diseases.

Abbreviations
dbGAP: database of Genotypes and Phenotypes; ADV: Advanced age-related macular degeneration; AMD: Age-related macular degeneration; CNV: Choroidal neovascularization; GA: Geographic atrophy; GCTA: Genomewide Complex Trait Analysis; GREML: Genomic-relatedness-based restricted maximum-likelihood; GRM: Genetic relationship matrix; GWAS: Genome-wide association study; h²: Narrow-sense heritability; h²w: Broad-sense heritability; IAMDGC: International Age-related Macular Degeneration Genomics Consortium; kb: Kilobasepairs; LD: Linkage disequilibrium; MAF: Minor allele frequency; PARIS: Pathway Analysis by Randomization Incorporating Structure; PC: Principal component; REMIL: Restricted maximum-likelihood; SDG: Statistical driver gene

Acknowledgements

The authors appreciate the contributions of the study participants. This work was completed with use of the High Performance Computing Resource in the Core Facility for Advanced Research Computing at Case Western Reserve University. The authors also acknowledge the contributions of the members of the International Age-related Macular Degeneration Genomics Consortium (IAMDGC): Lars G. Fritsche, Willmar IgI, Jessica N. Cooke Bailey, Felix Grassmann, Sebanti Sengupta, Jennifer L. Bragg-Gresham, Kathryn P. Burdon, Scott J. Hebbink, Cindy Wen, Mathias Gorski, Ivana K. Kim, David Cho, Donald Zack, Eric Souied, Hendrik P.N. Scholl, Elisa Bala, Kristine E. Lee, David J. Hunter, Rebecca J. Sardell, Paul Mitchell, Joanna E. Merriam, Valentina Cipriani, Joshua D. Hoffman, Tina Schick, Yara T.E. Lechanteur, Robyn H. Guymer, Matthew P. Johnson, Yingda Jiang, Chloe M. Stanton, Gabrielle H.S. Buitendijk, Xiaowei Zhan, Alan M. Kwong, Alexis Boleda, Matthew Brooks, Linn Gieser, Rinko Raknapriya, Kari E. Branham, Johanna R. Foerster, John R. Heckenlively, Mohammad I. Othman, Brendan J. Vote, Helena Hai Liang, Emmauelle Souzeau, Ian L. McAllister, Timothy Isaacs, Janette Hall, Stewart Lake, David A. Mackey, Ian J. Constable, Jamie E. Craig, Terrie E. Kitchner, Zhenglin Yang, Zhiguang Su, Hongrong Luo, Daniel Chen, Hong Ouyang, Ken Flagg, Danni Lin, Guanping Mao, Henry Ferreyra, Klaus Stark, Claudia N. von Strachwitz, Armin Wolf, Caroline Brandl, Guenther Rudolph, Matthias Olden, Margaux A. Morrison, Denise J. Morgan, Matthew Schu, Jeeyun Ahn, Giuliana Silvestri, Evangelia E. Tripon, Kyu Hyung Park, Lindsay A. Farrer, Anton Orlin, Alexander Bruckner, Mingyao Li, Christine A. Cucio, Saddek Mohand-Said, José-Alain Sahel, Isabelle Audo, Mustapha Benchabone, Angela J. Cree, Christina A. Rennie, Srinivas V. Govender, Shina Guhbi-Levi, Peter Campochiaro, Nathan Katsanis, Frank G. Holz, Frédéric Blond, Hélène Blanche, Jean-François Delenge, Robert P. Igo Jr., Barbara Truitt, Neal S.
The authors declare that they have no competing interests. Not applicable.

approved by institutional review boards as previously described [ 8]. All participants provided informed consent, and the study was

The genotype data analyzed during the current study were generated by

Availability of data and materials

The genotype data analyzed during the current study were generated by the IAMDGC as previously described [8] and are available through the database of Genotypes and Phenotypes (dbGAP; Accession: phs001039.v1.p1).

Ethics approval and consent to participate

The study participants were previously ascertained by IAMDGC cohorts as described [8]. All participants provided informed consent, and the study was approved by institutional review boards as previously described [8].

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Received: 29 January 2020 Accepted: 22 June 2020

Published online: 06 July 2020

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