Candidate gene studies of diabetic retinopathy in human

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Abstract Diabetic retinopathy (DR) is a multifactorial disease with complex pathophysiology. It is the main cause of blindness among the people in productive age. The purpose of this literature review is to highlight recent achievements in the genetics of diabetic retinopathy with particular focus on candidate gene studies. We summarized most of the available published data about candidate genes for diabetic retinopathy with the goal to identify main genetic aspects. We conclude that genetic studies reported contradictory findings and no genetic variants meet criteria of a diagnostic marker, or significantly elucidate the root of DR development. Based on these findings it is important to continue with the research in the field of DR genetics, mainly due to the fact that currently new possibilities and approaches associated with utilization of next-generation sequencing are available.

Keywords Diabetic retinopathy · DNA variants · Sequencing · Diabetes mellitus · Genetic studies

Introduction

Diabetes mellitus (DM) is one of the most significant health problems worldwide. It is a metabolic disorder in which elevated blood sugar levels are present as a result of the inability to produce a sufficient amount of insulin (type 1) or because of cellular insulin resistance (type 2). Both types of diabetes are associated with hyperglycaemia, oxidative stress, inflammation and macrovascular (coronary artery disease, atherosclerosis, hypertension and stroke) and microvascular complications such as retinopathy, nephropathy and neuropathy [1].

The number of patients with diabetes mellitus is rapidly increasing every year. Global mortality resulting from diabetes in adults was estimated to be 1.5 million deaths in 2012 (World Health Organization). It is estimated that there will be 418 million patients with impaired glucose tolerance and 380 million patients with T2DM by 2025 [2].

Diabetic retinopathy (DR) is a leading cause of visual impairment in patients at productive age. These alarming numbers highlight the necessity of optimization of diagnostic methods that will allow early identification of diabetic patients with significantly elevated risk of DR development that will help start optimal prevention and intervention. DR has an overall prevalence of 22–37% in individuals with known diabetes. It leads to damage of the retina microvasculature as a result of prolonged exposure to metabolic changes induced by diabetes. If left untreated, it may lead to blindness on account of continuous blood leakage due to the loss of retinal pericytes and fenestration [1]. DR is classified into two categories based on severity, namely less-severe nonproliferative diabetic retinopathy (NPDR) and severe proliferative diabetic retinopathy (PDR). The key changes of the retina in NPDR, as a result of hypoxia and venous bleeding, are microaneurysms, vascular leakage, hard
exudates, intraretinal microvascular abnormalities and cotton wool spots (Fig. 1). Retinal neovascularization induced by ischemia is the main characteristic of PDR [3].

The aetiology of this complex disease remains unclear and poorly understood. It is associated with both environmental and genetic factors. The possibility of developing and progression of DR is closely related to the duration of DM [1]. Almost all patients with T1DM and >60% of patients with T2DM are anticipated to have some type of retinopathy within the first 10 years of diabetes being diagnosed. The Diabetes Control and Complications Trial (DCCT) and United Kingdom Prospective Diabetes Study (UKPDS) clinical trials have also confirmed significant association between chronic hyperglycaemia and development and progression of DR, however, the fundamentals of how hyperglycaemia causes microvascular changes in the retina has not been fully elucidated [4]. Involvement of several biochemical pathways which can elucidate the role of hyperglycaemia in DR pathophysiology has been proposed, including activation of diacyl-glycerol (DAG)-PKC pathway, accelerated formation of advanced glycation end-products (AGE), increased polyol pathway flux, increased expression of growth factors (VEGF, IGF-1), haemodynamic changes, renin-angiotensin-aldosterone system (RAAS), leukostasis, subclinical inflammation, and oxidative stress that leads to increased expression of several proinflammatory genes (NF-kB, TGF-β, NOX4, Nrf2, etc.). Other risk contributors for DR development are dyslipidemia [5] and possibly blood pressure, but studies are contradictory about these risk modifiers [4, 6]. Pathways contributing to DR pathologies alongside with relevant genes are summarized in Fig. 2. In summary, increased vascular permeability, haemostatic abnormalities, endothelial dysfunction, increased tissue ischemia, angiogenesis and neovascularization is typical for overall DR pathophysiology [1].

**Genetic aspects of the diabetic retinopathy**

The above mentioned risk factors are not solely responsible for susceptibility to DR. Clinical studies have revealed considerable variations in the retinopathy onset and severity that cannot be fully explained by known risk factors such as the duration of diabetes, the level of glycemic control, or concomitant vascular disease [7]. For instance, some people might have DR even when they have good glycemic control and duration of DM is short. In contrast, other patients have poor glycemic control and prolonged duration of DM and yet may not develop DR. The probability of DR development also depends on the ethnicity; the Hispanics, the individuals of African descent and the Asians are more susceptible to DR [8, 9]. A study from 2008 reported retinopathy symptoms, including retinal microaneurysms, in nondiabetic patients with an optimal glucose level (glycosylated haemoglobin levels <5.0%) [10] and these microvascular changes were indistinguishable from lesions in diabetic DR. Therefore there is evidence that additional risk factors and genetic predispositions have a part in the pathogenesis of DR.
development and progression of DR and that these factors are independent of DM.

This is a solid confirmation of genetic contribution to the development and progression of DR. Over the past several years, progress has been made in identifying some of the susceptibility loci associated with DR through twin studies, family studies, candidate gene studies, linkage studies and small-scale GWAS (genome-wide association study). Twin and family studies have demonstrated that risk of DR emergence is three times higher for patients with a family history of DR than in patients without it for both T1DM and T2DM. Concordance is dramatically higher among monozygotic twins when compared to dizygotic twins [11]. One of the first twin studies has reported DR concordance of 68% in T1DM and 95% in T2DM [12]. Heritability score increases with the severity of DR, and has been estimated to be 18 and 52% for DR and PDR, respectively [13].

Candidate genes studies

Our knowledge of pathophysiology of DR allows us to propose possible candidate genes, which could play a role in the development and progression of DR. Candidate gene studies compare the frequency of a particular genetic variant in subjects with or without DR. This approach has revealed several genes with a possible key role in DR. These genes are part of different physiological and pathophysiological processes in organism, often associated with inflammation, such as RAAS (renin–angiotensin–aldosterone system), glucose induced pathways, remodeling of extracellular matrix (ECM), vascular endothelial dysfunction, and angiogenesis. It has been proposed that a great number of factors and genes with modest effect, as a part of different biochemical pathways, invoke pathological processes leading to DR.

Polyol pathway and its role in DR

Polyol pathway represents the main metabolic link between hyperglycaemia and damages caused by DM. Aldose reductase (ALR2) is the essential enzyme in the pathway. ALR2 converts glucose to sorbitol in an NADPH-dependent reaction. During the hyperglycaemia sorbitol accumulates in cells and induces osmotic stress and cellular damage. The above mentioned process leads to the destruction of retinal cells, microaneurysms, thickening of the basement membrane and loss of pericytes in animal models, which are also
| Gene symbol | Gene name                  | Function/cellular role                                                                 | Polymorphism                                                                 | Ch. | Type of DM | Population                                                                 | Comments                                                                                           | Ref.          |
|-------------|----------------------------|----------------------------------------------------------------------------------------|------------------------------------------------------------------------------|-----|-------------|-----------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------|---------------|
| AKR1B1/ALR2 | Aldose reductase gene      | Polyol pathway—conversion of glucose to sorbitol                                         | rs35839483 [(CA)n dinucleotide repeats]                                      | 7   | 1 and 2     | Chinese, Japanese, Indians, Chileans, Brazilians                           | z-2 microsatellite confers risk in all DR, z2 microsatellite against all DR                         | [20–27]      |
|             |                            |                                                                                        | rs759853 (c. C-106T)                                                        | 7   | 2           | Euro-Brazilian, Mainland Chinese, Han Chinese, Japanese                     | T allele protective against DR but according to some studies it is weak association                  | [25, 26, 28–30]|
|             |                            |                                                                                        | rs964083                                                                     | 7   | 2           | Australian                                                                  | Association with onset of diabetes                                                                   | [25]          |
| SDH         | Sorbitol dehydrogenase     | Polyol pathway—conversion of sorbitol to fructose                                       | rs2055858 (c. C-1214G)                                                      | 15  | 2           | Poland                                                                     | Weak associations; polymorphism possibly affect promoter activity                                   | [18, 19]     |
|             |                            |                                                                                        | rs3759890 (c. G-888C)                                                       | 15  | 2           | Japan, Poland, Caucasian-Brazilians                                        | Inconsistent finding, polymorphism possibly affect promoter activity                                 | [18, 31]     |
| ALDH2       | Mitochondrial aldehyde     | Polyol pathway—transformation from acetaldehyde to acetic acid, prevents creation of AGE| ALDH2*2                                                                      | 12  | 2           | Japanese                                                                   | Associated with protective effect against DR                                                        | [32]          |
| VEGF        | Vascular endothelial growth factor | Stimulation of angiogenesis and vasculogenesis | rs2010963 (c. C-634G) (c. C-460T)                                           | 6   | 2           | Japanese, Indian, Caucasian                                                | C allele confers risk for NPDR in T2DM                                                               | [33–41]      |
|             |                            |                                                                                        | rs25648                                                                      | 6   | 2           | Caucasian                                                                  | Possible association with DR                                                                         | [42, 43]     |
|             |                            |                                                                                        | rs1570360 (c. A-116G)                                                       | 6   | 2           | Multi-ethnic                                                               | T allele increase risk of DR but finding inconsistent                                               | [33, 34, 37] |
|             |                            |                                                                                        | rs309939                                                                     | 6   | 2           | Multi-ethnic                                                               | Inconsistent finding                                                                                 | [33, 34, 42] |
|             |                            |                                                                                        | rs35583934                                                                  | 6   | 1 and 2     | Multi-ethnic                                                               | T allele increase risk of DR but finding inconsistent                                               | [33–36]      |
|             |                            |                                                                                        | rs699947(c. A-2578C)                                                        | 6   | 2           | Multi-ethnic                                                               | (~ −2549) DEL increases risk but finding inconsistent                                               | [33]          |
|             |                            |                                                                                        | rs13207351 (c. A-152G)                                                      | 6   | 1 and 2     | Caucasian                                                                  | A allele increases risk but finding inconsistent                                                    | [33, 36, 40, 43–46]|
|             |                            |                                                                                        | rs735286 (c. C4618T)                                                        | 6   | 1 and 2     | Caucasian                                                                  | Associated with PDR in some of the studies                                                          | [34, 42, 46] |
|             |                            |                                                                                        | rs2146323 (c. C5092A)                                                       | 6   | 1 and 2     | Caucasian                                                                  | Haplopt-type-tagged SNP associated with severity of DR                                              | [42]          |
|             |                            |                                                                                        | rs833061 (c. C-1498T)                                                       | 6   | 2           | Chinese                                                                   | Inconsistent finding and weak association                                                            | [34, 37, 39, 46]|
|             |                            |                                                                                        | rs3025021                                                                   | 6   | 2           | Chinese                                                                   | Inconclusive                                                                                         | [33, 46]     |
|             |                            |                                                                                        | rs10434                                                                     | 6   | 1 and 2     | Caucasian                                                                  | G allele associated with blinding DR                                                                  | [33]          |
|             |                            |                                                                                        | rs833068                                                                   | 6   | 1 and 2     | Caucasian                                                                  | G allele confers risk in DR                                                                          | [33]          |
|             |                            |                                                                                        | rs833070                                                                   | 6   | 1           | Japanese                                                                   | Associated with early progression of DR but weak association                                          | [44]          |
|             |                            |                                                                                        | rs3025039 (c. C+936T)                                                       | 6   | 2           | Caucasian                                                                  | T allele increases risk                                                                               | [48]          |
| Gene symbol | Gene name | Function/cellular role | Polymorphism | Ch. | Type of DM | Population | Comments | Ref. |
|-------------|-----------|------------------------|--------------|-----|------------|------------|----------|------|
| bFGF/FGF2   | Basic fibroblast growth factor/fibroblast growth factor 2 | Stimulation of angiogenesis and tissue repair | rs1456044; rs308395; c. C-754G; c. T-553 A; c. T-834A | 4   | 2 | Multi-ethnic | A allele increases risk but weak associations | [15] |
| IGF-1       | Insulin-like growth factor 1 | Stimulation of cell growth and proliferation, inhibition of apoptosis | (CA)n | 4   | 2 | Southern Indian | 18-repeat of (CA) increases risk of DR | [51] |
| EPO         | Erythropoietin | Control of erythropoiesis, stimulation of proliferation, migration and angiogenesis in hypoxic cells | rs1617640, rs507392, rs551238 | 7   | 1 and 2 | Multi-ethnic, European American, Australian | TTA allele associated with PDR in European American, meta-analysis hasn’t found significant association; GCC haplotype associated with DR in Australian | [15, 52, 53] |
| RAGE        | Advanced glycosylation end product-specific receptor | Activation of pro-inflammatory genes | rs1800624 (c. T-374A); rs1800625 (c. T-429C); rs2070600 (p. G32S) | 6   | 2 | Indian, Chinese, African-Brazilian, Caucasian - Scandinavian, Caucasian, Indain, Danish, Indian, Chinese, Pakistani | Inconsistent finding and weak association, may be interacting with glycosylated hemoglobin; Inconsistent finding and weak association, functional studies show differences in polymorphic receptor activity | [15, 54–56] |
| ACEI        | Angiotensin-I converting enzyme | Component of the renin-angiotensin system—activation of angiotensin II | rs4646994 (c. G2350A)-INS/DEL at intron 16 | 17  | 1 and 2 | Caucasian - Slovene, Danish; Japanese, Multi-ethnic, Iranians, Chinese, Pakistanians | D allele possibly associated with DR in T2DM in Chinese, but inconsistent finding and weak association in other populations, associated with NPDR in Pakistanians | [62–66] |
| GSTT1       | Glutathione S-transferase T1 | Detoxifying enzyme—conjugation of reduced glutathione to a compounds | Null genotype | 22  | 2 | Caucasian - Slovenian | Greater risk of DR | [67] |
| GSTM1       | Glutathione S-transferase M1 | Detoxifying enzyme—conjugation of reduced glutathione to a compounds | Null genotype | 1   | 2 | Caucasian - Slovenian | Lower risk of DR | [67] |
| SOD2/MnSOD  | Mitochondrial manganese superoxide dismutase | Decrease of ROS production (transformation to to peroxide and oxide) | rs4880 (c. C47T, p. A16V) | 6   | 1 and 2 | Slovene (Caucasian), Finnish, Indian | C allele reduces risk of DR, not confirmed in Indian population | [15, 36, 56, 65, 68, 69] |
| eNOS3       | Endothelial nitric oxide synthases | Synthesis of nitric oxide (vasodilatation) | rs1338808 (27 VNTR intron 4 a/b); rs1799983 (c. C894T); rs41322052 (c. T-784C); rs2297518 (c. G-954C) | 7   | 2, 1&2 | Indians, West African, Caucasian - Brazilian, Caucasian - Brazilian, Danish, Multi-ethnic, Caucasian - Brazilian, Multi-ethnic, Caucasian | 4a allele protective effect against DR; G allele increases risk but weak association; Inconsistent finding and weak association | [15, 37, 70–73] | [15, 37, 50, 60, 72, 74] |

Table 1 (continued)
| Gene symbol | Gene name                      | Function/cellular role                                                                 | Polymorphism                                                                 | Ch. | Type of DM | Population                      | Comments                                                        | Ref. |
|-------------|--------------------------------|----------------------------------------------------------------------------------------|------------------------------------------------------------------------------|-----|-------------|----------------------------------|-----------------------------------------------------------------|------|
| RXRA        | Retinoid X receptor alpha      | Nuclear receptor—retinoic acid-mediated gene activation (antioxidants properties)     | rs3132300                                                                    | 9   | 1           | African American                 | Associated with progression of DR                               | [76, 77] |
| RXRG        | Retinoid X receptor gamma      | nuclear receptor—retinoic acid-mediated gene (antiproliferative effects)              | rs3818569                                                                    | 9   |             | Taiwanese                        | G allele associated with development of DR                       | [78] |
| UCP1        | Uncoupling protein-1           | Mitochondrial anion carrier protein (thermogenesis), protection against oxidative stress | rs1800592 (c. A-3826G)                                                      | 4   | 1 and 2     | Brazilian, Chinese, Danish       | G allele associated with increased risk of PDR                   | [60, 75, 79] |
| UCP2        | Uncoupling protein-2           | Mitochondrial anion carrier protein (thermogenesis), control of ROS production         | rs660339 (p. A55V, 45 bp INS/DEL)                                            | 11  | 1 and 2     | Brazilian                        | Risk factor for PDR                                             | [77, 80] |
| TLR4        | Toll-like receptor 4           | Pathogen recognition and activation of innate immunity                                 | rs10759931, rs1927914                                                        | 9   | 2           | Indian, Chinese                 | A,T alleles positively modulate the risk of DR, rs1927914 associated with susceptibility to DR in a Han Chinese population | [3, 55] |
| CFH         | Complement factor H            | Regulator of complement activation                                                    | rs800292 (p. J62V)                                                          | 1   | 2           | Chinese                          | Associated with DR                                              | [82] |
| CFB         | Complement factor B            | Regulator of complement activation                                                    | rs1048709                                                                    | 6   | 2           | Chinese                          | Associated with DR                                              | [82, 83] |
| MCP-1/CCL2  | Monocyte chemoattractant protein-1 | Cytokine—activation of monocytes, macrophages and lymphocytes | rs1024611 (c. A-2518G)                                                      | 17  | 2           | Chinese, Korean, Japanese       | G allele associated with susceptibility to DR and specifically PDR in Koreans | [84-86] |
| TGF-β1      | Transforming growth factor-beta 1 | Control of cell growth, proliferation, differentiation and apoptosis                   | c. T869C (p. L10P)                                                          | 19  | 2           | Multi-ethnic                     | Potential protect factor against DR                             | [15, 83] |
| ICAM1       | Intercellular adhesion molecule 1 | Stabilization of cell-cell interactions and facilitation of leukocyte endothelial transmigration | rs13306430                                                                  | 19  | 2           | Multi-ethnic                     | G allele confers protection                                      | [15, 77] |
| SLC2A1/GLUT1| Solute carrier family 2, member 1 | Transport of glucose across the plasma membranes                                     | rs841846 (c. A26177G)                                                       | 1   | 1 and 2     | African American, Malaysian     | Significant associations with severe DR, associated with progression of DR, not confirmed in Malaysian | [77, 94] |
| SLC2A11     | Solute carrier family 2, member 11 | Transport of glucose across the plasma membranes                                    | rs841853                                                                    | 1   | 1           | Malaysian, Multi-ethnic          | Weak association                                                | [77, 94] |
| SLC24A3     | Solute carrier family 24, member 3 | Sodium-calcium exchanger                                                             | rs2294895                                                                   | 20  | 1           | African American                 | Associated with progression of DR                               | [77] |

Table 1 (continued)

1 3
| Gene symbol | Gene name | Function/cellular role | Polymorphism | Ch. | Type of DM | Population | Comments | Ref. |
|-------------|-----------|------------------------|--------------|-----|------------|------------|----------|-----|
| PPARγ       | Peroxisome proliferator-activated receptor γ | Nuclear receptor—regulation of fatty acid storage and glucose metabolism, role in vascular permeability, inflammation, angiogenesis, neoangiogenesis, and insulin resistance | rs1801282 (c. C34G, p. P12A), rs10510419 | 3, 1 | Caucasian-Poland, Chinese, Danish, Multi-ethnic | Gallic acid confers protection against DR in Caucasian but finding inconsistent - protective effect against only PDR during T2DM in Pakistan population, not for Asian patients | [60, 79, 95, 96] |
| TCF7L2/TCF4  | Transcription factor 7-like 2 | Transcription factor for several genes (Wnt signaling pathway), vascular development | rs7901246, rs7901695, rs12255372 | 10 | Caucasian-Italian, Chinese, Multi-ethnic | Associated with DR, cardiovascular disease and coronary artery disease, rs7905146 associated with DR risk in Caucasian | [15, 79, 96–98] |
| OPG/OCIF    | Osteoprotegerin/osteoclastogenesis inhibitory factor | Cytokine receptor | rs2073618, rs3134069 | 8 | Caucasian-Slovenian | CA haplotype increase risk of DR | [99] |
| PAI-1       | Plasminogen activator inhibitor-1 | Serine protease inhibitor—inhibitor of plasminogen activation, tissue repair and remodeling | rs1799768 (4G/5G INS/DEL) | 7 | Indian, Caucasian, Euro-Brazilian, Multi-ethnic, Pakistani | 4G/5G allele increases risk but finding inconsistent, ethnicity discrepancies | [28, 55, 66, 100, 101] |
| MMP-2       | Matrix metalloproteinase-2 | Breakdown of extracellular matrix | c. C-1306T | 16 | Chinese | T allele associated with PDR | [7] |
| ANGPT1      | Angiopoietin 1 | Vascular development and angiogenesis | rs1283649 | 8 | African American | Significant associations with severe DR | [77] |
| APOE        | Apolipoprotein E | Transportation of lipoproteins, fatsoluble vitamins, and cholesterol | E2/E3/E4 | 19 | Mexican, Multi-ethnic | Inconsistent finding and weak association | [102, 103] |
| BBS2        | Bardet-Biedl syndrome 2 protein | Unknown function and link to DR | rs4784675 | 16 | African American | Significant associations with severe DR | [15] |
| CPVL/CHN2   | Carboxypeptidase, vitellogenic-like, chimerin 2 | Carboxypeptidase—unknown function/regulation of a cell growth, proliferation, and migration | rs39059, rs1002630 | 7 | Chinese | Increases risk of DR, significant in meta-analysis | [77, 104] |
| CTSH        | Cathepsin H | Lysosomal cysteine proteinase - degradation of lysosomal proteins, putative role in microcirculation changes | rs3825932 | 15 | Danish | T allele associated with reduced risk of progression to PDR | [60] |
| DRD2        | Dopamine receptor D2 | Dopamine receptor—regulation of vasodilation, aldosterone production and insulin secretion | rs7131056 | 11 | African American | Significant associations with severe DR | [77] |
| EDN1        | Endothelin-1 | Vasoconstriction | rs5370 (p. K198N) | 6 | Chinese | Reduced risk in Chinese | [77, 105] |
| ENPP1       | Ectonucleotide pyrophosphatase/phosphodiesterase 1 | Insulin resistance, interaction with integrin | rs1409181 | 6 | African American | Significant associations with severe DR | [77] |
| Gene symbol | Gene name | Function/cellular role | Polymorphism | Ch. | Type of DM | Population | Comments | Ref. |
|-------------|-----------|------------------------|--------------|-----|------------|------------|----------|------|
| ERBB3/HER3 | Human epidermal growth factor receptor 3 | Protein-tyrosine kinase—activation of downstream signaling pathways, unknown link to DR | rs2292239 | 12 | 1 | Danish | T allele associated with reduced risk of progression to PDR | [60] |
| FLT1/VEGFR1 | FMS-like tyrosine kinase 1/vascular endothelial growth factor receptor 1 | Protein-tyrosine kinase—control of cell proliferation and differentiation | rs62227 | 13 | 1 | African American | Associated with progression of DR | [77] |
| FRMD3 | FERM domain containing 3 | Maintaining cellular shape, putative TSG, unknown link to DR | rs10868025 | 9 | 2 | Chinese | Weak association with DR | [104] |
| HLA-B | Major histocompatibility complex, class I, B | Regulation of the immune system—presenting peptides on the cell surface | rs2523608 | 6 | 1 | African American | Significant associations with severe DR, associated with progression of DR | [77] |
| HTR1B | Serotonin receptor 1B | GPCR for serotonin—regulation of serotonin, dopamine, and acetylcholine release, putative regulator of retinal blood flow | rs1228814 | 6 | 1 | African American | Significant associations with severe DR | [77] |
| HTRA1/ARMS2 | HtrA serine peptidase 1/age-related maculopathy susceptibility 2 | Serine protease—regulation of insulin-like growth factors, putative regulator of cell growth and neovascularization | rs11200638, rs10490924 | 10 | 2 | Indian | Marginal association with DR | [55] |
| IL-10 | Interleukin-10 | Cytokine—pleiotropic effects in immunoregulation and inflammation | n. A-1082G | 1 | 2 | Indian | G allele is risk factor for PDR | [113] |
| INSR | Insulin receptor | Activation of the insulin signaling pathway | rs10500204 | 19 | 1 | African American | Associated with progression of DR | [77] |
| ITGA2B1 | Integrin α2β1 | Cell-cell and cell-extracellular matrix interactions | RFLP - Bgl II | 5/10 | 2 | Japanese, Caucasian | Risk factor for DR | [15, 106, 143] |
| ITGB5 | Integrin β5 | Cell-cell and cell-extracellular matrix interactions | rs9855359 | 3 | 1 | African American | Associated with progression of DR | [15] |
| MTHFR | Methylenetetrahydrofolate reductase | Remethylation of homocysteine to methionine | rs1801133 (c. C677T) | 1 | 2 | Japanese, Euro-Brazilian, Multi-ethnic, Turkish | Controversial findings, T allele possible increases risk of DR because of hyperhomocysteinemia | [28, 77, 107, 108] |
| NPY | Neuropeptide Y (p. L7P) | Vasorestriction, angiogenesis | rs16139 | 7 | 2 | Finnish | C allele increases risk but weak association | [144] |
| OLR1 | Oxidized low-density lipoprotein (lectin-like) receptor 1 | Recognition, internalization and degradation of oxidized low-density lipoprotein, putative regulator of Fas-induced apoptosis | rs2742115 | 12 | 1 | African American | Associated with progression of DR | [77] |
| Gene symbol | Gene name                                      | Function/cellular role                                                                 | Polymorphism                                      | Ch. | Type of DM | Population          | Comments                                      | Ref.                       |
|-------------|------------------------------------------------|----------------------------------------------------------------------------------------|-------------------------------------------------|-----|-------------|---------------------|-----------------------------------------------|----------------------------|
| PEDF/SERPINF1 | Pigment epithelium derived factor/serpin peptidase inhibitor, clade F member 1 (alpha-2 antiplasmin) | Antioxidative properties, inhibition of angiogenesis, neurotrophic factor (neuronal differentiation in retinoblastoma cells) | rs12150053, rs12948385, rs8697961, rs1126287     | 17  | 2          | Multi-ethnic        | Not associated with DR                       | [8, 15]                   |
| PON1        | Paraoxonase 1                                  | Cellular antioxidant— inhibition of HDL oxidation                                         | rs662 (p. Q192R), rs854560 (p. L55M)             | 7   | 2          | Multi-ethnic        | Inconsistent finding and weak association     | [15, 109]                  |
| PON2        | Paraoxonase 2                                  | Cellular antioxidant, hydrolytic activity—a putative role in defense responses to pathogenic bacteria | rs7493 (p. S311C), rs12026 (p. A148G)            | 7   | 1 and 2    | Multi-ethnic        | Inconsistent and weak association             | [77]                      |
| PROS1       | Protein S                                      | Co-factor for the antioxidant protease                                                | rs13062355                                       | 3   | 1          | African American   | Significant associations with severe DR      | [77]                      |
| PSMD9       | Proteasome 6S subunit, non-ATPase, 9            | Part of multicatalytic protease complex (proteasome)                                   | rs74421874, rs14259, rs3825172                    | 12  | 2          | Italian             | Associated with DR                            | [110]                     |
| ROBO2       | Roundabout, axon guidance receptor, homologue 2 | Axon guidance and cell migration, unknown link to DR                                   | rs10865559                                        | 3   | 1          | African American   | Significant associations with severe DR      | [77]                      |
| ROCK2       | Rho-associated, coiled-coil containing protein kinase 2 | Serine/threonine kinase—regulation of cytokinesis, smooth muscle contraction, the formation of actin stress fibers and focal adhesions, and the activation of the c-fos serum response element | p.T431N, p. R83K                                 | 2   | 1 and 2    | Turkish             | No association                               | [15, 111]                 |
| Romo-1      | Reactive oxygen species modulator 1            | Mitochondrial membrane protein—increase of the level of reactive oxygen species in cells | rs6060566                                        | 20  | 2          | Caucasian           | Independent risk factor for DR               | [112]                     |
| TF          | Transferrin                                    | Transportation iron from the intestine, reticuloendothelial system, and liver parenchymal cells to all proliferating cells in the body | rs3811647                                        | 3   | 1          |                                    | Associated with progression of DR            | [77]                      |
| TNF-α       | Tumor necrosis factor-alfa                     | Multifunctional pro-inflammatory cytokine (cell proliferation, differentiation, apoptosis, lipid metabolism, and coagulation) | rs361525 (c. G-238A), rs1800629 (c. G-308A), rs1799724 (c. C-857T) | 6   | 2          | Indian, Caucasian - Brazilians | AA genotype of rs361525 confers risk for pathogenesis of PDR in Indian, rs1800629 associated with PDR in Caucasian - Brazilians | [113–115]                 |
typical symptoms of DR in humans [14]. Three DR-associated ALR2 polymorphisms have been identified in different populations (see Table 1). The first polymorphism located at the 5′ end of the gene, the Z-2 allele of the (CA)n microsatellite located at the 5′ end of the gene increases risk of DR. In contrast, Z+2 and Z alleles show protective effect against DR [15]. Another polymorphism, rs759853, has shown association with DR where T allele confers protection against DR in T1DM [15, 16]. However, the use of AKR (aldo–keto reductase) inhibitors did not confirm expected results in clinical trials and they could not prevent progression of the disease [17]. But these clinical trials disregarded the genetic variants in the AKR2 gene which could have had negative impact on the function of AKR inhibitors. Another enzyme in the polyol pathway, sorbitol dehydrogenase (SDH) converts sorbitol into fructose in NAD+‑dependent reaction. Amano et al. found that SDH overexpression potentiated glucose toxicity to cultured retinal pericytes, thus leading to acceleration of pericyte loss, a typical trait of DR [18]. Polymorphisms rs2055858 and rs3759890 were identified in Polish, Japanese and Caucasian-Brazilian population and it is possible that they could affect the promoter activity of the SDH gene and have role in onset of DR [18, 19].

The mitochondrial aldehyde dehydrogenase 2 (ALDH2), expressed in vasculature, detoxifies reactive aldehydes formed from glucose and lipids, also prevents creation of AGE (advanced glycation end products) [32]. Morita et al. have reported a substantial relation between the ALDH2*2 allele and the incidence of DR in their study.

### Growth factors with role in DR

Vascular endothelial growth factor (VEGF) is one of the major factors in angiogenesis and influences vascular permeability of endothelial cells. VEGF is activated by microvascular changes induced by hypoxia during DM and also by hyperglycaemia [120]. Activation of VEGF leads to the destruction of the blood retinal barrier (BRB), the development of diabetic macular oedema and neovascularization typical for PDR. At the same time, elevated serum and vitreous levels of VEGF have also been described in eyes of patients with PDR [121]. Anti-VEGF therapies have led to the improvement of the patients’ condition and to the deceleration of retinal vessels proliferation [122]. Studies have revealed several polymorphisms in the VEGF promoter (rs210963, rs25648, rs1570360, rs3095039, rs35569394, rs699947, rs13207351, rs735286, rs2146323, rs833061, rs302502, rs10434, rs833068 and rs833070) with possible associations with DR [15, 33, 42, 44]. Rs210963 (–634C/G) has been associated with DR in Japanese and Indian populations [34, 35] whereas G allele of rs210963 has significant protective effect against NPDR in patients.
Abhary et al. have associated the GCC haplotype with DR in the Australian population [52, 53]. Interaction of various growth factors, cytokines, cell signalling molecules, and extracellular matrix are essential for angiogenesis during DR [125] while VEGF plays a crucial part [126] (Fig. 3).

Receptor for advanced glycation end products and cytokines

Hyperglycaemia causes nonenzymatic glycation of proteins and lipids and the creation of AGE. Accumulation of AGE leads to tissue damage by the formation of covalent cross-links between proteins, which alter structure and function of proteins. Another feature of AGE is its ability to interact with different surface receptors, such as the receptor for advanced glycation end products (RAGE). RAGE is an immunoglobulin and its activation leads to cytokine secretion. Cytokines accelerate the advance of diabetic complications by supporting proinflammatory processes and increasing endothelial permeability [127]. AGEs are found in the retinal vessels of diabetic patients where their levels correlate with those in the serum as well as with severity of retinopathy [128]. The c. T–374A (rs1800624), p. Gly82Ser (rs2070600) and c. T–429C (rs1800625) polymorphisms in the RAGE gene are associated with DR in Caucasians and with T2DM. Rs2010963 is also associated with higher risk of macular oedema in Japanese population [123]. There are constantly emerging studies identifying new polymorphisms in VEGF gene with possible connections to DR which underlines importance of this gene in the development of DR.

Other growth factors with a possible function in the pathology of DR are the basic fibroblast growth factor (bFGF) and insulin-like growth factor 1 (IGF-1). The bFGF is important for tissue repair and is angiogenic factor. Studies have revealed increased level of bFGF in patients with PDR and it seems to stimulate VEGF production. IGF-1 regulates the proliferation and differentiation of several cell types. Levels of intravitreal IGF-1 were found to be significantly increased in the eyes of patients with PDR compared to those of controls [124]. Variants identified to date are summarized in the Table 1.

Erythropoietin (EPO) plays an important role in stimulation of bone marrow stem cells, erythropoiesis, proliferation, migration, and angiogenesis in hypoxic vascular endothelial cells. Study has reported a elevated concentration of EPO in the vitreous of DM and PDR patients compared to controls [52]. There are two studies which have reported the association of rs1617640, rs507392, and rs551238 with the development of DR, but these studies report different findings. Tong et al. have determined the TTA haplotype as a risk contributor in European American population, whereas
Asian Indians [54, 57, 129, 130], but the association was not confirmed in Chinese [131].

Dysregulation of RAAS system

The rennin-angiotensin-aldosterone system (RAAS) is an endocrine system involved in the regulation of blood pressure and fluid balance. Patients with diabetes show dysregulation of RAAS system, namely angiotensin converting enzymes I and II (ACEI, ACEII) and angiotensin receptors which are upregulated in retina during PDR independently of blood pressure [132]. ACE converts angiotensin I (ATI) to angiotensin II (ATII) which mediates its haemodynamic effects through the angiotensin receptor ANGTR1 and ANGTR2. ATII in the eye regulates promotion of capillary growth, cell growth, intraocular blood flow and pressure, enhances vascular permeability, increases oxidative stress and via the expression of several growth factors including VEGF, IGF-1 and PDGF [15]. ACE inhibitors, angiotensin receptor blockers prevents neovascularization, reduce the incidence and progression of DR in T1DM. Studies have proposed that ACEII is also involved in PKC activation [133]. Meta-analysis suggested that ACE I/D polymorphism (insertion/deletion of a 287 bp Alu sequence in intron 16) may be associated with PDR [62].

The other polymorphisms modulating risk of DR

The retina is very sensitive to damage by oxidative stress. Oxidative stress is strongly implicated in the pathogenesis of DR, therefore the role of detoxifying enzymes, such as glutathione S-transferases (GST), was considered in the development of DR. Studies have shown that GSTT1-null genotype is found more frequently in the cases with DR in Caucasians patients with T2DM compared to controls, so the GSTT1-null genotype can be a risk factor for DR The individuals homozygous for the GSTT1-null allele had more generalized vasculopathy that leads to increased risk of sight threatening DR. In contrast, the GSTM1-null genotype may confer protection against development of DR in people with T2DM [67], but at the same time this polymorphism confers elevated risk for lung cancer [134]. There are reports that deficiency in GSTM1 leads to slower excretion of isothiocyanates. Isothiocyanates also suppress expression of VEGF which is the main inductor of retinal neovascularization in diabetes [67].

Oxidative stress induces a large amount of ROS and is assumed to damage the mitochondrial DNA. Mitochondrial manganese superoxide dismutase (MnSOD) prevents an excessive production of ROS by dismutation of superoxide radicals into hydrogen peroxide and hence defends the retinal endothelial cells from oxidative damage. The polymorphism rs4880 (c. C47T, p. A16V) affects a mitochondrial processing efficiency under oxidative stress and has been associated with DR in some studies [36, 68, 69].

Regarding PDR, presence of the 4a/4a genotype of the VNTR polymorphism for endothelial nitric oxide synthase (eNOS) has been associated with 3.4 times increased risk of PDR in Caucasian patients with T2DM [135]. In contrast, other studies have proposed that the 4a allele has a protective effect against DR [70, 71]. NO synthesized by eNOS is an endogenous vasodilator and has a role in induction of angiogenesis and regulation of VEGF expression. NO levels are significantly elevated in PDR patients relative to nondiabetic subject.

A study in 2008 revealed that retinoid-X receptor alpha (RXRA) possesses antioxidants properties and is associated with the development of DR [76]. Polymorphism rs3132300 has been linked with a progression of DR in T2DM in African American population. Also, polymorphism rs3818569 of the retinoid-X receptor gamma (RARG) has been found to be connected with an increased DR risk in the Taiwanese population [78].

Uncoupling protein 1 (UCP 1) is the mitochondrial inner membrane electron carrier that has a part in protection against oxidative stress. It has been proposed that UCP 1 and its product play role in insulin resistance when oxidative stress pathways are activated. SNP rs1800592, which is located in the promoter of the UCP1, has been shown to be associated with glucose homeostasis, adiposity and obesity, as well as changes in the body mass index (BMI) and body weight, resulting from metabolic disorders. UCP 1 has been implicated as a candidate marker for a risk factor of DR and the rs1800592 (c. A-3826G) polymorphism has been associated with PDR [75, 79]. Uncoupling protein 2 (UCP2) regulates production of reactive oxygen species (ROS) by mitochondria. Overproduction of ROS is associated with diabetic retinopathy (DR), thereby UCP2 gene polymorphisms can be involved in the development of this complication. rs660339 can be a relevant risk factor associated with PDR in both type 2 and 1 of diabetes [77, 80].

The inflammatory processes are a major part of the DR pathophysiology. They are often regulated by inadequate activation of members of the immune system. Toll-like receptor 4 (TLR4) takes part in the activation of a proinflammatory response by the ligand-dependent activation of the nuclear factor-kB (NF-κB) pathway. Any deregulation of TLR4 signaling due to single nucleotide polymorphisms (SNPs) in the extracellular domain of TLR4 may alter the ligand binding capacity and hence disturb the balance of pro- and anti-inflammatory cytokines [81]. It has been reported that rs4986790, rs4986791, rs10759931 and rs1927914 in TLR4 positively modulate the risk of DR [3, 81, 136].
There is increasing evidence from in vitro and in vivo studies that suggests a pathogenic role of the complement system in the development of diabetic angiopathy. In these studies, increased expression of several complement factors, namely, complement factor H (CFH), complement factor B (CFB), component 3 (C3), and component 5 (C5), has been observed in the vitreous of DR patients. CFH and CFB (an antagonist of CFH) contribute to the regulation of the activation of complement cascade. Polymorphism rs800292 (p.162V) in CFH affects protein-binding affinity with C3b and subsequently activation of the complement alternative pathway. A synergy effect between CFH rs800292 and CFB rs1048709 conferring a significantly increased risk for DR has been identified in the study of Wang [82].

Studies have reported significantly increased levels of monocyte chemotactic protein 1 (MCP-1) in aqueous and vitreous conditions in DR patients. MCP-1 has an ability to activate monocytes, macrophages and lymphocytes. Hyperglycaemia accelerates MCP-1 production in vascular endothelial cells and retinal epithelial cells which can lead to neovascularization and increased permeability of retinal vessels typical for PDR. Moreover rs1024611 polymorphism has been associated with DR in the Japanese, Korean, and Chinese populations [84–86].

Transforming growth factor β1 (TGF-β1) has an important role in angiogenesis, endothelial cell proliferation, adhesion and the deposition of extracellular matrix. The TGF-β1 gene may be involved in the development of DR through induction of angiogenesis and BRB breakdown. C. T869C (p. L10P) polymorphism has been associated with DR in the Japanese, Korean, and Chinese populations [84–86].

Intercellular adhesion molecule-1 (ICAM-1) has a major role in mediating the adhesion of circulating leukocytes to the blood vessel wall and transendothelial migration to the vascular intima. The increased retinal expression of ICAM-1 is thought to play a key role in leukostasis-mediated BRB breakdown, capillary occlusion and endothelial cell damage in DR [88]. Polymorphisms of ICAM-1 gene might have a role in modulation of its own gene expression but findings about p. K469E polymorphism (rs5498) are inconsistent across multiple studies [55, 88–91]. The G allele of rs13306430 could confer protection against DR in T2DM patients [15].

Solute carrier family 2 (SLC2A1), also known as facilitated glucose transporter (GLUT1) is expressed in endothelial cells of the BRB where SLC2A1 is the prevalent glucose transporter. Functional loss of BRB is typical for DR and studies have shown that patients with DR have high expression of GLUT1 in endothelial cells. SLC2A1 c. A26177G polymorphism has been associated with DR in a study of Ng [94].

The role of peroxisome proliferator-activated receptor γ (PPARγ) in DR pathogenesis has come to forefront mainly because of the protein’s role in vascular permeability, inflammation, angiogenesis, neovascularization, and insulin resistance, all of which contribute to the onset and severity of DR. However, the studies describing associations of PPARγ polymorphisms and DR have been inconsistent [15, 95].

Transcription factor 7-like 2 (TCF7L2/TCF4) is a key component in the regulation of fundamental processes such as vascular development. It has been found to mediate pathological neovascularization in PDR. Common variant rs7903146 in TCF7L2 has been reported to be strongly associated with T2DM and also with PDR in Caucasian [96]. An Italian study observed associations between TCF7L2 variants (rs7903146, rs7901695 and rs12255372) and DR, cardiovascular disease and coronary artery disease [97].

Osteoprotegerin (OPG), also called the osteoclastogenesis inhibitor factor (OCIF), is an important regulatory molecule in the vasculature. Rs2073618, rs3134069 polymorphisms have been linked with DR [99].

Plasminogen activator inhibitor-1 (PAI-1) is an inhibitor of plasminogen activation and is involved in tissue repair and remodeling. PAI-1 plays a crucial part in the regulation of intravascular fibrinolysis which is part of DR pathophysiology. Studies have investigated the connection between PAI-1 4G/5G and DR risk but findings have been inconsistent, maybe due to ethnicity discrepancies [15, 100, 101].

Matrix metalloproteinases (MMPs) are proteolytic enzymes that degrade extracellular matrix (ECM) components. MMPs also regulate cell proliferation, neovasculogenesis and tissue remodelling because degradation of the extracellular matrix (ECM) proteins of the basement membrane is necessary for endothelial cells to migrate, proliferate, and to form capillaries. Increased expression of MMP-2 may expedite degradation of the type IV collagen and the gap junction protein, accelerating the vascular complications of diabetes. It has been reported that c. C-1306T polymorphism seems to be genetic susceptibility factor for the development of DR [7].

In this review, we discussed candidate genes and polymorphisms with the highest genetic association with DR, or those most frequently analysed in studies in different population. The other genes and their polymorphisms associated with DR are summarized in Table 1 in alphabetical order. Studies concerning these genes have reported very weak or borderline associations, had small sample sizes and most of them have failed replication in other populations [137]. We did not include studies and polymorphisms that have been done on only one population and showed no associations with DR. None of the polymorphisms identified by candidate gene studies have achieved widespread acceptance as a marker of high risk of diabetic retinopathy. In part, this may be because of the complexity of DR which probably has more multifactorial, polygenic and environmental contributors to its pathophysiology.
In our review, we have included a number of genes in which multiple alleles or nearby SNPs have varying strengths of association or significance in relation to diabetic retinopathy. Possible explanation for this is variable association between the SNPs themselves with the causative change depended also on populations.

Another one of the possible reasons for the setback of the gene candidate approach is focusing on single SNPs when the linkage disequilibrium does not have to manifest [16]. It would be potentially useful to focus on haplotypes associated with DR instead of solely on SNPs. In addition, insufficient sample sizes to detect the modest effect of polymorphisms, incomplete coverage of variation in candidate genes and incorrect hypotheses about genes in the pathophysiology of DR are some of the reasons for controversial success of candidate genes studies. Based on these weaknesses of standard candidate gene studies, two recent studies have examined a higher number of candidate genes for DR in sample sizes larger than those used previously, in an approach that mimics a genome-wide approach [77, 138]. The first study, which examined 193 candidate genes with DR of type 1 diabetic African-Americans, found genetic associations in 13 genes with progression of DR and the polymorphisms are listed in Table 1 [77]. Identified genes are involved in pathways related to glucose metabolism, inflammatory processes, angiogenesis/vascular permeability, insulin signalling, retinal development, or blood pressure regulation. The second study, the Candidate gene Association Resource (CARe) has not confirmed connection between previously associated gene from numerous previous independent studies and DR [138]. The most interesting findings from this study are the variants in the P-selectin (SELP) (rs6128) and in the iduronidase (rs6856425) that have shown to be significantly linked with DR in European Americans, but were not seen in African-Americans, Hispanic Americans, or Asian Americans.

It is possible that the potential success of candidate gene studies lies in better characterization and definition of clinical phenotypes of DR, represented by specific patterns of severity and progression of DR. Only then, studies of candidate genes are worth pursuing, involving appropriately well-defined subgroups of patients [139]. Furthermore, it would be worthwhile to expand studies to those genes mutations in which are known to initiate hyperglycaemia [140] and indirectly lead to development of DM and DR.

**Conclusion**

DR remains to be one of the most complex, heterogeneous, multifactorial disorders in any genetic studies. It is one of the leading causes of blindness and visual impairment in the world and treatments options are limited. Research worldwide is focused on understanding the pathogenic mechanisms in DR with the key goal to prevent this disease and developing new drugs for treatments. There is a common consensus that the susceptibility to DR is contingent to a great amount of relatively common allelic variants with a modest effect, and how these genes interact among themselves and with environmental influences. Each of the allelic variants increased risk of DR by a small portion in overall susceptibility. The identification of genetic susceptibility loci for DR by genetic studies has not proved notably successful thus far, given the often contradictory and inconclusive results. It is obvious that the study of the DR genetics is still poorly developed and stands against numerous challenges. The most common approaches in studying this complex disease are insufficient for elucidating pathophysiology of DR. At the same time, most studies of new possible treatments disregarded genetic background of the patients, which could contribute to the treatment setback. Possible new NextGen sequencing methods and approaches based on interconnection of various omics, such as genomics, especially pharmacogenetics, transcriptomics, proteomics and metabolomics, will bring new breakthrough findings in the future. At this time, we can surely state that there is a long way ahead to fully understand this complex disease.

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**References**

1. Petrovič D (2013) Candidate genes for proliferative diabetic retinopathy. Biomed Res Int 2013:540416. doi:10.1155/2013/540416
2. van Dieren S, Beulens JW, van der Schouw YT, Grobbee DE, Neal B (2010) The global burden of diabetes and its complications: an emerging pandemic. Eur J Cardiovasc Prev Rehabil 17(Suppl 1):S3–S8. doi:10.1097/01.hjr.0000368191.86614.5a
3. Singh K, Kant S, Singh VK, Agrawal NK, Gupta SK, Singh K (2014) Toll-like receptor 4 polymorphisms and their haplotypes modulate the risk of developing diabetic retinopathy in type 2 diabetes patients. Mol Vis 20:704–713 (eCollection 2014)
4. Matthews DR, Stratton IM, Aldington SJ, Holman RR, Kohner EM (2004) Risks of progression of retinopathy and vision loss related to tight blood pressure control in type 2 diabetes mellitus: UKPDS 69. Arch Ophthalmol 122:1631–1640
5. Lim LS, Wong TY (2012) Lipids and diabetic retinopathy. Expert Opin Biol Ther 12:93–105. doi:10.1517/14712598.2012.641531
6. Chew EY, Ambrosius WT, Davis MD, Danis RP, Ganganaputra S, Greven CM et al (2010) Effects of medical therapies
on retinopathy progression in type 2 diabetes. N Engl J Med 363:233–244. doi:10.1056/NEJMoa1001288

7. Yang J, Fan XH, Guan YQ, Li Y, Sun W, Yang XZ et al (2010) MMP-2 gene polymorphisms in type 2 diabetes mellitus retinopathy. Int J Ophthalmol 3:137–140. doi:10.3980/j.issn.2222-3959.2010.02.10

8. Wong TY, Klein R, Islam FM, Cotch MF, Folsom AR, Klein BE (2010) Prevalence of diabetic retinopathy in the United States, 2005–2008. JAMA 304:649–656. doi:10.1001/jama.2010.1111

9. Zhang X, Saadine JB, Assis N, Rocha MN, Calliari LE, Longui CA, Monte O, de Melo MB (2007) Evaluation of AC(n) and C–106T polymorphisms of the aldose reductase gene in Brazilian patients with DM1 and susceptibility to diabetic retinopathy. Mol Vis 13:740–745

10. Santos KG, Tschiedel B, Schneider J, SOtouo K, Roisenberg I (2003) Diabetic retinopathy in Euro-Brazilian type 2 diabetic patients: relationship with polymorphisms in the aldose reductase, the plasminogen activator inhibitor-1 and the methylenetetrahydrofolate reductase genes. Diabetes Res Clin Pract 61:133–136

11. Kuo JZ, Wong TY, Rotter JI (2014) Challenges in elucidating the genetics of diabetic retinopathy. JAMA Ophthalmol 132: 96–107. doi:10.1001/jamaophthalmol.2013.5024.

12. Leslie RD, Pyke DA (1982) Diabetic retinopathy in identical twins. Diabetes 31:19–21

13. Looker HC, Nelson RG, Chew E, Klein R, Klein BE, Knowler WC et al (2007) Genome-wide linkage analyses to identify loci for diabetic retinopathy. Diabetes 56:1160–1166

14. Kinoshita JH (1990) A thirty year journey in the polyol pathway. Exp Eye Res 50:567–573

15. Abhary S, Hewitt AW, Burdon KP, Craig JE (2009) A systematic meta-analysis of genetic association studies for diabetic retinopathy. Exp Eye Res 86:647–652. doi:10.1016/j.exer.2008.01.009

16. Ng DPK (2010) Human genetics of diabetic retinopathy: current perspectives. J Ophthalmol. doi:10.1155/2010/172593

17. Mohamed Q, Gillies MC, Wong TY (2009) Management of diabetic retinopathy: a systematic review. JAMA 298:902–916

18. Amano S, Yamagishi SI, Koda Y, Tsuneoka M, Soejima M, Nakagawa K et al (2006) Prevalence of diabetic retinopathy in a multi-ethnic cohort in the United States. Am J Ophthalmol 142:446–455

19. Mohamed Q, Gillies MC, Wong TY, Klein R, Islam FM, Cotch MF et al (2007) Genome-wide linkage analyses to identify loci for diabetic retinopathy. JAMA Ophthalmol 132: 96–107. doi:10.1001/jamaophthalmol.2013.5024.

20. Ko BC, Lam KS, Wat NM, Chung SS (1995) An (A-C)n dinucleotide repeat polymorphism in the 5′-untranslated region of the aldose reductase gene is associated with diabetic retinopathy. Diabetes 44:727–732

21. Fujsiawa T, Ichikawa F, Yamada K, Miyagawa H, Uchiyashiki Y, Oniki K, Sakata M, Kajiwara A, Yoshida A, Jinnouchi H, Nakagawa K (2013) Association between aldehyde dehydrogenase 2 polymorphisms and the incidence of diabetic retinopathy among Japanese subjects with type 2 diabetes mellitus. Cardiovasc Diabetol 12:132. doi:10.1186/1475-2840-12-132

22. Abhary S, Burdon KP, Gupta A et al (2009) Common sequence variation in the VEGFA gene predicts risk of diabetic retinopathy. Invest Ophthalmol Vis Sci 50:5552–5558. doi:10.1167/iovs.09-3694

23. Awata T, Inoue K, Kurihara S, Ohkubo T, Watanabe M, Inukai K et al (2002) A common polymorphism in the 5′-untranslated region of the VEGF gene is associated with diabetic retinopathy in type 2 diabetes. Diabetes 51:1635–1639

24. Ulthra S, Raman R, Mukes BN, Rajkumar SA, Padmaja KR, Paul PG et al (2008) Association of VEGF gene polymorphisms with diabetic retinopathy in a south Indian cohort. Ophthalmic Genet 29:11–15. doi:10.1080/13880609.2010.501655

25. Kangas-Kontio T, Vavuli S, Kakko SJ, Penna J, Savolainen ER, Linnaamaa (2009) Polymorphism of the manganese superoxide dismutase gene but not of vascular endothelial growth factor gene is a risk factor for diabetic retinopathy. Br J Ophthalmol 93:14011406. doi:10.1136/bjo.2008.159012

26. Suganthalakshmi B, Anand R, Kim R, Mahalakshmi R, Karthi-rakash S, Namperumalsamy P et al (2006) Association of VEGF and eNOS gene polymorphisms in type 2 diabetic retinopathy. Mol Vis 12:336–341

27. Buraczynska M, Ksiazek P, Baranowicz-Gaszczyk I, Jozwiak L, Cebula K et al (2007) Association of the VEGF gene polymorphism with diabetic retinopathy in type 2 diabetes patients. Nephrol Dial Transplant 22:827–832

28. Szaflik JP, Wysocki T, Kowsalski M, Majsterek I, Borucza AI, Blasiak J, Szaflik J (2008) An association between vascular endothelial growth factor gene promoter polymorphisms and diabetic retinopathy. Graefes Arch Clin Exp Ophthalmol 246:39–43
40. Nakamura S, Iwasaki N, Funatsu H, Kitano S, Iwamoto Y (2009) Impact of variants in the VEGF gene on progression of proliferative diabetic retinopathy. Graefes Arch Clin Exp Ophthalmol 247:21–26. doi:10.1007/s00417-008-0915-3

41. Qiu M, Xiong W, Liao H, Li F (2013) VEGF–634G > C polymorphism and diabetic retinopathy risk: a meta-analysis. Gene 518:310–315. doi:10.1016/j.gene.2013.01.018

42. Churchill AJ, Carter JG, Ramsden C, Turner SJ, Yeung A, Qiu M, Xiong W, Liao H, Li F (2013) VEGF–634G > C polymorphism and diabetic retinopathy: a meta-analysis. PLoS One 884069. doi:10.1371/journal.pone.0084069.

43. Nakanshi K, Watanabe C (2009) Single nucleotide polymorphisms of vascular endothelial growth factor gene intron 2 are markers for early progression of diabetic retinopathy in Japanese with type 1 diabetes. Clin Chim Acta 402:171–175

44. Chun MY, Hwang HS, Cho HY, Chun HJ, Woo JT, Lee KW et al (2010) Association of vascular endothelial growth factor polymorphisms with nonproliferative and proliferative diabetic retinopathy. J Clin Endocrinol Metab 95:3547–3551. doi:10.1210/jc.2009-2719

45. Yang X, Deng Y, Gu H, Lim A, Altankhuyag A, Jia W et al (2011) Polymorphisms in the vascular endothelial growth factor gene and the risk of diabetic retinopathy in Chinese patients with type 2 diabetes. Mol Vis 17:3088–3096

46. Zeng Y, Dai F, Yang K, Tang Y, Xu M, Zhou Y (2015) Association between a vascular endothelial growth factor gene polymorphism (rs2146323) and diabetic retinopathy: a meta-analysis. BMC Ophthalmol 15:163. doi:10.1186/s12886-015-0155-3

47. Pororjan MD, Cătană A, Popp RA, Dumitrascu DL, Bala C (2015) Polymorphisms in the CTSH gene may influence the progression of diabetic retinopathy: a candidate-gene study in the Danish Cohort of Pediatric Diabetes 1987 (DCPD1987). Graefes Arch Clin Exp Ophthalmol 253:1959–1965. doi:10.1007/s00417-015-3118-8

48. Ng ZX, Kuppusamy UR, Poh R, Tajunisah I, Koay A, Fong K et al (2012) Lack of association between Gly82Ser, 1704G/T and 2184A/G RAGE gene polymorphisms and retinopathy susceptibility in Malaysian diabetic patients. Genet Mol Res 11:455–461. doi:10.4238/2012.March.12.2

49. Zhou JB, Yang JK (2010) Angiotensin-converting enzyme gene polymorphism is associated with proliferative diabetic retinopathy: a meta-analysis. Acta Diabetol Suppl 1:187–193. doi:10.1007/s00592-009-0160-1

50. Li Y, Ye Q, Hu Q, Shi Y, Xue C, Shi Y et al (2012) Association of angiotensin-converting enzyme gene polymorphism with diabetic retinopathy in the Chinese population. J Renin Angiotensin Aldosterone Syst 13:289–295. doi:10.1177/1470320311432187

51. Liang S, Pan M, Hu N, Wu YY, Chen H, Zhu J et al (2013) Association of angiotensin-converting enzyme gene 2350 G/A polymorphism with diabetic retinopathy in Chinese Han population. Mol Biol Rep 40:463–468. doi:10.1007/s11033-012-2081-2

52. Nikzamir A, Rashidi A, Esteghamati A, Nakhjavani M, Golmohammadi T, Khalilzadeh O (2010) The relationship between ACE gene insertion/deletion polymorphism and diabetic retinopathy in Iranian patients with type 2 diabetes. Ophthalmic Genet 31:108–113. doi:10.3233/DMA-2011-0863

53. Saleem S, Azam A, Masqood SI, Muslim I, Bashir S, Fazal N et al (2015) Role of ACE and PAI-1 Polymorphisms in the Development and Progression of Diabetic Retinopathy. PLoS One 10:e0144557. doi:10.1371/journal.pone.0144557

54. Cilenšek I, Mankoč S, Petrovič MG, Petrovič D (2012) GSTT1 null genotypes is a risk factor for diabetic retinopathy in Caucasians with type 2 diabetes, whereas GSTM1 null genotype might confer protection against retinopathy. Dis Markers 2184A/G RAGE gene polymorphisms and retinopathy susceptibility in Malaysian diabetic patients. J Diabetes 3:296–303. doi:10.1155/2013/264579.

55. Vanita V (2014) Association of RAGE (p.Gly82Ser) and MnSOD (p.Val16Ala) polymorphisms with diabetic retinopathy in T2DM patients from north India. Diabetes Res Clin Pract 104:155–162. doi:10.1016/j.diabres.2013.12.059

56. Kumaramanickavel G, Ramprasad VL, Sripiya S, Upaday NK, Paul PG, Sharma T (2002) Association of Gly82Ser polymorphism in the RAGE gene with diabetic retinopathy in type II diabetic Asian Indian patients. J Diabetes Complications 16:391–394

57. Yang Y, Wu Q, Li Y, Fan X, Hao Y, Sun H et al (2013) Association of the receptor for advanced glycation end products gene polymorphisms and circulating RAGE levels with diabetic retinopathy in Chinese population. J Diabetes Res 2013:264579. doi:10.1155/2013/264579.

58. Globocnik Petrovic M, Steblovnik K, Peterlin B, Petrovic D (2003) The –429T/C and –374T/A gene polymorphisms of the receptor of advanced glycation end products gene are not risk factors for diabetic retinopathy in Caucasians with type 2 diabetes. Klin Monbl Augenheilkd 220:873–876

59. Thorsen SU, Sandahl K, Nielsen LB, Broe R, Rasmussen ML, Peto T, Grauslund J, Andersen ML, Mortensen HB, Pociot F, Olsen BS, Broorsson C (2015) Polymorphisms in the CTSH gene may influence the progression of diabetic retinopathy: a candidate-gene study in the Danish Cohort of Pediatric Diabetes 1987 (DCPD1987). Graefes Arch Clin Exp Ophthalmol 253:1959–1965. doi:10.1007/s00417-015-3118-8

60. Ng ZX, Kuppusamy UR, Poh R, Tajunisah I, Koay A, Fong K et al (2012) Lack of association between Gly82Ser, 1704G/T and 2184A/G RAGE gene polymorphisms and retinopathy susceptibility in Malaysian diabetic patients. Genet Mol Res 11:455–461. doi:10.4238/2012.March.12.2

61. Zhou JB, Yang JK (2010) Angiotensin-converting enzyme gene polymorphism is associated with proliferative diabetic retinopathy: a meta-analysis. Acta Diabetol Suppl 1:187–193. doi:10.1007/s00592-009-0160-1

62. Li Y, Ye Q, Hu Q, Shi Y, Xue C, Shi Y et al (2012) Association between angiotensin-converting enzyme gene polymorphism and diabetic retinopathy in the Chinese population. J Renin Angiotensin Aldosterone Syst 13:289–295. doi:10.1177/1470320311432187

63. Liang S, Pan M, Hu N, Wu YY, Chen H, Zhu J et al (2013) Association of angiotensin-converting enzyme gene 2350 G/A polymorphism with diabetic retinopathy in Chinese Han population. Mol Biol Rep 40:463–468. doi:10.1007/s11033-012-2081-2

64. Nikzamir A, Rashidi A, Esteghamati A, Nakhjavani M, Golmohammadi T, Khalilzadeh O (2010) The relationship between ACE gene insertion/deletion polymorphism and diabetic retinopathy in Iranian patients with type 2 diabetes. Ophthalmic Genet 31:108–113. doi:10.3233/DMA-2011-0863

65. Saleem S, Azam A, Masqood SI, Muslim I, Bashir S, Fazal N et al (2015) Role of ACE and PAI-1 Polymorphisms in the Development and Progression of Diabetic Retinopathy. PLoS One 10:e0144557. doi:10.1371/journal.pone.0144557

66. Cilenšek I, Mankoč S, Petrovič MG, Petrovič D (2012) GSTT1 null genotypes is a risk factor for diabetic retinopathy in Caucasians with type 2 diabetes, whereas GSTM1 null genotype might confer protection against retinopathy. Dis Markers 3:296–303. doi:10.3233/DMA-2011-0863

67. Petrovič MG, Cilenšek I, Petrovič D (2008) Manganese superoxide dismutase gene polymorphism (V16A) is associated with diabetic retinopathy in Slovene (Caucasians) type 2 diabetes patients. Dis Markers 24:59–64

68. Tian C, Fang S, Du X, Ji C (2011) Association of the C47T polymorphism in SOD2 with diabetes mellitus and diabetic microvascular complications: a meta-analysis. Diabetologia 54:803–811. doi:10.1007/s00125-010-2004-5

69. Cheema BS, Kohli HS, Sharma R, Bhanasai A, Khullar M (2012) Endothelial nitric oxide synthase gene polymorphism and type
2 diabetic retinopathy among Asians Indians. Acta Diabetol 49:481–488. doi:10.1007/s00592-012-0437-7
71. Zhao S, Li T, Zheng B, Zheng Z (2012) Nitric oxide synthase 3 (NOS3) 4b/a, T-786C and G894T polymorphisms in association with diabetic retinopathy susceptibility: a meta-analysis. Ophthalmic Genet 33:200–207. doi:10.3109/13816810.2012.675398
72. Santos KG, Crispim D, Canani LH, Ferrugem PT, Gross JL, Roisenberg I (2012) Relationship of endothelial nitric oxide synthase (eNOS) gene polymorphisms with diabetic retinopathy in Caucasians with type 2 diabetes. Ophthalmic Genet 33:23–27. doi:10.3109/13816810.2012.620057
73. Chen Y, Huang H, Zhou JM, Doumatey A, Lashley K, Chen G et al (2007) Polymorphism of the endothelial nitric oxide synthase gene is associated with diabetic retinopathy in a cohort of West Africans. Mol Vis 26:2142–2147
74. Bazzaz JT, Amoli MM, Pravica V, Chandrasecaran R, Boulton AJ, Larjiani B, Hutchinson IV (2010) eNOS gene polymorphism association with retinopathy in type 1 diabetes. Ophthalmic Genet 31:103–107. doi:10.3109/13816810.2010.482553
75. Brondani LA, de Souza BM, Duarte GCK, Kliemann LM, Esteves JF, Marcon AS et al (2012) The UCPI-1 3826A/G polymorphism is associated with diabetic retinopathy and increased UCPI1 and MnSOD2 gene expression in human retina. Invest Ophthalmol Vis Sci 53:7449–7457. doi:10.1167/iovs.12-10660
76. Chai D, Wang B, Shen L, Pu J, Zhang X, He B (2008) RXR agonists inhibit high-glucose-induced oxidative stress by repressing PKC activity in human endothelial cells. Free Radic Biol Med 44:1334–1347. doi:10.1016/j.freeradbiomed.2007.12.022
77. Roy M, Hallman M, Fu Y, Machado M, Hanis CL (2009) Assessment of 193 candidate genes for retinopathy in African Americans with type 1 diabetes. Arch Ophthalmol 127:605–612. doi:10.1001/archophthalmol.2009.48
78. Hsieh CH, Pei D, Hung YJ, Hsiao FC (2011) Association between retinoid-X receptor-gamma genetic polymorphisms and diabetic retinopathy. Genet Mol Res 10:3545–3551. doi:10.4238/2011
79. Zhang Y, Meng N, Lv Z, Li H, Qu Y (2015) The gene polymorphism of UCPI1 but not PPARα and TCF7L2 are associated with diabetic retinopathy in Chinese type 2 diabetes mellitus cases. Acta Ophthalmol 93:e223–e229. doi:10.1111/aos.12542
80. Crispim D, Faqundes NJ, dos Santos KG, Rheinheimer J, Bouças AP, de Souza BM et al (2010) Polymorphisms of the UCPI2 gene are associated with proliferative diabetic retinopathy in patients with diabetes mellitus. Clin Endocrinol (Oxf) 72:612–619. doi:10.1111/j.1365-2265.2009.03684.x
81. Buraczynska M, Baranowicz-Gaszczyk I, Tarach J, Ksiazek A (2009) Toll-like receptor 4 gene polymorphism and early onset of diabetic retinopathy in patients with type 2 diabetes. Hum Immunol 70:121–124. doi:10.1016/j.huimimm.2008.12.003
82. Wang J, Yang MM, Li YB, Liu GD, Teng Y, Liu XM (2013) Association of CFH and CFB gene polymorphisms with retinopathy in type 2 diabetic patients. Mediators Inflamm 2013:748435. doi:10.1155/2013/748435
83. Liu L, Jiao J, Wang Y, Wu J, Huang D, Teng W et al (2014) TGF-beta1 gene polymorphism in association with diabetic retinopathy susceptibility: a systematic review and meta-analysis. PLoS One 9:e94160. doi:10.1371/journal.pone.0094160
84. Katakami N, Matsuiha M, Kaneto H, Matsuoka TA, Imanura K, Ishibashi F et al (2010) Monocyte chemoattractant protein-1 (MCP-1) gene polymorphism as a potential risk factor for diabetic retinopathy in Japanese patients with type 2 diabetes. Diabet Med Clin Pract 89:e9–e12. doi:10.1016/j.diabmed.2010.04.006
85. Jeon HJ, Choi HJ, Park BH, Lee YH, Oh T (2013) Association on monocyte chemoattractant protein-1 (MCP-1) 2518A/G with proliferative diabetic retinopathy in Korean type 2 diabetes. Yonsei Med J 54:621–625. doi:10.3349/ymj.2013.54.3.621
86. Dong L, Lv XY, Wang BJ, Wang YQ, Mu H, Feng ZL, et al (2014) Association of monocyte chemoattractant protein-1 (MCP-1) 2518A/G polymorphism with proliferative diabetic retinopathy in northern Chinese type 2 diabetes. Graefes Arch Clin Exp Ophthalmol 252:1921–1926. doi:10.1007/s00417-014-2651-1.
87. Beráník M, Kanková K, Benes P, Izakovicová-Hollá L, Znojil V, Hajeck D et al (2002) Polymorphism R25P in the gene encoding transforming growth factor-beta (TGF-beta1) is a newly identified risk factor for proliferative diabetic retinopathy. Am J Med Genet 109:278–283
88. Sun H, Cong X, Sun R, Wang C, Wang X, Liu Y (2014) Association between the ICAM-1K469E polymorphism and diabetic retinopathy in type 2 diabetes mellitus: a meta-analysis. Diabetes Res Clin Pract 104:46–49. doi:10.1016/j.diabres.2014.01.028
89. Liu L, Yu Q, Wang H, Zhang SX, Huang C, Chen X (2006) Association of intercellular adhesion molecule 1 polymorphisms with retinopathy in Chinese patients with type 2 diabetes. Diabet Med 23:643–648
90. Petrović MG, Osredkar J, Saraga-Babić M, Petrović D (2008) K469E polymorphism of the intracellular adhesion molecule 1 gene is associated with proliferative diabetic retinopathy in Caucasians with type 2 diabetes. Clin Experiment Ophthalmol 36:468–472
91. Zhu J, Gao SH, Chi XD, Zhang YS, Lai XZ, Zhong WL (2010) Relationship between diabetic retinopathy and K469E gene polymorphism of intracellular adhesion molecule-1. J Fujian Med Univ 44:190–193
92. Su X, Chen X, Lui L, Chang X, Yu X, Sun K (2013) Intracellular adhesion molecule-1K469E gene polymorphism and risk of diabetic microvascular complications: a meta-analysis. PloS One 8:e69940. doi:10.1371/journal.pone.0069940.
93. Yu Z, Li Y, Wu Y, Qu Y (2015) Association of ICAM-1 and HMGAl gene variants with retinopathy in type 2 diabetes mellitus among Chinese individuals. Curr Eye Res 30:1–5 (Epub ahead of print)
94. Ng ZX, Kuppusamy UR, Tajunishah I, Fong KCS, Chua KH (2012) Investigation of SLC2A1 26177A/G gene polymorphism via high resolution melting curve analysis in Malaysian patients with diabetic retinopathy. J Diabetes Complications 26:388–392. doi:10.1016/j.jdiacomp.2012.05.014
95. Tariq K, Malik SB, Ali SH, Maqsood SE, Azam A, Muslim I et al (2013) Association of Pro12Ala polymorphism in peroxisome proliferator activated receptor gamma with proliferative diabetic retinopathy. Mol Vis 19:710–717
96. Luo J, Zhao L, ChenAY, ZhangX, Zhu J, Zhao J et al (2013) TCF7L2 variation and proliferative diabetic retinopathy. Diabetes 62(7):2613–2617
97. Cicicacci C, Di Fusco D, Cacciotti L, Morganti R, D’Amato C, Novelli G et al (2012) TCF7L2 gene polymorphisms and type 2 diabetes: association with diabetic retinopathy and cardiovascular autonomic neuropathy. Acta Diabetol 50(5):789–799
98. Ding Y, Hu Z, Yuan S, Xie P, Liu Q (2015) Association between transcription factor 7-like 2 rs7903146 polymorphism and diabetic retinopathy in type 2 diabetes mellitus: a meta-analysis. Diab Vasc Dis Res 12:436–444. doi:10.1177/1479164115598274
99. Ramuš SM, Kumshe T, Petrović MG, Petrović D, Cilenšek I (2013) SNP 2073618 of the osteoprotegerin gene is associated with type 2 diabetes: association with diabetic retinopathy and cardiovascular disease. Biomed Research Internat. doi:10.1155/2013/364073
100. Nagi DK, McCormack LI, Mohamed-Ali V, Vudkin JS, Knowler WC, Grant PJ (1997) Diabetic retinopathy, promoter (4G/5G) polymorphism of PAI-1 gene, and PAI-1 activity in Pima Indians with type 2 diabetes. Diabetes Care 20(8):1304–1309
101. Zhang T, Pang C, Li N, Zhou E, Zhao K (2013) Plasminogen activator inhibitor-1 4G/5G polymorphism and...
retinopathy risk in type 2 diabetes: a meta-analysis. BMC Med. doi:10.1186/1741-7015-11-1
102. Santos A, Salguero ML, Gurrola C, Munoz F, Roig-Melo E, Pando- duro A (2002) The epsilon4 allele of apolipoprotein E gene is a potential risk factor for the severity of macular edema in type 2 diabetic Mexican patients. Ophthalmic Genet 23:13–19
103. Liew G, Shankar A, Wang JJ, Klein R, Bray MS, Couper DJ et al (2006) Apolipoprotein E gene polymorphisms are not associated with diabetic retinopathy: the atherosclerosis risk in communities study. Am J Ophthalmol 142:105–111
104. Hu C, Zhang R, Yu W, Wang J, Wang C, Pang C et al (2011) CPVL/CHN2 genetic variant is associated with diabetic retinopathy in Chinese type 2 diabetic patients. Diabetes 60:3085–3089. doi:10.2337/db11-0028
105. Li H, Louey JW, Choy KW, Liu DT, Chan WM, Chan YM et al (2008) EDN1 Lys198Asn is associated with diabetic retinopathy in type 2 diabetes. Mol Vis 14:1698–1704
106. Matsubara Y, Murata M, Maruyama T, Handa M, Yamagata N, Li H, Louey JW, Choy KW, Liu DT, Chan WM, Chan YM et al (2011) Proteasome modulator 9 gene is linked to diabetic nephropathy in Chinese type 2 diabetic patients. Diabetes 60:3085–3089. doi:10.2337/db11-0028
107. Yoshioka K, Yoshida T, Takakura Y, Kogure A, Uemekawa T, Toda H, Yoshikawa T (2003) No association between the MTHFR gene polymorphism and diabetic retinopathy in type 2 diabetic patients without overt nephropathy. Diabetes Care 26:1947–1948
108. Maeda M, Yamamoto I, Fukuda M, Nishida M, Fujitsu J, Nonen S, Igarashi T, Motomura T, Inaba M, Fujio Y, Azuma J (2003) MTHFR gene polymorphism as a risk for diabetic retinopathy in type 2 diabetic patients without serum creatinine elevation. Diabetes Care 26:547–548
109. Wang J, Yang MM, Rong SS, Ng TK, Li YB, Liu XM (2013) Association of paraoxonase gene polymorphisms with diabetic nephropathy and retinopathy. Mol Med Rep 8:1845–1851. doi:10.3892/mmr.2013.1710.
110. Paine SK, Sen A, Choudhari S, Mondal LK, Cowdhury IH, Basu A et al (2012) Association of tumor necrosis factor a, interleukin 6, and interleukin 10 promoter polymorphism with proliferative diabetic retinopathy in type 2 diabetic subjects. Retina 32:1197–1203. doi:10.1097/IAE.0b013e3281225f5f
111. Kanková K, Muzík J, Karásková J, Beránek M, Hájek M, Hajiak J, Skupien J, Bobrek J, Czogala M, Klupa T, Sieradzki J (2006) Clinical risk factors and the role of VDR gene polymorphisms in diabetic retinopathy in Polish type 2 diabetes patients. Acta Diabetol 43:114–119
112. Petrović MG, Kružliak P, Petrović D (2015) The rs6060566 of vitamin D receptor gene polymorphism and clinical characteristics on risk of diabetic retinopathy in Han Chinese type 2 diabetes patients. Gene 566:212–216. doi:10.1016/j.gene.2015.04.045
113. Cygankiewicz S, Sieradzka B, Malecki MT, Wolfok P, Skupien J, Bobrek J, Czogala M, Klupa T, Sieradzki J (2006) Clinical risk factors and the role of VDR gene polymorphisms in diabetic retinopathy in Polish type 2 diabetes patients. Acta Diabetol 43:114–119
114. Hong YJ, Kang ES, Ji MJ, Choi HJ, Oh T, Koong S-S, Jeon HJ (2015) Association between BsmI polymorphism in vitamin D receptor gene and diabetic retinopathy of type 2 Diabetes in Korean population. Endocrinol Metab (Seoul) 30:469–474. doi:10.3803/EnM.2015.30.4.469
115. Sesti LF, Crispim D, Canani LH, Polina ER, Rheinheimer J, Carvalho PS, Gross JL, Santos KG et al (2013) Lack of association between the Thr431Asn polymorphism in the platelet receptor for collagen. Blood 95:1560–1564
116. Kanková K, Muzík J, Karásková J, Beránek M, Hájek M, Hajiak J, Skupien J, Bobrek J, Czogala M, Klupa T, Sieradzki J (2006) Clinical risk factors and the role of VDR gene polymorphisms in diabetic retinopathy in Polish type 2 diabetes patients. Acta Diabetol 43:114–119
117. Zhong X, DU Y, Lei Y, Liu N, Guo Y, Pan T (2015) Effects of vitamin D receptor gene polymorphism and clinical characteristics on risk of diabetic retinopathy in Han Chinese type 2 diabetes patients. Gene 566:212–216. doi:10.1016/j.gene.2015.04.045
118. Cygankiewicz S, Sieradzka B, Malecki MT, Wolfok P, Skupien J, Bobrek J, Czogala M, Klupa T, Sieradzki J (2006) Clinical risk factors and the role of VDR gene polymorphisms in diabetic retinopathy in Polish type 2 diabetes patients. Acta Diabetol 43:114–119
119. Hong YJ, Kang ES, Ji MJ, Choi HJ, Oh T, Koong S-S, Jeon HJ (2015) Association between BsmI polymorphism in vitamin D receptor gene and diabetic retinopathy of type 2 Diabetes in Korean population. Endocrinol Metab (Seoul) 30:469–474. doi:10.3803/EnM.2015.30.4.469
120. Simó-Servat O, Hernández C, Simó R (2013) Genetics in diabetic retinopathy: current concepts and new insights. Curr Genomics 14:289–299. doi:10.2174/13892029113149990008
121. Sydorova M, Lee MS (2005) Vascular endothelial growth factor levels in vitreous and serum of patients with either proliferative diabetic retinopathy or proliferative vitreoretinopathy. Ophthalmic Res 37:188–190
122. Stewart MW (2012) Aflibercept (VEGF Trap-eye): the newest anti-VEGF drug. Br J Ophthalmol 96:1157–1158. doi:10.1136/bjophthalmol-2011-300654
123. Awata T, Kurihara S, Takata N, Neda T, Iizuka H, Okhuko T et al (2005) Functional VEGF C-634G polymorphism is associated with development of diabetic macular edema and correlated with macula retinal thickness in type 2 diabetes. Biochem Biophys Res Commun 333:679–685
124. Semeraro F, Cancarini A, dell’Omo R, Rezza S, Romano MR, Costagliola C (2015) Diabetic retinopathy: vascular and inflammatory disease. J Diabetes Res Suppl 2015:582060. doi:10.1155/2015/582060
125. Quazi Y, Maddula S, Ambati BK (2009) Mediators of ocular angiogenesis. J Genet 88:495–515
126. Abcouwer SF (2013) Angiogenic factors and cytokines in diabetic retinopathy. J Clin Cell Immunol. doi:10.4172/2155-9899.1000141
127. Tarr JM, Kaul K, Chopra M, Kohner EM, Chibber R (2012) Pathophysiology of diabetic retinopathy. ISRN Ophthalmol 2013:343560. doi:10.1155/2013/343560
128. Zong H, Ward M, Stitt AW (2011) AGES, RAGE, and diabetic retinopathy. Curr Diabetes Rep 11:244–252. doi:10.1007/s11892-011-0198-7
129. Lindholm F, Bakhtadze E, Sjogren M, Cilio CM, Agardh DC et al (2005) Functional VEGF C-634G polymorphism is associated with diabetic retinopathy or proliferative vitreoretinopathy. Ophthalmic Res 37:188–190
130. Ramprasad S, Radhia V, Mathias RA, Majumder PP, Rao MR, Rema M (2007) Ragge gene promoter polymorphisms and diabetic retinopathy in a clinic-based population from South India. Eye (Lond) 21:395–401.
131. JiXiong X, BILin X, MingGong Y, ShuQin L (2003) –429T/C and –374T/A polymorphisms of RAGE gene promoter are not associated with diabetic retinopathy in Chinese patients with type 2 diabetes. Diabetes Care 26:2696–2697
132. Wilkinson-Berka JL (2006) Angiotensin and diabetic retinopathy. Int J Biochem Cell Biol 38:752–765
133. Simó-Servat O, Hernández C, Simó R (2013) Genetics in diabetic retinopathy: current concepts and new insights. Curr Genomics 14:289–299. doi:10.2174/13892029113149990008
134. Matakova T, Sivonova M, Halasova E, Mistuna D, Dzian A, Berzincé P et al (2009) Gene polymorphisms of biotransforming...
enzymes (GSTs) and their association with lung cancer in the Slovakian population. Eur J Med Res Suppl 4:275–9
135. Cilenšek I, Mankoc S, Petrovic MG, Petrovic D (2012) The 4a/4a genotype of the VNTR polymorphism for endothelial nitric oxide synthase (eNOS) gene predicts risk for proliferative diabetic retinopathy in Slovenian patients (Caucasians) with type 2 diabetes mellitus. Mol Biol Rep 39:7061–7067. doi:10.1007/s11033-012-1537-8
136. Xu Y, Jiang Z, Huang J, Menq Q, Coh P, Tao L (2015) The association between toll-like receptor 4 polymorphisms and diabetic retinopathy in Chinese patients with type 2 diabetes. Br J Ophthalmol 99:1301–1305. doi:10.1136/bjophthalmol-2015-306677
137. Liew G, Klein R, Wong TY (2009) The role of genetics in susceptibility to diabetic retinopathy. Int Ophthalmol Clin 49:35–52. doi:10.1097/IIO.0b013e31819fe6d7
138. Sobrin L, Green T, Sim X, Jensen RA, Tai ES, Tay WT et al (2011) Candidate gene association study for diabetic retinopathy in persons with type 2 diabetes: the Candidate gene Association Resources (CARE). Invest Ophthalmol Vis Sci 52:7593–7602. doi:10.1167/iovs.11-7510
139. Cunha-Vaz J, Ribeiro L, Lobo C (2014) Phenotypes and biomarkers of diabetic retinopathy: the Weisenfeld award. Invest Ophthalmol Vis Sci 55:5412–5419. doi:10.1167/iovs.14-14884
140. Gašperiková D, Tribble ND, Staník J, Hučková M, Mišovicová N, van de Bunt M et al (2009) Identification of a novel β-cell glucokinase (GCK) promoter mutation (–71G>C) that modulates GCK gene expression through loss of allele-specific sp1 binding causing mild fasting hyperglycaemia in humans. Diabetes 58:1929–1935. doi:10.2337/db09-0070
141. Zhang HM, Chen LL, Wang L, Liao YF, Wu ZH, Ye F et al (2009) Association of 1704G/T and G82S polymorphisms in the receptor for advanced glycation end products gene with diabetic retinopathy in Chinese population. J Endocrinol Invest 32:258–262
142. Chen M, Lin W, Lu C, Chen C, Huang Y, Liao W, Tsai F (2014) Chimerin 2 genetic polymorphisms are associated with non-proliferative diabetic retinopathy in Taiwanese type 2 diabetic patients. J Diabetes Complications 28:460–463. doi:10.1016/j.jdiacomp.2014.04.009
143. Petrovic MG, Hawlina M, Peterlin B, Petrovic D (2003) BglII gene polymorphism of the α2β1 integrin gene is a risk factor for diabetic retinopathy in Caucasians with type 2 diabetes. J Hum Genet 48(9):457–460
144. Niskanen L, Voutilainen-Kaunisto R, Teräsvirta M, Karvonen MK, Valke R, Pesonen U, Laakso M, Uusitupa MIJ, Koulu M (2000) Leucine 7 to proline 7 polymorphism in the neuropeptide y gene is associated with retinopathy in type 2 diabetes. Exp Clin Endocrinol Diabetes 108(3):235–236