Retinopathy in subjects with type 2 diabetes at a tertiary diabetes clinic in Durban, South Africa: Clinical, biochemical and genetic factors

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Aim: To determine the prevalence of clinical and laboratory variables and genetic polymorphisms in association with diabetic retinopathy (DR) in subjects with type 2 diabetes attending a tertiary referral diabetes clinic in Durban, South Africa.

Methods: Cross-sectional study on 292 Indian and African patients with type 2 diabetes (71.5% women).

The presence of DR was determined by direct ophthalmoscopy. Clinical and laboratory data were collected and polymorphisms in the NOS3 (rs1722009, rs2070744, rs1799983) and VEGF (rs35569394, rs2010963) genes were determined.

Results: DR was present in 113 (39%) subjects. Those with DR were older (60.6 ± 9.6 vs. 55.4 ± 12.9 years, p = 0.005), had longer duration diabetes (18.5 ± 8.8 vs. 11.9 ± 9.2 years, p < 0.0001), higher HbA1c (9.2 ± 1.8 vs. 8.8 ± 1.7%, p = 0.049), serum creatinine (106.3 ± 90.2 vs. 75.2 ± 33.4 μmol/l), triglycerides (2.1 ± 1.2 vs. 1.9 ± 1.6 mmol/l, p = 0.042), proteinuria (72% vs. 28%, p = 0.001), and used more insulin (27% vs. 18%, p < 0.0001), anti-hypertensive (95% vs. 80%, p = 0.005), had longer duration diabetes (18.5 vs. 11.1 years, p = 0.0001), and used more insulin (39% vs. 22%, p < 0.0001), anti-hypertensive (95% vs. 80%, p = 0.005), and used more lipid-lowering therapy (70% vs. 56%, p = 0.023). There was no association between DR and any of the NOS3 or VEGF gene polymorphisms studied, although there were ethnic differences. After adjustment, diabetes duration (OR 1.05, 95% CI 1.01 - 1.08), presence of proteinuria (OR 4.15, 95% CI 1.70 - 10.10) and use of insulin therapy (OR 3.38, 95% CI 1.60 - 7.12) were associated with DR.

Conclusion: Hyperglycaemia, duration of diabetes and proteinuria are associated with DR in Indian and African patients in South Africa, whereas NOS3 and VEGF gene polymorphisms were not associated with DR.

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Introduction

Diabetic retinopathy (DR) is an important cause of visual loss and the progressively increasing global prevalence of diabetes suggests that there will be increasing numbers of persons affected by this complication. In terms of the global epidemiology of DR, a recent pooled data analysis, including an excess of 22,000 subjects from 35 different studies, reported that the age-standardized prevalence of any DR in subjects aged 20–79 years is 34.6%, proliferative diabetic retinopathy (PDR) 6.96% and diabetic macular edema (DMO) 6.81% [1]. This analysis did not include any studies from Africa and data on DR in African populations are sparse [2–4]. A systematic review of diabetes in sub-Saharan Africa between 1999 and 2011 reported that the prevalence of DR varied from 7% to 63% [3]. A previous study of African and Indian subjects with long-duration diabetes in KwaZulu-Natal, South Africa, showed that of 179 subjects with type 2 diabetes, 64.5% had DR, 47.9% non-proliferative DR only, 20.8% PDR and 26% had undergone laser photocoagulation therapy [5]. In this study, the subjects with DR had higher systolic blood pressure and longer diabetes duration than those without DR.

The major risk factors for the development of DR include disease duration, degree of hyperglycaemia, hypertension, obesity, dyslipidemia and genetic factors [6]. A number of candidate genes have been shown to be associated with DR, including polymorphisms in the endothelial nitric oxide synthase (eNOS) gene, the vascular endothelial growth factor (VEGF) gene, the aldose reductase (AKR1B1) gene and the angiotensin converting enzyme gene [7]. Furthermore, some studies have shown the intron 4a allele of the eNOS gene may be associated with a reduced risk of DR [8]. However, the influence of risk factors on the development of DR has not been studied in detail in populations in Africa. The aim of the current study was to examine the association between modifiable and non-modifiable risk factors and the development of DR in a group of
subjects with type 2 diabetes attending a referral diabetes clinic in KwaZulu-Natal, South Africa.

Patients and methods

Consecutive patients with type 2 diabetes attending a specialist referral diabetes clinic at Inkosi Albert Luthuli Central Hospital, Durban, South Africa, were enrolled. All patients attending the clinic were enrolled and there were no exclusion criteria. Each patient underwent clinical assessment including anthropometry and blood pressure measurement. Direct ophthalmoscopy through a dilated pupil was conducted by three experienced diabetologists. In subjects with proliferative DR, confirmation by an ophthalmologist was obtained. Retinopathy was graded as non-proliferative or proliferative. Non-proliferative changes included the presence of microaneurysms, venous beading, hard exudates, cotton wool spots and intra-retinal hemorrhages. Proliferative changes included neovascularization, intra-vitreous hemorrhages and fibrous retinal detachment [9]. Macular edema was not assessed. All patients provided written informed consent and the study was approved by the Biomedical Research Ethics Committee of the University of KwaZulu-Natal. Venous blood was collected for measurement of creatinine, glycated hemoglobin (HbA1c), lipid levels and for extraction of genomic DNA. All subjects had spot urine collected for the measurement of albumin and creatinine excretion and determination of urine albumin–creatinine ratio (ACR).

Assay methods

Creatinine was measured by the picric acid method, HbA1c by high performance liquid chromatography, cholesterol by enzymatic method, using cholesterol esterase and cholesterol oxidase conversion, followed by Trinder endpoint, triglycerides by the Fossati three step enzymatic reaction with Trinder endpoint and high density lipoprotein (HDL) by enzymatic reaction. Low density lipoprotein (LDL) was calculated with the Friedewald equation. Urine albumin excretion was measured by polyethylene glycol-enhanced immunoturbidimetric assay. Microalbuminuria was defined as ACR 2.5–25 mg/mmol creatinine in males and 3.5–35 mg/mmol creatinine in females. Proteinuria was defined as persistent dipstick positivity.

DNA was extracted from peripheral blood leucocytes using an in-house method involving sucrose lysis, phenol-chloroform extraction and ethanol precipitation.

Polymerase chain reaction (PCR) and gel electrophoresis was used to define the number of tandem repeat sequences in intron 4 (rs61722009) of the eNOS gene (NOS3). Allelic discrimination with real-time PCR (Taqman 3100, ABI) was used to determine the prevalence of −786 C/T (rs2070744) and 894 G/T (rs1799983) polymorphisms in the NOS3 gene and −634 G/C (rs2010963) and −2549 insertion-deletion (rs35569394) polymorphisms in the VEGF gene.

Statistical methods

In bivariate analysis of risk factors associated with retinopathy, Mann Whitney tests were used for ordinal data, Pearson’s chi square tests for categorical data and independent t-tests for quantitative normal data. Binary logistic regression analysis was used to determine factors associated with DR after adjustment for confounding. Statistical analysis was performed with SPSS version 15.

Results

The total study group included 292 subjects (Indian 175; African 117). Of these, 71.5% (n = 208) were women. The mean age was 57.5 ± 12 years, mean duration of diabetes 14.48 ± 9.59 years and mean HbA1c 9.06 ± 1.83%.

African subjects were older at diagnosis (45.4 ± 9.5 vs. 41.5 ± 12.0 years, p = 0.003), had a higher BMI (35.7 ± 6.5 vs. 30.0 ± 5.3 kg/m², p < 0.0001), higher SBP (138 ± 20 vs. 133 ± 15 mm Hg, p = 0.033), higher DBP (77 ± 10 vs. 74 ± 10 mm Hg, p = 0.025), lower total cholesterol (4.3 ± 1.0 vs. 4.6 ± 1.1 mmol/l, p = 0.019), lower triglycerides (1.7 ± 0.8 vs. 2.3 ± 1.8 mmol/l, p = 0.0003) and fewer were treated with statins (47.9% vs. 76%, p < 0.0001) [Table 1]. Fewer of the African subjects were current smokers (3.4% vs. 16.4%, p = 0.002). There was no difference between African and Indian subjects for age, number on anti-hypertensive therapy, number on insulin therapy, HbA1c, HDL cholesterol, LDL cholesterol, use of fibrates, serum creatinine and prevalence of microalbuminuria and proteinuria.

DR (non-proliferative and proliferative) was present in 113 (39%) of the subjects. Non-proliferative DR was present in 67 (23%) and PDR in 46 (15.8%) subjects. There was no difference in the prevalence of either non-proliferative or proliferative DR between African and Indian subjects (Africans, non-proliferative DR 19.7%, PDR 14.5%; Indians, non-proliferative DR 23.4%, PDR 16.0%, p = 0.996).

Subjects with any DR were compared to those with no DR for clinical, laboratory and genetic variables. Table 2 shows the clinical and biochemical variables. Both groups had a similar gender distribution and ethnic variation. When compared with subjects without DR, those with DR were older (60.6 ± 9.6 vs. 55.4 ± 12.9 years, p = 0.005) and had longer duration diagnosed diabetes (18.5 ± 8.8 vs. 11.9 ± 9.2 years, p < 0.0001). A higher proportion used insulin therapy (78% vs. 39%, p = 0.0001) and anti-hypertensive therapy (95% vs. 80%, p = 0.0003) and there were more smokers (58% vs. 42%, p = 0.046).

Subjects with DR had higher HbA1c (9.2 ± 1.8 vs. 8.8 ± 1.7% [77 vs. 73 mmol/mol], p = 0.049) and serum creatinine (106.3 ± 90.2 μmol/l vs. 75.2 ± 33.4 μmol/l, p < 0.0001). Microalbuminuria was present in fewer subjects with DR (38.5% vs. 61.5%, p = 0.001) whereas overt proteinuria was present in more subjects with DR (72% vs. 28%, p = 0.001).

Table 1: Clinical and laboratory characteristics of Indian and African subjects (n = 292)

| Gender | Indian | African | p-value |
|--------|--------|---------|---------|
| Male   | 61 (35) | 25 (21) | 0.02 |
| Female | 114 (65) | 92 (79) |         |
| Age (years) | 56.3 ± 12.0 | 59.2 ± 11.6 | 0.054 |
| Age at diagnosis (years) | 41.5 ± 12.0 | 45.4 ± 9.5 | 0.003 |
| Duration diabetes (years) | 14.9 ± 10.1 | 13.8 ± 8.9 | 0.4 |
| Smoking (%) | 29 (16.4) | 4 (3.4) | 0.002 |
| Insulin therapy (%) | 124 (71) | 83 (71) | 0.9 |
| Anti-hypertensive therapy (%) | 149 (85) | 108 (92) | 0.1 |
| Statin therapy (%) | 133 (76) | 56 (47.9) | <0.0001 |
| Body mass index (kg/m²) | 30.0 ± 5.3 | 35.7 ± 6.5 | <0.0001 |
| Systolic BP (mm Hg) | 133 ± 15 | 138 ± 20 | 0.033 |
| Diastolic BP (mm Hg) | 74 ± 10 | 77 ± 10 | 0.025 |
| HbA1c, (%) [mmol/mol] | 9.0 ± 1.7 [75] | 9.2 ± 2.0 [77] | 0.2 |
| Creatinine (μmol/l) | 84 ± 47 | 92 ± 83 | 0.4 |
| Microalbuminuria (%) | 65 (37) | 33 (28.2) | 0.2 |
| Proteinuria (%) | 21 (12) | 11 (9.4) | 0.5 |
| Cholesterol (mmol/l) | 4.6 ± 1.1 | 4.3 ± 1.0 | 0.019 |
| Triglycerides (mmol/l) | 2.3 ± 1.8 | 1.7 ± 0.8 | 0.0003 |
| HDL cholesterol (mmol/l) | 1.2 ± 0.3 | 1.2 ± 0.3 | 0.3 |
| LDL cholesterol (mmol/l) | 2.4 ± 1.0 | 2.3 ± 0.9 | 0.3 |
Table 2
Clinical and laboratory characteristics of subjects with and without diabetic retinopathy (n = 292)

|                          | Retinopathy | No retinopathy | p-value |
|--------------------------|-------------|----------------|---------|
| N = 113 (39%)            | N = 179 (61%)|
| Gender                   |             |                |         |
| Male                     | 34          | 50             | 0.6     |
| Female                   | 79          | 129            |         |
| Ethnic group             |             |                |         |
| Indian                   | 72          | 103            | 0.2     |
| African                  | 41          | 76             |         |
| Age (years)              | 60.6 ± 9.6  | 55.4 ± 12.9    | 0.005   |
| Duration diabetes (years)| 18.5 ± 8.8  | 11.9 ± 9.2     | <0.0001 |
| Smoking (%)              | 58.1        | 41.9           | 0.046   |
| Insulin therapy (%)      | 89          | 58             | 0.0001  |
| Anti-hypertensive therapy (%) | 96       | 80             | 0.0003  |
| Lipid-lowering therapy (%) | 70        | 56             | 0.023   |
| Body mass index (kg/m²)  | 31.6 ± 6.4  | 32.5 ± 6.3     | 0.2     |
| Systolic BP (mm Hg)      | 136.7 ± 17.7| 134.3 ± 17.03  | 0.2     |
| Diastolic BP (mm Hg)     | 72.3 ± 9.5  | 76.7 ± 10.0    | <0.0001 |
| HbA1c (%) [mmol/mol]     | 9.2 ± 1.8   | 8.8 ± 1.7      | 0.049   |
| Creatinine (μmol/l)      | 106.3 ± 90.2| 75.2 ± 33.5    | <0.0001 |
| Microalbuminuria (%)     | 38.5        | 61.5           | 0.001   |
| Proteinuria (%)          | 71.9        | 28.1           | 0.001   |
| Cholesterol (mmol/l)     | 4.6 ± 1.2   | 4.4 ± 0.9      | 0.1     |
| Triglycerides (mmol/l)   | 2.1 ± 1.2   | 1.9 ± 1.6      | 0.042   |
| HDL cholesterol (mmol/l) | 1.1 ± 0.2   | 1.2 ± 0.3      | 0.3     |
| LDL cholesterol (mmol/l) | 2.3 ± 0.9   | 2.3 ± 0.7      | 0.5     |

One hundred and seventy nine subjects were treated with statins (either simvastatin or atorvastatin), of whom 79 (44%) had DR and 100 (56%) had no DR. Only 6 subjects were treated with fibrates (1 African and 5 Indian subjects, all treated with bezafibrate).

When risk factors for DR were stratified according to ethnicity, there was no difference between African and Indian subjects.

Genetic data was available for 288 subjects (117 with DR and 171 with no DR). As shown in Table 3, none of the polymorphisms studied were associated with the presence of DR. Furthermore haplotypes of NOS3 were not found to be significantly different between those with and those without DR. Although the prevalence of polymorphisms at each of the five loci was found to vary between African and Indian subjects, analysis of NOS3 and VEGF gene polymorphisms within each ethnic group separately also failed to show any association with the presence of DR.

In logistic regression analysis, duration (years) of diagnosed diabetes (OR 1.05, 95% CI 1.01–1.08, p = 0.006), presence of proteinuria (OR 4.15, 95% CI 1.7–10.1, p = 0.002) and the use of insulin therapy (OR 3.4, 95% CI 1.6–7.1, p = 0.001) was associated with DR. There was no association with BMI, blood pressure, lipid levels or NOS3 or VEGF genotype.

Discussion

The current study has shown that subjects with DR were older, had longer duration diagnosed diabetes, higher HbA1c, higher serum creatinine and prevalence of proteinuria, more often treated with anti-hypertensive therapy and lipid-lowering therapy and more were smokers. The study failed to show an association with polymorphisms in NOS3 or VEGF genes.

Meta-analysis of the prevalence and risk factors for DR has shown that the overall global prevalence of any DR in subjects aged 20–79 years is 34.6% and that of PDR is 6.96% [1]. This report, however, did not include any studies from Africa. The prevalence of any DR in the current study (39%) is similar to the global estimate, although more subjects with PDR were identified in the current study (15.8%). Another study in an African population, conducted in Malawi, reported a similar overall prevalence of any DR to the current study (32.5%) [10]. The prevalence of retinopathy in the Diabcare Africa Study was lower than that recorded both in the current study and the Malawian study although the incomplete ascertainment of data (51% of subjects had an eye examination in the preceding 12 months), suggests that the prevalence may have been under-estimated [2].

The findings in the current study concur with the global meta-analysis that major risk factors for DR include diabetes duration and HbA1c. However, although HbA1c was significant in bivariate analysis, it failed to achieve significance in multivariate analysis when controlled for other risk factors. This may be partly accounted for by the cross-sectional nature of this study. The current study failed to confirm an association between DR and blood pressure and this is possibly explained by the higher proportion of those with DR receiving anti-hypertensive therapy. The measured blood pressures therefore, reflect the efficacy of drug therapy, thereby mitigating the influence of recorded blood pressure on DR. In addition, although data on the specific type of anti-hypertensive therapy were not collected, differences in the use of ACE-inhibitors and other anti-hypertensive agents may also partly account for this observation.

Although the pathogenesis of DR is multi-factorial, hyperglycemia is a major driver of the process, leading to alteration in capillary permeability and density [11–13]. The current study has shown a relationship between DR and hyperglycemia in that those with DR had longer duration of disease, higher HbA1c and more subjects with DR were treated with insulin than those without DR. The latter observation is probably a reflection of longer diabetes duration with greater reduction in beta-cell function necessitating treatment advancement to insulin therapy. This is, however, speculative, as data on indices of beta-cell function were not collected.

Table 3
Prevalence of NOS3 and VEGF gene polymorphisms in subjects with and without diabetic retinopathy (n = 288)

| Polymorphism | Retinopathy (N = 117) | No retinopathy (N = 171) | p-value |
|--------------|-----------------------|--------------------------|---------|
| NOS3 gene    |                       |                          |         |
| rs61722009   | aa                    | 17 (10)                  | 8 (7)   | 0.313   |
|              | ab                    | 64 (37)                  | 49 (44) |         |
|              | bb                    | 89 (51)                  | 50 (45) | 0.215   |
| rs2070744    | TT                    | 117 (67)                 | 69 (62) | 0.257   |
|              | TC                    | 51 (29)                  | 39 (35) |         |
|              | CC                    | 6 (4)                    | 3 (3)   | 0.514   |
|              | T                     | 285 (82)                 | 176 (80)| 0.650   |
|              | C                     | 63 (18)                  | 44 (20) |         |
| rs1799983    | GG                    | 137 (79)                 | 81 (73) | 0.329   |
|              | GT                    | 35 (20)                  | 28 (25) |         |
|              | TT                    | 2 (1)                    | 2 (2)   | 0.952   |
|              | G                     | 309 (89)                 | 190 (86)| 0.317   |
|              | T                     | 39 (11)                  | 32 (14) |         |
| VEGF gene    |                       |                          |         |
| rs35569394   | Deletion              | 86 (49)                  | 51 (46) | 0.651   |
|              | Insertion/deletion    | 65 (37)                  | 48 (43) |         |
|              | Insertion             | 23 (13)                  | 12 (11) | 0.675   |
| rs2010963    | GG                    | 83 (48)                  | 61 (55) | 0.283   |
|              | CG                    | 79 (45)                  | 42 (38) |         |
|              | CC                    | 12 (7)                   | 8 (7)   | 0.891   |
|              | G                     | 245 (70)                 | 164 (74)| 0.422   |
|              | C                     | 103 (30)                 | 58 (26) |         |
Previous studies have reported nephropathy to be associated with DR [11] and the current study found proteinuria to be more frequent in subjects with DR, whereas microalbuminuria was more frequent in those without DR. The association between proteinuria and DR is substantiated by the finding of higher serum creatinine in subjects with DR, suggesting perhaps more renal involvement (nephropathy). By contrast, some studies have found an association between microalbuminuria and DR [14]. A recent Chinese study found microalbuminuria to be associated with an increased risk of severe or proliferative retinopathy [15]. The reason for a higher frequency of microalbuminuria in subjects without DR in the current study is not known. It is possible that subjects with DR had longer duration of disease and more often had established nephropathy (shown by a greater frequency of proteinuria), whereas those without DR and shorter duration of disease, had markers of incipient nephropathy.

The Hoorn study found a positive relationship between cholesterol and triglyceride levels and DR in older subjects with type 2 diabetes [16]. The global meta-analysis also found an association between serum cholesterol and DMO [1]. The current study failed to show an association between total cholesterol, LDL cholesterol or HDL cholesterol and DR. However, lipid-lowering therapy was used more frequently in those with DR. Whether the efficacy of lipid-lowering therapy altered the relationship between serum cholesterol and DR is speculative. The finding that subjects with DR had higher triglyceride levels than those without DR is possibly a reflection of poorer metabolic control in those with DR.

Several inflammatory mediators have been shown to play a role in the development of DR, including VEGF, nitric oxide, tumor necrosis factor alpha and interleukin-1 beta [17]. The importance of the role of VEGF in the development of DR is underscored by the efficacy of anti-VEGF intra-ocular therapy in the management of DR and particularly in the management of DMO [12,18]. Furthermore, genetic factors have been shown to have an independent effect on the development of DR and polymorphisms in a number of candidate genes, including VEGF, NOS3 and aldose reductase (AKR1B1) have been studied [7]. A recent meta-analysis showed that the rs2010963 (−634 G to C) VEGF polymorphism was associated with DR in subjects with type 2 diabetes [19]. In a separate study, no significant association was found between the rs25648, rs1570360, rs3095039, rs35569394 and rs699947 VEGF polymorphisms and DR [7]. Some studies have shown a relation between NOS3 polymorphisms and DR, although meta-analysis has excluded most of the polymorphisms studied in this gene [7,20]. In agreement with the generally negative studies of candidate genes in DR, the current study failed to show a relationship between three NOS3 polymorphisms and two VEGF gene polymorphisms and DR. Although an independent genetic risk for the development of DR may exist, the effect size is likely to be small and require large study numbers to demonstrate positive associations.

Strengths of the current study include a multi-ethnic population, allowing comparison between groups and the inclusion of gene polymorphisms in the determination of factors associated with DR. Limitations include the relatively small numbers of subjects as well as the method of determination of DR, retinal photography may have improved the diagnostic assessment of DR.

In addition the findings, in this clinic-based study, may not be able to be extrapolated to the general population in view of the possibility of referral bias.

In conclusion, the current study has shown that in subjects with type 2 diabetes in KwaZulu-Natal, the well-established risk factors for DR are affirmed and a number of candidate genes failed to show an association with DR. These findings suggest that focus on control of hyperglycemia, blood pressure and serum lipids should remain the cornerstone of therapy, aiming at preventing the development of DR and other diabetes complications.

References

[1] Yau JW, Rogers SL, Kawasaki R, Lamoureux EL, Kowalski JW, Bek T, et al., for the Meta-Analysis for Eye Disease (META-EYE) Study Group. Global prevalence and major risk factors of diabetic retinopathy. Diabetes Care 2012;35:556–64.
[2] Sobngwi E, Ndong-Mbaye M, Boateng KA, Ramaiya KL, Njenga EW, Diop SN, et al. Type 2 diabetes control and complications in specialised diabetes care centres of six sub-Saharan African countries: the Diabcare Africa study. Diabet Med 2012;29:50–6.
[3] Stitt AW, Lois N, Medina RJ, Asamson P, Curtis TM. Advances in our understanding of diabetic retinopathy risk: a meta-analysis. Gene 2013;518:310.
[4] Glover SJ, Burgess PI, Cohen DB, Harding SP, Ho HWC, Zijlstra EE, et al. Prevalence of diabetic retinopathy, cataract and visual impairment in patients with diabetes in sub-Saharan Africa. Br J Ophthalmol 2012;96:156–61.
[5] Ahlary S, Hewitt AW, Burdon KP, Craig JE. A systematic meta-analysis of genetic association studies for diabetic retinopathy. Diabetes 2009;58:2137–47.
[6] Zhao S, Li T, Zheng B, Zheng Z. Nitric oxide synthase 3 (NOS3) 4b/a, t-786C and G894T polymorphisms in association with diabetic retinopathy susceptibility: a meta-analysis. Ophthalmic Genet 2012;33:200–7.
[7] Mvitu Mauka M, Longo-Mbenza B. Causes of visual disability among Central Africans with diabetes mellitus. Afr Health Sci 2012;12:193–7.
[8] Morlala AA, Pirie FJ, Gouws E, Amod O, Omar MA. Microvascular complications in South African patients with long duration diabetes mellitus. S Afr Med J 2001;91:987–92.
[9] Ding J, Wong TY. Current epidemiology of diabetic retinopathy and diabetic macular edema. Curr Diab Rep 2012;12:346–54.
[10] Derwa M, van der Aa J, Veldhuis R, van der Graaf Y, de Groen J, de Bruijn A, et al. Prevalence and risk factors for diabetic retinopathy: a meta-analysis. Ophthalmic Genet 2012;33:309–15.
[11] van Leiden HA, Dekker JM, Nijpels G, Heine RJ, Bouter LM, et al. Blood pressure, lipids, and obesity are associated with retinopathy: the Hoorn Study. Diabetes Care 2002;25:1320–5.
[12] Tang J, Kern TS. Inflammation in diabetic retinopathy. Prog Retin Eye Res 2011;30:343–58.
[13] van der Aa J, Veldhuis R, van der Graaf Y, de Groen J, de Bruijn A, et al. Prevalence and risk factors for diabetic retinopathy. The Beijing Communities Diabetes Study. 6. Retina 2012;32:322–9.
[14] van der Aa J, Veldhuis R, van der Graaf Y, de Groen J, de Bruijn A, et al. Prevalence and risk factors for diabetic retinopathy. The Beijing Communities Diabetes Study. 6. Retina 2012;32:322–9.
[15] van der Aa J, Veldhuis R, van der Graaf Y, de Groen J, de Bruijn A, et al. Prevalence and risk factors for diabetic retinopathy. The Beijing Communities Diabetes Study. 6. Retina 2012;32:322–9.
[16] Bandello F, Lattanzio R, Zucchiatti I, Del Turco C. Pathophysiology and treatment of diabetic retinopathy. Acta Diabetol 2013;50:1–20.
[17] Stitt AW, Lois N, Medina RJ, Asamson P, Curtis TM. Advances in our understanding of diabetic retinopathy. Clin Sci 2013;125:1–17.
[18] Newman DJ, Mattcock MB, Dawanay ABS, Kerry S, McGuire A, Yaqoob M, et al. Systematic review on urine albumin testing for early detection of diabetic complications. Health Technol Assess 2005;9:iii–iv.
[19] Xu J, Wei WB, Yuan MX, Yuan SY, Wan G, Zheng YY, et al. Prevalence and risk factors for diabetic retinopathy. The Beijing Communities Diabetes Study 6. Retina 2012;32:322–9.
[20] van der Aa J, Veldhuis R, van der Graaf Y, de Groen J, de Bruijn A, et al. Prevalence and risk factors for diabetic retinopathy. The Beijing Communities Diabetes Study. 6. Retina 2012;32:322–9.