Evaluation of microaneurysms as predictors of therapeutic response to anti-VEGF therapy in patients with DME

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Abstract

Administration of intravitreal anti-vascular endothelial growth factor (anti-VEGF) therapy is the first-line therapy for diabetic macular oedema (DME). However, some patients show no or insufficient response to repeated anti-VEGF injections. Therefore, it is necessary to identify factors that can predict this resistance against anti-VEGF treatment. Presence of microaneurysms (MAs) is a predictor of the development and progression of DME, but its relationship with the treatment response to the anti-VEGF agents is not well known. Therefore, we aimed to elucidate the relationship between the distribution of MAs and the response to anti-VEGF therapy in patients with DME. The number of MAs was measured before anti-VEGF therapy in each region using fluorescein angiography, indocyanine green angiography (IA), and optical coherence tomography angiography. Patients with DME were divided into the responder and non-responder groups after three loading phases. Differences in the distribution of MAs between the groups were investigated. Pre-treatment IA revealed more MAs in the nasal area in the non-responder group than in the responder group (10.7 ± 10.7 and 5.7 ± 5.7, respectively, in the nasal macula) (1.4 ± 2.1 and 0.4 ± 0.7, respectively, in the nasal fovea). Whereas, pre-treatment FA and OCTA could not reveal significantly difference between the groups. Detection of MAs in the nasal macula using pre-treatment IA may indicate resistance to anti-VEGF therapy. We recommend the clinicians confirm the presence of MAs in the nasal macula, as shown by IA, as a predictor of therapeutic response to anti-VEGF therapy in patients with treatment naive DME.

Introduction

Diabetic macular oedema (DME) is a disease that causes vision loss in patients with diabetic retinopathy. DME is a common complication of diabetic retinopathy caused by intraretinal fluid accumulation in the macular area [1]. Current treatments for DME include focal laser photocoagulation, steroid injection, vitrectomy, and intravitreal injection of anti-vascular endothelial growth factor (anti-VEGF). Anti-VEGF therapy is usually the first-line treatment
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Materials and methods

This report was derived from a prospective multicentre clinical study [Ranibizumab exploratory study on the evaluation of the local laser combination therapy to the non-reactive group to the diabetic macular oedema patients (RELAND study)] [UMIN000024208] [jRCTs061180035]. The inclusion criteria of the study were as follows: eyes must show definite retinal thickening due to DME as revealed by clinical examination using techniques such as...
slit-lamp examination and OCT, and at least one eye of each patient must meet all of the inclusion criteria and none of the exclusion criteria (Table 1).

Both eyes of study participants were included only if both of them were eligible for study entry. For safety reasons, both eyes of such participants were not injected with the drug on the same day as part of the initial treatment. In the analysis, both eyes were treated as two independent cases. The study protocols were approved by a certified review board at the Yamaguchi University Hospital (CRB6180002) as well as institutional review boards and ethics committees. All patients provided written informed consent before their enrolment. In the loading phase, three loading doses of 0.5 mg ranibizumab, an anti-VEGF agent, were administered as part of intravitreal therapy (IVT) (one dose/month, visits 1–3); subsequently, the eyes included in this study were assigned to the responder group or non-responder group depending on the BCVA and/or central foveal retinal thickness (CRT) compared to those before the loading phase (visit 1): responder group, BCVA improvement of ≥5 letters and/or CRT improvement of ≥20%; non-responder group, BCVA improvement of <5 letters and CRT improvement of <20% from visit 1 to visit 4. Non-response in this study was not considered an adverse event.

We analysed the data from patients who had also undergone IA and/or OCTA in the RELAND study. Age, sex, BCVA, CRT, intraocular pressure (IOP), FA, IA, and OCTA data before ranibizumab IVT and BCVA and CRT data at visit 4 were analysed. The number of MAs in each area of an anatomically modified ETDRS grid (Fig 1) was measured by three retina specialists (MW, RY, and KK).

FA and IA were performed using Heidelberg Retina Angiograph 2 (Heidelberg Engineering). Data of FA in early-phase and data of IA in late-phase were analysed. OCTA was performed using DRI-OCT Triton Plus (Topcon). Scan size was 6.0 mm × 6.0 mm; scan resolution was 320 × 320; repetition was 4 times per line; MAs were analysed in deep capillary plexus layer. To mask the patient’s characteristics, only images of FA, IA, and OCTA were presented to them.

Table 1. Inclusion and exclusion criteria.

| Inclusion criteria | Exclusion criteria |
|--------------------|--------------------|
| Willingness and the ability to provide signed informed consent | History of any anti-VEGF treatment for DME |
| Age ≥ 20 years | Persistent macula edema for ≥12 months |
| Type 1 or type 2 diabetes | Macular edema considered to be due to a cause other than diabetic macular edema |
| HbA1C of <10% within two months prior to the study entry | External ocular infection or suspected infection including conjunctivitis and chalazion |
| No cerebral vascular accident nor myocardial infraction within three months prior to the study entry | Severe intraocular inflammation including uveitis, active rubeosis and endophthalmitis |

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On the number of MAs in each area, the inter-rater agreement of the measurers on FA, IA and OCTA were 0.407025, 0.540321, 0.254121, respectively. Regardless of the inter-rater agreement, the average value of the three examiners was used as the MA number.

The primary objective of this study was to determine whether the number of MAs in each macula area of non-responder group is significantly higher than the responder group. The test of the null hypothesis that the number of MAs in both groups is equal was performed by the Welch’s t-test. The test was a two-tailed test with a significance level of 5%. The confidence interval is a two-tailed and the confidence coefficient is 95%. To supplement the primary analysis, the characteristics were also analyzed by the Welch’s t-test and the chi-square test.

Results

Characteristics of participants

Forty-eight patients were included in this clinical study, all of whom received ranibizumab IVT; 12 and 36 patients were assigned to the non-responder and responder groups,
respectively. Among the 36 patients in the responder group, 8 and 16 patients showed improvements in BCVA ($\geq$5 letters) and CRT ($\geq$20%), respectively, and 12 patients showed improvements in both BCVA and CRT (Fig 3).

The characteristics of the non-responder and responder groups in this study were as follows: age, 69.1 $\pm$ 4.3 and 66.8 $\pm$ 10.3 years, respectively; male-to-female ratio, 5:7 and 25:11, respectively; BCVA before the loading phase, 69.6 $\pm$ 14.3 and 66.7 $\pm$ 16.2 letters, respectively; BCVA after the loading phase, 67.3 $\pm$ 16.3 and 71.8 $\pm$ 14.0 letters, respectively; improvement in BCVA, $-2.3 \pm 4.4$ and $5.1 \pm 7.7$ letters, respectively; CRT before the loading phase, 373 $\pm$ 139 and 389 $\pm$ 129, respectively; CRT after the loading phase, 381 $\pm$ 140 and 253 $\pm$ 111 $\mu$m, respectively; improvement in CRT, $-4.7 \pm 23.3\%$ and $32.9 \pm 20.4\%$, respectively; IOP, 14.7 $\pm$ 3.5 and 14.8 $\pm$ 3.2 mmHg, respectively (Table 2). There were significant differences in CRT at visit 3 and visit 4 and improvements in BCVA and CRT between the groups. However, there were no significant differences in BCVA and CRT before the loading phase and BCVA at visit 4.

Fig 2. Detected MA of a patient. There are MAs in each the modified ETDRS grid areas. Red arrows indicate MAs.

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Relationship between MA and therapeutic response to ranibizumab IVT

There was no significant difference in the number of MAs detected using FA before the loading phase between the non-responder and responder groups (Table 3).

The number of MAs detected using IA was similar between the groups (Table 4). Surprisingly, the non-responder group showed a higher number of MAs, mainly in the nasal area, than did the responder group. In particular, there were significant differences in the nasal macula (non-responder group, 12.5 ± 10.6; responder group, 5.7 ± 5.7, \( P < 0.05 \)) and nasal fovea (non-responder group, 1.4 ± 2.1; responder group, 0.4 ± 0.7, \( P < 0.05 \)). There was no significant difference between the groups regarding the number of MAs detected by OCTA (Table 5).

Discussion

Anti-VEGF therapy for DME improves visual acuity and anatomical macular changes [25–29]. However, all patients with DME do not always respond to anti-VEGF therapy. Therefore, it is necessary to clarify the factors influencing the resistance to anti-VEGF therapy in patients with DME. MA involves the out-punching of retinal capillaries that are weakened by the loss of pericytes, which results in retinal focal haemorrhage [6, 7]. The presence of MAs is a predictor of DME development and progression [8, 10]. MAs detected by angiographies in patients with DME also contribute to the clinical response to anti-VEGF therapy [30, 31]. Indeed, the use of laser photocoagulation for leaking MAs was found to be effective for the treatment DME [1].
In this study, we examined the distribution of MAs by using multiple angiography techniques (FA, IA, and OCTA) and then investigated the relationship between the presence or distribution of MAs and the response to anti-VEGF therapy. Notably, MAs detected by IA in the nasal area contributed to resistance of the patients to three anti-VEGF loading doses. IA appears to

Table 2. Characteristics.

|                        | Non-responder | Responder | p value |
|------------------------|---------------|-----------|---------|
| Age (years; mean ± SD) | 69.1 ± 4.3    | 66.8 ± 10.3 | 0.48507 |
| Gender (no.; Male: Female) | 5: 7          | 25: 11    | 0.08519 |
| HbA1c (%)              | 7.5 ± 1.5     | 8.2 ± 1.2  | 0.344098|
| Pre-treatment BCVA at visit 1 (ETDRS letters; mean ± SD) | 69.6 ± 14.3   | 66.7 ± 16.2 | 0.585873|
| BCVA at visit 2 (ETDRS letters; mean ± SD) | 69.6 ± 14.2   | 69.2 ± 13.3 | 0.931332|
| BCVA at visit 3 (ETDRS letters; mean ± SD) | 69.1 ± 14.5   | 68.8 ± 13.4 | 0.956494|
| BCVA at visit 4 (ETDRS letters; mean ± SD) | 67.3 ± 16.3   | 71.8 ± 14.0 | 0.359004|
| Improvement BCVA (ETDRS letters; mean ± SD) | -2.3 ± 4.4     | 5.1 ± 7.7  | 0.003007|
| Pre-treatment CRT at visit 1 (μm; mean ± SD) | 373 ± 139      | 389 ± 129  | 0.725201|
| CRT at visit 2 (μm; mean ± SD) | 363 ± 145      | 295 ± 112  | 0.096851|
| CRT at visit 3 (μm; mean ± SD) | 387 ± 133      | 275 ± 129  | 0.0125176|
| CRT at visit 4 (μm; mean ± SD) | 381 ± 140      | 253 ± 111  | 0.0023279|
| Improvement CRT rate (%) | -4.7 ± 23.3    | 32.9 ± 20.4 | 0.00000286569|
| Pre-treatment IOP (mmHg; mean ± SD) | 14.7 ± 3.5     | 14.8 ± 3.2  | 0.879601|

Data are means ± SD, p value were calculated using Welch’s t-test.

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Table 3. Detected MA by FA.

|                     | Non-responder | Responder | p value |
|---------------------|---------------|-----------|---------|
| Macula Superior     | 17.0 ± 12.2   | 14.6 ± 9.0 | 0.487802|
| Inferior            | 12.4 ± 9.3    | 9.5 ± 6.6  | 0.307722|
| Lateral             | 14.5 ± 10.8   | 10.8 ± 7.5 | 0.271135|
| Nasal               | 13.2 ± 9.7    | 9.1 ± 7.2  | 0.183823|
| Parafovea Superior  | 2.5 ± 2.3     | 1.7 ± 1.8  | 0.240609|
| Inferior            | 1.7 ± 1.4     | 1.4 ± 1.7  | 0.479289|
| Lateral             | 1.4 ± 1.5     | 2.3 ± 3.0  | 0.215847|
| Nasal               | 1.6 ± 1.2     | 1.6 ± 1.8  | 0.942946|
| Fovea Superior      | 1.5 ± 1.5     | 0.8 ± 1.0  | 0.160108|
| Inferior            | 1.0 ± 1.0     | 0.9 ± 1.7  | 0.758671|
| Lateral             | 1.3 ± 1.3     | 0.7 ± 0.7  | 0.142024|
| Nasal               | 1.5 ± 2.2     | 0.5 ± 0.9  | 0.191712|

Data are means ± SD, p value were calculated using Welch’s t-test.

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accumulate at the permeable MA that is responsible for macular oedema [32]. As a supplement, there was no significant difference of MAs in nasal parafovea area between the two groups. The nasal parafoveal area might be too narrow to allow statistical significance. Whereas, MAs detected by IA especially contribute to macular oedema responsible for DME because IA-guided navigated focal laser photocoagulation is effective for the treatment of DME [32]. These results suggest that the presence of MAs detected by IA in the nasal area of the macula was associated with resistance to three anti-VEGF loading doses. We presented the multimodal images of representative cases in both group (Figs 4 and 5).

MAs are generally the earliest signs of clinically visible retinal damage and are a hallmark of DME. Accurate identification is important because their distribution and number provide prognostic information about the severity of DM and are also clinically relevant as a guide for the treatment of DME [18]. Indeed, the reduction of macular oedema is associated with the

Table 4. Detected MA by IA.

|                      | Non-responder | Responder | p value |
|----------------------|---------------|-----------|---------|
|                      | (N = 12)      | (N = 36)  |         |
| Macula               |               |           |         |
| Superior             | 12.3 ± 10.2   | 7.8 ± 7.0 | 0.158274|
| Inferior             | 8.1 ± 7.1     | 6.4 ± 6.3 | 0.427039|
| Lateral              | 9.0 ± 7.0     | 6.3 ± 5.8 | 0.223681|
| Nasal                | 12.5 ± 10.6   | 5.7 ± 5.7 | 0.0477773|
| Parafovea            |               |           |         |
| Superior             | 2.1 ± 2.3     | 1.6 ± 2.0 | 0.464580|
| Inferior             | 0.9 ± 0.9     | 1.5 ± 2.1 | 0.236424|
| Lateral              | 1.0 ± 1.1     | 1.4 ± 1.8 | 0.451470|
| Nasal                | 1.1 ± 1.1     | 1.2 ± 1.6 | 0.844735|
| Fovea                |               |           |         |
| Superior             | 1.0 ± 0.7     | 0.6 ± 0.8 | 0.0919663|
| Inferior             | 0.5 ± 0.8     | 0.7 ± 1.6 | 0.549018|
| Lateral              | 0.7 ± 1.0     | 0.3 ± 0.7 | 0.175130|
| Nasal                | 1.4 ± 2.1     | 0.4 ± 0.7 | 0.0391342|

Data are means ± SD, p value were calculated using Welch’s t-test.

Table 5. Detected MA by OCTA.

|                      | Non-responder | Responder | p value |
|----------------------|---------------|-----------|---------|
|                      | (N = 12)      | (N = 29)  |         |
| Macula               |               |           |         |
| Superior             | 6.3 ± 3.8     | 5.3 ± 4.4 | 0.424897|
| Inferior             | 5.3 ± 3.9     | 6.2 ± 3.8 | 0.540926|
| Lateral              | 4.5 ± 3.3     | 5.5 ± 2.9 | 0.41803|
| Nasal                | 4.3 ± 3.4     | 4.2 ± 3.0 | 0.988004|
| Parafovea            |               |           |         |
| Superior             | 1.5 ± 1.7     | 1.4 ± 1.2 | 0.655697|
| Inferior             | 0.9 ± 1.0     | 1.5 ± 1.0 | 0.141042|
| Lateral              | 0.9 ± 0.9     | 1.2 ± 1.2 | 0.430182|
| Nasal                | 0.9 ± 0.7     | 1.1 ± 0.9 | 0.546775|
| Fovea                |               |           |         |
| Superior             | 0.4 ± 0.7     | 0.6 ± 0.6 | 0.617404|
| Inferior             | 0.5 ± 0.5     | 0.5 ± 0.9 | 0.970334|
| Lateral              | 0.4 ± 0.5     | 0.6 ± 0.7 | 0.424587|
| Nasal                | 0.5 ± 1.1     | 0.4 ± 0.6 | 0.997254|

Data are means ± SD, p value were calculated using Welch’s t-test.
non-perfusion area, neovascularization, and number of MAs [33]. Macular capillary blood flow in patients with DME was found to be significantly lower than that in those with non-diabetes [34]. The nasal-temporal asymmetry of blood flow in the macular capillaries was significantly high in patients with DME. Furthermore, oxygen saturation is low on the nasal side of the macula [25]. The blood flow on the nasal side of the macula may be important for maintaining macular function and is highly sensitive to small changes in blood flow or oxygen saturation. MAs also localise to the nasal side of the macula and may interfere with normal nerve signalling in the papillary macular nerve fibre bundle, which results in indirect visual loss [35].

After the administration of three loading doses of the anti-VEGF agent, MAs detected on the nasal side of the macula using IA were not responsive to anti-VEGF therapy. The presence of large MA detected by IA in the nasal macula indicates that the DME condition is physiologically and anatomically severe, and such cases are likely to resist anti-VEGF therapy.

Accurate delineation of MA distribution, that is, location and number, is important for obtaining prognostic information about the severity of DME and for planning the treatment.

Fig 4. Detected MA of a non-responder case. 68-years-old male. There was a large MA (red arrow) surrounded blockings by hard exudates in nasal macula area. IA and FA could detect the MA; OCTA could not detect.

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of DME [12]. We investigated whether there were differences in the distribution of MAs between the responder and non-responder groups, using FA, IA, and OCTA. Interestingly, the MAs detected using IA, especially those on the nasal side of the macula, were predictive of resistance to three anti-VEGF loading doses, while those detected using FA and OCTA do not have predictive value in this regard. Theoretically, considering the molecular size, the sensitivity to detect MA is highest in the order of FA, IA, and OCTA. Because the molecular weight unit is 332, 775 and 64500, respectively. In this study, the total number of MAs in Tables 3–5 was consistent with the order. Also, we selectively analysed MAs in deep capillary plexus layer. Moreover, there were not monitoring centre to check the quality of the images. The noise in OCTA might have masked MA detection. Therefore, the detection sensitivities of OCTA in this study tended lower than other reports. Whereas, IA is useful for identifying leaking spots or larger capillaries in patients with DME [32, 36]. These previous data indicate that IA could characteristically help visualise MAs because of its low detection sensitivity. There is a report that the improvement of DME by anti-VEGF therapy correlated with the size of MAs [37]. Therefore, small MA may respond well to anti-VEGF therapy. In this study, MAs detected on

Fig 5. Detected MA of a responder case. 67-years-old female. There was less MAs in nasal area in spite of many MAs in superior and inferior area. Not only FA and OCTA but also IA could not detect noticeable MAs in nasal area.

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the nasal macula using IA was less than FA and OCTA in the responder group. This was because the responder group had many small MAs of detectable size for FA but not for IA. Moreover, MAs in the nasal macula is unlikely to develop for the reasons of the blood flow and oxygen saturation. However, when MAs develop large enough to be detected by IA, they may affect the effectiveness of treatment. Therefore, we assumed that IA, but not FA and OCT, is useful in predicting the response to anti-VEGF therapy. The number of cases might be too small to be significantly different in FA and OCTA. Whereas, MAs detected using IA are also detected at the same site in FA and OCTA. Therefore, more studies with a larger number of cases are needed to describe whether such MAs can be detected in FA and OCTA.

We have shown that the presence of MA on the nasal side of the macula detected using IA may be a biomarker of resistance to anti-VEGF therapy among patients with DME. The disruption of the blood-retinal barrier by VEGF contributes to the pathogenesis of DME [38]. Indeed, anti-VEGF agents are used as the primary treatment for DME, and these agents effectively improve macular oedema and vision in most patients with DME [25–29]. It is important to identify factors that predict resistance to anti-VEGF agents because the cost of anti-VEGF drugs is a major burden for patients with DME. There are many predictors of response to anti-VEGF therapy for DME, including pre-treatment CRT, HbA1c level [27], and presence of subretinal fluid, intraretinal cysts, and renal disease [28]. We focused on the relationship between the distribution of MAs and the reactivity of the treatment. Local retinal circulation is thought to have a great influence on the formation and distribution of MAs. Further investigation is needed to correlate the distribution of MAs between these predictors.

Among participant characteristics, there were no significant differences in BCVA from visit 1 to visit 4 (Table 2). There are studies in which some cases of DME did not show improvement despite the administration of intensive anti-VEGF therapy [4, 5]. Although criteria for therapeutic response to DME varied, these studies reported resistance to anti-VEGF therapy in 18–30% of DME eyes. Consistent with the data of previous studies, 25% (12/48) of DME eyes showed resistance in this study. Our results can also be explained by the fact that the pre-treatment BCVA was relatively better that that in other clinical studies [28, 39] and that the responder group showed lesser BCVA improvement than CRT improvement (Fig 3). The limitations of the study were the exploratory decision of the reactivity criteria and the unevenness of improvement in BCVA and CRT in the responder group.

In conclusion, detection on MAs in the nasal macular area using IA before anti-VEGF therapy of treatment naïve DME might indicate treatment resistance. We recommend the confirmation of the presence of MAs in the nasal macular area using IA to predict the therapeutic response to anti-VEGF therapy in treatment naïve DME eyes.

Supporting information

S1 Table. Minimal data set of characteristics. Sex, age, HbA1c, CRT, BCVA of each group are showed.
(XLSX)

S2 Table. Minimal data set of the number of detected MAs by FA. The number of MAs counted by three measurers in each area on FA.
(XLSX)

S3 Table. Minimal data set of the number of detected MAs by IA. The number of MAs counted by three measurers in each area on IA.
(XLSX)
S4 Table. Minimal data set of the number of detected MAs by OCTA. The number of MAs counted by three measurers in each area on OCTA.

(XLSX)

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