Diabetic Retinopathy Clinical Research Network: Diabetic retinopathy and diabetic macular edema

Red de Investigación de Retinopatía Diabética: Retinopatía diabética y edema macular diabético


Retina and vitreous department, Fundación Hospital Nuestra Señora de la Luz, IAP, Mexico City, Mexico

Abstract
The diabetic retinopathy clinical research network (DRCR.net), founded by the National Eye Institute, was formed in 2002. Currently it includes over 400 ophthalmologists in 115 clinical centers in 37 states of the United States. Since then, it has conducted several studies on various points of interest about diabetic retinopathy and diabetic macular edema. To date, the DRCR.net has published about 80 articles, including clinical trials (named with alphabetic letters), observational studies and various sub analyses derived from the primary studies. The results obtained provide retina specialists with substantial evidence regarding the treatment protocols classically proposed. Also, they evaluate new treatment guidelines and information concerning other important points about the disorders studied. The large amount of information provided by the protocols and the corresponding sub analyses make understanding them a difficult task. In this work, we intend to emphasize and summarize the essential points and conclusions of each of the protocols already published by the DRCR.net, highlighting their importance in clinical practice, which will result in better decisions for the individualized treatment of diabetic patients.

Key words: Clinical trial. Analysis. Diabetic retinopathy. Diabetic macular edema.
los puntos importantes y conclusiones de cada uno de los protocolos ya publicados por el DRCR.net, y remarcar su importancia en la práctica clínica, lo que resultará en mejores decisiones para el tratamiento individualizado de los pacientes diabéticos.

Palabras clave: Ensayo clínico, Análisis, Retinopatía diabética. Edema macular diabético.

Protocolo A

El Early Treatment Diabetic Retinopathy Study (ETDRS) mostró que el fotocoagulación focal o grid redujo el riesgo de pérdida visual moderada (MVL) causado por el edema macular diabético (DME) en 50% durante tres años después de su aplicación. La efectividad del laser no podría ser completamente explicada, aunque se sugirió que la cierre del filtrado causado por los microaneurismas podría ser uno de los mecanismos más importantes, al menos en parte. Posteriormente, algunos clínicos observaron una respuesta favorable en relación con la aplicación de focal/grid laser, no impactando realmente en los microaneurismas y, de igual manera, se observaron efectos adversos después de aplicaciones en áreas muy cerca de la fovea. Esto llevó al idea de generar un nuevo programa de tratamiento diferente al propuesto por el ETDRS.

Protocolo A comparó dos técnicas de fotocoagulación para el tratamiento del DME: la técnica modificada propuesta por el ETDRS (directo o grid laser) (grupo 1) versus una malla macular suave (MMG) técnica (grupo 2). El primer intenta reducir los efectos adversos observados por el espesor de las cicatrices en las zonas a distancia de la fovea. El segundo es una alternativa propuesta por el DRCR.net, que sugiere aplicar los puntos con menos potencia y más alejados de la fovea, con un espacio-interpunto mayor. Característicamente, los puntos deben aplicarse en el retinón sano, evitando la directa aplicación en los microaneurismas. Las técnicas de tratamiento se detallan en la tabla 1.

Se seleccionaron 263 pacientes (323 ojos) con no tratamiento DME para recibir uno de los dos programas de fotocoagulación. La agudeza visual (VA) se analizó y la correlación con el tiempo fue tomado con el Stratus 3.0. La fotografía fundus estándar del estudio ETDRS se tomó en la base y en cada visita (3,5 meses, 8 meses y 12 meses). La angiografía fluoresceína (FAG) se tomó en la base y a un año. La técnica de fotocoagulación se repetía en cada visita si el edema persistía.

El objetivo principal fue observar cambios en varios medidas de TD-OCT a 12 meses. Cambios en la agudeza visual corregida (VA) fueron considerados como objetivos secundarios.

As a result, at 12 months it was observed that in the eyes with central macular thickness (CMT) > 250 μm at baseline, CMT had a mean decrease of 88 μm in group 1 versus 49 μm in group 2 (p = 0.02). Macular volume decreased 0.8 mm$^3$ and 0.4 mm$^3$, respectively (p = 0.03). Mean change in BCVA was +0 letters and -2 letters, respectively (p = 0.10).

Lower effectivity was achieved with the MMG technique compared to the modified ETDRS technique$^1$. A subanalysis compared CMT with BCVA before and after treatment and found a modest correlation after treatment with laser photocoagulation. It is highlighted that VA measurement cannot be substituted even though TD-OCT is an important tool in the evaluation. Likewise, it showed that TD-OCT changes, both in the short and long term, are not predictors of VA changes$^5$.

When correlating TD-OCT with stereo fundus photographs to evaluate retinal thickness, a weak correlation with VA was observed, as well as a moderate correlation between TD-OCT and fundus photography in evaluating CMT and a slight superiority in the reproducibility, as well as a greater sensitivity of TD-OCT to detect retinal thickness changes$^3$.

It was also studied if the extension of DME evaluated by TD-OCT before treatment could explain the initial VA and predict the change in CMT or VA after treatment, but no answers were obtained with the subanalysis$^4$.

When evaluating the association between FAG characteristics and VA, TD-OCT and fundus photographs, they concluded that the baseline leakage area is associated with a reduced VA and increased thickness and volume values by TD-OCT at baseline, but no associations with changes at 12 months were found$^5$.

Protocolo B

En los noventa comenzó la investigación sobre el uso de varios agentes intravitréales que pueden tener un efecto beneficioso en diferentes patologías, incluyendo el DME. En 2001 y 2002, los primeros informes de casos aislados sobre el uso de triamcinolona intravitréal para el tratamiento del DME fueron publicados, sugiriendo algunas efectividad. Su uso comenzó a spreading according to surveys of the American Society of Retina Specialists. Since there was not a lot of evidence regarding its safety and effectiveness, it
was considered important to conduct a randomized clinical trial.

The study consisted in comparing triamcinolone with the then gold-standard proposed by the ETDRS for DME, focal or grid laser photocoagulation.

Patients with DME involving the fovea (n = 840) were randomized into 3 groups: 1) 1 mg of intravitreal triamcinolone, 2) 4 mg of intravitreal triamcinolone or c) focal/grid photocoagulation. The patients were treated at the beginning of the study and if necessary, retreated at each visit (with a 4-month interval). The triamcinolone groups could receive rescue laser in case of fulfilling failure criteria.

Those patients with visual impairment of 15 letters or more on two consecutive visits separated by 4 months (MVL) could receive an alternative treatment at investigators discretion (patients in the laser group could receive triamcinolone and those in the triamcinolone groups could receive laser). Results at 4 months showed a greater visual improvement in the 4 mg triamcinolone group compared to the other two; however, at one year of follow-up, there was no significant difference between the three groups.

At 2 and 3 years of follow-up (n = 306), there was a greater VA improvement in the laser group compared to the two triamcinolone groups. This difference was not considered attributable to the development of cataracts since the results were consistent when analyzing pseudophakic patients. In addition, CMT decrease was similar in the three groups. This study showed that focal/grid laser has greater long-term benefits compared to intravitreal triamcinolone in patients with DME, in addition to having less adverse effects (increased intraocular pressure and cataract progression)\textsuperscript{6,7}.

Protocol C

Some studies showed variations in retinal thickness due to OCT throughout the day in some patients with DME: it was higher in the morning and lower in the later hours of the day. Because OCT has become the standard tool for the evaluation of DME, it was considered

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**Table 1. Comparison of laser photocoagulation parameters for direct macular or modified grid technique proposed by the ETDRS and the mild macular grid technique by the DRCR.net.**

<table>
<thead>
<tr>
<th>Spot characteristics</th>
<th>Focal/grid photocoagulation (modified ETDRS technique)</th>
<th>Mild macular grid photocoagulation (MMG-DRCR.net)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Direct treatment</td>
<td>Directly treat all leaking microaneurysms in the thickened retinal area between 500 to 3,000 microns from the macular center (but not within 500 microns around the optic disc)</td>
<td>Does not apply</td>
</tr>
<tr>
<td>Color change of the microaneurysm with direct treatment</td>
<td>Not required, but at least a moderate gray burn, which should be evident on all microaneurysms</td>
<td>Does not apply</td>
</tr>
<tr>
<td>Size of spot with direct treatment</td>
<td>50 microns</td>
<td>Does not apply</td>
</tr>
<tr>
<td>Duration time with direct treatment</td>
<td>0.05 to 0.1 s</td>
<td>Does not apply</td>
</tr>
<tr>
<td>Grid treatment</td>
<td>Applied to all areas of diffuse leakage or non-perfusion within the described area of treatment</td>
<td>Applied to the entire described area for treatment (including non-thickened retina)</td>
</tr>
<tr>
<td>Area considered for grid treatment</td>
<td>500 to 3,000 microns superior, nasal and inferior from the macular center, 500 to 3,500 microns temporal to the macular center. At least 500 microns away from the optic disc</td>
<td>500 to 3,000 microns superior, nasal and inferior from the macular center, 500 to 3,500 microns temporal to the macular center. At least 500 microns away from the optic disc</td>
</tr>
<tr>
<td>Spot size with grid treatment</td>
<td>50 microns</td>
<td>50 microns</td>
</tr>
<tr>
<td>Duration time with grid treatment</td>
<td>0.05 to 0.1 s</td>
<td>0.05 to 0.1 s</td>
</tr>
<tr>
<td>Spot appearance with grid treatment</td>
<td>Barely visible (light gray)</td>
<td>Barely visible (light gray)</td>
</tr>
<tr>
<td>Inter-spot distance with grid treatment</td>
<td>Separation of two burn diameters</td>
<td>Total of 200 to 300 burns (2 to 3 burn diameters of separation)</td>
</tr>
<tr>
<td>Wavelength of focal or grid treatment</td>
<td>Green to yellow wavelengths</td>
<td>Green to yellow wavelengths</td>
</tr>
</tbody>
</table>
important to study its diurnal variation for a more accurate interpretation.

In this study (n = 156), the retinal thickness was measured using TD-OCT at the following time points: 8, 10, 12, 14 and 16 h. The average central subfield thickness (CST) at 8 h was 360 μm, with a decrease of 6% (13 μm) at 16 h. In 5 eyes (3%) there was a decrease of more than 25% of the retinal thickness. The retinal thickness change was significantly greater in eyes with the highest retinal thickness at 8 h. In 10% of the eyes there was a gain of more than 10 ETDRS letters and a loss of 10 letters in 6% of the eyes examined. According to the results of this study, it is considered that, although there may be a progressive decrease in retinal thickness throughout the day in patients with DME, these changes are small, occasional, and of little clinical relevance, so they should not be considered for scheduling measurements in clinical trials.

Protocol D

The vitreous humor has been implicated, at least in part, as a cause of macular edema (ME) by various physiological and mechanical mechanisms. Likewise, it was observed that the separation between the vitreous and the retinal surface resulted in a subsequent reduction of the retinal thickness in patients with DME, either by surgery or by spontaneous detachment. Therefore, the DRCR.net performed a prospective observational study (n = 87) to evaluate the visual and anatomical results (CMT by TD-OCT) after vitrectomy without concomitant cataract surgery in patients with DME associated with vitreomacular traction (VMT). In addition to vitrectomy, an epiretinal membrane (ERM) peeling was performed in 61% of the cases, internal limiting membrane (ILM) peeling in 54%, panretinal photocoagulation (PRP) in 40%, and triamcinolone injection at the end of the procedure in 64%. Mean baseline BCVA was 20/100 and baseline CMT was 491 μm. At 6 months, there was a mean CMT reduction of 160 μm and 68% of the patients had a reduction of at least 50% in thickness. Thirty-eight percent of the patients showed a gain of ≥10 letters, although 22% presented a loss of ≥10 letters. After 6 months no significant changes were observed.

These results suggest that it is beneficial to consider vitrectomy for the management of DME in eyes with MVL associated with VMT since most of the eyes showed a reduction in CMT and between 28 and 49% of the eyes showed significant visual improvement. It is worth mentioning that between 13 and 31% of the eyes showed significant visual worsening.

When analyzing the associated factors in those eyes that showed the greatest VA gain, it was observed that VA gain occurred in eyes with the worse initial BCVA and in those with an ERM peeling, so it is suggested that ERM peeling could be beneficial. The presence of vitreoretinal interface abnormalities is associated with a greater reduction in CMT but without significant differences in the final BCVA.

Protocol E

Injections of peribulbar corticosteroids were used for the treatment of ME secondary to uveitis, with favorable results, so their efficacy was suggested in DME. It was thought that peribulbar administration had fewer adverse effects than the intravitreal route, which would justify their use specifically in patients with DME and good vision, in whom the risks of intravitreal injections did not justify treatment application. Therefore, a pilot study (n = 129) was conducted to evaluate the effects of these drugs at the anterior and posterior subtenon level, with or without adjuvant photocoagulation, in patients with DME and good VA (≥ 20/40). Patients were randomized in 5 groups: 1) focal photocoagulation, 2) anterior subtenon injection of 20 mg triamcinolone, 3) anterior subtenon injection of 20 mg triamcinolone followed by focal photocoagulation after 4 weeks, 4) posterior subtenon injection of 40 mg triamcinolone and 5) posterior injection of 40 mg triamcinolone followed by focal photocoagulation after 4 weeks. The results at 34 weeks of follow-up did not show a significant difference in retinal thickness or VA between the 5 groups, and adverse effects attributable to the injections were reported, such as increased intraocular pressure and palpebral ptosis. In conclusion, peribulbar triamcinolone injections did not show any significant benefit in patients with DME and good VA.

Protocol F

The DRS and ETDRS studies showed that PRP reduces the risk of severe visual loss (<5/200 in 2 consecutive visits with a 4-month interval), so it became the gold-standard for proliferative diabetic retinopathy (PDR). But this treatment is not innocuous since several complications have been reported (e.g., onset or worsening of ME) and there is not a standardized number of PRP sessions.
The objective was to compare the effects of PRP on DME applied in a single session (group 1) versus 4 sessions (group 2) in patients with early PDR or severe non-proliferative diabetic retinopathy (NPDR) with relatively good VA (>20/32) and CMT < 300 μm by TD-OCT.

In 18%, preexisting ME was determined. During follow-up, retinal thickness tended to increase slightly compared to baseline levels, but CMT was slightly higher in group 1 on the day 3 (+9 versus +5; p = 0.01) and week 4 (+13 versus +5; p = 0.003), and at week 34, the difference was reversed, and the thickness was slightly increased in group 2 (+14 versus +22; p = 0.06). VA results were similar to CMT results. At 34 weeks, VA was slightly worse in group 2 (0 versus -2 letters; p = 0.006) compared to baseline. Despite the changes reported by TD-OCT, only one eye in each group required focal/grid laser.

As in other studies reported, the results did not show significant differences in the effect of treatment on VA. No adverse effects were identified in any group. In conclusion, PRP can be safely applied in a single session; however, this study does not provide enough evidence to show superiority when compared to several sessions.

Protocol G

ME is the main cause of MVL in diabetic patients. The diagnosis is clinical and is complemented by macular OCT. The indication to treat clinically significant DME is widely accepted, but the value of identifying subclinical DME is unknown. Subclinical DME with central involvement was defined as macular thickening by TD-OCT from ≥225 to ≤299 μm, but not observed on clinical examination.

In a first study carried out in 2008, the objective was to evaluate CMT by TD-OCT in diabetic patients without diabetic retinopathy (DR) or with mild NPDR (only microaneurysms), without central macular thickening (<225 μm) and with a VA better than 20/32 (n = 97) and compare it with reference values in a population without retinal disease. The average CMT was 201 ± 22 μm and was slightly higher in men. No other factors, apart from gender, were found associated with an increased central macular thickness. Concluding that in this group of patients, CMT does not differ from the reference values of a population without diabetes and that gender differences must be considered for thickness evaluation.

An observational study conducted in 2012 (n = 43) determined the progression rate from subclinical DME to clinically significant DME (CMT increase of 50 μm or CMT ≥300 μm) or DME requiring treatment. At 1 year, 16/43 eyes met the criteria for clinically significant ME. The probability of having a CMT increase at 1 year was 27% and at 2 years was 38%. It was demonstrated that subclinical ME is uncommon (4.8%), but that it has a higher percentage (1/4 to 1/2 to 2 years) of progression to clinically significant ME than that reported by the ETDRS (25% at 3 years). Monitoring of these patients is suggested because of their high risk of progression.

Protocol H

Bevacizumab is a humanized monoclonal antibody that inhibits all isoforms of vascular endothelial growth factor A (VEGF-A). It has been widely used off-label for the treatment of DME.

This was the first phase 2 study to evaluate the effect of bevacizumab in DME. One hundred and twenty-one patients were randomized to 5 groups: 1) baseline focal laser, 2) 1.25 mg bevacizumab at baseline and 6 weeks, 3) 2.5 mg bevacizumab baseline and at 6 weeks, 4) 1.25 mg bevacizumab baseline and sham reinjection (placebo) at 6 weeks, or 5) 1.25 mg bevacizumab baseline and at 6 weeks plus focal laser at 3 weeks. The first phase (12 weeks) evaluated the efficacy and the second phase evaluated safety up to 70 weeks. The primary objective was CMT by TD-OCT and BCVA at 24 weeks. Groups 2 and 3 had the greatest reduction in CMT at 3 weeks and improvement of 1 line of BCVA at 12 weeks, compared with the laser group alone, with no differences observed between groups 2 and 3. The combination of laser with bevacizumab showed no short-term benefits, but no adverse effects either. Regarding adverse effects, there was a case of endophthalmitis, two cases of acute myocardial infarction and one event of congestive heart failure. The conclusions were that bevacizumab reduces CMT more quickly while laser reduces it more slowly, but after 3 weeks the effect is similar. There is no short-term difference between both doses in CMT reduction, and the effect of bevacizumab has an initial peak that is stationary, or even decreases, at 3 and 6 weeks, suggesting that 6 weeks is a long time for a second dose. This study was the basis for the subsequent phase 3 studies conducted with bevacizumab.

Protocol I

The benefits of focal or grid laser for the treatment of DME demonstrated by the ETDRS in 1985 were verified.
by the DRCR.net (protocol B), remaining as the gold standard versus triamcinolone, after excluding the group of phakic patients who developed lens opacity. However, in those patients with a VA ≤ 20/40, approximately 20% of those treated with focal or grid laser presented a loss of ≥2 lines. This explains the need for new therapies that could have a greater benefit, such as monoclonal antibodies, either in monotherapy or as adjuvant therapy to photoacoagulation.

Protocol I is a multicentric clinical trial in which 854 study eyes were randomized from 691 participants with diagnosis of DME with central involvement. The main objective of the study was to evaluate the effect of ranibizumab or triamcinolone in combination with laser, compared with laser monotherapy, for the treatment of DME.

Subjects were randomized in 4 groups: 1) simulated intravitreal injection plus focal or grid early laser (within 7-10 days) (n = 293), 2) 0.5 mg ranibizumab plus early laser (n = 187), 3) 0.5 mg ranibizumab plus delayed laser (>24 weeks) (n = 188), or 4) 4 mg triamcinolone plus early laser (n = 186).

The treatment protocol for laser photoacoagulation was administered according to the modified technique of the original protocol proposed by the ETDRS, previously described and used in previous protocols of DRCR.net; while the treatment protocol with the intravitreal agents was carried out in the following way: from the beginning of the treatment until week 16, injections were applied every 4 weeks regardless of VA or macular thickness; from weeks 16 to 20 a monthly application was maintained, except when “success criteria” were met (VA> 20/20, CMT <250 μm); from week 24 to 48, patients with “success criteria” were categorized and treatment was delayed, those with improvement (VA gain of > 5 letters or improvement of CMT <10%) continued monthly injections. Those without improvement were assigned to treatment at the discretion of the investigator and, finally, those with failure (VA loss of 10 letters compared to baseline, CMT > 250μm and without improvement 13 weeks after completing the laser treatment) were assigned an “alternative” treatment at the discretion of the investigator. From week 48, visits were scheduled every 4 months, depending on the case. If treatment had been delayed in previous visits, the follow-up was extended to twice the time of the previous visit, with a maximum interval of up to 16 weeks.

The results from the first year showed the superiority of ranibizumab, either with an early laser (+9 ± 11) or with a delayed laser (+9 ± 12), for improving VA compared to a combined therapy of triamcinolone plus early laser (+ 4 ± 13) or laser monotherapy (+3 ± 13). The reduction in mean CMT by TD-OCT was similar for ranibizumab or triamcinolone. Likewise, in pseudophakic patients (n = 273), the visual improvement was comparable for both therapies.

After 3 years of follow-up, superiority was maintained in the ranibizumab groups with an early or delayed laser, and they presented a difference in mean VA of +2.9 letters in the delayed laser group compared to the early laser group. Likewise, the delayed laser group showed a greater percentage of patients with a gain of > 10 letters (56%) compared with the early laser group (42%).

The results at 5 years of follow-up showed proportions similar to the initial differences: ranibizumab with early (+7.2 letters) or delayed (+9.8 letters) laser showed a mean difference of -2.6 letters (p = 0.009). The subjects of the ranibizumab plus early or delayed laser group required an average of 8 to 9 injections during the 1st year, 3 to 4 during the 2nd year and 1 to 2 during the 3rd year, respectively. For years 4 and 5, the mean number of injections was 0 for both groups.

Likewise, those eyes assigned to the ranibizumab treatment groups had a lower rate of progression from severe NPDR to PDR (8 versus 42%) even when the injection was not applied monthly after week 12, compared to those assigned to the early laser monotherapy group. Therefore, it was also shown a beneficial effect of ranibizumab for the treatment of neovascularization, which suggests that indefinite monthly injections are not necessary to maintain these beneficial effects.

**Protocol J**

The worsening of pre-existing DME after PRP is a recognized adverse effect, although infrequent. VEGF has a role in the development and exacerbation of DME, so steroids and anti-VEGF drugs may reduce the exacerbation of pre-existing DME in patients with PDR and severe NPDR.

For this study, 345 eyes were randomized with a VA ≥ 20/320 and DME treated with focal/grid laser and with DR under treatment with PRP, into 3 groups: 1) baseline sham injection at 4 weeks, 2) 0.5 mg ranibizumab at baseline and 4 weeks, and 3) baseline triamcinolone and sham injection at 4 weeks, to evaluate the short-term effect on DME. At 14 weeks, the mean change in VA was -4 ± 14 letters, +1 ± 11 letters and +2 ± 11 letters in each group, respectively, with a greater proportion of eyes with improvement (> 10 letters) and a lower proportion with worsening (> 10 letters) in groups 2 and 3, respectively. The changes in retinal thickness
paralleled those of VA. Subsequent evaluations at 34 and 56 weeks only aimed to assess long-term safety, although it was observed that differences in VA and CMT did not persist until 56 weeks. This study concluded that the addition of triamcinolone or ranibizumab to focal laser plus PRP, is associated with an increased VA and a greater CMT decrease at 14 weeks but it was not able to determine the long-term benefit of this treatment20.

Protocol K

The criteria for retreating with focal/grid laser are an important aspect, and to some extent, there was limited information regarding the results obtained after a single laser session. The ETDRS and the DRCR.net established the criteria for retreatment (persistence of clinically significant ME, identification of one or more treatable lesions that could be the cause of visual loss) at 4-month intervals.

The objective of this study was to determine if CMT reduction associated with a single session of focal or grid laser would persist even after delaying treatment in patients with central involvement of DME after week 16. Treatment consisted in focal or grid laser photocoagulation in 122 eyes with clinically significant ME according to the modified ETDRS technique. From week 16, patients were followed-up every 8 weeks and laser treatment was delayed if they showed a gain of > 5 ETDRS letters or a CMT decrease > 10% compared to baseline.

At week 16, 47% showed a VA improvement. Of these, 48% (26 eyes) had a CMT ≥250 μm and were monitored until week 32. Only 11/26 eyes showed a CMT decrease of > 10% between week 16 and 32, with no need for retreatment.

This study demonstrated that 23 to 63% of the eyes that showed improvement but not a complete resolution of DME at 16 weeks after laser application, will continue to improve without needing retreatment until 32 weeks. This is useful evidence to assess a retreatment schedule21.

Protocol L

The ETDRS, in addition to providing solid evidence concerning DR and DME, established a standardized method for the assessment of manual optical refraction and thus BCVA (important research objectives). However, the methodology presents certain difficulties, among which can be highlighted the need for trained personnel, long time for the refraction and significant costs. An alternative to this, facilitated by technological advancement, could be autorefraction.

This study included prospectively 878 eyes of 456 participants from 26 centers. The main objective was to compare the VA obtained after performing automated refraction with different autorefractors versus the manual refraction (MR) performed according to the DCRR.net protocol in patients with visual loss secondary to DME with a wide range of VA (Snellen equivalent better than 20/400). The VA score was determined with the electronic VA test of the ETDRS. All the participants underwent the VA test on three occasions: on measurement with autorefraction and two additional measurements using manual refraction.

The spherical equivalent obtained was similar with both techniques, with a mean difference of 0.00 and a range of -1.75 to 1.13 diopters. But average VA was slightly better with manual refraction. The variability obtained between both techniques (automated refraction versus manual refraction) was considerably higher than between the two manual measurements (p <0.001) and was highly dependent on the autorefractor model. The smallest variability for automated refraction was obtained with the Topcon 8000 series.

The autorefractors used in this study were not able to replace manual refraction, mainly for clinical trials due to the high variability22.

Protocol M

Although it has been established through various studies that intensive glycemic control can reduce the frequency and progression of microvascular complications in individuals with diabetes mellitus (DM), it is clearly difficult to achieve, especially in the long term. Some authors point out obstacles that prevent optimal glycemic control, highlighting the lack of awareness about the potential damage to target organs, cultural aspects, as well as financial limitations, poor access to health services, depression, lack of perseverance and others23-26.

The DCRR.net carried out a study in 42 study centers. The objective was to determine whether the combination of personalized risk assessments and diabetes education in an ophthalmology clinic could improve glycemic control in people with diabetes, assessed by glycosylated hemoglobin (HbA1c) with a primary cut-off point at 1 year, extended to 2 years.

Diabetic patients were randomized to two groups: one that received the intervention and the other with control patients treated with standard care. Likewise,
each group was subdivided into those with more frequent than annual follow-up and those with annual follow-up. The patients in the intervention group received evaluation and personalized education, which consisted of: a) cut-off point for HbA1c, b) information on blood pressure and the severity of diabetic retinopathy (DR), c) personal risk for DR progression based on HbA1c and type of diabetes, d) a personalized report of the risk of kidney damage and retinopathy based on HbA1c, e) a chart with previous and recent reports of HbA1c, f) instructions to indicate the individual risks encountered and g) supplementary material for the management of diabetes. The patients in the control group were required to complete a minimum annual visit without receiving more education than usual in the ophthalmological visit.

The mean HbA1c report in patients with more frequent than annual follow-up was 8.3% in the control group (n = 502) and 8.6% in the intervention group (n = 488), with an annual change of -0.1% and -0.3%, respectively, while patients with annual follow-up had a mean HbA1c of 8.3% in the control group (n = 368) and 8.4% in the intervention group (n = 388), with a change of 0.0% and -0.1%, respectively. In conclusion, this study corroborated the difficulty of making a change in the personal behavior of patients, even with adequate education about the pathology and its possible complications, also without presenting differences between annual or more frequent follow-up27.

**Protocol N**

Vitreous hemorrhage secondary to PDR is considered one of the main indications for vitrectomy in the USA, carried out when the hemorrhage prevents the adequate application of PRP. Despite technological advances in retinal surgery, complications cannot be ignored. Some series of uncontrolled cases suggest that intravitreal antiangiogenics clear vitreous hemorrhage in the short-term so PRP can be applied without requiring vitrectomy.

This study evaluated the effect of intravitreal ranibizumab versus intravitreal saline to determine if ranibizumab increases the clearance rate of vitreous hemorrhage and thus avoids the need for vitrectomy. It was injected at baseline and at 4 and 8 weeks. At 16 weeks, which was the first cut-off point, the probability of vitrectomy was of 12% for the ranibizumab group and 17% for the saline group (p = 0.37). The main cause of vitrectomy between weeks 8 and 12 was persistent vitreous hemorrhage and, in a low proportion, progression to tractional retinal detachment (TRD). There was also a marked improvement in VA, a greater cumulative possibility of PRP completion and a lower recurrence rate of vitreous hemorrhage, suggesting a short-term biological effect of ranibizumab28.

Subsequently, in the 1-year analysis, the difference in VA did not persist and there were no relevant clinical differences between both groups, with a slightly greater possibility of PRP completion in the group treated with ranibizumab29.

**Protocol O**

It is important for clinical practice, as well as in the field of research, to know exactly what is the normal CMT in the eyes of individuals with DM but without DME and to differentiate it from those with DME by OCT.

In a previous study, it was observed that patients with DM without DR or with mild DR, as well as those with DR without DME, have thickness values similar to patients without DM and with apparently normal retinas. All this was determined by TD-OCT (OCT-3, Stratus).

They performed a first study that evaluated the macular thickness of people diagnosed with DM, without DR or with minimal DR using spectral-domain OCT (SD-OCT, Spectralis, Heidelberg Engineering, Inc.). Comparative measurements between the Spectralis and Stratus of the 9 ETDRS subfields were made, as well as a comparison with a reference database previously reported by the DRCR.net.

CMT mean (n = 122) was 270 ± 24 μm, with the central point in the range of 227 ± 25 μm, and the mean volume was 8.4 ± 0.4 mm³. Characteristically, men presented greater thicknesses and greater volumes. In contrast, the Stratus reported a mean CMT of 199 ± 24 μm (71 μm less than Spectralis) and an average volume of 6.7 mm³ ± 0.4 mm³ (1.7 mm³ smaller). The results were comparable with data obtained from healthy individuals without DM.

This study reported the mean thickness found in each of the 9 ETDRS macular field areas. Thicknesses ≥320 μm in men and ≥305 μm in women are proposed as gender-specific thicknesses to determine ME with central involvement of diabetic origin30.

The reproducibility of retinal thickness by TD-OCT and SD-OCT was evaluated, and equations were made to convert the measurements obtained by Stratus and determine if they are equivalent to SD. It was determined that there is a better reproducibility of the measurements with Spectralis compared to Cirrus or Stratus31. And when evaluating the reproducibility between CMT
obtained by Stratus or by Optovue RTVue, it was concluded that the variability of RTVue is similar to the one of Stratus or Cirrus²⁵.

The analysis provided by the network is useful for evaluating and comparing the results between trials and for the development of specific cut-off points of the different SD-OCTs available for subsequent clinical trials.

**Protocol P**

Even when cataract surgery is performed in diabetic patients routinely, there is limited data on VA and the behavior of CMT after said surgery in patients with DME.

This was a pilot, observational study with the main objective of assessing VA and the behavior of ME after cataract surgery in the eyes of patients previously diagnosed with DME with central involvement. They included 63 patients diagnosed with DME in at least 1 eye (CMT > 250 μm by TD-OCT, > 310 μm by SD-OCT). Thirty-five percent (n = 21) of the patients did not receive treatment for macular edema during the study. Forty-three percent (n = 26) received treatment before surgery (focal/grid laser, intravitreal triamcinolone within 4 months prior to surgery, intravitreal antiangiogenics within 2 months prior to surgery). Seven percent (n = 4) received intraoperative treatment and 42% (n = 25) received postoperative treatment. Of all patients who received treatment, 69% (n = 27) received intravitreal antiangiogenics.

At the end of the follow-up, at week 16, a change in the mean VA of +12 letters was observed (95% CI: +8 to +16), a gain of at least 4 lines was observed in 32% (95% CI: 20-45%) of the eyes, with a VA worsening of at least 2 lines in 10% (95% CI: 4-21%). Secondarily, a change in the mean CMT by OCT of -11 μm (95% CI: -11 to -28μm) at 16 weeks was observed. This study has the limitation of a small sample size associated with a heterogeneous treatment for DME; however, with the results obtained, it is estimated that at least half of the operated eyes that had previous diagnosis of DME may not show visual improvement. Due to the low level of evidence, it is not possible to obtain definitive conclusions in this study²⁵.

**Protocol Q**

It is widely accepted that DM increases the incidence of cataracts and may increase the risk of poor vision after surgery. On the other hand, cataract surgery has been reported as a risk factor for the incidence of DME or for the progression of DR.

The objective of this study was to determine the incidence of ME with central involvement after cataract surgery in patients with DR without pre-existing central macular thickening. This study enrolled 293 patients into 4 groups according to OCT: 1) without DME, defined as a central macular thickness less than the normal mean in all subfields; 2) “possible” DME in the central field, defined as a thickness between the mean and +2 standard deviations from the normal; 3) non-central “possible” DME, defined as a thickness of at least 1 non-central field between the mean and +2 standard deviations, and 4) “definitive” non-central DME, defined as at least one non-central field of at least +2 standard deviations.

The results were evaluated 16 weeks after surgery. It started with 329 participants; however, after adjusting for exclusion criteria and for the follow-up at 16 weeks, data from 261 eyes were analyzed. No eyes with previous DME developed EM with central involvement. Ten of the 97 (10%, 95% CI: 5-18%) eyes without central involvement prior to surgery developed central DME, 18 of the 147 (12%, 95% CI: 7-19%) eyes with “possible” central involvement developed central DME. The rate of development of DME with central involvement 16 weeks after cataract surgery differed according to the history of previous treatment for DME (p <0.001). Of the patients who had been previously treated for DME, 21% (95% CI: 14-30%) developed central ME, while 4% (95% CI: 29%) of those who did not have a history of prior treatment developed central ME. As for DR, 82% did not present changes, while 10% progressed. It was also found that those eyes with better VA and less severe DR at the beginning of the study had a lower incidence of central DME.

In conclusion, it was demonstrated that there is a higher possibility of developing DME with central involvement after cataract surgery in patients with non-central ME at baseline, as well as with history of previously treated DME with central involvement³⁴.

**Protocol R**

DME is the most common cause of MVL in patients with DR. The increase of several cytokines and other proinflammatory factors plays a role in DME explaining, at least in part, the pathophysiology of the disease. Non-steroidal anti-inflammatory drugs have been used for postoperative inflammation. Topical nepafenac is frequently used for the treatment of postoperative cystoid macular edema.
The main objective of this study was to evaluate the role of topical 0.1% nepafenac in eyes with non-central DME, defined as thickening within 3,000 μm from the center, without involving the fovea. Patients were randomized into 2 groups (n = 125). Group 1 (n = 61) received nepafenac 3 times a day. Group 2 (n = 64) received placebo. The initial follow-up was 12 months and treatment for DME was not allowed unless central involvement was observed.

The initial mean volume by TD-OCT was 7.8 mm³ and mean central thickness was 223 μm. The change at 12 months was -0.03 mm³ in group 1 and -0.02 mm³ in group 2, without differences when analyzed after adjusting for state of the lens and HbA1c. In group 1, 7 eyes (11%) developed central DME, while in group 2 nine eyes (14%) developed it. Of which, 5 eyes in group 1 and 3 eyes in group 2 required treatment.

The average change in VA was 0.09 letters and -0.15 letters, respectively, with a difference of 0.2 letters between the groups (p = 0.82). Among the side effects was a case of corneal lysis in one eye of the group treated with nepafenac, with a history of severe dry eye.

In conclusion, no benefit was found with the topical use of nepafenac for 12 months in terms of retinal thickness by OCT or VA. Nepafenac appears to be well tolerated, without showing differences in adverse effects compared to the control group.

Protocol S

PRP has been the standard treatment for PDR since the reports by the DRS and the ETDRS. Currently, with the advent of intravitreal antiangiogenic therapy, new alternative treatments have been proposed, based on the antiangiogenic effect of these drugs.

Protocol S evaluated the noninferiority of 0.5 mg ranibizumab (Lucentis, Genentech) versus PRP for the treatment of PDR with or without DME in patients without previous PRP. Participants were randomized (n = 305, 294 eyes) into 2 groups. Group 1 received PRP, completed in 1 to 3 visits, and group 2 received ranibizumab at baseline and every 4 weeks if they met the retreatment criteria, and could receive rescue laser if needed. In case of DME, group 1 could receive antiangiogenic treatment. In case of vitreous hemorrhage or tractional retinal detachment, vitrectomy was indicated.

The main result was a BCVA gain of +2.8 letters in group 1 compared to +0.2 in group 2 (p < 0.001) after 2 years of follow-up. Also, patients in group 1 showed increased visual field loss. Vitrectomy was performed in 15% of patients in group 1 and in 4% in group 2. More patients in group 1 developed DME (28 versus 9%) and 53% of patients in group 1 also received ranibizumab and only 6% of group 2 received PRP.

At 2 years there was no statistical difference in terms of activity or regression of neovascularization between both groups. There were no differences in the rates of retinal detachment, neovascular glaucoma or vitreous hemorrhage between both groups. The vitrectomy rate was significantly higher in the PRP group (15%) than in the ranibizumab group (4%) [36].

In a cost-benefit analysis in terms of monotherapy with ranibizumab or PRP for PDR, it was observed that it was more appropriate to initiate monotherapy with PRP for patients with PDR but without associated DME, and ranibizumab for those with DME at the time of diagnosis [37].

The 5-year results (n = 184) show that the average number of injections in group 1 was 7.9 and in group 2, 19.2. The mean change in VA was +10.5 versus +14.3 letters (p = 0.68), and the mean final BCVA in both groups was 20/25, which was consistent with the data at 2 years. Even though at 2 years group 1 had a greater loss of visual field, the decrease in peripheral visual field progressed in both groups during the 5 years of follow-up, with both 30-2 and 60-4 Humphrey visual field. The mean change in cumulative total score was -572 (635) dB versus -330 (645) dB, and the difference between groups was not statistically significant (p = 0.4). More than half of the eyes in group 1 were treated with ranibizumab for DME [38].

An important question that this protocol answers is whether treatment with ranibizumab poses a risk of developing serious complications in the absence of PRP after 5 years of follow-up. Few eyes in both groups developed neovascular glaucoma or neovascularization of the iris, and the incidence rate was similar. The overall rate of retinal detachment (mainly tractional) was highest in group 1 (15 versus 6%). At least half of the eyes in both groups developed some degree of vitreous hemorrhage, but only 42% in group 1 and 22% in group 2 required vitrectomy [39].

In conclusion, this protocol demonstrates that ranibizumab is not inferior to PRP in terms of final BCVA, so both therapies are adequate in the treatment of PDR, mainly for those with coexisting DME. When selecting a treatment, it is important to keep in mind individual factors of the patient, such as the possibility of close monitoring, costs and metabolic control status.

Protocol T

Among the wide variety of antiangiogenic agents available, aflibercept, bevacizumab, and ranibizumab...
are the most frequently used. All three have shown an important benefit and an adequate safety profile for the treatment of DME.

In this protocol, the three agents were compared. Patients (n = 660) were randomized into 3 groups: 1) 2.0 mg aflibercept, 2) 1.25 mg bevacizumab or 3) 0.3 mg ranibizumab. Doses were applied at baseline and then every 4 weeks (except if the response criteria were met during the first year), and every 4 to 16 weeks in the second year. The main objective was to evaluate the change in BCVA.

During the first year, improvement in VA was +13.3 with aflibercept, 9.7 with bevacizumab and 11.2 with ranibizumab, with a statistical but not clinically significant difference. When performing the analysis by VA ranges, in the group of VA 20/32 to 20/40 there was no statistical difference (p > 0.5), but if the initial VA was 20/50 or worse, aflibercept was better compared to the other two drugs (aflibercept versus bevacizumab [p <0.001], aflibercept versus ranibizumab [p = 0.003]) and no difference between ranibizumab and bevacizumab.

In the 2-year analysis, the results were similar. Patients with better initial BCVA achieved similar improvement with the three drugs. In those with an initial BCVA of 20/50 or worse, aflibercept induced the highest improvement in BCVA; however, the difference was lower than in the first year and it was not statistically significant compared to bevacizumab. The number of injections during the second year was less than in the first year.

The benefit of antiangiogenic agents has been demonstrated for the visual improvement and the decrease of CMT in eyes with DME. However, at least between 32 and 66% of the eyes treated with monthly antiangiotics for 6 months show persistent edema associated with low vision.

This study compared treatment with ranibizumab as monotherapy versus the addition of a dexamethasone implant in patients with persistent DME (DME treated with at least 3 loading doses of an antiangiogenic agent [bevacizumab, ranibizumab or aflibercept] within the 20 previous weeks).

Subjects were randomized (n = 116) in 2 groups: 1) 0.3 mg ranibizumab plus a 700 μg dexamethasone implant or 2) placebo implant plus 0.3 mg ranibizumab. Patients were evaluated every 4 weeks for 24 weeks and the need for retreatment was re-evaluated.

VA improvement was +2.7 letters in group 1 and +3 letters in group 2 (95% CI: -3.6 a 2.5; p = 0.73). A gain of ≥10 letters occurred in 22% of patients in group 1 and in 14% in group 2. CMT reduction was -110 μm in group 1 and -62 μm in group 2. In group 1, 29% developed intraocular hypertension compared to <1% in group 2 (p < 0.001). The total number of injections during the 24 weeks was 5.6 in group 1 and 5.7 in group 2.

In conclusion, adding dexamethasone to the antiangiogenic therapy did not show an additional visual improvement, while an additional decrease of the central macular thickness was determined when compared to the group treated with monotherapy. As it was reported in previous studies, the sustained-release dexamethasone implant predisposes to an increase in intraocular pressure, so its use should be monitored closely.

Protocols currently in development

Protocol V

The objective of the protocol is to evaluate the treatments for DME in eyes with very good vision. It will include eyes with DME by SD-OCT and a VA ≥ 20/25. Patients will be randomized to receive early treatment with antiangiotics (aflibercept and ziv-aflibercept) or early laser with delayed antiangiotics. And the main outcome will be to evaluate the proportion of eyes with a loss of ≥ 5 letters in 2 years.

Protocol W

This protocol consists of administering an intravitreal antiangiogenic (aflibercept) as a preventive therapy in patients with high risk of visual loss.

The primary objective is to evaluate the safety and efficacy of antiangiotics versus observation in eyes with severe NPDR and a VA ≥ 20/25 without DME and without previous treatment, as a preventive measure for the appearance of DME or PDR.

Protocol AA

It will evaluate peripheral DR lesions by ultra-widefield imaging (UWFI) and its association with the progression of DR severity. The objective is to assess if the evaluation of peripheral lesions by UWFI allows the evaluation of DR and to predict the progression rate over time,
compared with 7 standard field stereoscopic paired photos. Eyes with untreated NPDR and without macular edema will be included. Patients will undergo annual monitoring for 4 years to evaluate the relative risk of progression of two or more severity levels in groups with and without peripheral lesions using UWFI (Optos).

**Protocol AB**

This study will compare the treatment with intravitreal antiangiogenics (afiblercept) versus early vitrectomy for the treatment of vitreous hemorrhage secondary to PDR. The main objective is to compare the VA at 6 months after each of the interventions. In the antiangiogenics group, at least 2 monthly doses will be applied upon study initiation and later, vitrectomy may be performed up to 4 months later pro re nata in case of vitreous hemorrhage persistence. PRP may be applied if antiangiogenic treatment alone does not achieve disease control. In the vitrectomy group, surgery will be performed with intraoperative PRP in the first 2 weeks after randomization. Antiangiogenic injections may be applied in case of recurrent hemorrhage.

**Protocol AC**

It will consist in the administration of intravitreal aflibercept versus bevacizumab with delayed aflibercept for DME with central involvement. The objective is to compare the efficacy in terms of VA change. Patients in the bevacizumab group will be candidates for changing to aflibercept after 12, 16, 20 and 24 weeks if they meet the established criteria.

**Protocol AD (PROMINENT study)**

The objective is to evaluate whether treatment with pemfibrate (0.2 mg/12 h orally) compared to placebo reduces the rate of DR progression in type 2 diabetic patients with NPDR.

**Protocol GEN**

The objective of this study is to create a gene bank and to collect information about clinical phenotypes for the scientific community with the rationale that in the future this can provide enough information to evaluate susceptibility or genetic resistance regarding DR. Likewise, it is intended to determine variants that may impact on key biomarkers in the development of DME and neovascularization.

**Conclusion**

During the last few years, the DRCR network has provided clinical results on important research questions regarding the treatment of DME and, recently, of DR. There has been an answer to questions about treatments and technological innovations, either directly or indirectly, which otherwise would not have been answered by the industry or by other entities.

Antiangiogenic agents have been accepted globally as the first-line treatment in the management of ME and have shifted focal laser as adjuvant therapy in selected cases. Also, the application of grid macular laser is now in disuse. What currently is still under discussion is the management of PDR. It is well known the non-inferiority of antiangiogenics; however, their choice as first-line therapy is still controversial. In addition, steroids have shown certain benefit, although currently, they continue to be a second-line treatment for DME.

It should be noted that the conclusions provided by the network are the result of a collaboration between a group of clinical researchers who participate in academic centers, as well as in private practices, which, at least in theory, entails the ability to correctly extrapolate the results to daily clinical practice. However, we suggest that the results should be interpreted with caution always considering the individual characteristics of each patient, our environment and our main goal, which is to obtain the greatest benefit for patients using the selected therapy.

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**Conflicts of interest**

No conflicts of interest are reported by any author.

**Ethical disclosures**

The present work has been evaluated and authorized with the number AR-Ret-0015.

**Protection of human and animal subjects.** The authors declare that no experiments were performed on humans or animals for this study.

**Confidentiality of data.** The authors declare that no patient data appear in this article.
References


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