Intraocular vision-improving devices in age-related macular degeneration

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Abstract: Age-related macular degeneration (AMD) is a leading cause of blindness, especially in western countries, and will substantially burden society and the world's health care system. Patients with late AMD often accompany a progressive loss of central vision, which will heavily influence the quality of life and associated with increased risk of functional disability. The principal visual rehabilitation methods with low-vision magnifiers, such as hand or stand magnifiers, spectacles, and closed-circuit television, were cumbersome to use and cosmetically burdensome. Therefore, the development of intraocular vision-improving devices has become an attractive alternative to extraocular visual aids, and better life quality improvement has been reached among AMD patients. To evaluate each device’s safety and efficacy based on current research, we searched the Cochrane Library, PubMed using pre-reported search terms and keywords combined with both Mesh term and text words. We explored randomized clinical controlled trials, cohort studies, and case serial reports and summarizes three aspects: visual outcomes, safety outcome, and quality of life outcomes. There are four types of devices recommended for AMD patients illustrating in this article: an implantable miniature telescope (IMT), IOL-VIP System, EyeMax Mono, and Scharioth macula lens (SML). There is no doubt that these technological advancements would bring new hope for AMD patients. However, the lack of randomized controls, limited follow-up duration, and various visual acuity (VA) measurements in different studies would be difficult than IOL devices.

Keywords: Intraocular devices; implantable miniature telescope (IMT); EyeMax Mono; IOL-VIP system; Scharioth macula lens (SML)

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Introduction

Age-related macular degeneration (AMD), a leading cause of visual impairment in developed countries, affects the retina’s macular region, causing severe and progressive loss of central vision (1). On a global scale, the number of people with AMD is expected to be 200 million by 2020 and increasing to 300 million by 2040 (2). The estimated prevalence of late AMD in the three extensive population-based studies in European white people over 85 years old was 13.1% (3), and the 15-year incidence of late AMD was
6.8% (4). However, in Asia and Africa, the prevalence of late AMD was only 0.1–0.3% (2), and the 6-year incidence for early and late AMD is 4.2% and 0.2%, respectively (5). AMD occurs commonly in developed countries, potentially indicating that progression in demographic composition like aging, might play roles in the prevalence of the disease (6). Although AMD’s prevalence was lowest currently in Asia, the population still accounts for more than 60% and will project the most significant number of cases (2). Over the years, it would increase rapidly than other regions and heavily economic burdensome due to lack of income and profoundly impact the social and health care system worldwide.

One of the critical approaches to help patients with AMD was visual rehabilitation with low-vision magnifiers, like spectacles, hand-held telescopes, and magnifiers (7). However, there are some drawbacks. Firstly, those extraocular devices restricted the visual fields (VFs) and are limited to the relative movement between the eye and telescope, causing patients to move their heads to follow the image (8). Secondly, they are cosmetically burdensome and lead to motion sickness because of vestibular-ocular conflict (9).

Given the drawbacks illustrated above, improving visual acuity (VA) and quality of life among AMD patients remains an ongoing challenge. Intraocular implants, therefore, as a visual prosthetic device, have been developed as attractive alternatives. Nevertheless, it is unknown which devices are more effective and safer. The previous major review in 2017 has illustrated the optical fundamentals of IOLs for AMD and compared seven intraocular lenses (10). The purpose of this systematic review is to compare the efficacy and safety of different intraocular lenses for patients with AMD based on updated evidence. We present the following article following the PRISMA reporting checklist (available at http://dx.doi.org/10.21037/atm-20-5851).

Methods

Two researchers (WJ and MFF) initially searched the Cochrane Library, PubMed using pre-reported search terms and keywords combined with both Mesh term and text words. The full-text versions of any study were then screened independently and discussed when disagreements were exited to avoid article selection bias. The search term includes (Macular degeneration [MeSH] OR Macular degeneration) and (intraocular lens OR visual devices OR implantable IOL) OR EyeMax Mono OR oIAMD OR Scharioth lens OR Fresnel Prism OR IOL-VIP System OR Implantable Miniature Telescope OR Lipshitz macular implant. We searched randomized clinical controlled trials, cohort studies no matter prospective and retrospective, and case serial reports published from 2017 to 2020 April. However, for some devices that lack updates, we further summarize their characteristics in three aspects: VA outcomes, quality of life and functional outcomes, and safety outcomes. The language is specific to English, and the only intraocular lens or device in human patients affected by AMD was identified as eligible studies to include in our review. Extraction data included the necessary information (principle of the equipment, fundamental parameters), characteristics of patients (number of patients, age, assessment methods for AMD, and indication criteria), surgical operation (size of the incision, implantation position, timing for operation, binocular or monocular, contraindications and adverse events), preoperative evaluation (exercises, prefer retinal location (PRL) and approaches to measure VA), outcomes (short-time and long-time efficacy) and characteristics of the studies (title, publication year, country, and study design). This study’s primary outcome was the postoperative VA gain related to each device, and the secondary outcome was the evaluation of the quality of life, which is the original intention to design and promote those implantable visual devices.

Results

Intraocular magnifier telescope (IMT)

IMT is a device that designs for monocular use reduces the impact of the central vision blind spot. It projects the objects onto the larger area of the light-sensing retina that is not degenerated by AMD (11). It has comprised of high precision quartz glass wide-angle micro-optics mounted in a polymethyl methacrylate (PMMA) carrying device and, after removing the natural lenses, implanted in the capsular bag through a 10–12 mm limbal or scleral incision tunnel to extends through the pupil and acquire vertical clearance distance about 2.5 mm from the posterior cornea in case of damaging the posterior cornea endothelium The parameters of the device are 4.4 mm long, 3.6 mm in diameter, weigh 60 mg in aqueous and the overall haptic-to-haptic diameter of 13.5 mm. The lens aperture is 3.2 mm. There are two forms depends on different magnifications, namely the WA (wide-angle) 2.2x and the WA 3.0x (FDA 2010).

The most common approach used in IMT was a fixed focus Galilean telescope system. Two optical elements...
with high positive and negative power were used combined with the cornea embedded in the air to achieve higher magnification. The central VF of an operative eye would magnify 3× with a 20–24° field of view projected onto about 55° of the retina and the vision range, including near, intermediate, and distance. In contrast, the fellow eye is the sole eye for peripheral vision after surgery. However, the field of view is magnification at the expense of a significant VF reduction, which is not suitable for binocular implantation. Vision extensive rehabilitation for 6 months to 1 year were requested for all patients after surgery to be effectively using the device (12). With a low vision specialist’s help, the recipients would learn to reteach the brain to learn how to use each eye differently for a specific task.

The US FDA first approved in 2010 and reduced the potential candidates’ age limits from over 75 to over 65 years old in 2014. There are other limitations to FDA guidelines. The treatment candidate would have irreversible dry AMD, no longer for drug treatment, and no previous cataract surgery in the implanted eye (13). In 2020, the new models of IMT, namely WA 2.2×, WA 3×, WA DE 3×, NG IMT, and Tsert SI (NG SI IMT 3×), were CE marked and approved in European Union (Table 1). The new device (Figure 1), NG SI IMT 3×, is supplied as preloaded in the Tsert SI cartridge (Figure 2) what allows the implantation with a much smaller incision (SI) than with the previous model (Figure 3).

There were eight articles eligible and identified (14-21), and 274 patients contributed to the efficacy and safety analysis; among them, 174 patients were participated in follow-up studies at 18 and 24 months, whereas 63 patients eventually finished the follow-up study at 60 months.

Table 1 Comparison between WA-IMT model with NG (SI) system

| Parameters                      | WA-IMT          | NG (Si)  |
|---------------------------------|-----------------|----------|
| Magnification                   | 2.2×/2.7×       | Same     |
| Optics diameter (mm)            | 3.60            | Same     |
| AXL (height, mm)                | 4.40            | Same     |
| Overall diameter (mm)*          | 13.5            | 10.8     |
| Incision size (mm)              | 10–12           | 7.0–7.5  |
| Capsulorhexis size (mm)         | 7               | 6        |
| ECD loss                        | 23–25%          | 7.9%     |
| Corneal clearance-post op ACD (mm) | 2.5            | 3.5      |
| Sutures                         | 8–12            | 3–4      |
| Manipulation                    | High rate       | Almost none |
| Surgical duration (min)         | 60              | 25       |
| Surgical procedure              | For skilled surgeon | Simpler, less complicated |
| Patients feedback               | Few weeks PO    | Few days PO |

* include the haptic loop. ECD, endothelium cell density; ACD, anterior chamber depth.

Figure 1 NG SI IMT 3X. It was supplied by the VisionCare Ophthalmic Technologies Company.
VA outcomes

A prospective cohort study (14) includes 40 AMD patients (mean age was 77.1 years ±5.9), shows that both uncorrected distance visual acuity (UCDVA) and uncorrected near visual acuity (UCNVA) were improved by 0.3 Log MAR and 0.4 Log MAR respectively. Another 1-year clinical trial done by Lane (15) found out that 62% of participants gained three or more lines in both distances and near best-corrected visual acuity (BCVA), whereas 77% gained two or more lines. The most essential non-randomized multicenter case-control trial done by Hudson from 2006 to 2015 (16-18) shows that among 217 participants (mean age was 76 years old), 53% of operative eyes reached a 3-line or more improvement in BCVDA and BCNVA. For the 2-year follow-up trial, there are 174 patients participated, and 59.5% gained 3-line or more of BCVA. The increased percentage of VA improvement may occur due to the age of subjects may younger and satisfied with the operation results. In 2015, 63 patients were followed and divided into two groups based on their age. Group 1 (n=31) was included patients from 65 to 75 years old, and Group 2 (n=32) was 75 and over. In the younger group, 58% retained 3-line or more after five years of operation, and only 38% improved in the elder group. The mean BCDVA has improved 2.41±2.69 lines in all patients, lower than results at 2 years (3.2 lines). Only 1 patient (3.2%) had lost ≥2 lines of vision than 3 (9.4%) in group 2. Comparing the different models of IMT in patients at 1 and 2 years all show the 3× device had more considerable improvement in BCDVA than those implanted with the 2.2× device (3.6±1.9 versus 2.8±2.3 lines, P<0.001).

All research illustrated above shows that younger patients whose age is beyond 75 may have more motivation and thoughts to understand fully and participants to the vision of extensive rehabilitation, which is the primary role in improving visual acuity and used to the eyes have different tasks. Therefore, they would gain more improvement compared with the older one. It may notice that postoperative low-vision rehabilitation was painful for elderly patients and should be communicated with them ahead of the surgery.

Quality of life and functional outcomes

Three articles evaluate the quality of life improvement, which is one of the essential tasks for IMT implantation in patients with advanced or end-stage AMD. Activities of daily life (ADL) questionnaire applied in 2 articles (15,16), which represents activities of daily living grouped by two variables: type of activity (static or dynamic) and focusing distance (distance, intermediate, or near).

It shows a mean 14.1-point difference improvement (41.4 to 55.8, P<0.001) with statistically significant for all vision range for both static and dynamic dimensions at 1 year (16). Moreover, patients with younger age and more significant improvement in BCVA tended to score higher (15).
One study uses NEI VFQ-25 scores and shows the mean scores were improved by more than 7 points on 7 of 8 relevant subscales and correlated with a more significant point increase in BCDVA and BCNVA at least 2 lines improvement (P=0.0175) (15). However, the Peripheral Vision subscale decreased significantly from 67.6 to 62.9 (P<0.001), which is explainable that it would be a new experience to use one eye solely focus on the central area and the sensitivity in both eyes would be decreased (11). The effectiveness and cost-effectiveness of the IMT were evaluated by Brown (19) in 2011 by using quality-adjusted life-year (QALY) gain to quantify and be comparable. The incremental cost-utility ratios of $19,302/QALY for IMT implantation versus $16,045/QALY for fellow eyes that underwent intra-study cataract extraction. The results suggest that IMT is also well within the conventional limits of incremental cost-effectiveness.

Safety outcomes
Safety outcomes were mainly pointing at the rate of complications or adverse effects (AEs) at different periods of following time after surgery. Although the earliest 3 cases serial study done by Kaşkaloğlu in 2001 (20) fail to provide enough evidence to show VA improvements, they still bring about the possible complications like mild iritis and persistent bubbles existing inside the IMT that proved by studies based on larger populations. The most common AE illustrated by multiple types of research is inflammatory deposits on the device from 17.5% to 42.9%, and the rate increased as time goes by from 21% at 1 year to 25% at 2 years (15-17). Another common AE is Pigment deposits on the device from 10% at 1 year to 11% at 2 years (16,17). The treatment-needed intraocular pressure (IOP) increasing within 7 days, detected in 28% of participants (16). The rate of corneal edema, like conventional cataract surgery, was higher in the elderly (7.1% versus 4.3%) (18).

The IMT replaced were seen in 3.9% to 15% because of patient request, corneal edema, and the failure of the IMT manufacturer’s safety precautions to cause bubbles inside existing (14,17,18). The glare in bright light and reduced peripheral vision and depth perception reported being complained and drove dissatisfaction to the surgery outcomes (17). That remains that those improvements in the quality of manufacture and make strict patient selection criteria are the way to decrease the future incidence of such complications (14).

Although the design for IMT tried very hard to keep the device away from the posterior endothelium, the loss of endothelium cell density (ECD) was still reported by all researchers. The mean cell density decreased by 20% at 3 months, 13% to 25% at 1 year, and 29% at 2 years (17,18). At 5 years, the mean ECD loss was up to 35–40% among patients over 65 (18). The ECD loss may be correlated to baseline anterior chamber depth (ACD). One study shows that patients with 3.0 mm or more ACD are losing less endothelium cell counts at 2 years comparing with ACD less than 3.0 mm (17).

One ongoing study in IMT compares the safety and effectiveness of implanting the IMT in pseudophakia patients and is also estimated to be done in 2020 (21). In the Future, IMT clinical outcomes evaluation would need more studies to compare individuals randomized to the device versus non-implanted severed as controls.

IOL-VIP System (Intraocular Lens for Visually Impaired People) (Figure 4)

Another magnifier Galilean type telescope is the IOL-VIP System, launched in 2001 consists of two special lenses. One is a high biconcave IOL (about −66 D) embedded in the capsular acted as the eyepiece, determining the magnification. Another one is a high biconvex IOL (about +55 D) in the anterior chamber as an objective that can further modify the image. A new model called the IOL-VIP Revolution has also been developed since April 2012, implanted in the capsular bag, unlike the classic form (22). Both IOLs are made of polymethyl methacrylate and can filter the ultraviolet light. The diameter of the two optics is 5 mm, and the total length is 13 mm, which can offer an estimated magnification for a distance of 1.3. After standard phacoemulsification, the thick IOL (axial thickness max is 1.5 for AC IOL and 1.5 mm for in-the-bag IOL) need further enlarge the corneal incision to 7 mm to facilitate the implantation. There is a software that can perform a functional evaluation of the objective and subjective Preferred Retinal Locus (PRL) and BCVA, contrast sensitivity, and reading speed.

Meanwhile, it can also be used for training the PRL before and after the surgery. One study shows that the IOL-VIP® software would recognize more than 66% of patients whose PRL may too far from the fovea and/or not responding adequately to pre-surgical training ahead of the surgery so that none of the articles we found had reported an implantation removal (23). All patients were requested to undergo a 2-week preoperative training (12 30-min training sessions) and a 3-month postoperative rehabilitation
program (5 30-min training sessions per week for 12) to train and consolidate the PRL (24).

There are 5 articles fitting into the evaluation of the efficacy and safety of IOL-VIP, and 107 patients in total participated in observational studies without any controls (22-27). Except for 1 article mainly discussed the IOL-VIP® software (23), others fit into the evaluation of the efficacy and safety of IOL-VIP. The mean follow-up time from 6 to 50 months.

A new study done by 2019 showed that 12 AMD patients with mean age 72.3 were enrolled, and BCVA pre- and post-operation were 1.08±0.14 and 0.81±0.16 log MAR and statistically significant between the difference (P=0.05) (25). Another noncomparative interventional case series (26) depicts the same improvements (the difference was 0.69 Log MAR), and the main best-corrected clinical gain was 44%. Another case series study included 35 consecutive patients for 20 months (24). Besides the improvement in BCVA (difference was 0.51 Log MAR), they also consider the reading magnification gain and postoperative reading distance gain, which was ×6.2 and 7.66 cm, respectively. To further evaluate the rehabilitation program’s consequence, two groups were signed up by their preoperative VA. Patients with better preoperative VA tend to move their PRL to a more favorable position. In contrast, patients with low VA tend to have an unstable and peripheral PRL, which cannot be corrected by the training program. One retrospective study of 50 patients (age range between 36 and 85) with central scotoma for 4.2 years also detect the improvement in VA from 1.3 Log MAR to 0.68 Log MAR (27).

There is only 1 article (25) using VFQ-25 Scores for quality of life evaluation, and the difficulty with activities showed a statistically significant improvement from 39.4±16.3 to 43.8±15.8. Those improvements seen in scales may be related to the software using before the surgery, which can fitly evaluate patients’ residual visual function with an accuracy of 78% and prognosis for visual improvement based on simulation of the postoperative condition and lower the patients’ expectation. Therefore, a further study which including life quality evaluation and functional outcomes, would need to consider IOL-VIP System as more clinical applications.

The most common complication reported by studies is transient IOP increased after the operation. About 40% (4/10) of patients had increased IOP (24±3 mmHg) but can resolve in 1 week with the administration of topical timolol maleate 0.5% and corticosteroids (26). In one study, they do the preoperative iridotomy in all other cases after noticing 3 of the first 5 cases developed a pupillary block with IOP increased (24). Although 1 article done by Dag et al. (25) found that ACD was deeper post-operation from 2.7±0.3 to 3.1±0.7, the mean endothelium cell count was still significantly decreased postoperatively from 2,437±398 to 1,849±376, but lack of further information about the percentage among the subjects. Another study proved that endothelium cell loss appeared in 11.1% of subjects and 5.2% from 12 to 18 months (26). The incidence of PCO was 2.9–18% and can be managed by Nd: YAG laser capsulotomy (24,27). All in all, IOL-VIP System has a high satisfaction by AMD patients with software that can simulate the postoperative VA improvement and fewer complications reported. However, lacking clinical controls makes those conclusions less evidential and needs more information to evaluate its efficacy and safety further.

**iolAMD**

The hyperopic acrylic iolAMD is another Galilean telescope based implant that consists of two lenses designed to an optimized retinal image to all areas of the macular...
≤10° from fixation to 3 degrees away by a decentration of one of the IOLs and magnifies the image 1.25 to 1.3×. IOL 1 is a high-plus power lens (+63 D) implanted into the sulcus, which can gain depth of focus, whereas IOL 2 is a high-minus power lens (−49 D) embedded into the capsular bag. IOL 1 is 5 mm, and the overall diameter is 11.75 to 12.0 mm, with the haptic bent anteriorly. The optic of IOL 2 is 4 mm, overall length is 11.0 mm, and haptic vaulted posteriorly about 15°. After implantation, the distance recommended between optics of two lenses is 2 mm (28,29). The device can be folded, injected with a tip cartridge, and implant through a 3 mm non-suture incision without causing corneal astigmatism (30). The device’s optics generate transverse asphericity and maintain a breadth of focus across the macula (31) (Figure 5).

Two case series studies were included (29,31) with 15 patients participants, and none of the studies has controls and evaluates the quality of life-based on different scales. For one case series study (31), whose only evaluated 3 patients with AMD, the mean CDVA was improved 0.40±0.17 (from 0.08 to 0.5) and predicted CNVA was improved 0.49±0.13 (from 0.15 to 0.6) for 1 month. Moreover, at the end of 3 months after the operation, the mean CDVA was improved to 0.6, and the CNVA remained stable. Another prospective intervention pilot study with 12 patients followed for 4 months (29) shows that the mean decimal CDVA gain 67% (from 0.12 preoperatively to 0.20) and similar results in CNVA with less improvement about 50% (from 0.14 to 0.21). It also illustrated that the mean change in spherical equivalent (SE) was −1.5 D with a level of 0.5D of induced astigmatism, and it shows a myopic shift (from −0.1 D to −1.6 D after 4 months). Furthermore, the mean cylinder also improved. Microperimetry testing is also used to indicate a magnification effect and a deviation of images from the central of fovea up to 5° with fixation stability. Meanwhile, the mean threshold sensitivity increased from 7.2 to 20.2 dB. At baseline, the fixation points within a 2-degree circle and a 4-degree circle were 31% and 71%, respectively, whereas, after implantation, the number of points increased to 48% and 88%.

Most studies recorded the incidence rate of IOP increasing. One study did a precautionary intraoperative iridectomy among 9 patients (12 patients in total), and there is only one patient whose IOP arouse from 14 to 22 mmHg resolved by Nd:YAG laser (29). ECD reduced by 18% at 4 months, and the rate of corneal edema was 16.7%, also reported by one study (29). One patient’s IOLs in capsular bag vaulted anteriorly, which caused vision quality reduction and replaced with a smaller-size ordinary IOL. Except that, the other 11 patients checked by AS-OCT and found no evidence of gross deviation from the expected position (29).

Some demerits are existing in this device. First of all, two lenses combination restricted to eyes with previous cataract surgery and standard monofocal IOL implantation. Moreover, if the PRL changes with the progression of AMD, it would need further surgical manipulation, and patients usually use multiple PRLs for daily activities. A single restricted area would cause visual function damaging (30). Therefore, it is not offered by the manufacturer anymore today.

**EyeMax Mono**

According to the demerits of previous iolAMD devices, the advent of EyeMax Mono was approved in 2016 (32) and is similar to standard IOL, excepts for its optics that use hyperspherical design to increase the breadth of focus and the quality of the image supplied to all areas of the macula at ≤10° of retinal eccentricity and magnifies the image ×1.1 to ×1.2 (33). One of the most advanced parts is...
to provide a high-quality image with reduced blur in those areas where photoreceptor cell densities may still afford VA of 6/30 Snellen or better. It’s a new class of injectable, soft acrylic, extended macular vision IOLs (34) and permitting subjects to form single or multiple PRLs from their normal part of the macula as much as possible (35). There are two versions of the EyeMax mono: one is designed for capsular bag implantation following phacoemulsification. A second version is intended for sulcus implantation with standard, monofocal IOL that had already been implanted. The second version is a −3.5 D lens with a 5.0-mm hyperspherical optic component centered on a plate haptic component and delivers a modest hypermetropia correction. The overall diameter is 11.75 to 12 mm, and the haptic is vaulted posteriorly to reduce the risks associated with sulcal IOL implantation. The implant requires a modest postoperative refractive target of +2 D to +3 D for up to 1.2 D magnification (31). The EyeMax Mono can be binocularly implanted, and hypermetropia refractive target can be chosen to generate 10% to 20% magnification with spectacle correction in severe AMD cases (34) (Figure 6).

There were three consecutive cases of series studies included with 374 eyes. Robbie did the initial prospective interventional pilot assessment of the EyeMax Mono in 2018 (36), and 8 eyes of 7 patients with bilateral dry AMD underwent phacoemulsification and capsular bag implanted with 2 months follow-up postoperatively. Visual acuity improved for all patients, and mean CDVA improved from 0.93±0.22 preoperatively to 0.59±0.25, and the reading speed was reported a 57% improvement from 28±19 to 44±31 words per minute. Another prospective case series study included 22 pseudophakia eyes with 6 months follow-up (33). Both eyes of participants underwent small-incision sulcus implantation. The mean CDVA has improved ≥2 lines in all participants’ eyes. The evidence of improvement in CDVA over time from 0.90 Log MAR preoperatively to 0.51 Log MAR postoperatively at 6 months may suggest visual outcome improvement consists of a neuroadaptive process. Another consecutive case series of 244 eyes with mean age 80 years was conducted in 2018 and followed up for 3 months (37). Eighty-eight percent of cases reached the refraction target, and mean CDVA improved from 1.06 preoperatively to 0.71 postoperatively, equating to gain about 18 ETDRS letters. Mean CNVA was also improved from 1.36 to 0.88 postoperatively. Recently meta-analysis indicated that CDVA improvement with standard IOL implantation was only 6.5–7.5 ETDRS letters after 6–12 months of follow-up (38). In Austria, the first case report published in 2019 shows an 83-year-old Caucasian female with stable AMD in both eyes and did the operation on the left eye. It turns out after 6 months follow-up that her BCDVA improved from 0.2 to 0.5 (decimal), and BCNVA improved from 0.05 (decimal) to 0.2 (decimal) with no further long- and short-term complications (34).

The mean differences in endothelial cell counts were 143, and 0.8% (2/244) experienced steroid-induced IOP increasing at 4 weeks after surgery, which can be under controlled on topical hypotensive drugs and quite the treatment of steroid treatment (37). Another study showed that 9% of subjects with pseudophakia eyes experienced IOP elevated after the surgery. So far, there was no report about diplopia and lens misposition (31).

Based on the evidence provided by case series studies, cataract extraction with EyeMax Mono implantation performed safely. It can benefit the subject’s visual outcomes and optimized images across the macula. Other high-scale evidence levels with more subjects’ studies would need to evaluate the safety, efficacy fully, and, most importantly, the quality of life and functional outcomes of this device.

**Scharioth macular lens (SML)**

The fundamental optical approach for this new add-on macular lens, SML, is near triad reflex (39). When patients focus on near objects with a constricted pupil, and light rays are restricted mainly to the central optical area, that would have a magnification effect. Still, when patients focus on distance with dilated pupil, light rays will pass through the peripheral regions, and it would not be influenced. The central optic region provides relatively high refraction to achieve sharp vision in a range of 10 to 15 cm from the

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**Figure 6** EyeMax Mono. It was supplied by London Eye Hospital Pharma Company.
eye. Cones that for color vision and VA concentrated in
the fovea and Rods that are responsible for night vision,
sensitive motion detection, and peripheral vision are
abundant elsewhere in the retina. The reason why SML
does not distribute light away from the fovea since we
can detect motion and contours with non-foveal regions
of the retina expect reading in the dark so that the device
contributes to magnify the rest of the damaged fovea to
enable reading.

The Schirioth macula lens (SML) is a one-piece foldable
intraocular hydrophilic acrylic “add-on” lens with an overall
diameter of 13 mm and a 1.5-mm central portion with a
refractive power of +10 D solely in pseudophakia eyes. It is
implanted into the ciliary sulcus as an add on to the existing
in-the-bag IOL through an incision of 2.2 mm. It can be
implanted simultaneously with standard phacoemulsification
or at a late date (40). Wet AMD, iris neovascularization,
progressive glaucoma, zonulopathy, and shallow ACD are
not recommended for this device (39) (Figure 7).

There were 3 articles included in this review about
SML, with 91 patients participated for follow-up from 3 to
12 months (41-43). In 2019, a prospective multicenter
clinical trial done by Srinivasan reported the visual outcomes
following monocular SML implantation in pseudophakia
eyes with AMD (41). It includes 50 patients and a follow-up
for 12 months. They only had CDVA of 0.4 to 0.1 (decimal),
and CNVA can be improved about at least three lines when
tested with a +6.0 D at reading distance of 15 cm and +2.5
D 40 cm. The mean preoperative CNVA at 40 cm with
+2.5 D addition was 0.23±0.12 (decimal) preoperatively, and
after 1 year, the UNVA at 15 cm can improve to 0.57±0.33,
similar as screening test measurement preoperatively.
Furthermore, the mean CDVA remained unchanged at
1 year and confirmed that the SM does not affect the
distance VA. No intro- and post complications have been
reported. A retrospective study among 15 pseudophakia
patients for follow-up 3 months also improved the CNVA
and stable DVA (42).

Only one article presented in 2018 evaluated 26 patients’
quality of life by VFQ-25 after SML implantation (43)
showed 35% of patients achieved normal reading vision
with a gain of at least 3 lines and 28% overall improvement
on the questionnaire, especially for patients with legal
blindness.

Other approaches

A new foldable one-piece high-add IOL called LENTIS®
MAX, LS-313 MF80 (Teleon Surgical B.V, Berlin, Germany) (44) consisting of hydrophilic acrylates with a hydrophobic surface has been reported (Figure 8). IOL’s overall diameter is 11 and 6 mm optic, which can be implanted through a 2.2 mm incision. The new feature is a second additional near segment on the lens’s posterior surface, and the lens designed an aspheric biconvex that provides an additional power of 8.0 D, equating to 6.0 D at the spectacle plane. The design of the lens can magnify 1.5x at a distance of 30 cm and 3x at 15 cm. Moreover, a case report of an end-stage AMD patient implanted this high-add IOL was published and suggested improvement in both distances and near VA.

**Discussion**

There is no further evidence to evaluate the safety and efficacy of Fresnel Prism Intraocular Lens. As for Lipshitz macular implant (LMI) and LMI-SI, although there are also no further clinical studies, an ongoing randomized clinical trial to compare the OriLens intraocular telescope with standard low vision training in eyes with end-stage AMD (MIRROR 2017) (45). The results would expect in 2020. The essential characteristics and parameters about the lens discussed above had summarized in table implantable miniature telescope (IMT) was the only approved surgical device by FDA in the US and can provide central vision magnification at the cost of losing peripheral vision. Meanwhile, the incision for implantation is larger enough that it needs 6–8 sutures to close the wound and may cause severe astigmatism. IOL-VIP System and EyeMax Mono involves a smaller incision and be suitable for both phakic and pseudophakic eyes with similar magnification. Visual outcomes based on the restricted number of participants suggest that the VA improvement in EyeMax Mono is modest compared with IOL-VIP without extensive visual rehabilitation. The big breakthrough about EyeMax Mono was to provide a high-quality image in all macular areas extending up to 10° from the fovea in case of PRL changes or AMD progression. And the improvement in visual acuity had been proved to be case series of 244 subjects. There is no doubt that a case-control study with quality of life and function evaluation would conduct further.

Recently, a prospective multicenter clinical trial about the SML in previous pseudophakic eyes with AMD was published in 2019 (41) and provided more information about the visual and refractive outcomes. Although the SML does not offer distance magnification, it can magnify 2x for objects in a range of 10–15 cm from the eye and can be implanted through a 2.2 mm incision with a reduction of postoperative astigmatism.

For validity and generalizability of the studies included, VA outcomes vary in nature so that comparisons of efficacy and safety are difficult. No randomized controls in each IOL except IMT. Others are all case series with restricted subjects and limited follow-up duration that would not provide enough convincing evidence. Moreover, most of the studies did not limit the age range in the inclusion criteria, and long-term evidence and quality of life evaluation are insufficient in most of IOL. Due to the long and complicated rehabilitation after operations, understanding, and adequate expectations from patients, especially end-stage, should be informed ahead of time. In Table 2 we present characteristics of the significant intraocular devices available today for AMD patients.

All in all, those technological advancements in the treatment of AMD would bring new hope in VA and quality of life improvements to patients with no other treatment options previously. To date, more extensive, randomized, and long duration follow-up clinical trials are needed to further evaluation of safety and efficacy among different IOL based on other optic fundamentals. Based on the evidence so far, there are various strengths and limitations. Ophthalmologists’ options should value multiple aspects about the ocular anatomy that thick IOL may cause pupil block and increased IOP, living habits especially for reading and costs for individual AMD patients with low vision.
| Characterised in devices | IMT | IOP-VIP | iolAMD | EyeMax Mono | Scharioth macula lens (SML) | LMI-SI |
|--------------------------|-----|---------|-------|-------------|--------------------------|--------|
| Optical Fundamentals     |     |         |       |             |                          |        |
| Galilean type telescope; telephoto effect that enlarge images in the central VF |     |         |       | Single lens; hyperaspheric design increase the quality of the image of the macula at ≤10° of retinal eccentricity | Near Triad Reflex; central optic area providing high additional +10 D |        |
| Country                  | USA | Italy   | United Kingdom | United Kingdom | Switzerland | Israel |
| Magnification            | 3x  | 1.3x    | 1.25x to 1.3x | 1.1x to 1.2x | 2.0x        | 2.5x   |
| Material                 | Quartz glass | Polymethyl methacrylate (PMMA) | Hydrophobic acrylic | hydrophobic acrylic | Hydrophilic acrylic |     |
| Magnification            |     |         |       |             |                          |        |
| Diameter (mm)            | Telescope 4.4; haptic-to-haptic diameter: 13.5; lens aperture: 3.2 | Diameter 5; maximum AT for AC IOL 1.5; peripheral thickness for IOL in capsular bag 1.5; length 13 | IOL 2; optic 4, overall length 11; haptic vaulted posteriorly about 15°; IOL 1; optic 5, diameter 11.75–12 m; the distance between IOLs 2 | Optic 5; overall length 11.75–12; the haptic is vaulted posteriorly | Overall diameter 13 with 4 flex haptics; central optic diameter 1.5 | Diameter 5–6; central thickness 1.25 |
| Foldable                 | No  | No      | Yes   | Yes         | Yes          | No     |
| Lens Status              | Phakic | Phakic or pseudophakic eyes | Phakic eyes | Phakic or pseudophakic eyes | Pseudophakic eyes | Phakic or pseudophakic eyes |
| Incision size (mm)       | 10–12 | 7       | 3     | 2.8         | 2.2         | 5–5.5  |
| Implant position         | Capsular bag | Capsular bag and AC | capsular bag (IOL 2) and sulcus (IOL 1) | Capsular bag or sulcus | Giliar sulcus | Sulcus |
| Binocular/monocular       | Monocular | Binocular | Binocular | Binocular | Monocular | Binocular |
| Need PRL pick            | No  | Yes     | No    | No          | No          | No     |
| Rehabilitation needed     | Yes | Yes     | Yes   | No          | No          | Yes    |
| Pros                     | Greater magnification; works for both far and near; the only device with FDA approval | IOL-VIP Revolution can be used for wet AMD, glaucoma and degenerative myopia | Foldable, 3 mm sutureless incision, can dial the lens to another position if AMD expansion | Foldable, small sutureless incision, high-quality image in all areas of the macula extending up to 10° from the fovea in case of PRL changes or AMD progression. No extensive visual rehabilitation | Small incision, independent from lens status; no reduction of VF | Keep peripheral VF remain normal; high magnification |
| Cons                     | Big size, 12 or 7 mm sutured incision, endothelium cell loss, reduce the VF to 20–25° | Long surgery, 8 mm sutured incision, need prophylactic peripheral iridotomy | Relative unpredictability in the location of the lenses which could affect the final refraction | Relative unpredictability in the location of the lenses which could affect the final refraction | Does not provide distance vision magnification; magnification only achieved in a range of 10 to 15 cm from the eye | The risk of glare effects |

Note: https://www.medicontur.com/scharioth_macula_lens. Near Triad Reflex: miosis-accommodation-convergence. **AT, axial thickness; PRL, prefer retinal location.
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Footnote

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