Outcome comparison between wavefront-guided and wavefront-optimized photorefractive keratectomy: A systematic review and meta-analysis

Khaled M Hamam1,2, Mohamed I Gbreel1-2, Randa Elsheikh1,2, Amira Y Benmelouka2-4, Yassamine Ouerdane2-4, Amr K Hassan2-4, Aboalmagd Hamdallah5,6, Ahmed B Elshohry2-7, Anas Z Nourelden2-7, Ahmed T Masoud2-8, Asmaa A Ali2-9, Khaled M Ragab2,10, Ahmed M Ibrahim2,7

Photorefractive keratectomy (PRK) eye surgery is widely used for patients at risk for corneal ectasia to maintain an aspheric corneal shape. Wavefront-guided (WFG) ablation profile was designed to reduce pre-existing higher-order aberrations (HOA). We aimed to compare the corneal aberrations and visual outcomes between WFG and Wavefront Optimized (WFO) PRK in patients with myopia. Eight randomized clinical trials were included. We searched PubMed, Scopus, Web of Science and CENTRAL at March 2020, and updated the search in September 2020 using relevant keywords, The data were extracted and pooled as Mean Difference (MD) with a 95% Confidence Interval (CI), using Review Manager software (version 5.4). Pooled results showed no significance between Uncorrected Distance Visual Acuity (UDVA) and Corrected Distance Visual Acuity (CDVA) between both groups underwent WFG and WFO PPR after three months follow up (MD = -0.03; 95% CI: [-0.06, 0.00]; P = 0.07), (MD = -0.02; 95% CI: [-0.04, 0.01]; P = 0.22) respectively. Although, no significant difference between mean manifest cylinder after three and 12 months follow up, but the total MD for mean manifest cylinder difference was significantly lower with the WFG treatment method (MD = -0.12. 95% CI: [0.23-0.01], P = 0.03). This shows a slight advantage of the WFG over the WFO method. The visual performance showed similarity and excellent refractive outcomes in both WFG and WFO PRK. No significant statistical differences between the two approaches. On further comparison, there was a slight advantage of the WFG over the WFO method.

Key words: Higher-order aberration, meta-analysis, myopia, wavefront-optimized photorefractive keratectomy, wavefront-guided photorefractive keratectomy

Patients undergoing refractive surgery aim to achieve the best possible level of spectacle free vision.11 Although laser in situ keratomileusis (LASIK) is the most commonly selected refractive procedure for rapid postoperative recovery, PRK is still used for patients at risk for Corneal ectatic disorders and to avoid potential flap complications intraoperatively and in the future.12-13 However, UDVA can be improved by Myopic laser refractive surgery by producing an oblate cornea; visual quality is not enough post-surgery.14 In myopic Laser refractive surgery, because the laser beam enters the periphery, some parts are reflected, and the circular beam becomes elliptical resulting in a decrease in the effectiveness of the laser energy.15 Under-ablation of the peripheral cornea can be induced by these factors increasing HOA, especially spherical aberration.16

To overcome the problem of myopic laser refractive surgery, WFO ablation was designed to maintain an aspheric corneal shape by applying extra laser pulses to the peripheral cornea to diminish induction of spherical aberration.17 Another procedure, WFG ablation profile was designed to reduce pre-existing HOA.15 A few published studies compared WFG treatments with WFO treatments in patients undergoing LASIK17 and some reports17 show an improvement in the quality of vision after WFG treatment, but others11,13 showed no difference between the two profiles.18 WFO laser ablation profiles are designed to deliver additional treatment to the peripheral cornea in an attempt to preserve the naturally prolate shape of the cornea and minimize induction of higher-order aberrations while protecting the present aberrations of the eye.13 WFG treatments attempt to correct both lower- and higher-order aberrations requiring preoperative measurement of the eye’s aberrations using a Wavefront aberrometer.19

In this systematic review and meta-analysis, we aimed to compare the visual outcomes and corneal aberrations between WFO and WFG PRK and evaluate their effect on myopia for

1Faculty of Medicine, October 6 University, Giza, Egypt, 2Faculty of Medicine, University of Algiers, Algiers, 3Faculty of Medicine, Saad Dahlab University, Bliida, Algeria, 4Faculty of Medicine, South-Valley University, Qena, 5Faculty of Medicine Al-Azhar University, Damietta, 6Faculty of Medicine, Al-Azhar University, Cairo, 7Faculty of Medicine, Fayoum University, Fayoum, 8Faculty of Medicine, Assuit University, Assiut, 9Faculty of Medicine, El-Minia University, Minia, Egypt, 10International Medical Research Association (IMedRA)

Correspondence to: Dr. Mohamed Ibrahim Gbreel, Sharqia, Awlad-Sakr, Egypt. E-mail: mohamedgbreel207@gmail.com

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patient-perceived quality of vision to determine whether one treatment profile leads to more optimal vision than the other.

**Methods**

We followed the PRISMA statement guidelines\(^{[13]}\) during the preparation of this systematic review and meta-analysis and performed all steps according to the Cochrane handbook of systematic reviews of intervention.\(^{[14]}\) The PRISMA flow diagram for studies selected in the search process and eligibility assessment are summarized in Fig. 1.

**Literature search strategy**

We searched PubMed, SCOPUS, Web of Science and Cochrane CENTRAL through at March 2020, and updated the search in September 2020 for relevant clinical trials comparing optimized and guided photorefractive keratotomy. We used the following search strategy: (Wavefront-guided photorefractive keratectomy OR Wavefront-optimized photorefractive keratectomy) AND “Myopia”).

**Eligibility criteria and study selection**

We included studies that followed these criteria: (1) population are patients undergoing photorefractive keratotomy, (2) intervention and comparator are Wavefront-guided versus Wavefront-optimized photorefractive keratotomy operations, (3) study design: we included randomized clinical trials with no restrictions for languages. Besides, the references of included trials and relevant reviews were screened to ensure high-quality searching. We excluded animal trials, conference abstracts, non-randomized trials, and studies without relevant outcomes.

Two authors independently screened the title and abstract of all expected included studies following by full-text screening and then manually searching the references of the finally included papers eligible to meta-analysis.

**Data extraction**

Two independent authors performed the extraction step, and any disclosure among authors was resolved through discussion and referred to the study senior. We extracted data in a formatted data extraction excel sheet including (1) Summary as NCT, study design, population, Duration of treatment, and intervention [shown in Table 1]. (2) Baseline characteristics such as age, sex, mitomycin C use, Trefoil, and spherical aberration [shown in Table 2]. (3) Outcomes as UDVA, CDVA, mean manifest sphere, mean manifest spherical, and spherical equivalent.

**Quality assessment**

We used Cochrane’s risk of bias tool to perform the quality assessment; the tool is found in chapter 8.5 of the Cochrane

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**Figure 1:** The PRISMA flow diagram for included studies.

**Figure 2:** Risk of Bias assessment summary of the included studies.
### Table 1: Data summary

| Study ID.     | NCT       | Study Design                                   | Population | Duration of treatment | Intervention                                                                 | Comparison                                                                 |
|--------------|-----------|-----------------------------------------------|------------|-----------------------|------------------------------------------------------------------------------|---------------------------------------------------------------------------|
| Toy et al. 2016 | -         | Prospective, Randomize, Fellow eye-controlled clinical trial | 71 patients | 12 months follow up   | WFO-PRK using the WaveLight Allegretto Wave-400 Hz (Alcon)                   | WFG-PRK using the VISX CustomVue Star S4 IR                               |
| Sia et al. 2015 | -         | Prospective, randomized open-label study prospective, randomized clinical trial | 108 patients | (1,3,6,12) months follow up | Wavefront-guided                                                            | Wavefront-optimized treatment                                             |
| Ryan et al. 2017 | 1097525  | -                                             | 56 patients | (6,12) months follow up | Wavefront-guided (WFG)                                                      | Wavefront-optimized (WFO) photorefractive keratotomy (PRK) VISX CustomVueTM STAR S4 IRTM Excimer Laser with Active Track |
| Moshirfar et al. 2011 | -         | Prospective, single-masked, randomized, fellow-eye study | 23 patients | Three months follow up | WaveLight Allegretto system (Wavefront-optimized group), which utilizes the WaveLight® Allegretto 400 Hz Wave® Eye-Q Laser | Wavefront-guided PRK (VISX CustomVue Star S4 IR excimer laser system)       |
| Maurer et al. 2014 | -         | prospective randomized study                  | 27 patients | (1,3,6) months follow up | WFG PRK, WFG LASIK                                                           | WFG PRK and WFG LASIK                                                     |
| Lee et al. 2016 | 1135719   | prospective randomized study                  | 68 patients | 12 months follow up   | Wavefront-guided PRK (VISX CustomVue Star S4 IR excimer laser system)        | Wavefront-optimized PRK (WaveLight Allegretto Wave EyeQ 400 Hz excimer laser system; Alcon Laboratories, Inc., Fort Worth, TX) |
| He et al. 2015 | 1135719   | Prospective, randomized, fellow-eye controlled clinical trial | 71 patients | (1,3,6,12) months follow up | WFG PRK (Visx CustomVue Star S4IR excimer laser system; Abbott Medical Optics) | WFG PRK (WaveLight Allegretto Wave EyeQ 400 Hz excimer laser system; Alcon Surgical) |
| He et al. 2014 | 1135719   | Prospective, randomized, fellow eye controlled study | 71 patients | (1,3,6,12) months follow up | WFG PRK treatment by the VISX CustomVue Star S4 IR excimer laser system      | WFG PRK treatment by the WaveLight Allegretto Wave EyeQ 400 Hz excimer laser system |

**Figure 3:** (a) Uncorrected Distance Visual Acuity (UDVA) after 3 months follow-up. (b) Corrected Distance Visual Acuity (CDVA) after 3 months follow-up. (c) Spherical Equivalent after 3 months follow-up
handbook of systematic reviews of interventions 5.1.0. [5] This ROB assessment tool included the following domains of bias: selection, performance, detection, attrition, reporting biases, and other potential sources of bias. Author judgments fall into three categories: low, unclear, or high risk of bias for each domain. We used the quality assessment table provided in (part 2, Chapter 8.5) the same book. [5] We could not assess publication bias because of the small number of included studies, less than ten studies. [5]

Data synthesis

We used the mean difference (MD) to perform analysis of continuous outcomes, and risk ratio (RR) to analyze dichotomous outcomes. The analysis was performed using (Review Manager Software, version 5.4) under a fixed-effect model in all outcome data. Statistical heterogeneity between studies was assessed by observation of the graphs and measured by Chi-square test and I-squared (I^2) test for the degree of the heterogeneity. We conducted subgroup analysis according to postoperative follow-up of UDVA to stratify the surgical efficacy on UDVA.

Results

Literature search

The initial search resulted in 374 articles from 4 databases: 77 articles from PubMed, 56 articles from Cochrane, 93 articles from Scopus, and 138 articles from Web of Science. Of these 374 articles. We excluded 108 articles due to duplication. 266 articles underwent title, and abstract screening, and 241 were excluded because they did not follow our PICO criteria. The remaining 25 articles underwent full-text screening. A total of eight papers were finally included for the final qualitative analysis and six articles for quantitative analysis.

Quality assessment

Based on the Cochrane risk of bias tool that is found in chapter 8.5 of the Cochrane handbook of systematic reviews of interventions 5.1.0, [5] none of the eight studies had the selection, attrition, or reporting bias. 2 studies had performance bias as the participants and personnel were not fully blinded, and 1 study had a different type of bias. (Risk of bias summary shown in Fig. 2).

Outcomes

1. UDVA

After 3 months follow up, the pooled studies showed no significant difference in UDVA of 20/20 in the eyes underwent WFG and WFO PRK (MD = -0.03; 95% CI: [-0.06, 0.00]; P = 0.07). Pooled results were homogenous (I^2 = 0%, P = 0.93) [8,17]. Fig. 3a.

2. CDVA

The pooled studies showed no significant difference in CDVA of 20/20 in the eyes underwent WFG and WFO PRK after 3 months follow up (MD = -0.02; 95% CI: [-0.04, 0.01]; P = 0.22). Pooled results were homogenous (I^2 = 0%, P = 0.70) [8,17]. Fig. 3b.

3. Spherical Equivalent

The pooled studies showed no significant difference in the spherical equivalent of 20/20 in the eyes underwent WFG and WFO PRK after 3 months follow up (MD = 0.04; 95% CI: [-0.51, 0.58]; P = 0.89). Pooled results were homogenous (I^2 = 0%, P = 0.93) [8,17]. Fig. 3c.
Figure 4: Follow up assessment of UDVA
4. Follow up assessment of UDVA

The pooled studies showed no significant difference in UDVA of 20/10 in the eyes underwent WFG and WFO PRK after 6, 12 months follow-up (RR = 0.56, 95% CI: [0.26, 1.22], P = 0.15). Pooled results were homogenous (I² = 50%, P = 0.16). [18,19] UDVA of 20/15 in the eyes underwent WFG and WFO PRK after 1, 3, 6, 12 months follow-up (RR = 1.06, 95% CI: [0.93, 1.22], P = 0.37). Pooled results were homogenous (I² = 41%, P = 0.15). [17-20] UDVA of 20/20 in the eyes underwent WFG and WFO PRK after 1, 3, 6, 12 follow-up (RR = 1.03, 95% CI: [0.96, 1.11], P = 0.35). Pooled results were homogenous (I² = 48%, P = 0.10). [17-20] UDVA of 20/25 in the eyes underwent WFG and WFO PRK after 6, 12 months follow-up (RR = 1.00, 95% CI: [0.98, 1.02], P = 1.00). Pooled results were homogenous (I² = 0%, P = 1.00). [18,19] UDVA of 20/30 in the eyes underwent WFG and WFO PRK after 1, 3, 6, 12 months follow-up (RR = 1.01, 95% CI: [0.93, 1.09], P = 0.89). Pooled results were homogenous (I² = 0%, P = 0.94). [17-19] UDVA of 20/40 in the eyes underwent WFG and WFO PRK after 1, 3, 6, 12 follow-up (RR = 1.01, 95% CI: [0.93, 1.09], P = 0.89). Pooled results were homogenous (I² = 0%, P = 0.89). [17-19] UDVA of 20/50 in the eyes underwent WFG and WFO PRK after 6, 12 months follow-up (RR = 1.00, 95% CI: [0.98, 1.02], P = 1.00). Pooled results were homogenous (I² = 0%, P = 0.94). [17-19] UDVA of 20/60 in the eyes underwent WFG and WFO PRK after 1, 3, 6, 12 months follow-up (RR = 1.00, 95% CI: [0.98, 1.02], P = 1.00). Pooled results were homogenous (I² = 0%, P = 0.94). [17-19] UDVA of 20/70 in the eyes underwent WFG and WFO PRK after 6, 12 months follow-up (RR = 1.00, 95% CI: [0.98, 1.02], P = 1.00). Pooled results were homogenous (I² = 0%, P = 0.94). [17-19] UDVA of 20/80 in the eyes underwent WFG and WFO PRK after 1, 3, 6, 12 months follow-up (RR = 1.00, 95% CI: [0.98, 1.02], P = 1.00). Pooled results were homogenous (I² = 0%, P = 0.94). [17-19] UDVA of 20/90 in the eyes underwent WFG and WFO PRK after 6, 12 months follow-up (RR = 1.00, 95% CI: [0.98, 1.02], P = 1.00). Pooled results were homogenous (I² = 0%, P = 0.94). [17-19] UDVA of 20/100 in the eyes underwent WFG and WFO PRK after 1, 3, 6, 12 months follow-up (RR = 1.00, 95% CI: [0.98, 1.02], P = 1.00). Pooled results were homogenous (I² = 0%, P = 0.94). [17-19] UDVA of 20/125 in the eyes underwent WFG and WFO PRK after 1, 3, 6, 12 months follow-up (RR = 1.00, 95% CI: [0.98, 1.02], P = 1.00). Pooled results were homogenous (I² = 0%, P = 0.94). [17-19]

5. Mean Manifest sphere

The pooled studies showed no significant difference in the mean manifest sphere after 3 months follow up (MD = 0.09%, 95% CI: [-0.46, 0.63], P = 0.76), and after 12 months (MD = 0.14, 95% CI: [-0.46, 0.70], P = 0.62). [8,17,21] Pooled results were homogenous at 3 and 6 months (I² = 0%, P = 0.99), (I² = 0%, P = 0.90) respectively. The total mean difference is 0.11 D for WFG vs WFO (95% CI: [0.28, 0.5], P = 0.57). The pooled results were homogenous (I² = 0%, P = 0.89).

6. Mean manifest cylinder

The pooled studies showed no significant difference in the mean manifest cylinder after 3 months follow up (MD = 0.09, 95% CI: [-0.23, 0.05], and after 12 months (MD = -0.16, 95% CI: [-0.33, 0.09]).
A total of 96 patients out of 164 patients in the WFG group had Mitomycin C applied to their eyes.[18] Comparing the change in HOAs from baseline to treatment groups, the change in coma was 0.2 microns in the WFG group (0.12 SD) and 0.21 microns in the WFO group (0.12 SD). Changes in other types of HOAs, such as trefoil, and spherical aberration were 0.16 microns (0.09 SD) for WFG and 0.17 microns (0.11 SD) for WFO group in trefoil and spherical aberration and 0.1 microns (0.12 SD) for WFG and 0.11 microns (0.14 SD) for WFO group in spherical aberration.[9]

**Discussion**

We established a systematic review and meta-analysis to compare the visual outcomes and corneal aberrations between WFG and WFO PRK in patients with myopia. The data from our analysis estimated that there is no significant difference between WFG and WFO in UDVA, CDVA, mean manifest sphere and spherical equivalent but the total of MD of manifest cylinder after three and 12 months follow-up was significantly lower with the WFG ablation profile. To our knowledge, we have comprehensively reported the outcomes following WFG and WFO PRK covering up to 12 months after surgery.

The wavefront-based technique has better clinical results and visual performance when compared to conventional therapy.[10,12,22] WFO and WFG photoablation techniques also reduce the postoperative occurrence of HOAs comparing to other conventional therapies.[23]

He et al. found no significant induction of HOAs after WFO vs WFG therapies. Still, WFG had a significant advantage over WFO in postoperative UDVA, CDVA, residual refractive errors, and contrast sensitivity.[9] Moshifar et al. investigated the three months outcomes following WFG and WFO ablation profile in PRK and found that both platforms produced equivalent refractive safety, predictability, and change in spherical aberration; but WFG PRK yielded slightly better results in contrast sensitivity.[14] Sia et al. demonstrated that both techniques have comparable excellent refractive efficacy, safety, predictability, and stability profiles.[28]

Yu et al. observed 100 patients randomized to WFG treatment or WFO LASIK treatment in both eyes with questionnaires which showed no difference between the two groups in terms of patient satisfaction.[29] Patients were observed for six months and questioned about the level of glare, halos, light sensitivity, fluctuations of vision, ghost images, and starbursts they experienced. They again found no differences between the two treatment groups except perhaps an increase in ghost images in the WFG group at six months. Similar to previous studies, symptoms increased from baseline at the first month but returned to preoperative levels from three months onward. This reversal in the symptoms was most likely due to epithelial remodelling.[24]

Although WFG treatment has the advantage of precisely individualized correction of higher-order aberrations at the time of surgery, small gains in visual acuity, predictability, and HOAs compared to WFO in laser-assisted LASIK, both methods of PRK have been shown to result in comparable visual results.[8] It is important to be able to predict the change in corneal curvature induced for a given amount of refractive correction on both treatment modalities for preoperative planning and future intraocular lens selection. Regarding future cataract surgery, after myopic laser photorefractive keratectomy, corneal power calculations are typically overestimated, causing the selection of underpowered intraocular lenses, and subsequently, the hyperopic surprise may occur.[10,12,27-34]

WFG and WFO treatment profiles differ such that while WFG was developed to consider pre-existing HOAs, WFO was designed to consider HOAs induced by conventional treatment, particularly spherical aberration.[18] Yet again, WFO treatment may be preferred if it has overall equivalent outcomes to WFG treatment because WFO can be more easily performed without needing to analyze aberrometry data.[16] Several studies have compared the quality of vision after WFG and WFO treatments in patients undergoing LASIK.[11,12,35,37] Some have shown no difference between the two profiles.[36,37] However, other studies suggested improved results with WFG treatment.[34,37]

**Quality of evidence**

We included six Randomized Clinical Trials (RCTs) to the quantitative analysis constituting a strong level of evidence. All steps were performed according to the Cochrane handbook and PRISMA checklist.[14,15] By measuring the quality level, the included studies are ranged from low, moderate to high quality.

The main strength and limitation of our study

This is the most comprehensive report to date on outcomes following WFG and WFO PRK covering up to 12 months postoperative refractive outcomes. All outcomes were homogenous under the fixed-effect model. It is important to note that our review was limited by studies reporting UDVA at different times following the procedure, some studies reported UDVA at three months, others at 6, while others at 12, and some studies had strict enough follow up to report it on all those time periods. Also, we have included a limited number of trials.

**Conclusion**

Results of the visual performance show that eyes achieve similar and excellent refractive outcomes in both WFO and WFG PRK. There are no significant statistical differences between the two approaches. Only slightly better results, lower surgical side effects, and higher patient satisfaction was noticed in WFG treated eyes, giving it a slight advantage over the WFO method. Larger sample sizes and more studies are required to show equivalency or superiority of WFG compared to WFO effectively. We recommend further, and more extensive studies are conducted with stricter follow up and more diverse follow-up checklists to determine the best guidelines for myopia PRK treatment.

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**Conflicts of interest**

There are no conflicts of interest.

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