Development of ocular hypertension secondary to tamponade with light versus heavy silicone oil: A systematic review

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Aim: The intraocular silicone oil (SO) tamponades used in the treatment of retinal detachment (RD) have been associated with a difference ocular hypertension (OH) rate. To clarify, if this complication was associated to use of standard SO (SSO) versus heavy SO (HSO), we performed a systematic review and meta-analysis of comparative study between two kind of SO (standard or light vs. heavy) for the treatment of RD and macular hole, without restriction to study design. Materials and Methods: The methodological quality of two randomized clinical trials (RCTs) were evaluated using the criteria given in the Cochrane Handbook for Systematic Reviews of Intervention, while three non-RCTs were assessed with the Newcastle–Ottawa Scale and Strengthening the Reporting of Observational Studies in Epidemiology checklists. We calculated Mantel–Haenszel risk ratio (RR) with 95% confidence intervals (95% CIs). The primary outcome was the rate of patients with OH treated with SSO compared to HSO. Results: There were a higher number of rates of OH in HSO compared to SSO. This difference was statistically significant with the fixed effect model (Mantel–Haenszel RR; 1.55; 95% CI, 1.06–2.28; P = 0.02) while there was not significative difference with the random effect model (Mantel–Haenszel RR; 1.51; 95% CI, 0.98–2.33; P = 0.06). Conclusion: We noted a trend that points out a higher OH rate in HSO group compared to SSO, but this finding, due to the small size and variable design of studies, needs to be confirmed in well-designed and large size RCTs.

Key words: Glaucoma, heavy silicone oil, light silicone oil, ocular hypertension, retinal detachment, standard silicone oil

The silicone oils (SOs) are intraocular tamponades diffusely used in the treatment of retinal detachment (RD) and macular hole.\[1-5\] The ocular hypertension (OH) has been also reported in literature after pars plana vitrectomy (PPV) and SO injection, with an incidence ranging from 3% to 40% of cases.\[14-16\] Since there is the need to support the retina in any quadrant nowadays, there are two kind of SOs available: (i) Standard SO (SSO), that has lower density than water, providing a good support for the superior retina; (ii) heavy SO (HSO), that has a heavier density than water, making it able to provide an effective postoperative tamponade of the inferior quadrants. HSO is a mixed compound obtained by adding SO to the semifluorurate (alkane or ether) in a miscible ratio. The presence of semifluorinate should make the compound more unstable and prone to induce emulsification and increase of intraocular pressure (IOP). Although using of HSO in case of inferior retinal breaks or macular hole, reduces the RD’s recurrence\[13\] and provides to high success rate of macular hole closure,\[12\] several case series report a different rate of increase of postoperative IOP, with an incidence ranging from 14% to 30.7% of cases.\[13-15\] The OH represents an important risk factor for both SSO and HSO, which can lead to the development of secondary glaucoma.\[16,17\] To clarify, if this complication was associated to the type of SO, several studies have been conducted,\[18-22\] nevertheless they have reduced statistical power and discordant results. Therefore, a systematic review was needed to clarify this issue.

Materials and Methods

We searched MEDLINE, EMBASE, Scopus and Google Scholar for studies that comparing the OH rate in patients treated with HSO with those treated with SSO for RD. Any study comparing HSO with SSO for RD, without restriction to study design or language, was included. The keywords used in the search were “SO”, “HSO” and “RD and SO”. We performed the final search on December 23, 2013.

Three authors (VR, FS, MRR) independently screened for potential relevance the titles and abstracts of studies to identify those that fulfilled the inclusion criteria: Patient treated with SO, vitrectomy surgery with tamponades, comparison on the complications of HSO versus SSO, and IOP.

Disagreements or doubts were resolved by discussion. We obtained in full, any study that potentially meets the inclusion criteria based on the title, abstract or both and assessed these studies against the inclusion criteria.

The following data were extracted independently by three reviewers (VR, FS, MRR): Administrative details (authors, year of publication), details of participants (number, setting, baseline characteristics by group), details of the study (study design and type), details of IOP after surgery, outcome descriptions and outcomes measures in each group with reasons.
The data abstracted for dichotomous variables are the number of eye with OH. When additional data were needed, we contacted the corresponding author of each study by E-mail in order to access further information.

Three authors (VR, FS, MRR) independently assessed studies fulfilling the review inclusion criteria for methodological quality. For randomized clinical trials (RCTs), we used the criteria given in the Cochrane Handbook for Systematic Reviews of Intervention.[23] The risk of bias was assessed in individual studies across six domains: Random sequence generation, allocation concealment, blinding of participants and personnel (performance bias), blinding of outcome assessors (detection bias), incomplete outcome data, selective outcome reporting. We categorized these judgments as “low risk”, “high risk”, or “unclear” risk of bias.

For observational studies, we used two commonly adopted checklists, the Newcastle–Ottawa Scale (NOS) and the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) checklists, from which two reading grids were adapted.[24-26] The NOS is a nine-point scale that assigns points on the basis of the process of selection of the cohorts or the cases and the controls (0–4 points), the comparability of the cohorts or the cases and the controls (0–2 points), and the identification of the exposure and the outcomes of study participants (0–3 points). The NOS was developed to assess the quality of nonrandomized studies for the purpose of incorporating quality assessments in the interpretation of meta-analytic results. This scale is recommended by the Cochrane Non-randomized Studies Methods Working Group available at the electronic address (http://www.ohri.ca/programs/clinical_epidemiology/oxford.htm). STROBE is an international, collaborative initiative of epidemiologists, methodologists, statisticians, researchers and editors involved in the conduct and dissemination of observational studies. The STROBE statement consists of a checklist of 22 items, which relate to the title, abstract, introduction, methods, results and discussion sections of articles. Eighteen items are common to cohort studies, case-control studies and cross-sectional studies and four are specific to each of the three study designs. The STROBE statement provides guidance to authors about how to improve the reporting of observational studies and facilitates critical appraisal and interpretation of studies by reviewers, journal editors and readers.

We assessed statistical heterogeneity using $\tau^2$, Cochran’s $Q$ and the $P$ statistic. The $P$ statistic describes the percentage of total variation across trials that are due to heterogeneity rather than sampling error.[27,28]

Dichotomous outcomes (e.g. rate of patients with OH) are presented as risk ratios (RRs) with 95% confidence intervals (CIs). The software used was Review Manager (RevMan, Version 5.3, The Nordic Cochrane Centre, The Cochrane Collaboration, Copenhagen, Denmark, 2014). In case of no heterogeneity ($P = 0$), studies were pooled using a fixed effect model. Where values of $P$ were $> 0$, a random effects analysis was undertaken.[23,29] In addition, both the random effects model and the fixed model will be used as a sensitivity analysis for evaluating the possible bias effects of smaller studies.

Where values of $P > 75\%$ indicated a very high level of heterogeneity, we refrained to pool data from different studies and we undertook a narrative overview.

The surgical techniques were generally consistent with minor variation in different studies.

Surgery was performed with monitored anesthesia care and a retrobulbar block. In both groups, the surgical procedure included a standard 3-port PPV. During vitrectomy, the vitreous base was thoroughly removed. Epiretinal traction removal, internal limiting membrane peeling and relaxing retinotomies were performed, when necessary. The retinal periphery was inspected for retinal breaks, and any peripheral break found was treated with cryocoagulation or endolaser photocoagulation. A fluid–air exchange procedure was then performed with humidified air. At the end of the surgical procedure, SSO or HSO was injected. The iridotomies were performed when necessary and located according to the density of the oil used. At the end of surgery, the surgeons aimed to reach a complete filling and the eyes appeared clinically completely filled by the compound. Suture of the sclerotomies followed. In all cases, the surgery was not combined with scleral buckle placement.

The outcome measure was the incidence of OH at 1-month after surgery, calculated as the number of patients with IOP $> 21$ mmHg with or without medical therapy in each study.

**Results**

Three thousand one hundred eighty-eight studies were identified through a literature search. The most of these studies were excluded because no-comparative studies, unqualified intervention and unqualified outcome measure. Only five studies comparing the OH rate after PPV for RD using SSD or HSO were included [Fig. 1]. The random effects model will be used and a fixed effect model will be considered as a

![Diagram](image_url)
sensitivity analysis for evaluating the possible bias effects of smaller studies.

**Study characteristics**

A total of 441 eyes in 5 included studies were included in this meta-analysis. The 50.3% of participants were female. Sample sizes in these studies ranged from 30 to 180 eyes. Three studies were carried out in Italy, and one study was carried out in Turkey and United Kingdom each. All of the studies included OH rate after PPV for RD [Table 1].

There were 2 RCTs and 3 retrospective case–control studies. In the RCT of Kocak and Koc,[19] there was no information on random sequence generation and outcome concealment. Both studies were open label, but the outcome measure probably was not affected by blinding status. The two studies were judged as low risk of bias for the remaining items.

Methodological quality of 3 retrospective case–control studies was evaluated using NOS and STROBE. The quality of the studies, according NOS, was high: All the studies met eight of the nine study quality criteria. The only unmet item was related to comparability criteria since the studies did not report on statistical methods used to adjust for confounding. According STROBE checklist, almost all of case–control studies reported the scientific background, described the setting and locations, included a clear description of the inclusion criteria and give a clear definition of outcomes. No one provided sufficient information on how the sample size and described measures to reduce potential sources of bias; likewise, all the reports did not describe the approach to dealing with missing data or to controlling for confounding factors. All the studies described the statistical methods, but an assessment of confounding (e.g., odds ratios, RRs) was not reported. The selected studies described the characteristics of participants in detail and reported numbers of outcome events.

The pooled data are presented as Mantel-Haenszel RR and 95% CI with a fixed effect model [Fig. 2a] and random effect model [Fig. 2b] for the OH rate. The heterogeneity test shows lack of heterogeneity ($\chi^2$, 4.41; df 4; $P = 0.35$); however, since $I^2$ were $>0 (=9$). Both the random effects analysis and the fixed effect analysis are presented. Moreover, the fixed effect model is considered as a sensitivity analysis for evaluating the possible bias effects of smaller studies. There were a higher number of rates of OH in HSO compared to SSO. The authors did not observe any pupil block mechanisms in these series.[18] Kocak and Koc observed OH in SSO group in 7 cases (23%) in the early postoperative period, in 1 case (3%) by the 1st month, whereas in HSO group the OH was present in 7 eyes (22%) in the early postoperative period, in 4 eyes (13%) by the 1st month.[19] Mete et al. reported that 2 eyes (11%), treated with SSO, developed OH that was controlled with topical beta-blocker treatment. In the group treated with HSO, 5 eyes (29%) developed OH that in one case was refractory to topical treatment and required further ciliary body cyclo-photoagulation. No statistically significant differences were found between the two groups ($P = 0.238$; Fisher’s exact test).[20] Romano et al. stated that the long-term increased IOP depends on an open-angle mechanism rather than a closed-angle mechanism, since they did not report any case of pupillary block associated with SSO or HSO.[21] They stated that the persistence of a raised pressure could be related to the presence of emulsified oil in the anterior chamber and in the trabecular meshwork, which persists after the removal of SO.[21]

Wong et al. showed that the IOP is higher in eyes treated with HSO at day 1 and between 7 and 14 days postoperatively ($P = 0.05$ and 0.01, respectively), whereas by the 4th week, the difference of IOP between the two groups was not significant ($P = 0.17$). At 4 weeks, IOP higher than 30 mmHg was still present in 9 eyes (12.7%) in HSO group and in 1 eye (1.8%) in the SSO group. However, according to the authors, the raised IOP in HSO group was more difficult to treat.[19]

The OH due to an intraoperative overfilling is still a risk factor, but easily managed by the surgeon.[30]

Our meta-analysis considered only cases of OH in open-angle, frequent complication following PPV and SO

| First author, country, year publication | Study design | Patients | Number of patients enrolled in the arms (SSO/HSO) | Patient with OH (SSO vs. HSO) |
|----------------------------------------|-------------|----------|-----------------------------------------------|-------------------------------|
| Avitabile, Italy, 2011                 | Randomized control study | RD with macular hole and posterior staphyloma in highly myopic eyes | 15/15                         | 2/15 versus 4/15 |
| Kocak, Turkey, 2013                    | Randomized control study | RD in patients with inferior breaks | 30/31                         | 1/30 versus 4/31 |
| Mete, Italy, 2011                      | Retrospective case-control | RD with myopic macular hole | 17/25                         | 5/17 versus 4/25 |
| Romano, Italy, 2010                    | Retrospective case-control | RD | 105/75                                      | 12/105 versus 12/75 |
| Wong, UK, 2009                         | Retrospective case-control | RD | 57/71                                        | 13/57 versus 30/71 |

RD: Retinal detachment, SSO: Standard silicone oil, HSO: Heavy silicone oil, OH: Ocular hypertension
injection, ranking second to cataract as a late complication of SO. At least in theory, the use of HSO, mainly for the presence of semifluorinated alkanes, has been considered to be higher risk factor than SSO for postoperative OH. The semifluorinated alkanes in the beginning have been only used as intraoperative tamponades to keep the retina stable during the surgery and as silicone solvent to remove SO remnants from vitreous cavity. The main problem associated with semifluorinated alkanes was the presence of emulsification, as for perfluocarbon, liquid in up to 100% in the postoperative of all cases. The emulsification was thought to be responsible for inflammatory reaction due to foreign body response to the emulsified droplets. The stability of the semifluorinated alkanes is significantly improved adding SO to the semifluorinated, and keeping at same time a density higher than water. The mixed compound generated, HSO, is currently used as postoperative intraocular tamponades, however, the current literature still report OH in the postoperative period. The results of this meta-analysis, according to the random model, lack of statistical significance. Nevertheless, it was possible to observe a trend for lower OH rate in SSO group [Fig. 2b]. This tendency is confirmed by the results of the fixed model showing a statistically significant difference in OH rate between the SSO and HSO groups.

Other risk factors associated with an increased OH rate after PPV, rather than the presence of semifluorinated alkanes in the compound, are the lens status, angle pathology, preexisting glaucoma, steroid response, axial myopia, overfilling, migration of SO into the anterior chamber, surgical procedures commonly used in conjunction with PPV, systemic diseases. The surgeon should also consider, in the planning of the vitreoretinal surgery, that several studies confirmed the strong relationship between high myopia and glaucoma. In noncomparative studies a significant association of diabetes mellitus with OH after SO injection was found. Other authors reported that OH rate was independent from systemic conditions such as diabetes mellitus. In our comparative studies no one of authors found any systemic association with OH rate.

Nowadays the clinical management of eyes with OH after PPV and SO injection is medical therapy.

Jackson et al. observed that when SO was present in anterior chamber, high IOP usually ensued and the OH could be difficult to treat. In the comparative studies the most of patients are treated with medical therapy only 1.5% (7/441 eyes) required to glaucoma surgery. Since the trabeculectomy has a limited role and success rate, besides being technically difficult, the surgical therapy with Ahmed glaucoma valve implantation with inferior placement have gained wide acceptance in recent years. Previous studies have shown that Ahmed glaucoma valves have a low complication rate, especially, if compared with other glaucoma implants. Furthermore in the 1991 was proposed a new surgical technique for the pars plana insertion of a glaucoma implant, instead of anterior chamber, for the
management of neovascular glaucoma.[30] This modification can potentially cause significant anterior segment complications. Recently Maris et al.[31] in a retrospective study did not report differences in IOP and complication after implantation of the Ahmed glaucoma valve in posterior or in the anterior chamber. There are several important limitations to this meta-analysis, including the lack of randomized control trials, the inclusion of retrospective studies, with all the limits that they have, variety of surgeons, techniques and instruments used. In summary further well-designed and large size RCTs are necessary to confirm our results.

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