Short-Term Use of Dexamethasone/Netilmicin Fixed Combination in Controlling Ocular Inflammation After Uncomplicated Cataract Surgery

Aldo Caporossi¹
Giovanni Alessio¹
Francesco Fasce³
Giorgio Marchini⁴
Antonio Rapisarda⁵
Vincenzo Papa⁶

¹Department of Ophthalmology, Policlinico Universitario A. Gemelli IRCCS Università Cattolica del Sacro Cuore, Roma, Italy; ²Ophthalmology Unit, Dipartimento di Scienze mediche di base, Neuroscienze e Organi di Senso Università di Bari, Azienda Ospedaliera Policlinico Consorziale, Bari, Italy; ³Ophthalmology Unit, IRCCS San Raffaele, Milano, Italy; ⁴Ophthalmology Unit, Ospedale Policlinico G.B. Rossi, Università di Verona, Verona, Italy; ⁵Ophthalmology Unit, Azienda Ospedaliera Garibaldi, Catania, Italy; ⁶Medical Affairs SIFI SpA, Catania, Italy

Purpose: To evaluate the short-term anti-inflammatory effect of dexamethasone/netilmicin fixed combination in the management of ocular inflammation after cataract surgery.

Patients and Methods: Open-label, randomized, active-controlled, clinical study conducted in 6 sites in Italy; 238 patients were randomized 2:1 to dexamethasone/netilmicin (dexa/net, n=158) or betamethasone/chloramphenicol (beta/chl, n=80). Treatment started the day of surgery and continued 4 times daily for 7 days. The primary efficacy parameter was the anterior chamber (AC) flare. The percentage of patients displaying none or mild (ie, only barely detectable) AC flare was defined as “efficacy rate”, whereas the percentage of patients showing a decrease of AC flare score from baseline was defined as “percentage of responders”. Additional parameters evaluated were AC cells, conjunctival hyperaemia, corneal and lid oedema, symptoms of ocular discomfort, visual acuity, and intraocular pressure. Dexa/net was considered effective if the efficacy rate was not inferior (by means of 97.5% confidence interval) to that of beta/chl.

Results: After 7 days of treatment, no AC flare was observed in 92.8% (dexa/net) and 92.3% (beta/chl) of patients, whereas no AC cells were observed in 91.5% (dexa/net) and 93.6% (beta/chl) of patients, respectively. The “efficacy rate” was 100% in both groups, whereas the “percentage of responders” was 94.1% in the dexa/net and 93.6% in the beta/chl group. The p-value to reject the null hypothesis of inferiority was <0.001. Other efficacy parameters confirmed both treatments as highly effective, despite their difference in steroid content (2 mg/mL for beta/chl vs 1 mg/mL for dexa/net). IOP and visual acuity at the end of the study were comparable. Two cases of allergic conjunctivitis were considered adverse events and were both related to dexa/net.

Conclusion: Short-term use of dexa/net fixed combination is safe and effective in the control of post-operative inflammation following uncomplicated cataract surgery.

Keywords: cataract surgery, fixed combination, netilmicin, chloramphenicol

Introduction
Cataract surgery the most common surgical medical procedure performed annually across the European Union states in approximately 5 million individuals.¹ After cataract extraction, a varying degree of post-surgical inflammation may occur. Although such inflammation is, in most cases, self-limited, the use of anti-inflammatory agents can rapidly resolve it and prevents serious complications.
resulting from uncontrolled inflammations (as cystoid macular edema).\textsuperscript{2} Accordingly, the use of topical corticosteroids and/or non-steroidal anti-inflammatory drugs remains the mainstay of post-surgical management of these patients. A recent study funded by the European Society of Cataract and Refractive Surgeons (ESCRS) showed that a combination of a non-steroidal anti-inflammatory drugs and dexamethasone reduces the risk for developing cystoid macular oedema after cataract surgery.\textsuperscript{3} Apart ocular inflammation, endophthalmitis is the most important complication of cataract surgery. Its severity depends on the virulence and the quantity of inoculated pathogens. \textit{Coagulase-negative Staphylococci} and \textit{S. Aureus} are the most common pathogens isolated in this infection.\textsuperscript{4} Moreover, during the past decade a major concern was related to the high prevalence of infections due to methicillin-resistant \textit{S. Aureus} (MRSA) and methicillin-resistant \textit{Coagulase-negative Staphylococci} (MRCoNS). A preoperative antisepsis of the periocular area with topical povidone–iodine and the use of intracameral cefuroxime are actually considered the standard for endophthalmitis prophylaxis.\textsuperscript{5,6} However, cefuroxime is not effective versus methicillin-resistant strains.\textsuperscript{8} The use of post-operative topical antibiotics is controversial, but they are frequently used in the real life to sterilize the ocular surface and prevent any access of microorganisms inside the eye.\textsuperscript{5,7} Topical antibiotics can be administered either alone or in combination with corticosteroids. Steroid/antibiotic fixed combinations have several advantages over the use of single components, as better compliance, lower costs, and reduction of the potential wash-out effect.\textsuperscript{8} The choice among different products depends on the bacterial susceptibility to the antibiotic and the type and strength of the steroid included in the formulation. A steroid/antibiotic fixed combination containing 1 mg/mL dexamethasone and 3 mg/mL netilmicin is available in several EU and non-EU countries under the trade name of Netildex (SIFI SpA, Italy) and exhibits a fast and effective control of ocular inflammation, either post-operative or not.\textsuperscript{8-10} Since the use of dexamethasone after cataract surgery is established,\textsuperscript{11} the product’s added value is the presence of netilmicin, a wide spectrum antibiotic also covering methicillin-resistant strains ensuring a complete sterilization of the ocular surface in the immediate post-surgery period.\textsuperscript{12-14} In this study, a short treatment with such steroid-antibiotic combination was tested in patients who underwent to uncomplicated cataract surgery.

**Patients and Methods**

**Trial Design**

This was a multicenter, open, randomized, active-controlled, parallel group, Phase IV clinical study conducted in 6 sites in Italy at Bari, Milano Verona, Catania, and Rome (2 centres). The main objective of the study was to evaluate the effectiveness of 1 mg/mL dexamethasone plus 3 mg/mL netilmicin (dexa/net) ophthalmic solution treatment for 7 days in the control and treatment of post-surgical ocular inflammation following cataract surgery. A control group of patients was treated with 2 mg/mL betamethasone plus 5 mg/mL chloramphenicol (beta/chl) eye suspension. The study protocol was approved by the institutional review board at each research center. All participants provided written informed consent and the study was conducted according to the Declaration of Helsinki and Good Clinical Practice guidelines. The trial is registered at the European Union Clinical Trials Register (EudraCT) no. 200600330513.

**Participants**

Any subject older than 40 years, able to give an informed consent and scheduled for having micro-incisional cataract surgery was eligible for the study. Subjects were considered not eligible if they had: 1) history of ocular inflammatory diseases, ocular herpes infection, iritis, uveitis or Sjogren’s syndrome, 2) concomitant ocular pathologies, 3) intraocular pressure >24 mmHg; 4) previous ocular surgery in the affected eye in the previous 12 months; 5) previous laser treatment in the affected eye in the previous 6 months; 5) use of any ocular medication within 14 days prior to study entry; 6) any ocular infection within month prior to study entry; 7) known or suspected allergy to ophthalmic preservatives, aminoglycosides and steroids. Two hundred thirty-eight consecutive patients (116 males and 122 females) were screened and enrolled. Mean age was 72 years (range 42–90 years). Age and gender were well-matched among groups of treatment. Patients disposition, according to the CONSORT diagram, is displayed in Figure 1.

**Interventions**

Two hundred and thirty-eight patients were randomized, using a computer-generated list, in a 2:1 ratio to dexa/net (n=158) or beta/chl (n=80). The different packaging of two products did not allow a double-masked approach. Treatment started the day of surgery (day 0) administering...
1 drop in the conjunctival sac of the operated eye and continued 4 times a day for 7±1 day. Afterwards investigators could continue the anti-inflammatory treatment according to patient conditions and their clinical routine. Control visits were scheduled at day 1 and day 7 (±1) after surgery.

Outcomes
The examination at each visit included best corrected visual acuity, slit-lamp examination, ophthalmoscopy and applanation tonometry. The primary efficacy parameter evaluated was the anterior chamber (AC) flare scored from none to severe using a 0 to 3-point scale (Table 1). Other

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### Table 1 Scoring Systems Used in the Study

| Score | Anterior Chamber Flare       | Anterior Chamber Cells | Conjunctival Hyperemia              | Lid and Corneal Edema | Symptoms (Burning/Tearing/Pain) |
|-------|------------------------------|------------------------|-------------------------------------|-----------------------|---------------------------------|
| 0     | none                         | none                   | none                                | none                  | none                            |
| 1     | mild (barely detectable)     | mild (1 to 10 cells)   | mild (some vessel injected)         | mild                  | mild (present but not distressing) |
| 2     | moderate (iris and lens detail clear) | moderate (11 to 50 cells) | moderate (diffuse injection, individual vessels discernible) | moderate | moderate (distressing but not interfering with daily life) |
| 3     | severe (iris and lens details not visible and fibrin in the anterior chamber) | severe (> 50 cells) | severe (intense injection, individual vessels not easily discernible) | severe | severe (very distressing and interfering with daily life) |

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parameters of analysed included: AC cells, conjunctival hyperaemia, corneal and lid oedema, and symptoms of ocular discomfort (pain, burning, and tearing). All these clinical variables were also graded from none to severe using a 0 to 3-point scale (Table 1).\textsuperscript{15} Safety variables monitored during the study were adverse events, intraocular pressure (IOP), and visual acuity.

**Statistics**

Statistical analysis was performed by an expert statistician using the SAS system version 9.2. The sample size estimate was based on a one-sided test of equality of paired proportions of the primary end point. This was estimated to be 96% in the control group treated with dexamethasone for 10 days. Considering that in this trial treatment duration was 7±1 day, the estimated percentage of patients with AC flares was reduced to 90%. Based on this assumption, 219 evaluable patients were required to detect at 15% difference between treatments to have a 90% chance (delta = 10%) of statistically proving efficacy, in a non-inferiority design study. To allow for about 10% subjects drop-out, a final sample size of 238 patients was planned. Statistical evaluation was performed as previously described for similar studies.\textsuperscript{8,15} Listing, tables, graphs, and statistical output were generated using the SAS software. Briefly, the main efficacy parameter evaluated was AC flare at day 7 after surgery. The “efficacy rate” (primary efficacy parameter) was defined as the percentage of patients displaying none or mild AC flare (ie, score 0 or 1), whereas the “percentage of responders” (secondary efficacy parameter) was defined as the percentage of patients showing a decrease of AC flare score from baseline and patients scoring 0 at both baseline and at Day 7 after surgery. Dexa/net would be declared effective, if it can be shown that the response rate is not inferior (by means of 97.5% confidence interval) to that of beta/chl. Statistical analyses were performed on predefined subsets. The Full Analysis Population (FA) consisted of subjects operated and treated for at least 5 days and that reached the end point visit (Day 7 after surgery). The Per-Protocol (PP) Population consisted of all subjects of the FA population without any relevant protocol deviations. The main efficacy parameters (“efficacy rate” and “percent of responders”) were analysed in both FA and PP populations, whereas all other parameters were analysed in the FA population only. Safety was analysed in all subjects who received at least one dose of drug [Intention-to-Treat (ITT) population]. For other efficacy parameters, intraindividual score differences were calculated, assuming a zero difference if the respective post-dose values were not available. These score differences were compared between treatments by the Mann–Whitney U-test or the Wilcoxon test for independent samples. All secondary efficacy parameters were presented by descriptive analysis. All safety results are presented with their descriptive analyses only. Adverse events (AEs) were coded using the MedDRA dictionary (version 14).

**Results**

As shown in Figure 1, the ITT population consisted of 238 patients (dexa/net=158 and beta/chl=80). Seven patients were excluded due to a protocol violation and therefore the FA population consisted of 231 patients (dexa/net=153 and beta/chl=78); the PP population included 190 patients fully adherent to the protocol (dexa/net=126 and beta/chl=64). Patients were well-matched among groups regarding baseline characteristics.

Main efficacy results are displayed in Tables 2 and 3. After 7 days of treatment most of patients had no signs of inflammation in the anterior chamber. The main efficacy parameters (“efficacy rate” and “percentage of responders”) were comparable between groups. The p-value to reject the null hypothesis of inferiority was p<0.001 (Table 2). Specifically, at the end point visit no AC flare

**Table 2 Primary Efficacy Outcomes**

|                         | Dexa/Net | Beta/Chl | P values (Rejection of the Null Hypothesis) | One Side 97.5% CI |
|-------------------------|----------|----------|-------------------------------------------|------------------|
| **Efficacy rate**       |          |          |                                           |                  |
| FA                      | 100%     | 100%     | /                                         | /                |
| PP                      | 100%     | 100%     | /                                         | /                |
| **Responders**          |          |          |                                           |                  |
| FA                      | 94.1%    | 93.6%    | <0.001                                    | −0.061 +∞        |
| PP                      | 92.9%    | 92.2%    | 0.0002                                    | −0.073 +∞        |

Notes: Efficacy rate= percentage of patients displaying at the study endpoint (day 7 after surgery) none or mild anterior chamber flare (ie, score 0–1). Responders= percentage of patients showing at the study endpoint (day 7 after surgery) a decrease of anterior chamber flare score from baseline or a score 0 at both visits (baseline and day 7). FA=full analysis population (dexa/net: n=153; beta/chl: n=78). PP= per-protocol population (dexa/net: n=126; beta/chl: n=64)
was observed in 92.8% of patients treated with dexa/net and 92.3% of patients treated with beta/chl. In addition, no AC cell was detected in 92.8% of patients treated with dexa/net and 92.3% of patients treated with beta/chl (Table 3). Furthermore, all additional efficacy parameters evaluated (conjunctival hyperaemia, corneal oedema, and lid oedema) confirmed that both treatments were equally effective in reducing ocular inflammation (Table 3). Same conclusion can be draw also for ocular symptoms (tearing, burning and pain) (Table 4).

During the study, non-serious AEs occurred in 2.1% of patients (5/238). These AEs occurred in the group treated with dexa/net. Three events (subconjunctival hemorrhage, retinal hemorrhage, and capsular rupture) were considered not related to treatment but to the surgical procedure, whereas two cases of allergic conjunctivitis were considered related to the use of dexa/net. Final IOP was 14.3±2.3 mmHg (mean ±SD) in the group treated with dexa/net and 14.1±2.3 mmHg in the group treated with beta/chl. Final visual acuity was 8.7 ±2.0 decimals (mean ±SD) in the dexa/net group vs 8.9 ±1.7 in the beta/chl group.

Discussion
Cataract surgery is the most common surgical intervention. Despite actual techniques have reduced considerably signs

**Table 3 Additional Efficacy Outcomes**

|                                      | Day 1 |        | Day 7 |        | p-value Difference Between Groups (Wilcoxon Rank-Sum Test) | p-value Difference Within Group (Pratt-Wilcoxon Test) |
|--------------------------------------|-------|--------|-------|--------|-----------------------------------------------------------|------------------------------------------------------|
|                                      | Dexta/Net | Beta/Chl | Dexta/Net | Beta/Chl | Dexta/Net | Beta/Chl |
|                                      | N (% )     | N (% )     | N (% )     | N (% )     |                                                      |                                                      |
| **Full Analysis Population, N**      | 153 | 78 | 153 | 78 | 0.385 | <0.0001 |
| **Anterior chamber flare score**     | 0=None | 53 (34.7%) | 34 (43.6%) | 142 (92.8%) | 72 (92.3%) | 0.941 | <0.0001 | 0.0499 |
|                                      | 1=Mild   | 92 (60.1%) | 38 (48.7%) | 11 (7.2%) | 6 (7.7%) |
|                                      | 2=Moderate | 8 (5.2%) | 6 (7.7%) | 0 | 0 |
|                                      | 3=Severe | 0 | 0 | 0 | 0 |
| **Anterior chamber cells score**     | 0=None,  | 85 (55.6%) | 47 (60.3%) | 140 (91.5%) | 73 (93.5%) | 0.4594 | <0.0001 | <0.0001 |
|                                      | 1=Mild   | 57 (37.2%) | 22 (28.2%) | 13 (8.5%) | 5 (6.5%) |
|                                      | 2=Moderate | 11 (7.2%) | 9 (11.5%) | 0 | 0 |
|                                      | 3=Severe | 0 | 0 | 0 | 0 |
| **Conjunctival hyperaemia**          | 0=None | 75 (49.0%) | 38 (48.8%) | 126 (83.3%) | 70 (89.7%) | 0.128 | <0.0001 | 0.0499 |
|                                      | 1=Mild   | 69 (45.1%) | 36 (46.1%) | 22 (14.4%) | 8 (10.3%) |
|                                      | 2=Moderate | 9 (5.9%) | 4 (5.1%) | 4 (2.6%) | 0 |
|                                      | 3=Severe | 0 | 0 | 1 (0.6%) | 0 |
| **Lid oedema**                       | 0=None | 125 (81.7) | 70 (89.7) | 150 (98.0) | 76 (97.4) | 0.852 | <0.0001 | 0.0499 |
|                                      | 1=Mild   | 28 (18.3) | 7 (8.9) | 2 (1.3) | 2 (2.5) |
|                                      | 2=Moderate | 0 | 1 (1.2) | 1 (0.6) | 0 |
|                                      | 3=Severe | 0 | 0 | 0 | 0 |
| **Corneal oedema**                   | 0=None | 60 (39.2%) | 31 (39.7%) | 148 (96.7%) | 75 (96.1%) | 0.082 | <0.0001 | 0.0499 |
|                                      | 1=Mild   | 70 (45.7%) | 32 (41.1%) | 5 (3.3%) | 3 (3.9%) |
|                                      | 2=Moderate | 20 (13.1%) | 14 (17.9%) | 0 | 0 |
|                                      | 3=Severe | 3 (2.0%) | 1 (1.3%) | 0 | 0 |
of post-surgical inflammation and the risk of endophthalmitis, patients are often treated in the post-operative period with topical steroids and antibiotics (frequently in a fixed combination) as well as with non-steroidal anti-inflammatory drugs.2–7

The aim of this study was to evaluate the effectiveness of a short treatment with a steroid/antibiotic fixed combination containing 1 mg/mL dexamethasone and 3 mg/mL netilmicin in the control of post-operative inflammation. Results have shown that after 7 days more than 90% of patients have no sign of inflammation in the anterior chamber (AC flare and AC cells = score 0) and that the others have only a negligible level of inflammation (score 1). Accordingly, also signs of ocular surface inflammation were absent (score 0) at the study endpoint (7 days) in most patients (82% for conjunctival hyperaemia, 97% for corneal oedema and 98% lid oedema). The control treatment used in the study was another steroid-antibiotic fixed combination containing 2 mg/mL betamethasone and 5 mg/mL chloramphenicol. It is interesting to note that both treatments had a comparable anti-inflammatory efficacy rate despite the double dose of steroid present in beta/chl. This finding suggests, therefore, that a short-term treatment with 1 mg/mL dexamethasone is appropriate to manage post-surgical inflammation and that a stronger and extended exposure to steroids is not necessary. This ensures an effective control of post-operative inflammation and, at the same time, reduces the risk of developing corticosteroid-related side effects and avoids an overuse of antibiotics.

In our opinion, the main benefit of dexa/net over other available steroid-antibiotic fixed combinations is related to the antibiotic component. Indeed, netilmicin has a wide-spectrum activity (which also includes methicillin-resistant strains)12–14 and a negligible toxicity for the ocular surface.16 Since post-surgical infections are due to microorganisms resident in the ocular surface, mainly Coagulase-negative Staphylococci (in particular S. epidermidis) and S. Aureus,17 it is critical to use antibiotics highly effective on these germs. If not eradicated, these bacteria may enter inside the eye starting the day of surgical procedure until corneal incision is fully closed. For this reason, antibiotic treatment, when used, should starts the same day of surgery. Netilmicin is able to sterilize lid margins and conjunctiva when given before cataract surgery.14,18 Moreover, the bacterial flora on the ocular surface isolated from patients

| Table 4 Ocular Symptoms |
|-------------------------|
| Day 1 | Day 7 | p-value Difference Between Groups (Wilcoxon Rank-Sum Test) | p-value Difference Within Group (Pratt-Wilcoxon Test) |
|      |        |                          |                          |
|      | Dexa/Net | Beta/Chl | Dexa/Net | Beta/Chl | Dexa/net | Beta/chl |
| Full Analysis Population, N | 153 | 78 | 153 | 78 | 0.402 | 0.0030 | 0.0029 |
| Tearing |        |        |        |        |        |        |        |
| 0=None | 119 (77.8%) | 55 (70.5%) | 135 (88.2%) | 68 (87.1%) | 0.402 | 0.0030 | 0.0029 |
| 1=Mild | 32 (20.9%) | 22 (28.2%) | 16 (10.5%) | 10 (12.8%) |
| 2=Moderate | 2 (1.3%) | 1 (1.3%) | 2 (1.3%) | 0 |
| 3=Severe | 0 | 0 | 0 | 0 |
| Burning |        |        |        |        |        |        |        |
| 0=None, | 117 (74.0%) | 64 (82.1%) | 140 (91.5%) | 73 (93.5%) | 0.652 | <0.0001 | 0.0030 |
| 1=Mild | 38 (24.1%) | 15 (19.2%) | 13 (8.5%) | 5 (6.5%) |
| 2=Moderate | 3 (1.9%) | 1 (1.3%) | 0 | 0 |
| 3=Severe | 0 | 0 | 0 | 0 |
| Pain |        |        |        |        |        |        |        |
| 0=None | 137 (89.5%) | 63 (80.8%) | 145 (94.8%) | 77 (98.7%) | 0.032 | 0.070 | 0.0007 |
| 1=Mild | 16 (10.5%) | 13 (16.7%) | 7 (4.6%) | 1 (1.3%) |
| 2=Moderate | 0 | 2 (2.5%) | 0 | 0 |
| 3=Severe | 0 | 0 | 1 (0.6%) | 0 |
undergoing cataract surgery is highly susceptible to netilmicin and much less to other antibiotics.\textsuperscript{14} It is important to highlight that, regardless their source, \textit{S. Aureus} and \textit{Coagulase-negative Staphylococci} isolates are often characterized by methicillin resistance (MR) and multiresistance (MDR) to several classes of antibiotics; this finding is even more common in older patients, as those operated for cataract.\textsuperscript{19} Netilmicin is effective on methicillin-resistant \textit{S. Aureus} (MRSA), methicillin-resistant \textit{Coagulase-negative Staphylococci} (MRCoNS) and MDR \textit{Coagulase-negative Staphylococci},\textsuperscript{12–14} whereas these microorganisms display a high resistance to fluoroquinolones.\textsuperscript{12,13,19}

As a potential limitation of the study, AC inflammation was measured, for practical reasons, by slit-lamp examination rather than by a laser flare and cell meter. Even if the scoring system used to measure flare and cells by slit-lamp examination is subjective and semi-quantitative, yet it corresponds to the actual daily routine of practice. Moreover, a consistency of results obtained with bio-microscopy and laser flare and cells meter measurements has been described.\textsuperscript{20} In summary, a short-term use of dexa/net fixed combination is effective as beta/chl in the control of post-operative inflammation following uncomplicated cataract surgery.

Data Sharing Statement

The raw data of the study used may be released upon request to the corresponding author.

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Author Contributions

All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal: gave final approval of the version to be published; and agree to be accountable for all aspects of the work.

Disclosure

VP is an employee of SIFI SpA (Italy). AC and AR are retirees. All remaining authors have no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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