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#### LETTER

# Toxic anterior segment syndrome and intracameral injection of cefuroxime axetil

Susanne Gardner<sup>1</sup> Peter Barry<sup>2</sup> Luis Cordovés<sup>3</sup>

<sup>1</sup>European Society of Cataract and Refractive Surgery, Dublin, Ireland; <sup>2</sup>Ophthalmic Surgery, St Vincents University Hospitals Group, Dublin, Ireland; <sup>3</sup>Retina and Vitreous Section, Ophthalmology Service, Hospital Universitario de Canarias, Spain

Correspondence: Susanne Gardner 1266 W Paces Ferry Road, Atlanta, GA 30327, USA Email susanneg@bellsouth.net

http://dx.doi.org/10.2147/OPTH.S91943

#### Dear editor

We read with concern the findings of toxic anterior segment syndrome in patients receiving intracameral injection of cefuroxime axetil, as described in the report by Çakir et al<sup>1</sup> in the March 2015 issue of *Clinical Ophthalmology*. Notably, the authors state that the form of cefuroxime used to extemporaneously compound cefuroxime for intracameral injection was cefuroxime axetil and that 17 cases of toxic anterior segment syndrome ensued. With over one million doses of intracameral cefuroxime administered to date, and with extensive clinical experience,<sup>2</sup> the unfortunate scenario described by the authors has not been realized elsewhere.

Based on information offered in the report, the root of the problem may lie in the inadvertent, or inappropriate, use of the axetil form of the antibiotic. Use of cefuroxime axetil does not conform with the guidelines published by the European Society of Cataract and Refractive Surgery,<sup>3</sup> and does not reflect the form of cefuroxime used in Aprokam<sup>®</sup>, the approved injectable form of cefuroxime that has recently become commercially available in Europe and in Turkey.

The axetil form of cefuroxime could be obtained from various sources, but is a form of cefuroxime not intended for injection. The axetil form of cefuroxime is used in, for example, tablets or oral suspensions where the drug particles do not remain in solution, but in a solid form or suspension. Cefuroxime axetil is an ethyl ester of cefuroxime that requires hydrolysis by esterases in the body to yield free cefuroxime. The axetil moiety of cefuroxime axetil is then metabolized to acetic acid and acetaldehyde.

Aside from the improper use of the axetil form of cefuroxime, and any effects from metabolic byproducts in the eye, concerns also surround the extemporaneous compounding procedure itself. One assumes the axetil form would have had to be placed into a liquid vehicle, and sterilization attempted prior to injecting into the anterior chamber of the eye. This procedure poses challenges in terms of accurate dose determination from a suspension and also sterilization of the suspension, both of which would likely interfere with accurate assessment of any dose actually delivered.

Once inside the eye, both the particulate matter of the axetil form and the metabolic acetic acid and acetaldehyde byproducts of axetil could have been noxious to the eye, in addition to uncertainties surrounding the delivered dose and sterility. Each of these factors, excluding the cefuroxime base, would themselves likely produce toxic anterior segment syndrome after intracameral injection. We hope our communication to the corresponding author is received and that such practices are discouraged in the region.

This report again underscores the hazards of extemporaneous compounding of products for injection into the eye. In answer to the clinical demand for a commercially

Clinical Ophthalmology 2015:9 1865-1867

© 2015 Gardner et al. This work is published by Dove Medical Press Limited, and Licensed under Creative Commons Attribution — Non Commercial (unported, v3.0) permission from Dove Medical Press Limited, provided the work is properly attributed. Permissions beyond the scope of the License are administered by Dove Medical Press Limited. Information on how to request permission may be found at http://www.dovergersc.om/permissions.php available cefuroxime product, Aprokam was approved by the European Medicines Agency, with an indication for intracameral injection as prophylaxis for endophthalmitis after cataract surgery.

Aprokam does not use cefuroxime axetil, but does use the proper base for formulation into an injectable product. We thank the authors for bringing attention to the potential hazards of improper extemporaneous compounding as described in this publication.

## Disclosure

The authors report no conflicts of interest in this communication.

### References

- 1. Çakir B, Celik E, Aksoy NÖ, et al. Toxic anterior segment syndrome after uncomplicated cataract surgery possibly associated with intracameral use of cefuroxime. *Clin Ophthalmol.* 2015;17:493–497.
- Behndig A, Montan P, Stenevi U, Kugelberg M, Lundström M. One million cataract surgeries: Swedish Cataract Registry 1992–2009. J Cataract Refract Surg. 2011;37:1539–1545.
- Barry P, Cordovés L, Gardner S. ESCRS guidelines for prevention and treatment of endophthalmitis following cataract surgery: data, dilemmas and conclusions, 2013. Available from: www.escrs.org. Accessed June 14, 2015.

## **Authors' reply**

Burçin Çakır Erkan Celik Nilgün Özkan Aksoy Özlem Bursalı Turgay Uçak Erdinç Bozkurt Gursoy Alagoz

Sakarya University Education and Research Hospital, Sakarya, Turkey

Correspondence: Burçin Çakır Sakarya University Education and Research Hospital, Adnan Menderes Street, Sakarya 54000, Turkey Tel +90 532 660 5657 Email b\_koklu@yahoo.com

## **Dear editor**

We are grateful to Gardner et al for their attention and interest in our paper. In our study, we investigated possible causes of an outbreak of toxic anterior segment syndrome step by step and finally we focused on the use of cefuroxime axetil at the end of surgery. This drug was not produced for intracameral use, and dilution problems have been reported previously.<sup>1,2</sup> While we had our suspicions about the agent and changed from cefuroxime axetil to moxifloxacin, Aprokam<sup>®</sup> was not commercially available in our country.

Despite the European Society of Cataract and Refractive Surgery (ESCRS) recommendations, use of intracameral cefuroxime has not been fully adopted. An ESCRS survey of member ophthalmic surgeons conducted in 2012 reported that the most common reasons for not using intracameral cefuroxime during cataract surgery were: the unavailability of a country/clinic-specific protocol, unavailability of an approved preparation, and concern over the risk of dilution errors.<sup>3</sup> These major concerns are directly addressed by Aprokam, which, as per the UK National Health Service National Patient Safety Agency injectable risk assessment proforma, is likely to be classified as a lower-risk product. Aprokam is currently the only product licensed for prophylaxis of postoperative endophthalmitis, and received approval from the European Medicines Agency in 2012.

In summary, we agree with the concerns of Gardner et al about the extemporaneous compounding of products delivered into the eye, and we recommend use of approved commercial and single-unit products to avoid the risk of dilution errors and contamination.

## Disclosure

The authors report no conflicts of interest in this communication.

### References

- Lockington D, Flowers H, Young D, Yorston D. Assessing the accuracy of intracameral antibiotic preparation for use in cataract surgery. *J Cataract Refract Surg.* 2010;36(2):286–289.
- Yoeruek E, Spitzer MS, Saygili O, et al. Comparison of in vitro safety profiles of vancomycin and cefuroxime on human corneal endothelial cells for intracameral use. J Cataract Refract Surg. 2008;34(12):2139–2145.
- 3. Barry P. Adoption of intracameral antibiotic prophylaxis of endophthalmitis following cataract surgery: update on the ESCRS Endophthalmitis Study. *J Cataract Refract Surg.* 2014;40:138–142.

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