

Rayner Intraocular Lenses Limited

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www.rayner.com

01 January 2016

Ref: 2015-02

To whom it may concern,

Field Safety Notice (FSN)

Healthcare professionals and stock control managers are advised that Rayner Intraocular Lenses Limited is voluntarily recalling certain Hydrophilic Acrylic Single Use Intraocular Lens system packs that are in distribution.

Internal quality checks have revealed that certain products released to market may contain a higher than usual level of residual polishing compound (aluminium oxide) that is used in the manufacturing process of intraocular lenses (IOLs).

Rayner maintains extremely high levels of product quality and has set stringent internal limits for residual aluminium oxide levels. There is a possibility that certain products released to market may contain levels that slightly exceed these limits.

As a precaution, and because clinically significant levels of aluminium oxide have, on rare occasions been linked to cases of Toxic Anterior Segment Syndrome (TASS) in published literature ⁽¹⁾, Rayner Intraocular Lenses Limited has initiated a voluntary recall of these products.

Product Name and Model Number	Batch Number	Sphere Power	Cylinder Power
Superflex Aspheric 920H	105E76367	+20.5D	
Superflex Aspheric 920H	105E76370	+21.0D	
C-flex Aspheric 970C	105E76411	+24.0D	
C-flex Aspheric 970C	105E76417	+25.0D	
C-flex Aspheric 970C	105E76419	+26.0D	

Product sold to Ireland:









Product Name and Model Number	Batch Number	Sphere Power	Cylinder Power
Superflex Aspheric 920H	105E76490	+10.0D	
Superflex Aspheric 920H	105E76492	+12.0D	
Superflex Aspheric 920H	105E76513	+13.5D	
Superflex Aspheric 920H	105E76515	+15.0D	
Superflex Aspheric 920H	105E76516	+15.5D	
Superflex Aspheric 920H	105E76517	+17.0D	
Superflex Aspheric 920H	105E76548	+20.5D	
Superflex Aspheric 920H	105E76551	+21.0D	
Superflex Aspheric 920H	105E76554	+21.0D	
Superflex Aspheric 920H	105E76568	+6.0D	
C-flex Aspheric 970C	105E76574	+26.0D	
C-flex Aspheric 970C	105E76576	+27.0D	
T-flex Aspheric 623T	105E76607	+14.5D +4.0D	
Superflex Aspheric 920H	105E76658	+16.0D	
Superflex Aspheric 920H	105E76659	+17.0D	
Superflex Aspheric 920H	105E76660	+17.5D	
T-flex Aspheric 623T	105E76683	+17.5D	+3.0D
Superflex Aspheric 920H	105E76713	+13.0D	

Product Name and Model Number	Batch Number	Sphere Power	Cylinder Power
Superflex Aspheric 920H	115E76725	+17.5D	
Superflex Aspheric 920H	115E76879	+7.0D	
T-flex Aspheric 623T	115E77242	+18.0D	+2.0D
Superflex Aspheric 920H	115E77266	+5.0D	

Information for stock control managers and healthcare facilities:

All listed batches (Lots) should be immediately quarantined in the first instance and then returned as promptly as possible (please refer to instructions below).

Rayner Intraocular Lenses Limited offers free replacement, reimbursement or substitution.

Information for Healthcare professionals and surgeons:

If a lens with an indicated serial number has already been implanted, it is recommended that as a precaution, you monitor for symptoms of TASS post operatively for up to a month in case of late onset presentation.

If cases of TASS do arise, please promptly report these cases to us and / or directly to your Health Authority.

Return contact details:

Return the completed response form and any product for replacement, reimbursement or substitution to:

FAO: Vigilance Department Rayner Intraocular Lenses Limited The Ridley Innovation Centre 10 Dominion Way Worthing BN14 8AQ United Kingdom

The completed form may also be returned to Rayner via e-mail to <u>feedback@rayner.com</u> or by fax to +44 (0) 1903 258901.

Rayner Intraocular Lenses Limited's Customer Commitment

Rayner Intraocular Lenses Limited sincerely apologises for any inconvenience this action may cause you. Replacement, reimbursement or substitution will be issued to you at the earliest opportunity.

Rayner Intraocular Lenses Limited is committed to ensuring that our products are manufactured to the highest standard and wish to inform you that we take all such matters extremely seriously.

Notification to Competent Authorities

By copy of this letter, Rayner Intraocular Lenses Limited wishes to inform you that the National Competent Authority (NCA) has been notified.

Should you have any questions regarding this field action, please do not hesitate to contact me or your Rayner representative.

Yours faithfully,

Eleanor Rees Regulatory Vigilance Manager Rayner Intraocular Lenses Limited

Ref:

¹Don Calogero, MS et al; Evaluation of Intraocular Reactivity to Metallic and Ethylene Oxide Contaminants of Medical Devices in a Rabbit Model; *Ophthalmology 2012;119:e36–e42*

Rayner Intraocular Lenses Limited Field Safety Notice Response Form

Device Name/Model	LOT Number	Implanted		Returned to Rayner			
		Yes 🗌	No 🗌	Yes 🗌	No 🗌		
		Yes 🗌	No	Yes 🗌	No 🗌		
		Yes	No	Yes 🗌	No 🗌		
		Yes 🗌	No	Yes 🗌	No		
		Yes 🗌	No 🗌	Yes 🗌	No 🗌		
		Yes 🗌	No 🗌	Yes 🗌	No 🗌		
		Yes 🗌	No 🗌	Yes 🗌	No 🗌		
		Yes 🗌	No 🗌	Yes 🗌	No 🗌		
		Yes 🗌	No 🗌	Yes 🗌	No 🗌		
Name and title of person completing form:							
Facility name:							
I have read and understood the contents of this Field Action.							
I have notified all affected persons of this Field Action.							
Signature:		Date:					

E-mail to feedback@rayner.com

Fax to +44 (0) 1273 324623 for the attention of: Rayner Vigilance Department.