

Flexible,  
robust  
solutions  
from BSI

An Ophthalmic  
Medical Device  
Notified Body



# Expertise and experience

Giving clarity to compliance

Updated May 2017

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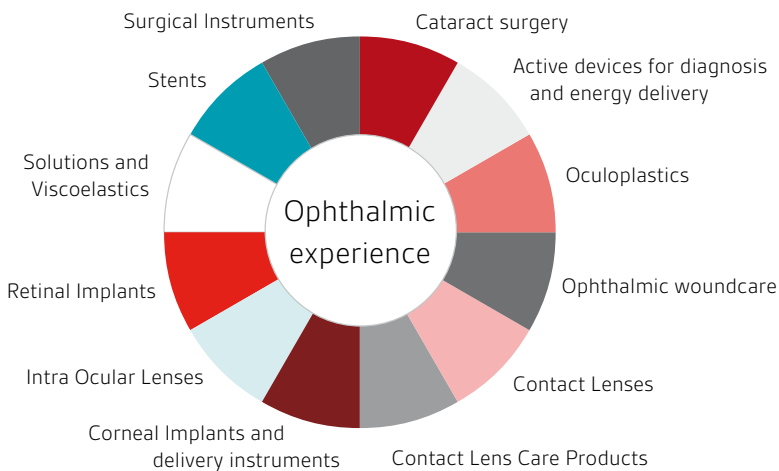
...making excellence a habit.™



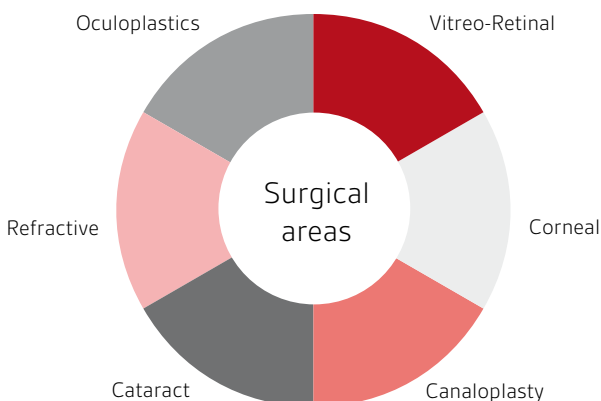
## Unrivalled expertise from the premier Ophthalmic Notified Body.

BSI is proud of the technical expertise we can provide in Ophthalmics. Our team has experience gained from both industry and regulatory roles. We have direct experience with manufacturers working with a wide range of ophthalmic medical devices.

**BSI's Ophthalmic experience includes a wide range of devices including:**



**Covering the following surgical areas...**



## Three unique reasons to make BSI your Ophthalmic Notified Body

### Experience and expertise:

BSI focuses on excellence, thereby reducing your corporate risk.

### Bespoke service:

BSI offers a range of review services, giving you a greater level of flexibility as well as predictability.

### Market Access:

Our efficient review services means your product reviews won't slow down your launch plans, helping you to stay ahead of the competition.

# Efficient and experienced services into global markets – giving clarity to compliance

## Be prepared

In the competitive medical device market place, ensuring that product development meets all regulatory requirements is essential. Understanding and consideration of the complicated clinical and regulatory requirements early in the product lifecycle could ensure your company gains the competitive advantage. Consolidated clinical and regulatory planning will assist your company to maximise resources and minimise time to market.

## CE-Excellence

Clients work with us because we understand the challenges medical device manufacturers face in getting compliant products to market efficiently and safely.

Our CE-Excellence review services deliver the efficiency you need to be competitive while maintaining confidence through a robust review. Explore the options below:

- CE-Standard: Our standard service reviews are completed by experienced BSI Product Experts, giving you confidence in the review.
- CE-Dedicated FastTrack: This service allows you to schedule your technical documentation review with a dedicated BSI Product Expert.
- CE-Onsite FastTrack: This review is conducted at your premises; a BSI Product Expert visits the facility for a period of time. This allows dynamic communications and opportunities for immediate responses.

## Worldwide access

Our expertise offers a wide range of proven regulatory and quality management programs that work together for full international compliance. Our QMS solutions include: ISO 13485, ISO 9001, ISO 14001 and many more.

BSI is a recognized Certification Body in Australia, Brazil, Canada, Hong Kong, Japan, Malaysia and Taiwan, and is a recognized MDSAP Auditing Organization for all participating Regulatory Authorities.

## Seamless transfer to BSI

If you decide to transfer your certification to BSI, we can offer a seamless exercise with comprehensive support and the absolute minimum level of disruption. We have expertise encompassing the full range of industry sectors and management system standards.

## Certification support

Throughout the certification process and beyond we can continually support you by providing:

- Expert training courses:
  - In-house for your company
  - Public courses, see website for the latest schedule
- Regulatory Updates, helping you plan for the future
- Free webinars
- Access to relevant standards

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**Please Note:** our programs do not guarantee CE marking certificate will be issued within a certain amount of working days, but are based on completing the review process with either a positive or negative recommendation.

BSI CE-Dedicated FastTrack and CE-Onsite FastTrack are not available for devices utilizing animal tissue or containing human blood derivatives or medicinal substances.



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# Global expertise



## Certification services

CE marking  
ISO 13485 QMS Auditing  
MDSAP Auditing – Australia, Brazil, Canada, Japan and USA  
Japan PMD Act  
Brazil INMETRO 60601 auditing and combined INMETRO, ISO 13485 and CE marking auditing  
Hong Kong CAB  
Malaysia CAB  
Taiwan TCP

## Training courses

CE marking for AIMD, MDD and IVD  
ISO 13485 QMS  
MDSAP: Fundamentals and readiness  
Introduction to risk management for medical devices  
CE marking Medical Devices with Software  
Compiling and Maintaining Technical Files and Design Dossiers  
Clinical Evaluation for Medical Devices  
Device - Drug Combinations: Practical Guidance on Borderline Issues and the Consultation Process  
Process Validation for the Medical Device industry  
Post Market Surveillance and Vigilance  
Medical Devices Utilizing Material of Animal Origin.  
Global Market Access courses

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**Your resource in worldwide compliance: Call BSI today on **1300 730 134** or visit **[bsigroup.com/en-au](https://www.bsigroup.com/en-au)** – to start your journey**

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