bsi.

Want Everything you need to to be an know about working with BSI external as an external **IVD** expert **IVD** expert for BSI?

...making excellence a habit."

What is BSI Medical Devices?

BSI The Netherlands (2797) is a leading Notified Body; we review medical devices to ensure that they conform to the requirements of the European Directives and Regulations. BSI UK (0086) is a UK Approved Body able to provide conformity assessments under the new UKCA scheme.

BSI performs conformity assessments to ensure devices are safe and perform as intended

What is an external IVD expert?

BSI's team of experts has a broad range of clinical, scientific and technical expertise, allowing us to review most aspects of these devices.

However, we sometimes need additional expertise; this is where your specialist scientific knowledge and experience would be invaluable, helping us to ensure that patient safety remains the focus of BSI product reviews. If you are interested in supporting BSI in assessing IVD medical devices for conformity in the European Union and Great Britain, we want to hear from you.

What are the areas we need additional expertise in?

These are some of the areas where we would value your contributions:

Histocompatibility Molecular Biochemisti immunogenetics **É Bacteriolo** Pathology and Virology Parasitology mmunology histopathology

What are the benefits to you?

- Be involved with innovation
- Support product review to ensure patient safety
- Work alongside a specialist team

Key requirements before applying:

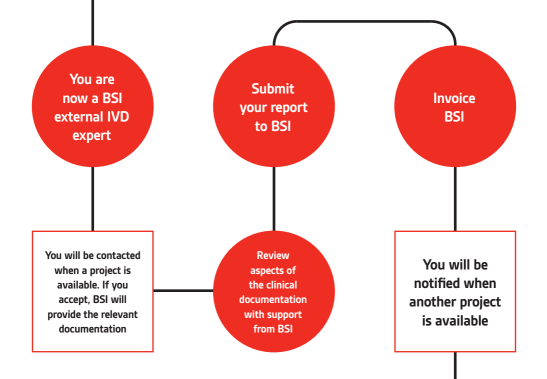
- You will be a senior scientist or clinician
- You will be educated to degree level or higher
- You will have at least four years' professional experience, including a significant period in a clinical environment



We review your education and experience

We include you on our secure database of external IVD experts. We will provide training in medical device legislation as required

* This means you agree to complete timely, unbiased reviews within your area of expertise Sign our Consultancy and Impartiality Agreements*



Interested in learning more?

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