

Synthetic Biomaterials

Technical Work Area 40

Project 6

Comparability in the thermal stability of a protein antibiotic

Objectives

The aim of this study is to validate the thermal stability of a pharmaceutically relevant protein antibiotic in native and near native environments by chiroptical spectroscopy, e.g. Circular Dichroism (CD) spectroscopy.

Background

There is an increasing shift in industry towards protein biologics – 30% of total sales and 50% of top 100 pharma sales by 2022 – with oncology and infectious diseases accounting for the most significant activity.

Proteins undergo structural transitions in their response to changes in while their activity environments. depends on conformational stability and reversibility. These require routine monitoring to support QA in product Regulatory agencies development. emphasise the need for higher-order structural characterization using CD spectroscopy to ensure comparability of biological products for pharmaceutical development (e.g. ICH Q5E).

However, achieving reproducibility in measurements is hampered by the lack of high-order reference materials and methods to benchmark the performance of protein biologics. This is further supported by international comparability studies performed to date (e.g. CCQM

P59, P59.1), which revealed the need for suitable reference materials to ensure the repeatability of CD measurement results. The proposed biologic is a protein antibiotic that folds in aqueous media into a highly stable tertiary structure exhibiting complete reversibility under thermal denaturation conditions.

Standardisation needs

The pre-standardisation needs focus on:

- far-UV CD spectra comparability between laboratories
- transition temperature evaluation upon thermal denaturation
- comparability in cooperativity and reversibility of the transition
- instrument performance validation and measurement variabilities as a function of temperature and concentration

Relevant Standards

ISO Guide 35 Reference materials

ISO 15189: 2012, BS ISO 29301: 2010

ISO 15194: 2009

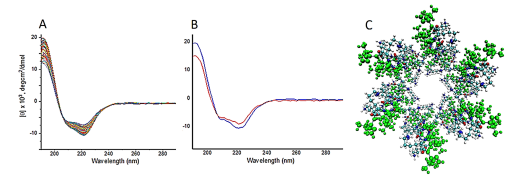
EU 2017/746: IVD regulation

ICH Q5E: guidelines for comparability of biological products.

Relevant Committees

ISO/TC 276 - Biotechnology ISO/TC 229 - Nanotechnologies CCQM CIPM – CAWG, PAWG

Call for Participation



(A) CD spectra of the antibiotic at set time points during thermal denaturation showing cooperativity of unfolding. (B) CD spectra before and after the thermal denaturation showing the reversibility of folding. (C) Model antimicrobial pore formed

Work Programme

- the antibiotic will be distributed to participants as a lyophilised powder
- protocols for sample preparation and measurements will be disseminated
- collected CD spectra will be compared with full uncertainty evaluations

Second stage analysis

 procedures developed will be repeated by a smaller group of participants to determine the effect of test parameters on the repeatability and reproducibility of the measurement results.

Knowledge Transfer

International round-robin tests, good practice guidelines, peer-review publications and presentations in conferences and standardisation venues.

Status

Study In progress since February 2019.

Additional Volunteers Welcome

Participants fund their own study in the project

For more information on participation, please contact:

Project Leader

Dr. Jascindra Ravi

National Physical Laboratory, UK

jascindra.ravi@npl.co.uk

TWA Chair **Dr. Max Ryadnov**National Physical Laboratory, UK

max.ryadnov@npl.co.uk

www.vamas.org

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