

 Contenido archivado el 2022-07-06

ENATRANS Webinar - The regulation Ecosystem for Nanomedicine

The Eu-funded project ENATRANS organises this webinar to inform about the regulation Ecosystem for Nanomedicine and give practical advice about what is required and who can help at the different steps of the Nanomedicine regulation process.

 1 Diciembre 2017 - 1 Diciembre 2017

 Germany



© ENATRANS 2017

The translation of any new medical invention into a final product is a highly regulated and complex process. To bring a new medical product or device on the market it has to go through a long and complex process involving many authorities at regional, national and international level. To successfully manage this process it is mandatory to study and understand the different parts of the process in detail.

The webinar gives an overview of the broader nanomedicine ecosystem and issues to be aware of such as regulation, pricing and reimbursement systems by providing all data about these steps in place in Europe, at the EU, and Member States level. The provided information, links and downloads will serve as a guide about what is required in which order and who can help at the different steps of the process.


Free-of-charge registration at: <https://zoom.us/meeting/register/8c53975e94f9846b34538d7d4481ef37> 

Palabras clave

Nanomedicine, Nanobiomedical, regulation, translation, nanotechnology for healthcare

Colaborador

Aportado por

VDI/VDE Innovation + Technik GmbH
Germany 

Proyectos conexos



ENATRANS

Enabling NAnomedicine TRANSlation

28 Junio 2023

PROYECTO

Última actualización: 27 Noviembre 2017

Permalink: <https://cordis.europa.eu/event/id/152469-enatrans-webinar-the-regulation-ecosystem-for-nanomedicine/es>

European Union, 2025