



Read more about how we are strengthening clinical trials in Denmark and the EU here

Clinical Trials Denmark Invests to Remain the EU's #1

With a new ambitious life science strategy in place, the Danish Medicines Agency and the Danish National Center for Ethics are setting the direction for the development of regulatory framework that contributes to **Denmark's position** as a leading life science nation.

Our focus



Fast Review Process:

Especially for early phase trials.



High-Level Expertise: Developing regulatory framework to align with the latest technology and research.



Pragmatic Approach:

Without compromising patient safety.



Responsibility: As a leading Reporting Member State (RMS), Denmark takes responsibility for the process from start to finish.



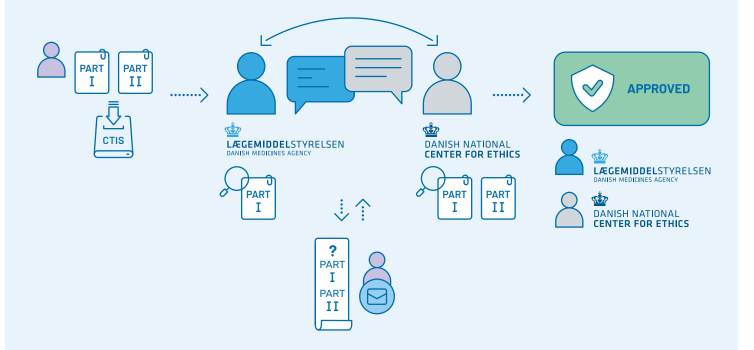
Transparency: Ensuring clear, well-reasoned, and consistent requirements in Denmark and the EU.



Accessibility: Being a reliable sparring partner for all stakeholders.

Did you know that Denmark is the country in the EU conducting the most clinical trials with medicinal products per million inhabitants?

Through collaboration, we ensure an optimal approval process



In Denmark, applicants have easy access to guidance from both authorities.

The aim is that mononational trials receive response within 31 days in Denmark - and even faster for early phase trials.

Contact



Danish Medicines Agency

Clinical Trials Unit T +45 44 88 91 23 E kf@dkma.dk

laegemiddelstyrelsen.dk



Medical Research Ethics Committees

Danish National Center for Ethics T +45 72 21 66 77

E dketik@dketik.dk

researchethics.dk