



From concept to market-ready medical devices

Without unnecessary complexity

Bringing a medical device to market requires more than engineering. Development, regulatory requirements, and production must work together.

Ergo-Tec connects these steps.

Either as a complete solution or in selected phases.

What you gain

- Fewer interfaces and less coordination effort
- Clear responsibilities across all project phases
- No need to build your own production setup
- A structured path from concept to series production
- A partner who actively drives implementation

How your product moves forward

Concepts designed for production

Your requirements are translated into technical solutions that can be manufactured efficiently.

Early prototypes for faster decisions

Functional prototypes allow testing and validation before moving into production.

Regulatory requirements integrated from the start

Documentation and compliance are considered early in the process.

Components that fit together

Materials and parts are selected to meet technical and regulatory needs.

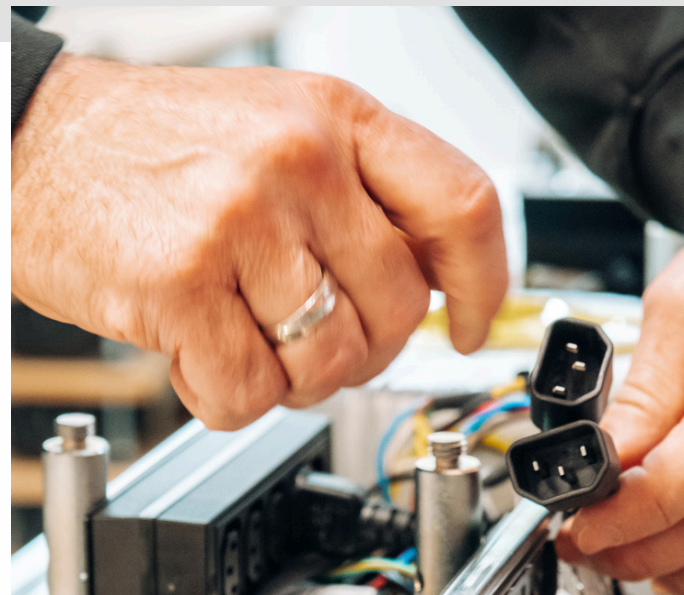
Reliable series production

Assembly and manufacturing follow defined processes for mid-volume series.



When it makes sense to work with Ergo-Tec

- You require additional components or systems to complete your product
- You want to develop and manufacture with one partner
- You have a product but no production capabilities
- You are preparing a start-up for market entry
- Your internal production is limited or highly complex





Clear collaboration – your know-how stays protected

You define requirements and specifications.
Ergo-Tec implements them in a structured way.

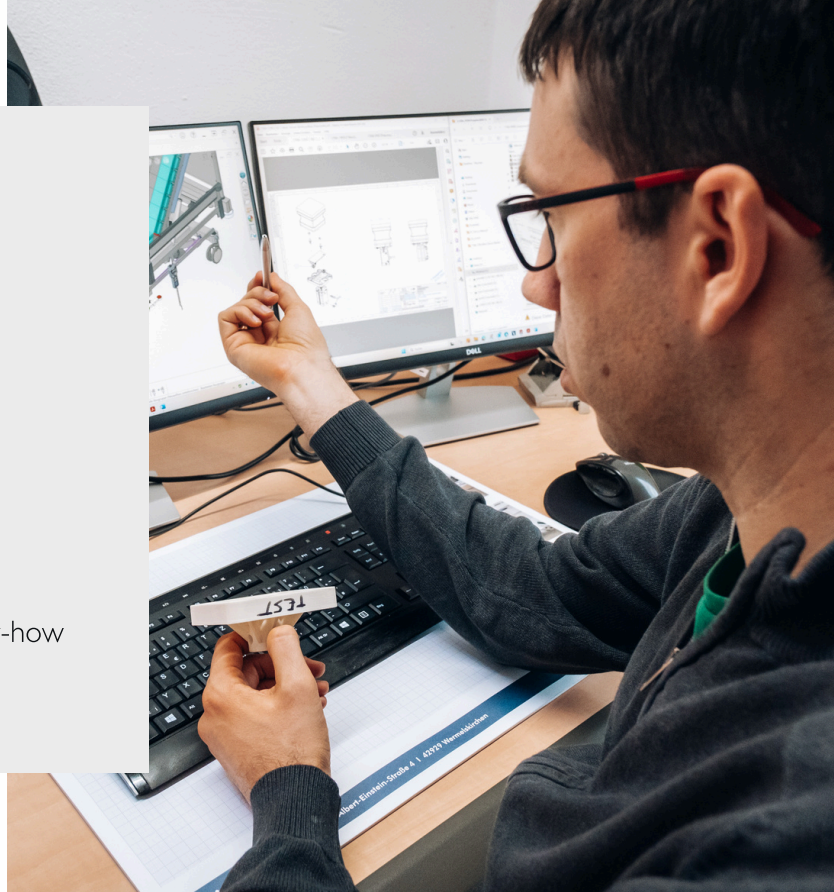
All results remain transparent and traceable – your know-how stays with you.

Experience that keeps projects moving

With over 25 years in medical device development and manufacturing, Ergo-Tec understands how to align development, regulatory preparation, and production. Our certification according to DIN EN ISO 13485 ensures structured processes and consistent quality throughout all project phases. This reduces delays and avoids late-stage changes.

Let's talk about your project

If you are planning to develop or scale a medical device, we support exactly where you need it.



Focus: Patient positioning systems

Our key focus lies on electrically adjustable systems such as:

- examination and treatment chairs
- operating tables
- hospital beds
- specialized systems such as dysphagia chairs



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