



## Effectum Medical AG

assists in the development of innovative medical devices and technologies throughout the certification and life cycle management process. Acting as legal manufacturer and by offering our Plug & Play (e)QMS, we speed up your time to market and ensure compliance and excellence in every stage of development.



**E F F E C T U M**  
**M E D I C A L**

# AI/ML DEVICE CERTIFICATION



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# ARTIFICIAL INTELLIGENCE (AI) MACHINE LEARNING (ML) DEVICE CERTIFICATION

## AI and ML Device Certification for Healthcare Navigate AI/ML Integration with Confidence

Incorporating Machine Learning (ML) and Artificial Intelligence (AI) into medical devices brings transformative potential, but it also requires navigating a complex regulatory environment. Effectum Medical provides expert guidance to ensure your devices meet all relevant compliance standards, including the European AI Act, MDR, and IVDR.



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## COMPLIANCE SERVICES

### Our Comprehensive AI/ML Compliance Services

#### AI Readiness & Compliance Checks

- **Gap Analyses & AI Readiness:** We review your documentation to identify any gaps, ensuring it aligns with AI Act requirements.
- **Dual or Single Regulation Assessments:** Tailored checks to meet both AI Act and MDR/IVDR compliance, providing peace of mind at every stage.

#### CE Certification Support

- **Expert Guidance:** Our team assists in achieving CE certification, offering specialized support for ML/AI-based software to help you navigate the approval process efficiently.

#### Technical Documentation Templates

- **Streamlined Preparation:** Access customizable templates that simplify the preparation of technical files, helping you maintain compliance while saving time.

#### AI Act Compliance Solutions

- **Harmonized Compliance Strategy:** Our established processes ensure that your systems meet both AI Act and MDR/IVDR requirements efficiently and effectively.



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## LIFECYCLE SUPPORT FOR ML/AI DEVICES

From development through to post-market monitoring, we support your ML/AI device across its entire lifecycle:

- **Pre-market Conformity Assessments:** Determine if a notified body conformity assessment is necessary or if internal controls will apply. Our team will also assist in preparing technical documentation in line with both AI Act and MDR/IVDR standards.
- **Clinical Evaluation Support:** We help align clinical evaluations with ML/AI-specific regulatory requirements.
- **Continuous Monitoring Solutions:** Ensure your device remains compliant as it evolves in real-world use, meeting ongoing monitoring and regulatory requirements.

### Preparing for the Future of Healthcare AI Compliance

Developing a robust ML/AI device can take 1-2 years, with conformity assessments adding another year. To meet the August 2027 deadline for high-risk applications under the EU AI Act, it's critical to start early.

Let us guide you through each step to ensure readiness for this evolving regulatory landscape.



## WHY CHOOSE EFFECTUM MEDICAL?

### Why Choose Effectum Medical?

- **Dual Expertise:** Specialists in AI regulation and medical device compliance.
- **Efficiency:** Streamlined processes that align with AI Act and MDR/IVDR.
- **Future-Proofing:** Preparedness for upcoming regulatory changes, ensuring your product's compliance and adaptability.

Don't let regulatory requirements stall your healthcare AI innovations. **Partner with Effectum Medical** to turn compliance into a growth opportunity and build patient trust.

**Contact us** for a personalized consultation to prepare your ML/AI models for the European market.



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## SUPPORT OPTION



### **Outsourced Legal Manufacturer:**

Streamline market access and compliance with our trusted legal manufacturer services. Need speed? Ask about our fast-track option!



### **Plug & Play (e)QMS:**

Accelerate your launch with a ready-made EN ISO 13485, MDR/IVDR, and FDA-compliant quality management system.



### **Plug-and-Play eQMS Bootcamp:**

Flexible training to empower self-management of your QMS.



### **Rent an Expert:**

Add quality management and regulatory expertise to your team as needed.



### **Regulatory Opinions:**

Access expert insights for global compliance and market strategy.



### **AI/ML Device Certification:**

Certification services for AI and machine learning-based medical devices.



### **CH-REP (Swiss Authorized Representative):**

Facilitate Swiss market access with local representation. Visit [Effectum CH-REP AG](https://www.effectummedical.com/CH-REP) for details.

For more information visit: [www.effectummedical.com](https://www.effectummedical.com)



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