

“All Non-European manufacturers of medical equipment who wish to market their products in the EU market under their own name must designate a Regulatory Authorized Representative in order to meet CE requirements.”

The European medical device market represents an annual sales volume in excess of 40 Billion Euros. This means, it has become the second largest world-wide, after the United States.

There are, as always, certain obstacles and hurdles to be overcome before the sale of medical devices into Europe. **MEDDEVICE can help you to** easily gain access to this immense opportunity by assisting you to register for CE-marking and by acting as your Authorized Representative as defined by the European Medical Devices Directives (MDD 93/42/EEC with rev. 2007/47/EC).

MEDDEVICE opens the door to Europe providing you with the best representation possible in the ever-changing environment of EU legislation.

