



Unlock the World with Penn Pharma

Getting your IMPs and Commercial products into the EU has never been simpler with Penn Pharma's PharmacEUtical Portal.

Penn's single site operations facility is based in the UK, which makes it ideally located to assist non-EU clients import and distribute pharmaceutical and clinical trial material throughout the EU and the world. With all operations being on one site, it creates a seamless integrated service which aids project communication and enables us to maintain a personal approach when dealing with our clients' projects.

Penn is MHRA approved and is FDA inspected. Penn operates an ISO based, mature quality management system which is regularly inspected by both the Medicines and Healthcare Products Regulatory Agency (MHRA) and the United States Food and Drug Administration (FDA).

With the advent of the new European Clinical Trials Directive (2001/20/EC) in May 2004, Penn was one of the first companies in Europe to obtain the new MA-IMP licence.

Penn has temperature controlled Storage & Distribution and Supply Management warehouses and a global distribution network to assist you with the importation, storage and distribution of your clinical trials materials to anywhere in the world.



With 6 full-time dedicated QPs, lab support, controlled drug and cold chain distribution networks, we can assist you in accessing the EU and Global patient population. Our FDA history combined with MHRA regulation and fully cGMP compliant services ensures that your products will not only access the EU but will be of the highest standards possible. Full retesting and stability studies are available for commercial products no matter how big or small the volume.

You're in good hands with Penn.

Penn Pharma

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For more information

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WITH YOU EVERY STEP OF THE WAY