



What we do?



Expertise in Process Research/Development and scale-up (non-GMP and cGMP).



Technology driven organization with a flexible approach. Founded by scientists with proven industrial track record.



Research partner for Innovator/Generic Pharmaceutical and Biotech companies across the globe



OUR SERVICES:



CONTRACT DEVELOPMENT

We offer a complete service including:

- regulatory strategy evaluation
- formulation development
- •scale-up
- manufacturing process validation
- methods development and validation
- •ICH stability test





From design to finished product, we support our customer's project all the way.



OUR SERVICES:



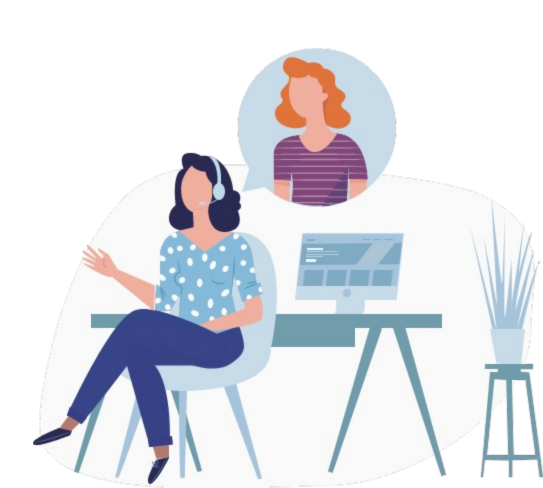
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AS LICENSING OUT:

- •We out-license our registration dossiers, our registration dossiers, for the partners to register and apply for a Marketing Authorisation in their country with their trademark.
- •We are also willing to follow the registration procedure up to the granting of the Marketing Authorisation on behalf of our partner (in any) and then transfer the related MA once obtained directly to our partner.

In both cases, the partner becomes the MA holder and finished product manufacturer.



We grant professional regulatory support from a preapproval to the post-approval stage for the full lifecycle of any product.





We put patients first and help people around the world live better and healthier life':

Choose quality and innovation - choose us.

Choose quality and innovation - choose us. With us, you can be sure of the quality and effectiveness of our products.



Make the right choice for your health and the health of your loved ones - choose our company.

All Products Worldwide:

Alimentary tract & metabolism



Dermatologicals



Antiinfective agents

Anti-neoplastic &





Blood & blood forming organs

Cardiovascular system



Musculo-skeletal system

Genito urinary system & sex hormones

Nervous system



Sensory organs



Respiratory system

Systemic hormonal agents



OTC



OUR PRODUCTION FACILITY IN UNITED KINGDOM



Noumed House Shoppenhangers Road Maidenhead, SL6 2RB United Kingdom

Phone: +44 (0) 20 3399 8900









An MHRA accredited Manufacturing Facility based in Maidenhead, UK.

Noumed's manufacturing facility has the capability to manufacture various solid dosage forms which include plain and coated tablets, and hard gelatin capsules. The facility is equipped with R& D suites to aid New Product development, undertaking feasibility and optimization trials and for manufacturing small batches for clinical trials etc.

All production takes place in isolated ISO-Class 8 / Class D certified rooms in accordance with cGMP and FDA guidelines. Additionally, all airflows are regulated through the use of airlocks and pressure differentials.







Manufacturing is supported with in-process storage areas and material handling equipment to improve efficiency. The facility has 3 suites for tablet manufacturing and 2 suites for capsule manufacturing.

The tablet manufacturing suite has 3 independent granulation areas supported by tablet compression machines and coating pans. The 3 granulation suites are equipped with a Vacuum Transfer System.

The 2 capsule manufacturing suites are fitted with automatic capsule filling lines. The manufacturing facility is equipped with a range of cage blenders from 50 litres to 1200 litres, and has a dedicated area for formulation/product development."

OUR PRODUCTION FACILITY IN TURKEY

Cerkezkoy

The World Medicine facility in Cerkezkoy, in the province of Tekirdag, was completed in 2020. The facility is our flagship investment, producing a range of pharmaceuticals including liquid sterile forms, lyophilized sterile forms, solid forms, semi-solid forms, aerosol inhalers and non-sterile liquid forms. Equipped with machinery which exceeds world standards using the latest technology, the facility is controlled with systems that meet the strictest production standards in the international arena in terms of hygiene and security.

Providing services with an annual production capacity of 1.3 billion boxes (13.6 billion semi-products/year), our factory operates on an area of 50,000 m2 with competent staff with their finger on the pulse of global dynamics, who possess extensive production knowledge and who work to a quality policy of product and service perfection.





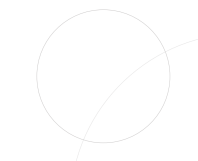


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Non Sterile Liquid and Semi-Solid Section Production	n Total 350 million boxes/year
Syrup Filling	75 million bottles/year
Screw Cap, Filling	100 million bottles/year
Snap On Filling	125 million bottles/year
Nasal Spray Filling	75 million bottles/year
Suppository Filling	200 million pieces/year
Pomade Tube Filling	40 million tubes/year
Sterile Liquid and Lyophilized Injectable S Total Production	boxes/year
Small Sized Vial Liquid Filling	150 million vials/year
Small Vial Lyophilisation	35 million vials/year
Large Sized Vial Liquid Filling	25 million vials/year
Ampoule Filling	400 million ampoules/year
Muti-Doses Eye Drop Filling	100 million bottles/year
Mono-Dose Eye Drop Filling	150 million pieces/year
Blisters (Vials / Ampoule)	220 million blisters/year
Solid Section Total Production	500 million boxes/year
Tablet	10 billion tablets/year
Capsule Filling	1,6 billion capsules/year
Bottle Powder Filling	60 million bottles/year
Sachet Filling	250 million sachet/year
Stick Filling	200 million stick/year
Blister (Tablet / Capsule)	1 billion blister/year



CONTACT US:





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