



Manufacturer & Exporters of

WHO GMP CERTIFIED COMPANY Bulk Drugs & Drug Intermediates

About Us

Established in the year 1996, We, Vaikunth Chemical Pvt. Ltd., take immense pride in introducing ourselves as a prominent manufacturer, supplier and exporter of a wide range of bulk drugs and drug intermediates. A more than 25 years of experience and expertise has strengthened our morale and has motivated us to offer innovative solutions in drugs to our valued clients worldwide.

Vaikunth has made successful forays in Manufacturing of Bulk Drugs and Drug Intermediates.

The company has carved a niche for itself with its dynamic and strong technical team. We take pride in developing and commercially manufacturing of our molecules for our esteemed clients.



MANUFACTURER & EXPORTERS OF BULK DRUGS & DRUG INTERMEDIATES



ESTABLISHED IN THE YEAR 1996

Got Manufacturing License for Bulk Drugs and Drug Intermediates. Form 25. L/C No. G/25/1996

1996

2009

GMP CERTIFICATE

Company receive GMP certificate from FDCA for Bulk Drug Manufacturing. Certificate No. S-GMP/0910148

EXPANSION

Amalgation of plot number C1B/407/10 &11 with existing Plot No. 408/4 & 5

2020

REAPPROVAL OF WHO GMP CERTIFICATE

Company receive Reapproval of WHO GMP certificate for Bulk Drug Manufacturing. Certificate No. 23044037

2017

WHO GMP CERTIFICATE

> Company receive WHO - GMP certificate for Bulk Drug Manufacturing. Certificate No. 20011804

2023



Vision

To forge a company that utilises its scientific and technological capabilities combined with strategic research to deliver manufacturing competencies.



Quality Policy

Vaikunth is committed to manufacturing highest quality Active Pharmaceutical Ingredients and its Intermediates by strict adherence to WHO current Good Manufacturing Practice and Quality Management System to enhance our customer satisfaction.

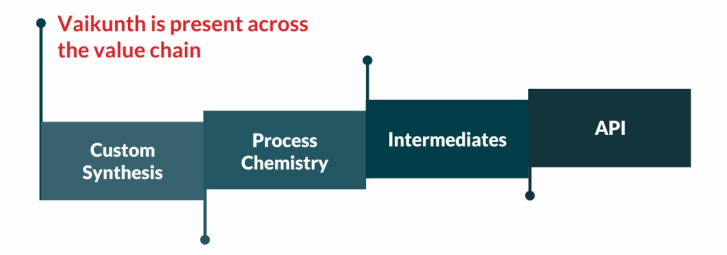


EHS PHILOSOPHY

Vaikunth accords highest priority to carry out the business in a sustainable manner, ensuring safety of their employees and protection of environment.

Business Highlights





Strategy in Place for Next Growth Phase

Strengthen position as a speciality high and generic player with best in class regulatory and development capabilities.

Long term supply contract and new product development to be the key growth drivers.

Integrated Capabilities

Strong end-to-end capabilities with presence across the entire value chain of APIs and Drug intermediates made by strong manufacturing and regulatory capabilities along with front-end presence across markets.

Diversified Geographical Presence

Vaikunth has global presence in over 40 countries by exporting the high quality APIs & Drug intermediates.

Strong Operating Performance

Growth going forward to be driven by development of new product segment & supplies to customers.

Strong Manufacturing Infrastructure

API manufacturing facilities with approvals from leading agencies providing global competitive edge.

Manufacturing Facilities are in compliance with WHO GMP, GLP, ISO 9001:2015, ISO 14001:2015 & Halal.

Excellent Regulatory & R&D Capabilities

Strong track record of regulatory documents fillings.

Strong capability of R&D so that we can provide customers with a variety of special products from grams to kilograms and tons.

The capability of R&D is our core competitiveness.

Product List

Bulk Drugs APIs

PRODUCT NAME	CAS NO.	GRADE	THERAPEUTIC CATEGORY
Cetirizine Hydrochloride	83881-52-1	IP, BP, EP, USP	Antihistaminic
Levocetirizine Dihydrochloride	130018-77-8	IP, USP	Antihistaminic
Metoclopramide Hydrochloride	54143-57-6	IP, BP, EP, USP	Antiemetic
Tolperisone Hydrochloride	3644-61-9	JP, IH	Muscle Relaxant
Trifluoperazine Hydrochloride	440-17-5	IP, BP, EP, USP	Nueroleptic
Metoclopramide	364-62-5	BP, EP	Antiemetic
Ethopabate	59-06-3	BP, EP, USP	Antiprotozoal
Diethylamine salicylate	4419-92-5	BP, EP	Anti inflammatory
Aminosalicylic acid	65-49-6	USP	Antituberculosis
Trazodone Hydrochloride	25332-39-2	BP, USP	Antidepressant
Chlorzoxazone	95-25-0	USP	Muscle Relaxant
Prochlorperazine	58-38-8	USP	Antipsychotic / Antiemetic
Prochlorperazine Maleate	84-02-6	IP, BP, USP	Antipsychotic / Antiemetic
Valproic Acid	99-66-1	IP, USP	Antiepileptic / Anticonvulsant
Sodium Valproate	1069-66-5	IP, BP, EP, USP	Antiepileptic
Divalproex Sodium	76584-70-8	IP, USP	Antiepileptic / Anticonvulsant







Intermediates

PRODUCT NAME	CAS NO.	USED FOR API MANUFACTURING	
P-Chloro Benzophenone	134-85-0		
P-Chloro Benzhydrol	119-56-2		
P-Chloro Benzhydryl Chloride	134-83-8	Cetirizine Hydrochloride	
P-Chloro Benzhydryl Piperazine	303-26-4		
Cetirizine Base	83881-51-0		
Para Amino Salicylic Acid	65-49-6		
4-Acetyl Amino Salicylic Acid	50-86-2		
"Methopabate or 4-Acetyl Amino-2- Methoxy Benzoic Acid Methyl Ester"	4093-29-2		
"5-Chloro Methopabate or Methyl-4- Acetamido-5-Chloro-2-Methoxybenzoate"	4093-31-6	Metoclopramide Hydrochloride	
"Metoclopramide Base or 4-Amino-5-Chloro-N-[2-(Diethylamino) Ethyl]-2- Methoxybenzamide"	364-62-5		
N,N-Diethyl Ethylene Diamine	100-36-7		
(-)-4-Chlorobenzhydrilamine	163837-57-8	Lovacetivizina Dibudra ablasi da	
(-)-P-Chloro Benzhydryl Piperazine	300543-56-0	Levocetirizine Dihydrochloride	
Methyl-4-Amino-2-Hydroxy Benzoate	4136-97-4	Ethopabate	
Methyl-4-Acetamido-2-Hydroxy Benzoate	4093-28-1	Europubato	

APIs Under Development

PRODUCT NAME	CAS NO.	CATEGORY
Ezetimibe	163222-33-1	Antihyperlipidemic
Fexofenadine	83799-24-0	Antihistamine
Elagolix Sodium	832720-36-2	GnRH receptor antagonist

Manufacturing Facility



























































300+ 40+ 25 +CUSTOMERS **COUNTRIES YEARS**



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