TMDA/DMC/MRE/F/016 Rev #:02



THEUNITEDREPUBLICOFTANZANIA



MINISTRYOFHEALTH

TANZANIAMEDICINES ANDMEDICALDEVICESAUTHORITY

PUBLIC ASSESSMENT REPORT FOR ATORVAS 10 (ATORVASTATIN CALCIUM 10 MG) FILM COATED TABLETS

Version number 01 3rd January, 2023

TMDA Headquarters, Plot No. 56/1, Block E, Kisasa B Centre, Swaswa Road, P. O. Box 1253, Dodoma – Tanzania, Telephone: +255 (26) 2961989/2061990/+255 (22) 2450512/2450751/2452108, Email: <u>info@tmda.og.tz</u>, Website: <u>www.tmda.go.tz</u> Toll free: 0800110084

1. Introduction

ATORVAS 10 is a generic medicine of Lipitor (Atorvastatin calcium 10 mg). ATORVAS 20 is a Lipid modifying agents medicine belonging to HMG-CoA-reductase inhibitors group. ATORVAS 10 is a selective, competitive inhibitor of HMG-CoA reductase, the rate-limiting enzyme responsible for the conversion of 3-hydroxy-3-methyl-glutarylcoenzyme A to mevalonate, a precursor of sterols, including cholesterol. Triglycerides and cholesterol in the liver are incorporated into very low-density lipoproteins (VLDL) and released into the plasma for delivery to peripheral tissues. Low-density lipoprotein (LDL) is formed from VLDL and is catabolised primarily through the receptor with high affinity to LDL (LDL receptor). Atorvastatin lowers plasma cholesterol and lipoprotein serum concentrations by inhibiting HMG-CoA reductase and subsequently cholesterol biosynthesis in the liver and increases the number of hepatic LDL receptors on the cell surface for enhanced uptake and catabolism of LDL. Atorvastatin reduces LDL production and the number of LDL particles. Atorvastatin produces a profound and sustained increase in LDL receptor activity coupled with a beneficial change in the quality of circulating LDL particles. Atorvastatin is effective in reducing LDL-C in patients with homozygous familial hypercholesterolaemia, a population that has not usually responded to lipid-lowering medicinal products

ATORVAS 10 is approved in Tanzania for use in adults, adolescents and children aged 10 years or older with primary hypercholesterolaemia including familial hypercholesterolaemia

Registration number	TAN 21 HM 0106		
Brand name	ATORVAS 10		
Generic name, strength and	Atorvastatin calcium 10 mg		
form			
ATC classification	C10AA05, Lipid modifying agents, HMG-CoA-		
	reductase inhibitors		
Distribution category	POM		
Country of origin	India		
Associated product	ATORVAS 20 (Atorvastatin calcium 20 mg)		
Marketing Authorization Holder	Sunberg Life Sciences PVT. Limited, No.15, Gopalakrishna Road, T. Nagar, Chennai 600 017. Tamil Nadu. India Telephone: +91 44 2345 2030-34 Telefax: +91 44 2345 2036, E-Mail: vijay@softgelhealthcare.com		
Local Technical	Moraf Pharmaceuticals Ltd,		
Representative	P.O. Box 21323, Dar es Salaam,		

1.1 Product details

Tanzania.		Tanzania.
-----------	--	-----------

1.2 Assessment procedure

The application for registration of ATORVAS 10 was submitted on 22nd November, 2017. The product underwent full assessment. Assessment was completed in four rounds of evaluation. ATORVAS 10 was registered on 29th March, 2022.

1.3 Information for users

Visual description of the finished	Orange coloured circular slightly biconvexed		
product	film coated tablets and scored in the middle		
	on one side		
Primary packing material	3 × 10's Alu - Alu Blister Pack		
Secondary packing materials	Carton box		
Shelf-life and storage condition	24 months		
	Store below 30°C. Protect from light and		
	moisture		
Route of administration	Oral		
Therapeutic indications	Atorvastatin is indicated as an adjunct to diet		
	for reduction of elevated total cholesterol		
	(total-C), LDL-cholesterol (LDL-C),		
	apolipoprotein B, and triglycerides in adults,		
	adolescents and children aged 10 years or		
	older with primary hypercholesterolaemia		
	including familial hypercholesterolaemia		
	(heterozygous variant) or combined (mixed)		
	hyperlipidaemia (corresponding to Types Ila		
	and IIb of the Fredrickson classification) when		
	response to diet and other no		
	pharmacological measures is inadequate.		
	Atorvastatin is also indicated to reduce Total-		
	C and LDL-C in adults with homozygous		
	familial hypercholesterolaemia as an adjunct		
	to other lipid-lowering treatments (e.g. LDL		
	apheresis) or if such treatments are		
	unavailable. Prevention of cardiovascular		
	disease Prevention of cardiovascular events		
	in adult patients estimated to have a high risk		
	for a first cardiovascular event, as an adjunct		

	to correction of other risk factors
--	-------------------------------------

2. Labelling and product information

Summary of product characteristics

The SmPC included all the relevant information to ensure correct and safe use of the medicine by healthcare providers. The complete SmPC can be accessed <u>here</u>.

Package insert/leaflet

The package insert is confirmed to be derived from the SmPC and contains sufficient data for the end user. Since the product is POM that is intended for long term use the package insert contains full prescribing information as per SmPC.

Container labels

The product label information is presented in English. Details in the secondary pack label include:

Brand name: ATORVAS 10 (Atorvastatin calcium 10 mg) film coated tablets

Composition: Atorvastatin calcium 10 mg, Lactose, Microcrystalline Cellulose, Povidone K30, Calcium Carbonate, Croscarmellose Sodium, Magnesium Stearate, Opadry White, Sunset Yellow Lake, Dichloromethane, Isopropyl Alcohol

Pack size: 3 × 10's

Manufacturing details: <batch number, manufacturing date, expiry date>

Storage conditions: Store below 30°C. Protect from light and moisture

Manufacturer address: The Madras Pharmaceuticals, No.137B, Old Mahabalipuram Road, Karapakkam, Chennai - 600 096, Tamil Nadu, India.

Unique identifier: NA

Special warnings/precautions or instructions for use: <include warnings or IFU if applicable>

The details of the primary pack include:

Brand name and strength: ATORVAS 10 (Atorvastatin calcium 10 mg) film coated tablets

Manufacturing details: <batch number, manufacturing date, expiry date>

Name of manufacturer: The Madras Pharmaceuticals

The content of the primary and secondary labels was aligned to the requirements of the Part V of the Compendium: Guidelines on Format and Content of Product Labels for Medicinal products. The label contains sufficient information for proper identification of the medicine and post marketing follow up of the product.

Describe any approved deviation to the requirements and the justification for the deviation.

Mock labels are appended as annex I

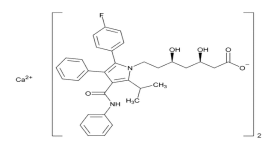
3. Scientific discussion

Quality of Active Pharmaceutical Ingredient(s)

Information on quality of the API was submitted in form of Full details

General properties

Atorvastatin Calcium API is compendia in USP/BP/JP Molecular formula: $C_{66}H_{68}CaF_2N_4O_{10}.3H_2O$ Chemical name: (3R, 5R)-7-[2-(4-fluoropheilyl)-3-phellyl-4- (phenylcarbamoyl)-5-(propan-2-yl)-IH-pyrrol-lylJ-3, 5 dihydroxyheptanoate calcium Trihydrate Structure:



Critical physico-chemical properties of the API were very slightly soluble in water. acetonitrile; slightly soluble in ethanol (95 percent), freely soluble in methanol.

<u>Manufacture</u>

The API manufacturing site, DSM Sinochem Pharmaceuticals India Private Limited, Bhai Mohan Singh Nagar, Toansa, Tehsil: Balachaur; District: Nawanshahr - 144 533 Punjab, India was noted to comply with WHO GMP requirements as evidenced by the GMP certificate issued by Food and Drug Authority, Punjab, Pariwar Kalyan Bhawan, Sector 34-A Chandigarh. Atorvastatin Calcium API is manufactured by chemical synthesis using conventional techniques. Sufficient controls of quality of materials and in-process checks were employed throughout the manufacturing process.

Specifications

The API specifications were set as per in-house standards and ICHQ3A. The parameters monitored during quality control are: Appearance, solubility, identification,

optical rotation, related substances, heavy metal, water, assay, particle size distribution. Compliance to these specifications were established via batch analysis data and stability studies.

Stability and container closure system

The re-test period of Atorvastatin Calcium API is 48 months when packed in LDPE bag which in turn placed in HDPE drum and stored at 30°C.

Quality of the Finished Pharmaceutical Product

Formulation

ATORVAS 10 is a White coloured circular slightly biconvexed film coated tablets and scored in the middle on one side packed in Alu - Alu Blister. ATORVAS 10 contains Atorvastatin calcium and other ingredients listed here after Lactose, Microcrystalline Cellulose, Povidone K30, Calcium Carbonate, Croscarmellose Sodium, Magnesium Stearate, Opadry White, Sunset Yellow Lake, Dichloromethane, Isopropyl Alcohol. The quantities of all ingredients are confirmed to be in line with the recommendations of Handbook of Pharmaceutical Excipients, Edition 8th in terms of function and quantities. Ingredient, Lactose is of safety concern therefore appropriate warnings were included in the product label.

Manufacture

The finished product was manufactured at The Madras Pharmaceuticals, No.137B, Old Mahabalipuram Road, Karapakkam, Chennai 600 096. India. The compliance of the site to TMDA GMP standards was confirmed through site inspection on 5th June 2014.

Specifications

The FPP is compendia in BP/USP. The manufacturer controls the quality of the finished product as per in-house and ICHQ3B requirements. The parameters monitored during quality control are: Appearance, identification, average weight, uniformity of weight, disintegration time, dissolution, uniformity of content, related substance, assay, microbial limit test, residual solvents. Compliance to the standard was established using batch analysis data and stability data.

Stability and container closure system

Stability studies were conducted on three batches of the finished product stored at $30^{\circ}C \pm 2^{\circ}C/75\% \pm 5\%$ RH for 24 months and $40^{\circ}C \pm 2^{\circ}C/75\% \pm 5\%$ RH for 6 months. Based on the stability data presented, the approved shelf-life is 24 months when stored in Alu – Alu blisters at below $30^{\circ}C$.

Safety and efficacy information

The biowaiver was approved based on additional strength.

ATORVAS 10 fulfilled the criteria for waiving an in-vivo bioequivalence study as per relevant TMDA guidance. Dissolution profiles of ATORVAS 10 (Atorvastatin calcium 10 mg) film coated tablets was compared to < ATORVAS 20 (Atorvastatin calcium 20 mg) film coated tablets .less than 85% of the labelled amount of Atorvastatin calcium had dissolved in all three media. Therefore, necessitating calculation of similarity factor f2, which was noted to be above 50.

4. Benefit-Risk Assessment and Conclusion

On basis of the data submitted, the current state of knowledge and compliance to Good Manufacturing Practice, the benefit of the product outweighs the risks associated with its use when used in accordance to the summary of product characteristics. ATORVAS 10 is recommended for registration.

5. Post-approval updates

Variation applications

Reference number	Date submitted	Change requested	Recommendation	Granting date

Feedback from pharmacovigilance, post marketing surveillance and enforcement activities

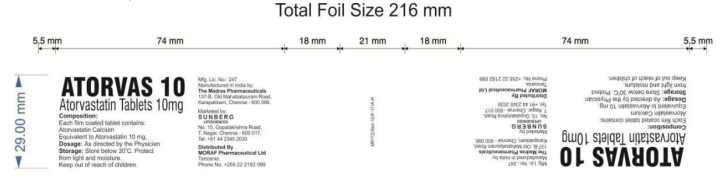
Type of feedback	Impact	Response

Re-registration applications NA

PART 5: CHANGE HISTORY

Version number	Date	Description of update	Section(s) Modified	Approval date

Annex I: Mock up label



ATORVAS 10 Atorvastatin Tablets 10mg

Composition:

Each film coated tablet contains: Atorvastatin Calcium Equivalent to Atorvastatin 10 mg. **Dosage:** As directed by the Physician **Storage:** Store below 30°C. Protect from light and moisture. Keep out of reach of children.

200%

Mfg. Lic. No.: 247 Manufactured in India by: **The Madras Pharmaceuticals** 137-B, Old Mahabalipuram Road, Karapakkam, Chennai - 600 096.

Marketed by: SUNBERG LIFESCIENCES

No. 15, Gopalakrishna Road, T. Nagar, Chennai - 600 017. Tel: +91 44 2345 2030

Distributed By MORAF Pharmaceutical Ltd Tanzania. Phone No. +255 22 2182 099



LBH - 108x22x47 mm