

TMDA/DMD/MCIE/F/002
REV.# 01



THE UNITED REPUBLIC OF TANZANIA
MINISTRY OF HEALTH



TANZANIA MEDICINES AND MEDICAL DEVICES AUTHORITY

CORDEN PHARMA GmbH, PLANKSTADT, GERMANY
PUBLIC GMP DESK ASSESSMENT REPORT

Date: March, 2025



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Part 1: General information about the company

1.1 Manufacturer's details	
Name of manufacturer/ Applicant	Corden Pharma GmbH - Germany/ Astrazeneca PLC – United Kingdom
1.2 Inspected site Details	
Name & physical address of inspected manufacturing site	Name: Corden Pharma GmbH Physical address of the site: Otto-Hahn-Strasse 1, Plankstadt, Baden, Wuerttemberg, 68723, Germany 24hrs contact; +49 6202 99-1166
Name of Unit/ block/ workshop number inspected	The company has three (3) buildings for the production of bulk products; Buildings 1, 2, and 3 whereas facility 3 is dedicated for the manufacturing of veterinary products
1.3 Inspection details	
Date of desk review	23 rd July, 2024
Date of last inspection by the SRA, WHO-PQ or EAC / SADC for production line applied at TMDA	This facility was inspected on 12-06-2024 by <i>Regierungspraesidium Tuebingen Leitstelle Arzneimittelueberwachung Baden - Wuerttemberg</i> by a competent authority of Germany, GMP Certificate No. DE_BW_01_GMP_2024_0174 (Human medicinal products) and DE_BW_01_GMP_2024_0175 (veterinary products) was issued which is valid for three years from date of inspection. These certificates can be traced in the EudraGMP database
1.4 Brief report of the activities undertaken at the site	
Summary of the activities performed at the site	The facility is involved in • Manufacturing, packaging and batch certification of general pharmaceutical

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	<p>products for human use and veterinary medicinal products inform of non-sterile solid dosage form (tablets, capsules); special requirements for medicinal products (OSD) with high potent active substances</p> <ul style="list-style-type: none"> Quality control testing (physical/chemical and microbiology: non-sterility) of general pharmaceutical products for human use and veterinary medicinal products inform of non-sterile solid dosage form (tablets, capsules)
Production lines applied at TMDA	Hormonal products for human use in the form of tablets

Part 2: Review of submitted documentary evidence

2.1. Site master file (*describe consistency of SMF as per requirements*)
Contents and layout of Site master file No. 03 B43 SOP003034 41 presented complied with requirements stipulated in TFDA GMP Regulations, 2018

2.2. Provide list of all regulatory inspections carried out in the past three years.

This facility was inspected on 12-06-2024 by *Regierungspraesidium Tuebingen Leitstelle Arzneimittelueberwachung Baden - Wuerttemberg* by a competent authority of Germany, GMP Certificate No. DE_BW_01_GMP_2024_0174 (Human medicinal products) and DE_BW_01_GMP_2024_0175 (veterinary products) was issued which is valid for three years from date of inspection. This certificate can be traced in the EudraGMP database

2.3. Manufacturing license and GMP permit granted by the local National Medicines Regulatory Authority (NMRA).

This facility was inspected on 12-06-2024 by *Regierungspraesidium Tuebingen Leitstelle Arzneimittelueberwachung Baden - Wuerttemberg* by a competent authority of Germany, GMP Certificate No. DE_BW_01_GMP_2024_0174 (Human medicinal products) and DE_BW_01_GMP_2024_0175 (veterinary products) was issued which is valid for three years from date of inspection. This certificate can be traced in the EudraGMP database



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- 2.4. Valid GMP certificate issued by WHO listed authority and/or that from WHO prequalification and Regional Harmonization Initiatives/AMA (whichever is applicable) for inspection carried out within the past three years for production line(s) applied at TMDA.

Not applicable, see section 2.2 of this report

- 2.4.1. Name of SRA/WHO-PQ/RECs

Regierungspraesidium Tuebingen Leitstelle Arzneimittelueberwachung Baden – Wuerttemberg

- 2.4.2. Dates of inspection
23rd June, 2024

- 2.4.3. Scope of inspection / List of compliant production line

Hormonal products for human use in the form of tablets

- 2.4.4. A confirmation by the senior QA representative that a full WLA audit covering the product(s) has been performed and all matters dealt with and attest to the authenticity of the information

A confirmation by senior QA representative that a full SRA audit covering casodex production line has been performed and that all matters have been dealt with and attest to the authenticity of the information, was not provided. However, this may be waived as the GMP compliance of the site has been confirmed by the GMP certificate accessed from Eudra GMP website

- 2.5. Regulatory Actions against the facility that were taken in the past three (3) years.

The facility confirmed that there were no high-risk market complaints for the past three years, it was also noted that there are no complaints received from the Tanzania Market for the products as well for the past three years.

- 2.6. PQR(s) of the concerned product(s) (If products have not been registered).

This is waived because the product of concern has been registered by TMDA, this implies that PQR was assessed during the dossier assessment

- 2.7. Real time and Accelerated Stability studies under Zone IVb conditions (*If products have not been registered*).



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This is waived because the product of concern has been registered by TMDA, this implies that PQR was assessed during the dossier assessment

2.8. Review of Aseptic processing and filling validation protocols and reports (*for sterile products*).

This is waived because the manufacturing facility applied for and was GMP certified by SRA for manufacturing of hormonal products for human use in the form of tablets

2.9. Review of validation master plan; policy on validation qualification and calibration (If no product has been registered).

This is waived because the product of concern has been registered by TMDA, this implies that PQR was assessed during the dossier assessment

2.10. Market complaints in the last three years for products applied at TMDA

The facility confirmed that there were no high-risk market complaints for the past three years, it was also noted that there are no complaints received from the Tanzania Market for the products as well for the past three years.

Part 3: Conclusion

Based on the desk assessment and evidence(s) provided Corden Pharma GmbH, Otto-Hahn-Strasse 1, Plankstadt, Baden, Wuerttemberg, 68723, Germany is considered to be operating at an acceptable level of compliance with the requirements of the Tanzania Food, Drugs and Cosmetics (Good Manufacturing Practice Enforcement) Regulations, 2018 for manufacturing of hormonal products for human use in the form of tablets.

This TPIR will remain valid until 22nd July, 2027, provided that the facility will remain compliant following any inspections conducted in the period.