

***SUMMARY OF AXIS CLINICALS LIMITED AND REGULATORY  
INSPECTIONS AND APPROVALS  
(November - 2023)***

**INTRODUCTION:**

AXIS Clinicals Limited is a Contract Research Organization for clinical research to support bioequivalence studies and Phase Trials. AXIS Clinicals Limited offers services for Medical writing, Study site support, Bioanalysis, Data management and Statistical services as a one stop full time global CRO to serve the pharmaceutical industry by keeping pace with latest techniques, procedures by ensuring scientific validity of all research projects and complying with global regulations.

AXIS Clinicals Limited is headquartered in Hyderabad, India with operations and facilities in China, Mexico and USA to support the global sponsors. In India, AXIS is having 2 facilities to support bioequivalence studies. Further details were provided under the Table “Legal Identity of the facilities”.

**Key Features:**

- AXIS Clinicals Limited has a total of 9 study areas that have an overall capacity of 330 beds.
- The bioanalytical lab located at head office has equipped with 26 LC-MS/MS and 2 ICP-OES and 1 ICP-MS to support bioanalytical services.
- Pharmacokinetic analysis and Biostatistical analysis of data will be carried out by using WinNonlin and SAS software's.
- A Clinical Research (CR) and Clinical Data Management (CDM) departments will support phase trails in hospitals and Data management by using Oracle clinical.
- AXIS Clinical Reference Laboratory is approved by NABL as per ISO 15189:2012 and accredited.
- AXIS Clinicals Limited has an Independent Quality Assurance Team and Common Quality Management System across all facilities to have uniform data / conduct of the studies / Documentation / final report preparation.

AXIS Clinicals Limited formerly known as Trident Life Sciences Limited and the scope of the change is only name of the company and has been approved by the Registrar of Companies, Government of India and regulatory agencies like Drug Controller General of India (DCGI) etc.

**LEGAL IDENTITY OF THE FACILITIES:**

Facility	Address	Scope	Bed capacity	USFDA FEI & DUNS Number
Head office (ACL-01)	AXIS Clinicals Limited 1-121/1, Survey No. 66 & 67 (Part), Miyapur, Serilingampally, Hyderabad, Telangana, 500049, India.	Clinical, Clinical Reference Lab, Bioanalytical, Pharmacokinetic and Statistical services and Phase trails (Clinical Research), Clinical Data Management (CDM).	192 beds in 4 study areas with bed capacities 62,58,36, 36	FEI No: 3006649033, DUNS No: 86-346-1407
ACL-02  (Operations at this unit were closed from February- 2017)	AXIS Clinicals Limited, Plot No. 33 to 35, Alluri Sitaramaraju Nagar, Opp. J.P.N Nagar, Miyapur, Hyderabad, Telangana, 500049, India.	Clinical Phase	128 bed capacities in 4 study areas with bed capacities of 28,36,28 and 36	FEI No: 3010297993, DUNS No: 65-062-3114
ACL-03	AXIS Clinicals Limited, Atharva, Opp. Rajpath club, S.G highway, Bodakdev, Ahmedabad- 380054, Gujarat- India	Clinical Phase	138 bed capacities in 5 study areas with bed capacities of 36,15,15, 36 and 36.	FEI No: 3012359023, DUNS No: 65-100-6277

**Note:** The bio-analytical, Pharmacokinetic and biostatistical services will be supported from head office (ACL 01) located in Hyderabad for all the facilities (ACL-01, ACL-02 & ACL-03).

**REGULATORY INSPECTIONS AND ACCREDITATIONS:**

AXIS Clinicals Limited conducted bioequivalence studies to support marketing authorization for various regulated markets including US FDA, UK-MHRA, AEMPS-Spain, INFARMED-Portugal, ANSM-France, Turkey-MOH, WHO, DCGI, Health Canada, Thai-MOPH, NPRA-Malaysia, GCC Gulf, MCC-South Africa, MHSD-RoK Kazakhstan etc. for various sponsors. Our facility has been audited and approved by various global regulatory agencies and accreditation bodies and the details were described in the following table.

**US-FDA REGULATORY INSPECTIONS:**

S.No.	Year	Inspection dates	Scope	Facility	Outcome	Receipt of EIR	Remarks
1	2008	05 <sup>th</sup> -14 <sup>th</sup> Mar 2008	All Phases	ACL-01	Form 483 issued	Yes	Nil
2	2008	27 <sup>th</sup> -31 <sup>st</sup> Oct 2008	All Phases	ACL-01	Form 483 issued	Yes	Nil
3	2009	26 <sup>th</sup> -30 <sup>th</sup> Oct 2009	All Phases	ACL-01	No Form 483	Yes	Nil
4	2010	31 <sup>st</sup> May-11 <sup>th</sup> Jun 2010	Clinical Phase	ACL-01	No Form 483	Yes	Nil
5	2011	14 <sup>th</sup> -16 <sup>th</sup> Feb 2011	Clinical Phase	ACL-02	No Form 483	Yes	Nil
6	2011	16 <sup>th</sup> -21 <sup>st</sup> Mar 2011	Bioanalytical	ACL-01	Form 483 issued	Yes	Nil
7	2011	10 <sup>th</sup> -14 <sup>th</sup> Oct 2011	Clinical Phase	ACL-01	No Form 483	Yes	Nil
8	2012	01 <sup>st</sup> -05 <sup>th</sup> Oct 2012	Clinical Phase	ACL-01	Form 483 issued	Yes	Nil
9	2013	06 <sup>th</sup> -08 <sup>th</sup> Aug 2013	All Phases	ACL-01	No Form 483	Yes	Nil
10	2014	10 <sup>th</sup> -20 <sup>th</sup> Mar 2014	Clinical Phase	ACL-02	No Form 483	Yes	Head office address captured in EIR
11	2016	16 <sup>th</sup> -18 <sup>th</sup> Mar 2016	Bioanalytical (GMP) Phase	ACL-01	Form 483 issued	No	Analytical services with ICP-OES
12	2016	06 <sup>th</sup> -10 <sup>th</sup> Jun 2016	Bioanalytical Phase	ACL-01	Form 483 issued	Yes	Nil
13	2016	11 <sup>th</sup> -21 <sup>st</sup> Jul 2016	Clinical Phase	ACL-01	No Form 483	Yes	Nil
14	2016	11 <sup>th</sup> -25 <sup>th</sup> Aug 2016	Clinical Phase	ACL-02	No Form 483	Yes	Nil
15	2018	21 <sup>st</sup> -25 <sup>th</sup> May 2018	Clinical Phase	ACL-03	No Form 483	Yes	Unannounced Inspection
16	2018	27 <sup>th</sup> – 30 <sup>th</sup> Aug 2018	Bioanalytical Phase	ACL-01	No Form 483	Yes	Unannounced Inspection
17	2018	29 <sup>th</sup> Oct – 05 <sup>th</sup> Nov 2018	Clinical Phase	ACL-01	No Form 483	Yes	Nil
18	2022	07 <sup>th</sup> – 31 <sup>st</sup> Mar 2022	Bioanalytical Phase	ACL-01	No Form 483	No	Remote Audit
19	2022	25 <sup>th</sup> – 28 <sup>th</sup> Apr 2022	Clinical Phase	ACL-01	No Form 483	Yes	Close-out letter received.
20	2022	02 <sup>nd</sup> – 05 <sup>th</sup> May 2022	Clinical Phase	ACL-03	No Form 483	Yes	Close-out letter received.

**US-FDA REGULATORY INSPECTIONS – CR STUDIES (PATIENT POPULATION STUDIES):**

S.No.	Year	Inspection dates	Scope / Therapeutic Indication	Facility	Outcome	Receipt of EIR	Remarks
1	2012	28 <sup>th</sup> May to 01 <sup>st</sup> Jun 2012	Clinical Phase / Epilepsy	AXON Hospital, Hyderabad, Telangana, India	No Form 483	No	Nil
2	2012	04 <sup>th</sup> to 08 <sup>th</sup> Jun 2012	Clinical Phase / Epilepsy	St.Theresa's General Hospital, Hyderabad, Telangana, India	Form 483 issued	No	Nil
3	2016	28 <sup>th</sup> Nov to 02 <sup>nd</sup> Dec 2016	Clinical Phase / IBS with Constipation	M.V.Hospital & Research Centre, Lucknow, Uttar Pradesh, India	Form 483 issued	No	Nil
4	2017	13 <sup>th</sup> to 17 <sup>th</sup> Feb 2017	Clinical Phase / IBS with Constipation	Inamdhari Hospital, Pune, Maharashtra, India	No Form 483	No	Form FDA 484 issued
5	2017	27 <sup>th</sup> Feb to 3 <sup>rd</sup> Mar 2017	Clinical Phase / IBS with Constipation	Gandhi Hospital, Hyderabad, Telangana, India	Form 483 issued	No	Nil
6	2017	17 <sup>th</sup> to 21 <sup>st</sup> Apr 2017	Clinical Phase / IBS with Constipation	Samvedana Hospital, Varanasi, Uttar Pradesh, India	No Form 483	No	Form FDA 484 issued
7	2017	06 <sup>th</sup> to 10 <sup>th</sup> Nov 2017	Clinical Phase/ Psoriasis / Rheumatoid Arthritis	Rathi Hospital, Ahmedabad, Gujarat, India	No Form 483	Yes	Form FDA 484 issued
8	2017	13 <sup>th</sup> to 17 <sup>th</sup> Nov 2017	Clinical Phase / Schizophrenia	Ratandeep Multispeciality Hospital, Ahmedabad, Gujarat, India	No Form 483	No	Form FDA 484 issued
9	2019	04 <sup>th</sup> to 08 <sup>th</sup> Feb 2019	Clinical Phase / Iron Deficiency Anemia	KRM Hospital and Research centre, Lucknow, UP-226010, India.	Form 483 issued	No	Nil
10	2019	05 <sup>th</sup> to 07 <sup>th</sup> Mar 2019	Diarrhea caused by Giardia lamblia	The Grant Government Medical College, Mumbai Central, Mumbai, Maharashtra, India	No Form 483	Yes	Nil
11	2019	09 <sup>th</sup> to 19 <sup>th</sup> Jul 2019	Clinical Phase / Iron Deficiency Anemia	Bodyline Hospitals Pvt. Ltd., Ahmedabad, Gujarat, India	Form 483 issued	No	Form FDA 484 issued
12	2019	11 <sup>th</sup> to 14 <sup>th</sup> Sep 2019	Clinical Phase / Schizophrenia	Shri Hathkesh Healthcare Foundation, Junagadh, Gujarat, India	No Form 483	No	Form FDA 484 issued
13	2019	16 <sup>th</sup> to 19 <sup>th</sup> Sep 2019	Clinical Phase / Iron Deficiency Anemia	Baroda Medical College, Vadodara, Gujarat, India	No Form 483	No	Form FDA 484 issued
14	2019	23 <sup>rd</sup> to 26 <sup>th</sup> Sep 2019	Clinical Phase / Iron Deficiency Anemia	Nirmal Hospital Pvt.Ltd, Surat, Gujarat, India	No Form 483	No	Form FDA 484 issued

15	2019	23 <sup>rd</sup> to 27 <sup>th</sup> Sep 2019	Clinical Phase / Iron Deficiency Anemia	ACSR Govt. Medical College, Nellore, Andhra Pradesh, India	Form 483 issued	No	Form FDA 484 issued
16	2019	04 <sup>th</sup> to 08 <sup>th</sup> Nov 2019	Diarrhea caused by Giardia lamblia	Supe Heart and Diabetes Hospital and Research Centre Nashik, Maharashtra, India.	Form 483 issued	No	Nil
17	2020	02 <sup>nd</sup> to 06 <sup>th</sup> Mar 2020	Schizophrenia	Anand Multispeciality Hospital and Research Centre, Gandhi Nagar, Gujarat, India	Form 483 issued	No	Form FDA 484 issued
18	2021	05 <sup>th</sup> Mar 2021 to 02 <sup>nd</sup> Apr 2021	Asthma	Lotus Multispecialty Hospital Beside Swastik School, Opposite Baliyadev Temple, Motera Stadium Road Ahmedabad, Gujrat, India	Virtual inspection for study related document verification		
19	2021	14 <sup>th</sup> Mar 2021 to 23 <sup>rd</sup> Apr 2021	Asthma	Shree Hospital, S. No. 7/3/B Gulmohor Society Kharadi, Pune 411014, Maharashtra, India	Virtual inspection for study related document verification		
20	2022	18 <sup>th</sup> to 22 <sup>nd</sup> Apr 2022	Vulvar and Vaginal atrophy	Anand Surgical Hospital Pvt., Ltd. Memco Cross Road, Naroda Road, Ahmedabad, Gujarat, India	No Form 483	No	Nil
21	2022	29 <sup>th</sup> Aug to 02 <sup>nd</sup> Sep 2022	Vulvar and Vaginal atrophy	Ojas Multispecialty Hospital, Pune, Maharashtra, India-412101	No Form 483	No	Nil

**OTHER REGULATORY INSPECTIONS:**

S.No.	Year	Inspection dates	Scope	Facility	Outcome	Remarks
<b>ANVISA- Brazil</b>						
1	2007	06 <sup>th</sup> -10 <sup>th</sup> Aug 2007	All Phases	ACL-01	Approved	Nil
2	2009	12 <sup>th</sup> -16 <sup>th</sup> Oct 2009	All Phases	ACL-01	Approved	Nil
3	2011	05 <sup>th</sup> -06 <sup>th</sup> Aug 2011	All Phases	ACL-01	Approved	Nil
4	2012	17 <sup>th</sup> -19 <sup>th</sup> Oct 2012	All Phases	ACL-01	Approved	Nil
5	2013	05 <sup>th</sup> -08 <sup>th</sup> Aug 2013	All Phases	ACL-01	Approved	Nil
6	2014	02 <sup>nd</sup> -04 <sup>th</sup> Jun 2014	All Phases	ACL-01	Approved	Nil
7	2015	02 <sup>nd</sup> -04 <sup>th</sup> Mar 2015	All Phases	ACL-01	Approved	Nil
8	2017	07 <sup>th</sup> – 11 <sup>th</sup> Aug 2017	All Phases	ACL-01	Approved	Validity till 03 <sup>rd</sup> Jan 2019
<b>UK-MHRA</b>						
1	2009	24 <sup>th</sup> -27 <sup>th</sup> Feb 2009	All Phases	ACL-01	Approved	GCP Certificate Issued
2	2011	17 <sup>th</sup> -18 <sup>th</sup> Oct 2011	Clinical Phase	ACL-01	Approved	Certificate Issued
3	2014	01 <sup>st</sup> -05 <sup>th</sup> Sep 2014	All Phases	ACL-01	Approved	Nil
4	2018	04 <sup>th</sup> – 08 <sup>th</sup> Jun 2018	All Phases	ACL-01	Approved	Inspection includes the review of Patient Population study.
5	2021	29 <sup>th</sup> Nov – 08 <sup>th</sup> Dec 2021	All Phases	ACL-01 & ACL-03	Approved	Remote Audit.



**OTHER REGULATORY INSPECTIONS:**

S.No.	Year	Inspection dates	Scope	Facility	Outcome	Remarks
<b>ANSM-France</b>						
1	2009	02 <sup>nd</sup> -03 <sup>rd</sup> Jun 2009	Clinical Phase	ACL-01	Accepted Compliance	Nil
<b>MOH-Turkey</b>						
1	2011	18 <sup>th</sup> -19 <sup>th</sup> Dec 2011	All Phases	ACL-01	Approved	GCP Certificate issued
<b>Thailand-BLQS-GLP Inspection</b>						
1	2014	21 <sup>st</sup> -25 <sup>th</sup> Jan 2014	Bioanalytical	ACL-01	Approved	GLP Certificate issued
<b>WHO</b>						
1	2015	17 <sup>th</sup> -20 <sup>th</sup> Mar 2015	All Phases	ACL-01	Approved	Nil
2	2017	18 <sup>th</sup> -21 <sup>st</sup> Dec 2017	All Phases	ACL-01	Approved	Nil
<b>GCC-DR (Gulf Central Committee for Drug Registration; Member States: Bahrain, Kuwait, Oman, Qatar, Saudi Arabia, United Arab Emirates)</b>						
1	2016	15 <sup>th</sup> -16 <sup>th</sup> Feb 2016	All Phases	ACL-01 & 02	Approved	Nil
<b>MCC (Medicines Control Council) – South Africa</b>						
1	2017	25 <sup>th</sup> – 27 <sup>th</sup> Mar 2017	Clinical Phase	ACL-01	Approved	Nil
<b>NPRA (National Pharmaceutical Regulatory Agency) – Malaysia</b>						
1	2018	18 <sup>th</sup> – 22 <sup>nd</sup> Jun 2018	All Phases	ACL-01	Approved	Validity till 20 <sup>th</sup> Sep 2023
2	2023	22 <sup>nd</sup> - 26 <sup>th</sup> May 2023	All Phases	ACL-01	Successfully Completed	Nil
<b>INFARMED (Portuguese National Authority of Medicines and Health Products)</b>						
1	2018	18 <sup>th</sup> – 20 <sup>th</sup> Jun 2018	Clinical Phase	ACL-03	Approved	Nil
<b>AEMPS (Agencia Española de Medicamentos y Productos Sanitarios) – Spain</b>						
1	2018	17 <sup>th</sup> – 21 <sup>st</sup> Dec 2018	Clinical Phase	ACL-03	Approved	Nil

**DCGI INSPECTIONS:**

S.No.	Year	Inspection dates	Scope	Facility	Approval Date	Remarks
1	2006	16 <sup>th</sup> Dec 2006	All phases	ACL-01&02	08 <sup>th</sup> Feb 2008	Approved
2	2010	13 <sup>th</sup> -14 <sup>th</sup> Jul 2010	All phases	ACL-01&02	Nil	Compliance submitted & accepted
3	2013	18 <sup>th</sup> -20 <sup>th</sup> Sep 2013	All phases	ACL-01&02	11 <sup>th</sup> Nov 2014	Approved
4	2015	16 <sup>th</sup> -17 <sup>th</sup> Oct 2015	All phases	ACL-01&02	16 <sup>th</sup> Mar 2016	Approved
5	2016	19 <sup>th</sup> Apr 2016	Clinical Phase	ACL-03	01 <sup>st</sup> Aug 2016	Approved
6	2016	22 <sup>nd</sup> -23 <sup>rd</sup> Aug 2016	All phases	ACL-01	Nil	Nil
7	2017	08 <sup>th</sup> Jun 2017	All phases	ACL-01	09 <sup>th</sup> Aug 2017	Approved (Validity till 08 <sup>th</sup> Aug 2020)
8	2019	17 <sup>th</sup> Jul 2019	Clinical Phase	ACL-03	12 <sup>th</sup> Sep 2019	Approved (Validity till 11 <sup>th</sup> Sep 2024)
9	2020	03 <sup>rd</sup> to 04 <sup>th</sup> Jun 2020	All phases	ACL-01	23 <sup>rd</sup> Jul 2020	Approved (Validity till 22 <sup>nd</sup> Jul 2025)

**Note:** Details of DCGI inspections for facility approval for conducting bioequivalence studies and clinical research with a notification/ inspection report/approval were listed here.

**ACCREDITATIONS:**

NABL (National Accreditation Board for Testing and Calibration Laboratories)						
S.No.	Year	Inspection Dates	Type of Visit/Audit	Facility	Outcome	Remarks
1	2006	27 <sup>th</sup> Dec 2006	Pre-Assessment	ACL-01	Pre-Assessment Approved	Nil
2	2007	15 <sup>th</sup> -16 <sup>th</sup> Jan 2007	Assessment	ACL-01	Approved	Nil
3	2008	28 <sup>th</sup> -29 <sup>th</sup> Jun 2008	Surveillance	ACL-01	Approved	Nil
4	2009	07 <sup>th</sup> -08 <sup>th</sup> Mar 2009	Reassessment	ACL-01	Approved	Nil
5	2010	Aug 2010	Desk top audit	ACL-01	Approved	Nil
6	2011	11 <sup>th</sup> -12 <sup>th</sup> Jun 2011	Reassessment	ACL-01	Approved	Nil
7	2012	Aug 2012	Desk top audit	ACL-01	Approved	Nil
8	2013	11 <sup>th</sup> -12 <sup>th</sup> May 2013	Reassessment	ACL-01	Approved	Nil
9	2014	Jul 2014	Desk top audit	ACL-01	Approved	Nil
10	2015	30 <sup>th</sup> -31 <sup>st</sup> May 2015	Reassessment	ACL-01	Approved	Nil
11	2016	01 <sup>st</sup> Aug 2016	Desk top audit	ACL-01	Approved	Nil
12	2017	27 <sup>th</sup> & 28 <sup>th</sup> May 2017	Reassessment	ACL-01	Approved	Nil
13	2018	21 <sup>st</sup> Jun 2018	Desk top audit	ACL-01	Approved	Nil
14	2019	14 <sup>th</sup> & 15 <sup>th</sup> May 2019	Reassessment	ACL-01	Successfully Completed	Nil
15	2020	25 <sup>th</sup> Jul 2020	Desk top audit	ACL-01	Approved	Nil
16	2021	10 <sup>th</sup> & 11 <sup>th</sup> Jul, 2021	Reassessment	ACL-01	Approved	Nil
17	2022	Aug 2022	Desk top audit	ACL-01	Approved	Nil