
CLINICAL RESEARCH / HEALTHCARE EDUCATION, TRAINING & CONSULTING SERVICES

www.cliniminds.com

Vision

**TO ESTABLISH WORLDCLASS INSTITUTION TO
OFFER WIDEST RANGE OF HIGH QUALITY
CLINICAL RESEARCH, HEALTHCARE AND
PHARMACEUTICAL PROGRAMS AND BUSINESS
SOLUTIONS**

Cliniminds Background

- Tenet Health Edutech Pvt. Ltd. was established in 2004 by a team of professionals from Clinical Research, Healthcare & Pharma industry.
- Cliniminds offers wide range of educational programs and training solutions for the clinical research, pharma and healthcare industry..
- At present we offer over 23 educational and professional training programs in pharma and clinical research using classroom; online and corporate workshops mediums. Programs are backed up by the high quality copyright content and faculty with global perspective.
- Cliniminds is Headquartered at Delhi, and have branches in Bangalore, Hyderabad, Mumbai, Ahmedabad, Cochin, Bhopal and Vijaywada. New branches are being set up in other cities as well.
- Also offer E-Learning solutions for the healthcare & pharma companies and CROs.

Key Activities

- Classroom Professional Educational Programs in the field of Clinical Research & Pharma in India – Diploma & Certificate Programs.
- Online/Distance Learning – Global market
- Corporate Training Solutions
- E-Learning Solutions for Corporates
- Consulting Services
- Medical & Scientific Writing

OBJECTIVE OF PRESENTATION

- Objective of this presentation is to seek International Certification / Accreditation or Co-Certification of Cliniminds' Clinical Research training programs.
- Offer some of your clinical research and pharma industry training and educational programs in India.
- Explore other options of some of Cliniminds programs at your institution.

BACKGROUND

- ⊕ Clinical research jobs are amongst most sought after jobs segment in the healthcare industry with over 40% new top job offerings
- ⊕ Major gap in Demand & Supply of trained manpower – An unmet need
- ⊕ With the shift of work to India; expected market size of \$2 billion and entry of new players, over 50,000 professionals will be required in 2012.
- ⊕ Current demand is over 12,000 professionals per year.
- ⊕ Example : Qunitiles Corp, USA have alone hired over 2,500 new employees in the last 3 years.

- ⊕ The current facilities are able to train only 4,000 professionals.
- ⊕ Employers required well trained professionals.
- ⊕ Attractive Salaries Offered & High Annual Salary Growth.

OPPORTUNITY

⊕ Global Clinical Research Training Size	\$2 billion
⊕ Market Size in 2005-06	\$200 million
⊕ Estimated Market Size in 2009	\$600 million
⊕ 2012 Projected Industry Spending on CRO Services and Investigator Grants	\$2 billion
⊕ Number of CROs - current	80
⊕ Number of MNC & Indian Pharmacos in Clinical Research	70
⊕ Clinical Research Market Growth (Annual)	40%
⊕ Full time & Site Staff Required in 2012	50,000
⊕ New Protocols received by Indian Regulator everyday	20

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- ⊕ Patients / Subjects Required in 2012 350,000
 - ⊕ Over 3,000 new Certified Investigators will be required in 2008 and over 12,000 by 2012

INDIA HUB OF GLOBAL CLINICAL RESEARCH

- ⊕ India is one of the top 3 countries where companies plan to spend the most R&D dollars over the next 3 years.
- ⊕ Favoured destination ahead of countries like Israel, Philippines, Canada, China, Ireland & Russia in terms of Overall Climate (*Gartner Report, January 2003*)
- ⊕ Powerhouse in R&D (*e.g. GVK Wyeth Global R&D Deal*)
- ⊕ Over 60 CROs offer Phase I to IV trials complying with ICH-GCP guidelines.
- ⊕ Over US\$500 million foreign investment expected in the next 18 months
- ⊕ Over 200 hospitals serving as sites for clinical trials, India is emerging as one of the fastest recruiter of subjects across the world.
- ⊕ Some of the top medical/technical universities in Asia

INDIA HUB OF CLINICAL OF CLINICAL RESEARCH

- ⊕ The clinical community is populated with English speaking, western-trained graduates
- ⊕ Sophisticated technological infrastructure
- ⊕ 100 million plus English speaking / trained professionals (Largest outside US)
- ⊕ Over 2 million science post graduates and growing...
- ⊕ Large pool of treatment naïve patients from multiethnic and multiracial backgrounds
- ⊕ Better patient recruitment, retention and compliance
- ⊕ Participants generally benefit, as the trials conducted in India, mostly in phase II - IV, provide improved care and COST savings as procedures and drugs are provided at no charge

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- ⊕ Cost effective operations
 - ⊕ Higher GMP/GLP/GCP Compliance
 - ⊕ Maximum number of approved GMP plants outside USA
 - ⊕ Excellent quality management, Technology and infrastructure
 - ⊕ Increasing presence of all Pharma majors, CROs & also in-house CROs set up by leading pharma companies
 - ⊕ Strong IT industry availability of IT skilled manpower

INDIA'S VITAL CLINICAL STATISTICS

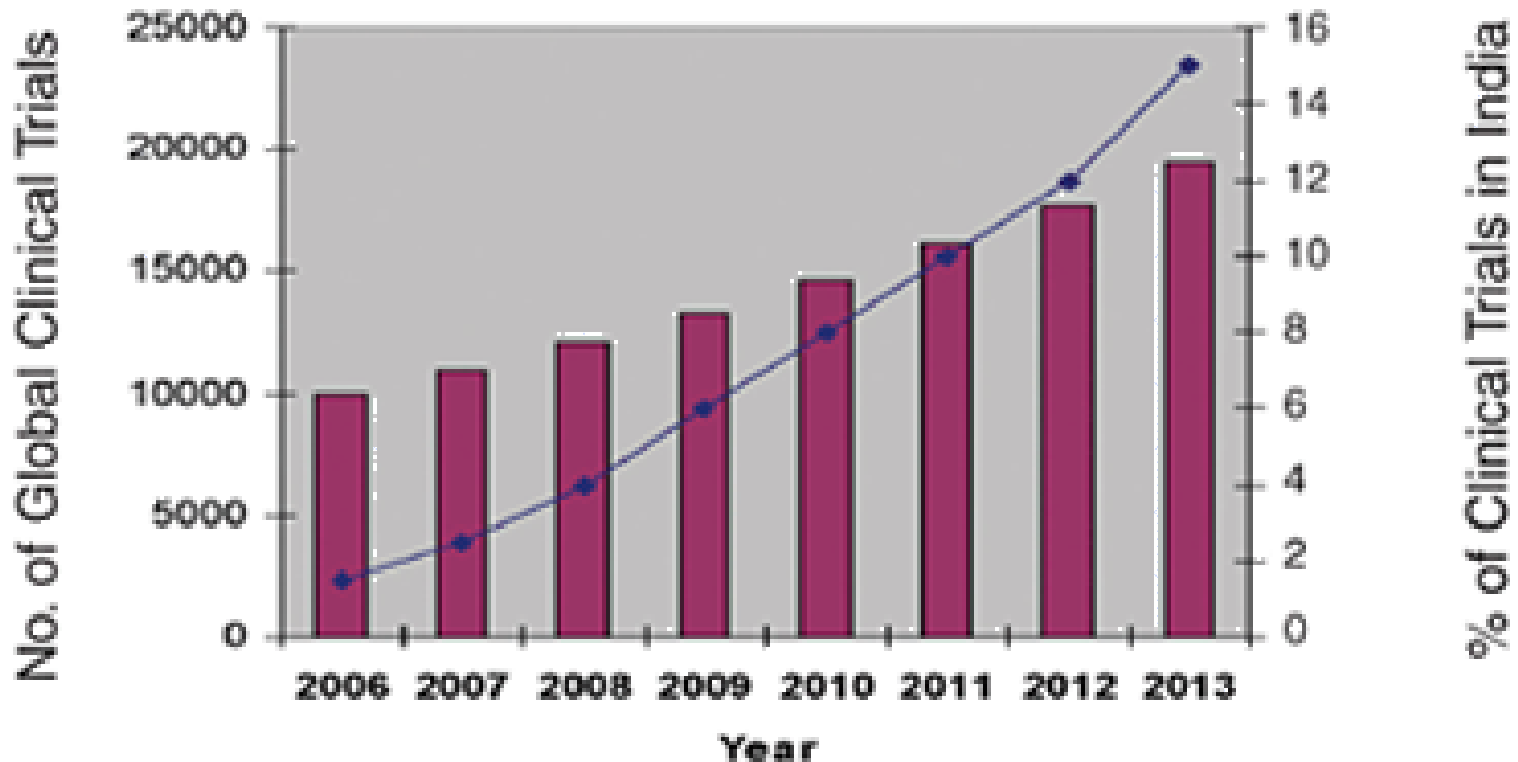
- ⊕ Cancer: 3 million
- ⊕ Diabetes: 34 million
- ⊕ HIV: 8-10 million
- ⊕ Epilepsy: 8 million
- ⊕ Hypertension: 150 million
- ⊕ Schizophrenia: 1 million
- ⊕ Asthma: 40 million
- ⊕ Alzheimer's: 1.5 million
- ⊕ Cardiac-Related Deaths: 2 million
- ⊕ Recruits for genetic studies
- ⊕ 600,000 practicing physicians
- ⊕ 14,000 hospitals
- ⊕ 700,000 beds
- ⊕ 17,000 medical graduates per year

REGULATORY FRAMEWORK

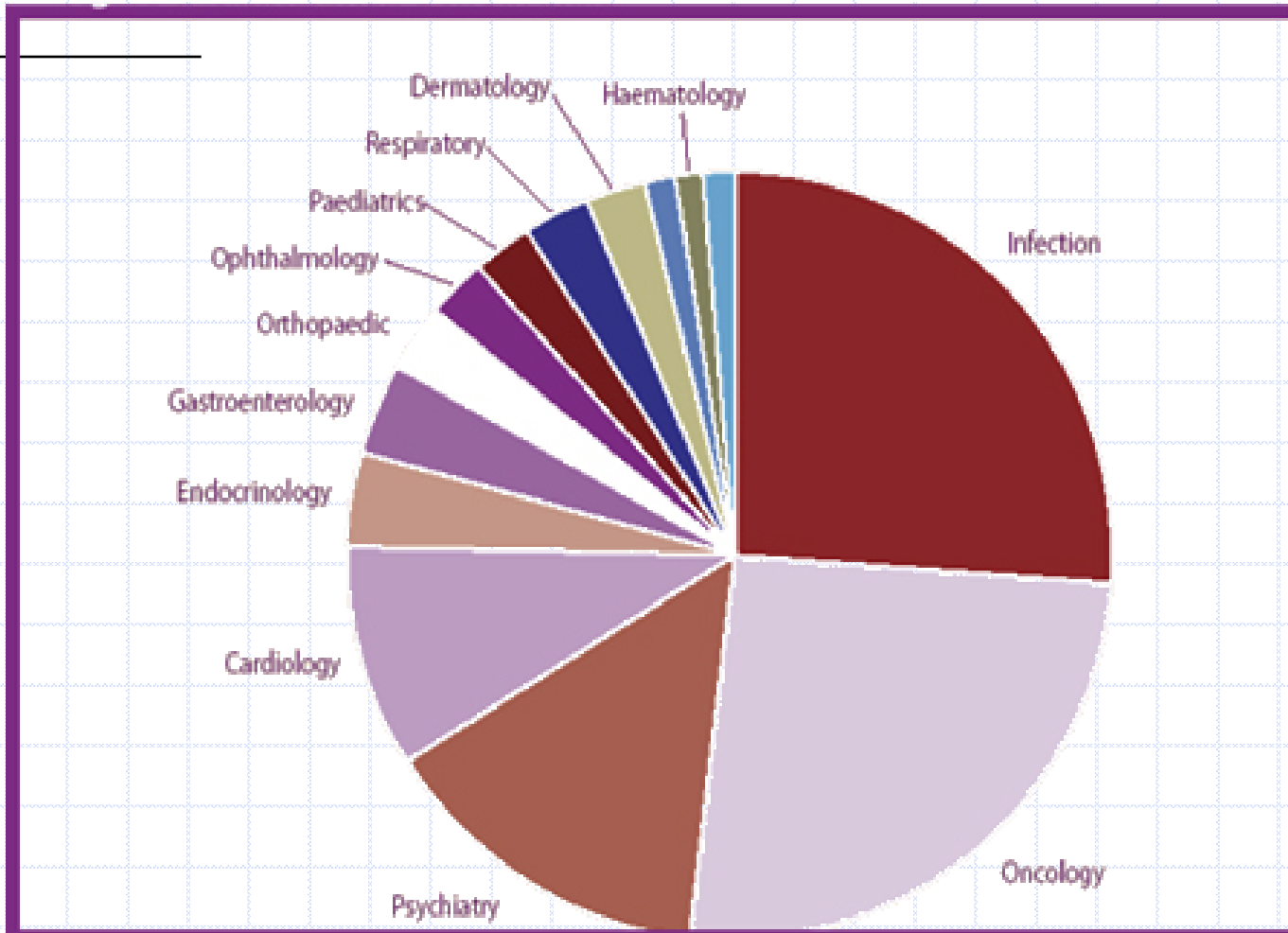
- ⊕ Positive Regulatory Environment – Protocols Approved by DCGI / Schedule Y
- ⊕ CDSCO (Central Drugs Standard Control Organisation) to regulate Clinical Research
- ⊕ Further strengthening of environment by setting up National Drug Authority
- ⊕ Intellectual property protection
 - As of January 2005, recognize product patents from 1995 to present
- ⊕ Clinical Trial Protocol Approval Time is reducing
- ⊕ Duty Free Import of Clinical Trial Supplies
- ⊕ Easier Drug Importation Procedure
- ⊕ ICMR Guidelines on the Safety of Human Subjects
- ⊕ **USFDA OFFICE IN INDIA**

Percent of Global Trials in India

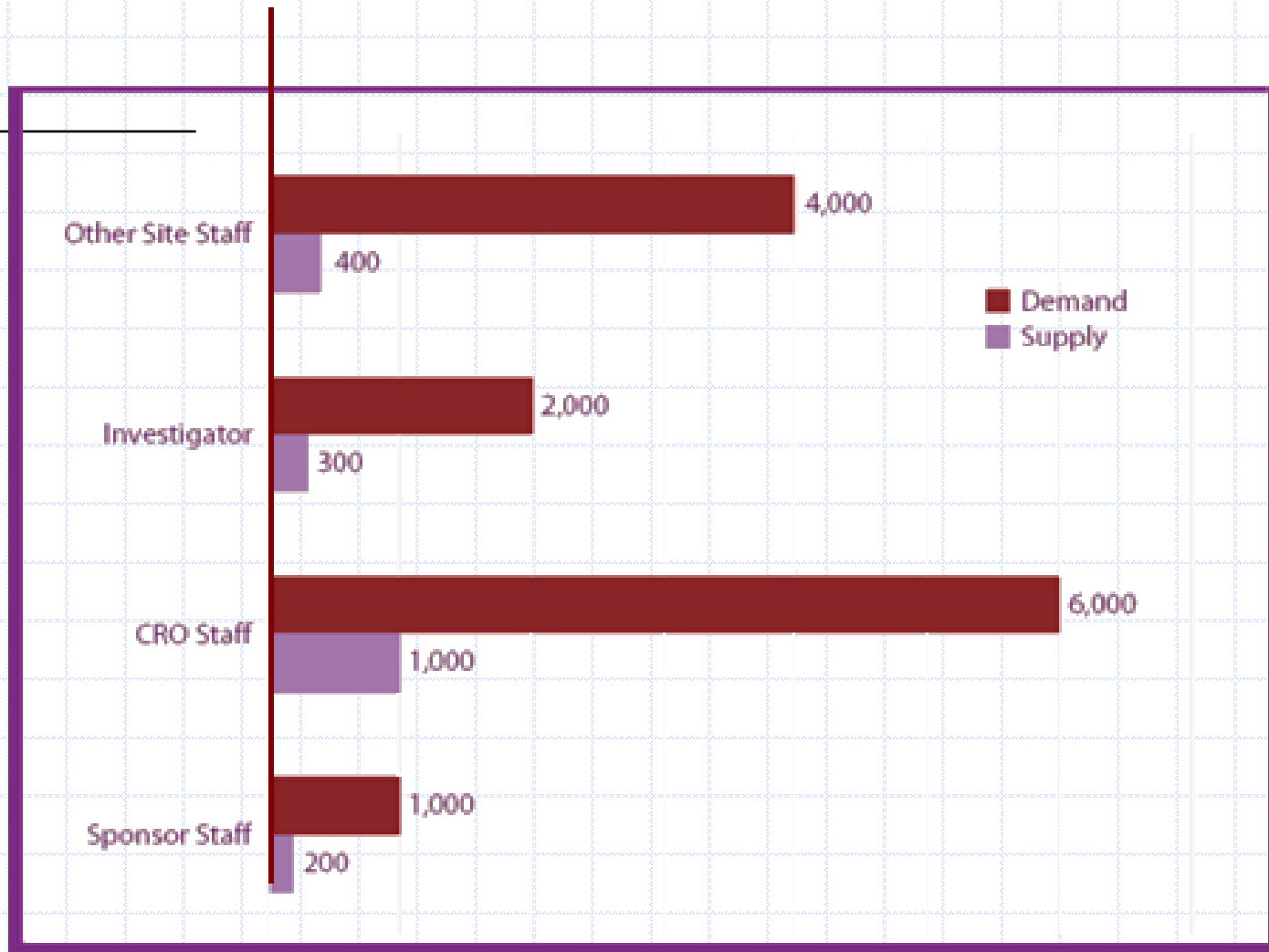
Number of Global Trials and % In India



Clinical Trials Done in India



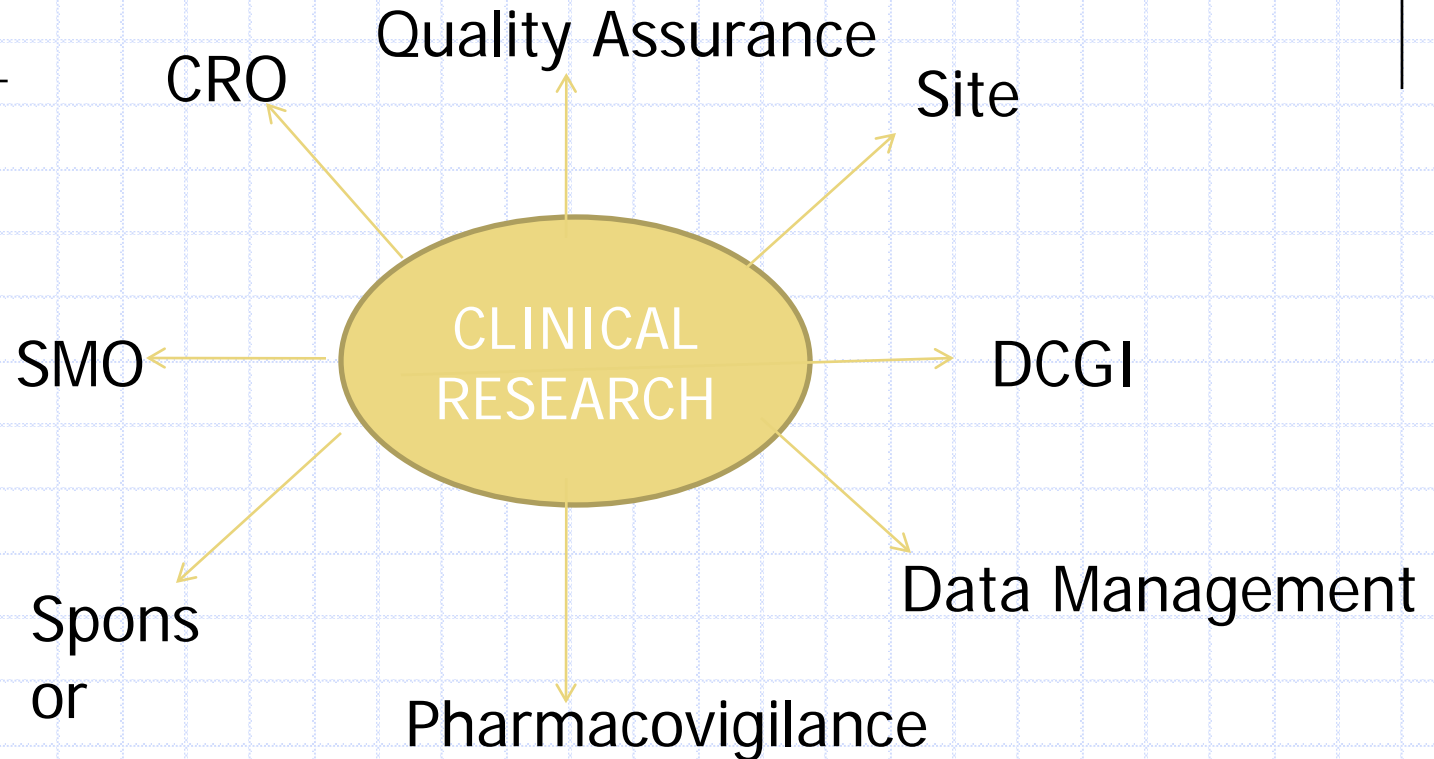
Demand Supply Gap in India



CR Career Pathway

- ⊕ Pharmaceutical Companies
- ⊕ Clinical CROs (Contract Research Organizations)
- ⊕ BA/BE Centers
- ⊕ SMOs (Site Management Organizations)
- ⊕ Data Management CROs
- ⊕ IT Companies in Healthcare / Clinical Domain
 - EDC Service Providers
 - Central Laboratories
 - Packaging & Labeling & Contract Manufacturers
 - Investigator & Site Staff
 - Training Centers.

Functions in Career Pathways



Career in CROs / Pharma / Biotech Companies

- ⊕ Clinical Trial Assistant (CTA)
- ⊕ Clinical Research Associate (CRA)
- ⊕ Senior CRA
- ⊕ Clinical Team Leader
- ⊕ Project Manager
- ⊕ Senior Project Manager
- ⊕ Manager Medical & Regulatory
- ⊕ Manager –Safety / Patents
- ⊕ Manager Quality Assurance
- ⊕ Medical Director
- ⊕ Associate Director –Clinical
- ⊕ Associate Director –Projects
- ⊕ Director –Business Development
- ⊕ Director / Head (Clinical Operations)
- ⊕ General Manger / CEO / President



Phase I / II / III / IV Trial
Project Management
Drug Development Planning
Monitoring
Source Data Verification
Safety Reporting
Regulatory Approval
QA Audits
Business Development

Career in SMO

- 
- ⊕ Clinical Research Coordinators (CRC) / Study Coordinators
 - ⊕ Principal Investigators / Co-Investigators
 - ⊕ Medical Monitors
 - ⊕ Project Manager / Senior Project Manager
 - ⊕ Manager Medical & Regulatory
 - ⊕ Manager Quality Assurance
 - ⊕ Manager –Business Development
 - ⊕ Medical Director
 - ⊕ Associate Director –Clinical
 - ⊕ Associate Director –Projects
 - ⊕ Director / Head (Clinical Operations)
 - ⊕ General Manager / CEO / President

Career opportunities in DM

- 
- ⊕ Data Entry Operator
 - ⊕ Data Validation Executive
 - ⊕ QA Executive
 - ⊕ Data Manager
 - ⊕ QA Manager
 - ⊕ Statistical Programmer
 - ⊕ Statistician
 - ⊕ Data Reviewer
 - ⊕ Data Base Designer
 - ⊕ Medical Writer
 - ⊕ Head –Data Management

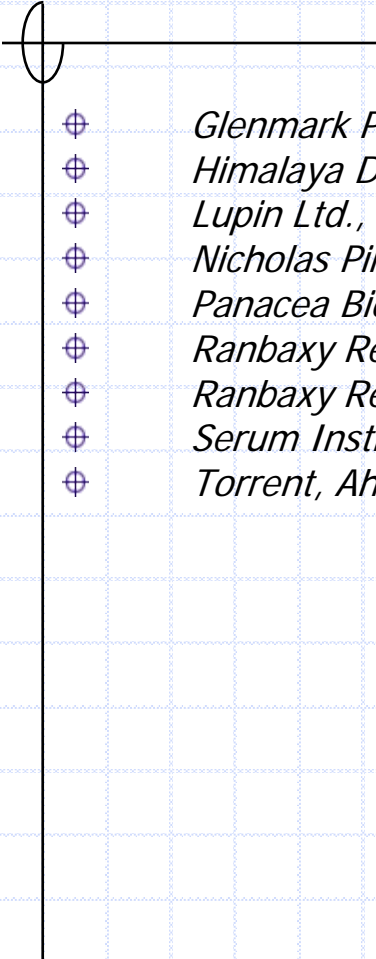
Key Cities in India for Clinical Research

- ⊕ Delhi & NCR Region
- ⊕ Mumbai
- ⊕ Pune
- ⊕ Ahmedabad
- ⊕ Vadodara
- ⊕ Hyderabad
- ⊕ Bangalore
- ⊕ Chennai
- ⊕ Cochin
- ⊕ Trivandrum
- ⊕ Chandigarh
- ⊕ Bhopal
- ⊕ Indore
- ⊕ Coimbatore
- ⊕ Visakhapatnam

KEY PHARMA COMPANIES IN CLINICAL RESEARCH

- ⊕ *Abbott, Mumbai*
- ⊕ *Chiron, Mumbai*
- ⊕ *Astra Zeneca Pharma India Ltd, Bangalore*
- ⊕ *Astra Zeneca Foundation, Bangalore*
- ⊕ *AventisPasteur, Delhi*
- ⊕ *Pfizer Ltd, Mumbai*
- ⊕ *Pfizer Biometrics, Mumbai*
- ⊕ *Altana(Zydus), Mumbai*
- ⊕ *Eli Lilly, Delhi*
- ⊕ *Boston Scientific, Delhi*
- ⊕ *Hospira, Delhi*
- ⊕ *Merck ,Delhi*
- ⊕ *Sanofi Aventis Syntho Lab, Mumbai*
- ⊕ *GSK, Glaxo SmithKline Pharmaceuticals Ltd, Mumbai*
- ⊕ *Novartis International Clinical Development Center, Mumbai*
- ⊕ *Novartis Pharma, Mumbai*
- ⊕ *Roche, Mumbai*
- ⊕ *Sandoz, Mumbai*
- ⊕ *Wyeth, Mumbai*
- ⊕ *BMS, Mumbai*
- ⊕ *Novo Nordisk, Bangalore*
- ⊕ *Lundbeck, Bangalore*
- ⊕ *Eisai Pharmaceuticals, Mumbai*

- ⊕ *LG Life Sciences, Delhi*
- ⊕ *Bayer, Mumbai*
- ⊕ *GE, Delhi*
- ⊕ *Johnson & Johnson, Jansenn Cilag, Mumbai,*
- ⊕ *Cordi Baxter, Delhi*
- ⊕ *BD Biosciences, Delhi*
- ⊕ *Bharat Biotech, Hyderabad*
- ⊕ *Bharat Serum, Mumbai*
- ⊕ *Cadila Pharmaceuticals*
- ⊕ *Cipla, Mumbai*
- ⊕ *Emcure, Pune*
- ⊕ *Fulford India Mumbai*
- ⊕ *Indus Biotherapeutics, Ahmedabad*
- ⊕ *IPCA, Mumbai*
- ⊕ *Shreya Biotech, Pune*
- ⊕ *Shantha Biotechnics Pvt. Ltd. Hyderabad*
- ⊕ *Sun Pharma, Mumbai*
- ⊕ *Torrent Pharmaceutical Ltd, Gandhi nagar,*
- ⊕ *USV Ltd. Mumbai*
- ⊕ *Wockhardt, Mumbai*
- ⊕ *Zydus Cadilla, Ahmedabad*
- ⊕ *Biocon, Bangalore*
- ⊕ *Cadila Pharmaceuticals, Ahmedabad*
- ⊕ *Intas Pharmaceuticals Ltd., Ahmedabad*

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- ⊕ *Glenmark Pharmaceuticals Ltd., Mumbai*
 - ⊕ *Himalaya Drugs ,Bangalore*
 - ⊕ *Lupin Ltd., Pune*
 - ⊕ *Nicholas Piramal,Mumbai*
 - ⊕ *Panacea Biotech, Delhi*
 - ⊕ *Ranbaxy Research Laboratories, Delhi*
 - ⊕ *Ranbaxy Research Laboratories, Gurgaon*
 - ⊕ *Serum Institute of India, Pune*
 - ⊕ *Torrent, Ahmedabad*

KEY CROs IN INDIA

Quintiles
Synchron
Lambda
Genpact
Siro Clinpharm
I-Gate
Reliance Clinical Services
PPD
Onmnicare
ICON
Clin Trac
PharmaNet
Pharm-Olam
Aizant
Lotus Labs
Vimta
GVK
BioServe Clinical Research
Apothecaries
Clinsys
Fortis
Kendle
Bioassay
Clinworld
Perinclinical
Quest Life Sciences

Ahmadabad / Bangalore
Ahmedabad /Bangalore
Ahmedabad
Delhi/Bangalore
Mumbai
Mumbai
Mumbai
Mumbai
Bangalore
Bangalore
Bangalore
Bangalore
Bangalore
Hyderabad
Bangalore
Hyderabad
Hyderabad
Hyderabad
Delhi
Delhi/Noida
Delhi/Noida
Delhi/Gurgaon
Baroda
Bangalore
Mumbai

WHY CAREER IN CLINICAL RESEARCH

- ⊕ Fastest growing segment in the healthcare / pharmaceutical industry
- ⊕ Excellent Opportunity to Develop Combination of Technical and Management Skills
- ⊕ Part of the Global Growth Opportunity
- ⊕ Wider Job Horizon
- ⊕ Rapidly Growing Opportunities...and growing (internal and external)
- ⊕ Attractive Compensation and future growth
- ⊕ Higher Job Satisfaction
- ⊕ Continuous Training Opportunities
- ⊕ International Opportunities

CHALLENGES FOR HUMAN RESORUCES

- ⊕ Very rapid growth in number and size of companies
- ⊕ Spectrum of skills required is in scarcity
- ⊕ Lack of specialization
- ⊕ Limited pool of experienced people
- ⊕ Employee Retention
- ⊕ Continuous Training

Cliniminds - Key Achievements

- ⊕ Established 23 programs with extensive, high quality course content and faculty: Online; E-Learning; Distance Learning & Classroom. More industry oriented programs currently in pipeline.
- ⊕ Presence in 8 key cities of India – Delhi; Mumbai; Hyderabad; Bangalore; Ahmedabad, Cochin, Bhopal and Vijaywada. 4 more cities to start soon.
- ⊕ Program content is targeted towards all major global markets and address the global regulatory environment. Overseas students are growing from Americas; Europe; Middle East; Africa and other countries.
- ⊕ Developed feature rich state-of-the-art user friendly ONLINE LEARNING SYSTEM. System now used by major pharma companies for in-house training.
- ⊕ ***Have been regularly conducting clinical research training programs for National Institute of Health, USA in the Indian market.***

Key Achievements

- ⊕ Awaiting the accreditation of our programs by Government of India.
- ⊕ Top of the Line Faculty & Industry Driven academic Council.
- ⊕ Co-certified Training Programs with **NIH USA**, MSD, Bioserve, Max Healthcare and other companies.
- ⊕ **Key Clients for Training Workshops and Online Training Programs**
: NIH USA; MSD; Fresenius Kabi, Germany; Dabur; Novartis; Quintiles; Ranbaxy; GSK; Bioserve; Max Healthcare; Johnson & Johnson; Panacea Biotec
- ⊕ Strong & Multiple industry tie ups / collaborations to provide practical orientation
- ⊕ Trained over 1,000 professionals, with increasing number of international students for online programs from North America, South America, Europe and Middle East.
- ⊕ Accreditation / Certification from Pharmaceutical Society of India & ISO 9001:2000 by JAS ANZ.

Key Achievements

- ⊕ Conducted several industry workshops for leading corporates, viz. NIH, MSD (Merck), Novartis, Biocon, Quintiles, Bioserve, Ranbaxy, Panacea, Max Healthcare, Max Neeman, Apollo Hospitals, Fresenius Kabi, Merck & Co., Panacea Biotec, Asian Clinical Trials, Quintiles.
- ⊕ Very High student satisfaction levels and excellent placements.
- ⊕ Experience in conducting multi location training programs using Videoconferencing technology.

Training Workshops Offered

- Over 35 Workshops for major Indian and MNCs have been conducted in the following areas :
 - Conduct & Management of Clinical Trials
 - Monitoring of Clinical Trials
 - ICH GCP Workshop on Clinical Research
 - Quality Assurance in Clinical Research
 - Auditing & Inspections of Clinical Trials
 - Pharmacovigilance
 - Regulatory Affairs
 - Medical & Scientific Writing
 - Ethics in Clinical Research
 - Roles & Responsibilities of Investigators
 - Conduct of Cancer Clinical Trials
 - Conduct & Management of BA/BE Studies
- Duration of these workshops have been from 1 – 3 days.
- Workshops are also blended with Online Learning System for follow up training.
- Programs can be customised to suit your needs.

Certifications offered by Cliniminds

- ⊕ Advanced Post Graduate Diploma in Clinical Research & Pharmacovigilance
- ⊕ Advanced Post Graduate Diploma in Clinical Research & Regulatory Affairs in Pharma & C R
- ⊕ Advanced Post Graduate Diploma in Clinical Research & Regulatory Affairs in Pharma & C R-Class Room
- ⊕ Advanced Post Graduate Diploma in Clinical Research & Clinical Data Management & Biostats +SAS
- ⊕ Advanced Post Graduate Diploma in Clinical Research & Clinical Data Management & Biostats +SAS-Class Room
- ⊕ Advanced Post Graduate Diploma in Clinical Research
- ⊕ Diploma in Clinical Research
- ⊕ Post Graduate Diploma in Pharmacovigilance
- ⊕ Certificate Program in Conducting & Managing Bioequivalence & Bioavailability Studies
- ⊕ Certificate Program in Conducting & Managing Clinical Trials for Cancer Patients

Certifications offered by Cliniminds

- ⊕ Post Graduate Diploma in Clinical Data Management & SAS
- ⊕ Certificate Program For Clinical Trial Investigators & Site Personnel
- ⊕ Certificate Program in Monitoring of Clinical Trials
- ⊕ Post Graduate Diploma in Clinical Trials Management
- ⊕ Post Graduate Diploma in Clinical Research for Nurses
- ⊕ Certificate Program in Quality Assurance in Clinical Research
- ⊕ Post Graduate Diploma in Regulatory Affairs
- ⊕ Post Graduate Diploma in Biostatistics
- ⊕ ICH GCP Certificate Program In Clinical Research For Medical Practitioners
- ⊕ Integrated Post Graduate Diploma In Clinical Research & Pharmacovigilance
- ⊕ Post Graduate Diploma in Pharmacovigilance - Class Room
- ⊕ Integrated Post Graduate Diploma in Clinical Research & Pharmacovigilance - Class Room
- ⊕ Post Graduate Diploma in Medical and Scientific Content Writing
- ⊕ Post Graduate Diploma in cGMP – Good Manufacturing Practices
- ⊕ Certificate Program in Clinical Trial Regulations & GCP in Europe & UK
- ⊕ Certificate Program in Clinical Trials Auditing & Inspection
- ⊕ Post Graduate Diploma in Clinical Trials Management - Class Room
- ⊕ Post Graduate Diploma in Clinical Data Management & SAS - Class Room

Cliniminds Students

1. Cliniminds students are from the medical, dental, homeopathic, pharmacy, biotechnology, biochemistry, microbiology and other life sciences background with Advanced Post Graduate Diploma in Clinical Research; Post Graduate Diploma in Clinical Trials Management; Pharmacovigilance; Regulatory Affairs; Data Management & SAS; Monitoring.
2. All students are provided hands on practical training on regulatory affairs; clinical trials conduct & management; ethics; subject recruitment & retention; roles & responsibilities; essential documents, viz. protocol, CRF, ICF, Trial Master File and other documents, monitoring; project planning & management, medical & scientific writing; drug safety, data management, basics of bio stats and other areas.
3. Large number of CROs, pharma companies and hospitals have recruited number of clinical research students from Cliniminds.
4. There are large number of overseas students from North America, Europe, Middle East, South America

ONLINE PROGRAMS

⊕ Cliniminds Online Learning Systems has been designed to meet the needs of both individual students and corporates. Large number of companies have used Cliniminds system for internal training, viz. Ranbaxy; Quintiles|GSK; Bioserve; Asian Clinical Trials, etc.

⊕ Following are the key features :

- Ready to use and user friendly
- Bulk user license with control panel
- Customisation / mix and match from Cliniminds Programs.
- System allows you to upload some of your internal training documents for users. Only accessible for your own teams
- Users account could be created or blocked in no time
- Realtime Records of Training for HR / Training manager to monitor the progress of the users
- Online Evaluation
- Online data can not be saved, download or copy/pasted. Printing could be allowed.
- Effective for site staff / investigators
- Fully protected.
- 99.999% uptime
- Cost effective
- Confidentiality.

Our Capabilities / Strengths

- ⊕ Top of the line industry experts and faculty with extensive industry experience
- ⊕ Expertise in content development
- ⊕ Strong academic team and faculty
- ⊕ Strong practical training and placement assistance to students
- ⊕ Over 20 years of experience in advising, setting up and managing education, healthcare and clinical research businesses.
- ⊕ Experience in setting up turnkey projects in the clinical research & healthcare sector
- ⊕ Promoters come with strong Industry experience and clear vision for the clinical research and healthcare industry
- ⊕ Strong regulatory experience
- ⊕ Strong understanding of marketing and distribution



TESTIMONIALS

The sessions were particularly informative for me, who was not versed in the Indian regulations.

I think the Schedule Y information was of particular interest to me.

It was of particular benefit to the group that Cliniminds focused on GCP, Tox studies, Pre-Clin studies, and Phase I-II-III. The assessment at the completion of the presented materials was great. It is always a must to assess competency as part of a training. All speakers were great presenters and had very good talks.

QA Specialist

NIH

US Government



It's a great privilege for me to give a testimonial about my training institute. I can say that it was a perfect time for me to enhance my skills for widening my opportunities in Clinical Research. I appreciate the study materials because it was a blend of both simplicity and authenticity covering every nook and corner pertaining to my specialization. The coaching classes cum workshops I attended, was completely Industry oriented and professionally executed. I can surely say to the new aspiring candidates, that the courses provided by Cliniminds will be the first of its kind where you get the real touch of the Clinical Research Industry and new energetic start.

Dr. Praveen. S

Clinical Research Coordinator

Max Neeman International

New Delhi



Its really my pleasure and proud to say that I am a student of Cliniminds. This institution helped me a lot to shape up my career beyond my expectation. I am very thankful to the institution to made me to procure a good job in good company. The classes were really in practical sense, which helped me, a lot in this industry to cope up with the work environment. I recommend this institution for those who aspires their careers in Clinical Research.

Dr.Monika Tyagi



I joined Cliniminds in March,08 batch of Advanced P.G.Programme for Clinical Research.It was a very interesting, informative course. In addition,I also attended 3 workshops organized by Cliniminds on ICH-GCP, Quality Assurance and Pharmacovigilance.The workshops were of very good quality in terms of content and faculty.Overall I have found the staff very helpful and the material provided of good quality.I joined Max Neeman International in April,2008 as a CRC.Cliniminds actively helped me to get the job.I have now been promoted to Drug Safety Manager-Medical Monitoring.I sincerely thank Cliniminds and wish best of luck to the institute for the future.

Dr. Sutirtha Mukhopadhyay
Manager – Drug Safety - Icon



I was a part of the **Cliniminds 2008** batch for APCR at the NEW DELHI. The programme helped me in knowing the Clinical Research field which was relatively new to me, the interactive session and the periodic projects and tests helped us to grow our knowledge; the teachers were helpful. The institute helped me in getting a Placement at MAX NEEMAN INTERNATIONAL while I was doing the course, and by the time I completed the course, I was promoted. The regular counseling helped me in planning my career. ***I WISH THE BEST OF LUCK TO ALL THE CURRENT AND FUTURE STUDENTS OF CLINIMINDS.***

Dr SAURABH SAXENA B.D.S P.G.D
Clinical Research Associate MAX NEEMAN
INTERNATIONAL Max House, 1st Floor,
1, Dr. Jha Marg, Okhla-III New Delhi-
110020



It would be injustice if I failed to appreciate my courses (Certificate program in Clinical Research ; Pharmacovigilance ; & Regulatory Affairs) from Cliniminds. . The courses made me industry-ready, and also gave me the edge that others do not have. The study manuals for all the courses were quite comprehensive and easy to understand and in my opinion, it covered all aspects, while also giving ample knowledge about the subject. I would really like to thank Cliniminds for making me understand about the basics of Clinical Research Industry and delivering the right knowledge in a structured manner. All in all it was a great experience and one that I would be ready to undergo any time again.

Dr. Sandeep Bhatia

Medical Advisor

Sanofi-Aventis



Thank you for organizing an ICH GCP Workshop at Max Healthcare at such a short notice.

I am pleased to inform you that the workshop was very successful, and have completely met out objectives.

We are pleased with the content quality and the faculty.

We would be glad to use you training services for our future Clinical Research Training requirements.

Dr. Saroj Kumar Sabath
Manager-Clinical Research
Max Healthcare



I am really glad to inform you that I have joined MAX NEEMAN as CRC from 16th March 09.

I am really thankful to you for recommending my name to Max Neeman.

It was great learning in Cliniminds from eminent and experienced teachers from Clinical Research field.

Sincere thanks for all your support and encouragement.

Dr. Anuja Mathkari

I rate it as best one, because here in our company we have online training sessions just same as u have, but it is somewhat different more student friendly and never encountered any problems interms of difficulty in navigation and completion.

I salute the designer of site and content of study material, i also recommend all my colleagues about the course.

I owe to your cliniminds for ignite the minds of students by such a awesome course.

Waiting for your foster certification in course.

D.Raghavendra,
Quality Assurance Specialist,
Quality & Compliance
Management
GSK
Gurgaon, India.

I have taken the course "Integrated Post Graduate Diploma In Clinical Research and Pharmacovigilance" during the period of May - July, 2009. Having found it myself on-line, I found it very important to be sure the site was reliable. From the very first steps of my on-line interaction with the course providers I put whatever initial uncertainties I had aside. The authors were committed, prompt, responsive and engaged. They would take all of my inquiries seriously and usually get back to me within 24 hours, which was more than understandable due to the time difference. This remained true from the transfer of advance payment up to the moment I was issued the graduation certificate. Above all though, it was the concept that caught my attention in the first place. The name of the course matched my area of interest and once I got my access rights I learned that its content lived up to it as well. Being put together in a comprehensive way, it also provided a user-friendly learning and knowledge testing interface and manageable timelines. I can recommend the course to anyone who is looking for the value for their money. The absence of any of the international English language evaluation requirement also saves money and time, provided one is already familiar with the specifics of the English used by the international clinical trials community, which no English course can be of much help with anyway.

Tomas Novak,

PharmD July 9, 2009 Prague,
Czech Republic"

Very good faculty, very interactive and positive in accepting comments, queries and answering to their best capability and knowledge.

The content of presentation was adequate and simple for understanding to freshers as well.

I liked the institute and wish to enroll myself for future workshops and in the ongoing courses.

Rashmi Kulshrestha

Associate Director Regulatory
Affairs

Ranbaxy Laboratories Ltd.



I found the workshop on ICH –GCP conducted at the Cliniminds highly interesting and praiseworthy for the fact that those two days of brainstorming sessions gave me a detailed insight into various aspects of Clinical Research imperative for doctors and its growing demand in Pharmaceutical industry. The faculty at the Cliniminds is highly skilled in their job and good enough to solve my queries. I would like to wish the Cliniminds the best of wishes in all their endeavors towards developing skilled and motivated Clinical Researchers I found the workshop on ICH –GCP conducted at the Cliniminds highly interesting and praiseworthy for the fact that those two days of brainstorming sessions gave me a detailed insight into various aspects of Clinical Research imperative for doctors and its growing demand in Pharmaceutical industry. The faculty at the Cliniminds is highly skilled in their job and good enough to solve my queries. I would like to wish the Cliniminds the best of wishes in all their endeavors towards developing skilled and motivated Clinical Researchers

Dr. Manish Mahajan
M.D. Deptt. of Biophysics
A.I.I.M.S.

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