The Clinical Development Services Agency (CDSA) organized a training programme on Good Clinical Practice (GCP) supported by the National Biopharma Mission (NBM) on the 28th and 29th January 2020. GCP is an international ethical and quality standard for the design, conduct, performance, monitoring, auditing, recording, analyses and reporting of clinical trials that provides assurance that the rights, safety and well-being of human research participants are well protected, data and reported results are credible and accurate. This makes creating awareness about GCP mandatory when we talk about innovation in medicine and healthcare. The launch of the New Drugs and Clinical Trials Rules, 2019 on March 19th 2019, opens up a new set of opportunities to accelerate innovations in India. Poor understanding of the current regulatory requirements mixed with inadequate compliance to best global practices may lead to delays and failures which can be significantly reduced through awareness programmes. The workshop was thus designed with a goal to create awareness and understanding of GCP to enable compliance with the current guidelines and regulatory requirements. The session today had about 70 participants from academia, industry, ethics committees’ members, researchers, CROs and pharma companies in the audience.

**Contact Persons:** Dr. Siuli Mitra, THSTI, Faridabad, Haryana; Ms. Vandana Chawla (CDSA coordinator for the workshop)

Email ID: smitra@thsti.res.in