



ADI Intellect Provides Expert Medical Communication Solutions & 2D & 3D Animation

Delivering the Highest Quality Regulatory and Medical
Documents Spanning the Drug Development Continuum

Why Choose ADI?

The demand for medical writing has increased considerably in recent years. There are several reasons; more research studies are being conducted today in the biomedical field, pharmaceutical companies are discovering more new drugs, and diagnostic and therapeutic companies are developing innovative medical devices.

Concomitantly, regulatory and scientific documents need to be produced for submission to regulatory authorities during their approval process in addition to a plethora of supporting documents aimed at educating a diverse audience of healthcare professionals and lay people. It is a formidable challenge.

Simply put, we thrive in this environment.

ADI Intellect Advantages

- Thoroughly Research Project
- Slavish Adherence to Accuracy & Details
- Think Logically
- Work Collaboratively
- Excellent Communication & Time Management Skills
- Audience Centric Approach

35+ Years
in Regulatory & Medical Writing
Experience in 20+ Therapeutic Areas

Regulatory Writing

Pharmaceutical companies understand that first-to-market is paramount in gaining an advantage over their competitors. Regulatory writers are an indispensable part of the drug approval process.

The regulatory writer must interpret clinical trial data correctly and summarize the statistical findings to produce timely, clear and concise documents that will pass regulatory scrutiny. Our regulatory medical writers are trained in the latest regulatory requirements.

They possess a detailed understanding of the International Conference on Harmonization (ICH) guidelines, regulatory style requirements, tone, and preferred terminology.

Our regulatory writers are experts who consistently meet and surpass our client's needs. They will expertly guide you through all phases of your clinical trial to inform the targeted audience about the drug under study.

Regulatory Writing Portfolio

- > All documents comply with ICH E3 requirements
- > Deliverables include clinical study reports, appendices, trial protocols, report and informed consent forms
- > Investigator brochures and briefing packages for meetings with regulators
- > Package inserts
- > Pharmacovigilance reports e.g. annual safety reports
- > Periodic adverse drug experience reports (PADER)
- > Patient safety narratives
- > Common Technical Document (CTD) Summaries
- > Clinical Evaluation Report (CER) for medical devices
- > Site-specific and country-specific reports

Clinical Study Design

We will design every aspect of your clinical trials. Our years of experience and knowledge combined with our medical animation resources, will showcase your clinical trial results in the clearest and most compelling way. This will maximize your chance of gaining acceptance from the FDA. In turn, allowing you to move forward with other clinical trials and product development plans.

Your clinical trial will be created by experienced professionals who can maximize the probability of acceptance by regulatory bodies.

You do not want corporate objectives for your clinical trial to flounder or fail before the trial can get off the ground.

Clinical Study Objectives

- > Clearly describe clinical trial objectives and methodology
- > Identify key data to be collected for Clinical observations
- > Conduct patient profiling including number needed
- > Assure safety and efficacy of all aspects of the trial
- > Strict adherence to regulatory requirements including protocol
- > Assure quality and statistical analysis of trial data

2.5 Million Hours
in Clinical Trial Intelligence
Spanning 150,000 Clinical Trials

Medical Writing

New information is constantly being added to the field of medicine via an ever-increasing pipeline of drug discovery and development, expanding clinical experience, and medical device innovation. This information needs to be effectively communicated to different audiences; physicians and other healthcare professionals, patients and consumers, investors, and regulators.

Medical writing is the discipline of writing scientific documents. Medical writers may not be the original scientists who do the actual research but work with the physicians/scientists involved in the generation of data and help present the information in the most effective manner.

The importance of good medical writing cannot be ignored as science depends on clear and concise reporting. Meticulous research can appear flawed if it is poorly presented.

Medical Writing Portfolio



Marketing
Communications
Literature



Clinical Trials,
Drug Intelligence
& Report Writing



Medical & Scientific
Education



Bibliographic
Searches



Manuscripts
& Publications



Patient
Narrative



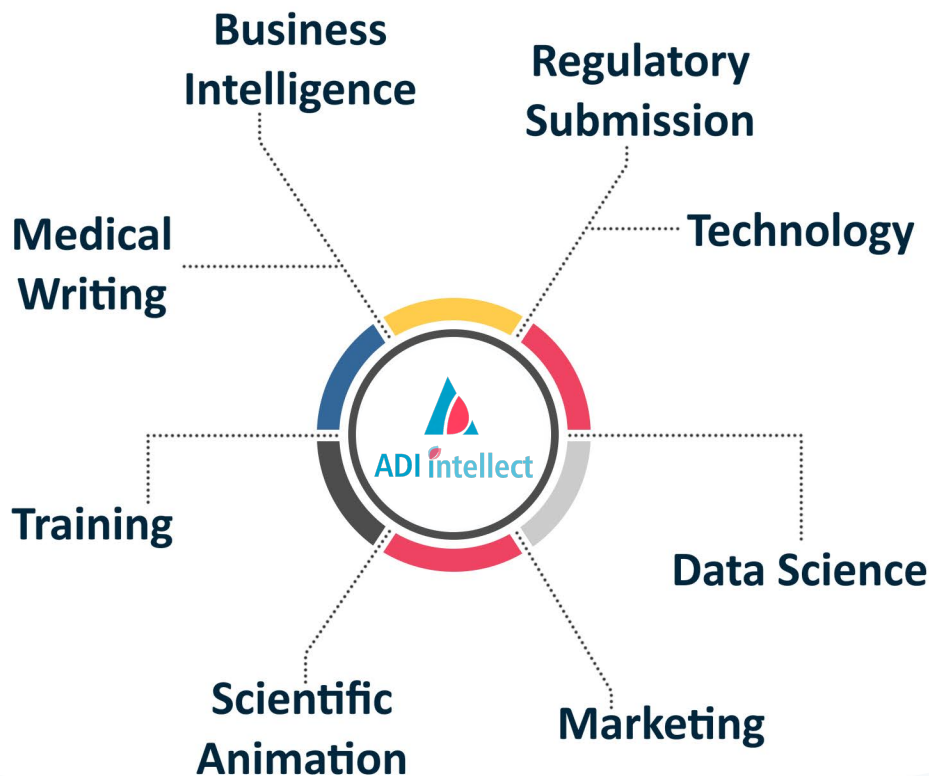
Our Companies

The ADI Intellect offers both horizontal and vertical services for various industry segments. Our offerings help industry leaders and departments make faster decisions through our **Business Intellegience** services, reach out to their audience with our **Animation**, allow their teams to be more productive quickly through our **Interactive Learning**, acquire the best technology performance through our **IT Service**, obtain cutting edge Data analytics through our **BIG DATA** technology, benefit from our **Process Outsourcing** to maximize business focus and cost benefits, abet the drug development process through our **Clinical Trial Services**, and facilitate new drug development utilizing our **Drug Repurposing** assistance.

What started with a small group of 7 team members has now expanded to a team of hundreds of expert professionals; ADI continues to transcend geographical limits to win over global associates.

The ADI's incredible growth rate, more than 50% p.a., reflects well on its leadership, the entire team's dedication, and its commitment to excellence. Entrepreneurial spirit defines ADI; performing everything with utmost precision, motivated by an ever-innovative approach.

Our management team shares our success with renowned medical doctors and researchers, creative animators and designers, and innovative technical staff. As a result, the ADI is recognized worldwide as best in class in knowledge process outsourcing, new drug delivery processes, high end tools for drug development, and contract research and consultation. Every day is a new and welcome challenge at ADI. Come challenge us!



ADI Intellect

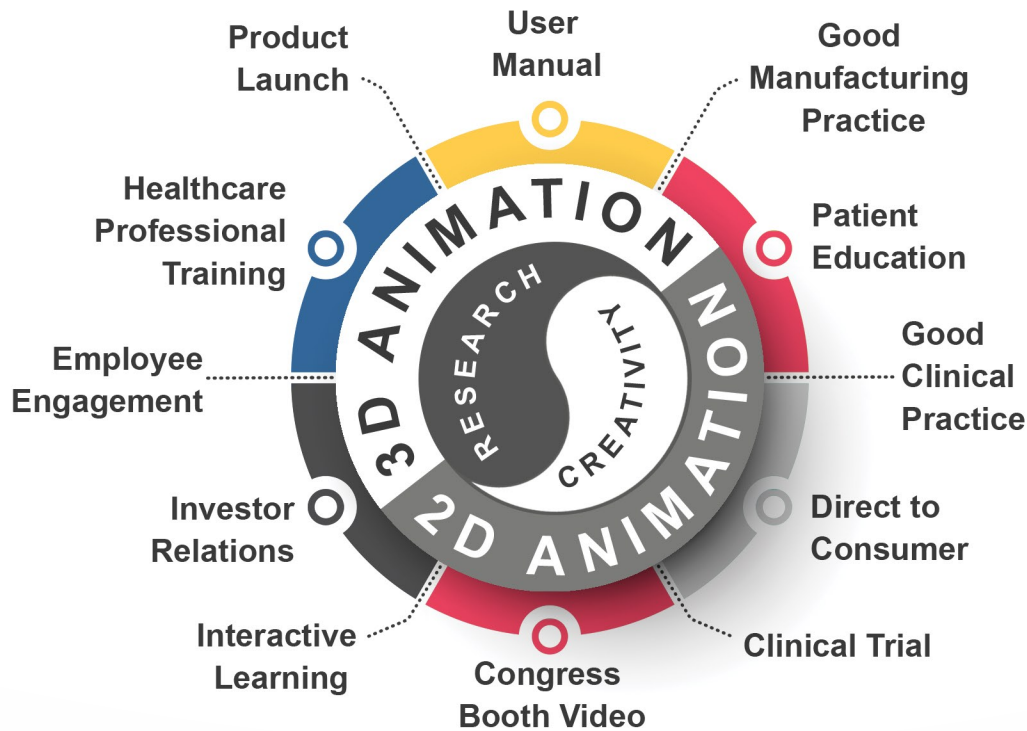
At ADI, our dedicated clinical research professional offers simplified communication solutions to the specific needs of healthcare organizations, ensuring compliance with regulatory standards both internally and externally. Our approach is audience-centric, evidence-based, and simplified, utilizing infographics and 3D graphic illustrations for better understanding. With our unparalleled expertise, we seamlessly translate complex medical concepts into simple and engaging content, equipped to ignite innovation and more clarity in the dynamic world of Med-Tech.



2D & 3D **Animation**

ADI intellect **Promotes Deeper**
Understanding of Your Products

Before, During & After Development We Will Convey
Your Drug's Benefits with the power of Medical Animation.



Our Process & Methodology

- Audience Centric Approach
- Research Based Content Development
- Library of Human Anatomy Models & Illustrations
- Designed by Team of Subject Matter Experts

5000+ Minutes
of Animation Content Created
Experience in 20+ Therapeutic Areas



We Couple **Passion** with
Technical **Proficiency** to Accelerate
Drug Commercialization.

Contact **Us**



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