

Research Acronyms

Adverse Event of Special Interest	AESI
Adverse Event	AE
Adverse Drug Reaction	ADR
Central Institutional Review Board	CIRB
Collaborative IRB Training Initiative	CITI
Code of Federal Regulations	CFR
Clinical Laboratory Improvement Amendments	CLIA
Clinical Research Assistant	CRA
Clinical Research Coordinator	CRC
Case Report Form	CRF
Clinical Trial Agreement	CTA
ClinicalTrials.gov	CT.gov
Code of Federal Regulations	CFR
Collaborative IRB Training Initiative	CITI
Confidential Disclosure Agreement	CDA
Conflict of Interest	COI
Coverage Analysis	CA
Contract Research Organization	CRO
Data Management Plan	DMP
Data and Safety Monitoring Board	DSMB
Data and Safety Monitoring Committee	DSMC
Department of Health & Human Services	DHHS
Electronic Case Report Form	e-CRF
Electronic Data Capture	EDC
Electronic Medical Record	EMR
Federal Wide Assurance	FWA
Financial Disclosure Form	FDF
Food and Drug Administration	FDA
Food and Drug Administration	FDA
Genetic Information Nondiscrimination	GINA
Good Clinical Practice	GCP
Health Insurance Portability and Accountability Act	HIPAA
Human Exposure Review Committee	HERC
Human Research Protection Program	HRPP
Investigational Brochure	IB

Institutional Biosafety Committee	IBC
Informed Consent Form	ICF
Independent Ethics Committee	IEC
Institutional Review Board	IRB
International Conference on Harmonization	ICH
Investigational Device Exemption	IDE
Investigational Drug Service	IDS
Investigational New Drug Application	IND
Investigational Product	IP
Investigator's Brochure	IB
Legally Authorized Representative	LAR
Manual of Procedures	MOP
Materials Transfer Agreement	MTA
Non-Disclosure Agreement (UCSD agreement)	NDA
New Drug Application	NDA
National Institutes of Health	NIH
Office of Grants Administration	OCGA
Office of Clinical Trials Administration	OCTA
Office of Human Research Protection	OHRP
Office of IRB Administration	OIA
Office of Sponsored Programs	OSP
Pharmacodynamics	PD
Pharmacokinetics	PK
Principal Investigator	PI
Protected Health Information	PHI
Quality Assurance	QA
Quality Control	QC
Radiation Safety Committee	RSC
Serious Adverse Event	SAE
Standard Operating Guidelines	SOG
Sub-Investigator	Sub-I
Unanticipated Problem Report	UPR