



## **IRB Statement of Compliance**

IntegReview IRB, an independent IRB located in Austin, Texas, was established in 1999. The IRB offers five meetings (Monday-Friday) each week for review of U.S., Latin American and Japan research sites, as well as certain global educational research projects. We provide quality review by experienced individuals and perform quality assurance/control on all submission and approval documentation. Designed to accelerate the IRB process without compromising accuracy or the protection of human subjects, IntegReview IRB delivers study documents to the investigator in an efficient and timely manner.

IntegReview IRB is committed to meeting rigorous standards for quality and maintaining sound policies and procedures involved in the protection of human research participants. IntegReview IRB was initially awarded full accreditation of its human research protection program (HRPP) by the Accreditation of Human Research Protection Programs, Inc.<sup>®</sup> (AAHRPP) in June 2007. AAHRPP accreditation will expire in September of 2020.

Written standard operating procedures govern IntegReview IRB for initial, continuing, full board, expedited and exempt review of clinical, social-behavioral, education and evaluation research studies. IntegReview IRB complies with the regulations as defined in the United States Food and Drug Administration (FDA), Code of Federal Regulations, Title 21, Parts 50, 54, 56, 312 and 812, International Conference on Harmonisation (ICH) Guidelines for Good Clinical Practices, E6, the Department of Health and Human Services (DHHS) regulations as identified in the Code of Federal Regulations, Title 45, Part 46, Title 34, Parts 98 and 99, other regulations as applicable, as well as local and state laws.

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