

Newsletter



Implementation of a Quality Culture at the Research Site

A quality culture is reflected in the consistency of the site's daily processes. Standardization of procedures and effective team communication help minimize errors and ensure protocol compliance. Two things to consider while implementing this approach are:

SOPs and Quality Assurance

Standard Operating Procedures form the foundation of the site's quality system. Proper implementation and periodic review allow clinical practice to remain aligned with regulatory and operational study requirements.

Ongoing Preparedness for Monitoring and Audits

A proactive approach to quality enables the site to maintain continuous compliance, reducing risk during monitoring visits, audits, and regulatory inspections.

Interested in collaborating or learning more about our programs? Visit <https://alliance.rcm.upr.edu/iscorerc/>

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Congratulations!

Our congratulations to trainee Brenda A. Guzman on the successful completion of the Clinical Research Coordinator Development Program this February. This accomplishment represents a meaningful milestone in her professional journey and highlights a strong commitment to growth within the field of clinical research.



ISCORE-RC CRCDP Program Experience

By Brenda A. Guzman

"My experience in the ISCORE-RC Clinical Research Coordinator Development Program has been very enriching. I have gained a greater understanding of the importance of the study coordinator's role within a research team. This program has strengthened my knowledge of good clinical practices, research ethics, and proper protocol management. The way the course is designed allowed me to apply what I learned to real-life situations. I really liked the combination of online learning with the hands-on training component, as it allowed me to apply what I was learning almost immediately. This experience has been a great contribution to my professional development and has reinforced my commitment to clinical research focused on the well-being of participants. I feel more confident, organized, and prepared to coordinate clinical studies with responsibility and professionalism."