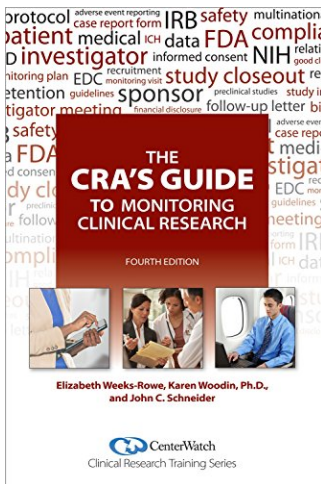


[PDF] The CRA's Guide To Monitoring Clinical Research, Fourth Edition

Elizabeth Weeks-Rowe, Karen E. Woodin Ph.D.,
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The CRA's Guide to Monitoring Clinical Research, fourth edition, continues to be a vital resource for both novice and experienced CRAs seeking to learn more about the field of monitoring or to better understand their roles and responsibilities as the industry becomes more global and technologically focused. This edition includes helpful tips and strategies, checklists, personal experiences, traveling tips, key takeaways and exercises. With new and updated chapters on the evolving CRA roles and responsibilities, monitoring for device and biologic trials, risk-based monitoring, globalization of studies, EMR, web-driven data collection, the sub-PI role, and more,

The CRA's Guide is a must-have training and educational tool that you'll refer to time and time again.

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