Study Teams Must Manage Visits in OnCore: In the coming months, a greater emphasis will be placed on managing OnCore study visits. This means the study staff must mark an OnCore visit as Occurred, Missed, or N/A within two business days of the visit’s occurrence. To assist study teams, the OnCore team will send ‘late visit’ reports to study coordinators. If billing questions should arise because visits are not managed within time limits, study teams will receive inquiries from the Clinical Billing Office.

Contact the OnCore team for more information.

Facilities & Support You Need: The CCTS can connect you to specialized capacities and expertise, like extended stays for healthy research participants and nutrition-related research support.

Contact the CCTS to learn more.

Enhancing the Diversity of Clinical Trial Populations - Eligibility Criteria, Enrollment Practices, and Trial Designs Guidance for Industry: Toward health equity, it is essential that clinical trials enroll diverse populations with a broad range of characteristics (e.g., sex, ancestral background, age, comorbid conditions, socioeconomic determinants). Important guidance was published by the FDA last November.

Read it here.

Thank you for reading,

UAB Clinical Trials Administration Committee (CTAC)

Do you have news or updates to share with the clinical trials community at UAB?

Send us an email!