Educational Pre-Conference Workshop/In-Conference Workshop Descriptions

Pre-Conference Workshops
Short courses on topical methods or issues related to clinical trials and are scheduled for 4 hours. The focus is on education and training and will include hands-on work and plenty of time for questions and discussion. Please include a bullet point description of how the workshop will be structured (e.g., 10-min presentation followed by 30-mins of small group work with hands-on use of software etc.). Pre-Conference Workshops are offered at an additional cost to the attendees and will be held on the Sunday prior to the SCT Annual Meeting. Submissions with faculty from diverse backgrounds and affiliations are strongly encouraged. The Primary Contact/Author can edit the submission in the portal until the associated submission deadline as indicated on the portal. All session proposals, abstracts, etc. submitted by the designated submission deadline will be considered the final versions. No additional editing can be made after the indicated submission deadlines.

In-Conference Workshops
Interactive sessions on a method or topic related to trials. In-Conference Workshops are not intended to be purely didactic but will have plenty of time for discussion and may include small-group work, hands-on use of tools and software, troubleshooting and "ask-the-expert" time. In-Conference Workshops are scheduled for 90 minutes. Please include a bullet point description of how the tutorial will be structured (e.g., 10-min presentation followed by 30-mins of small group work with hands-on use of software etc.). In-Conference Workshops will be Monday, Tuesday, and Wednesday in conjunction with the SCT Annual Meeting. Submissions with faculty from diverse backgrounds and affiliations are strongly encouraged. The Primary Contact/Author can edit the submission in the portal until the associated submission deadline as indicated on the portal. All session proposals, abstracts, etc. submitted by the designated submission deadline will be considered the final versions. No additional editing can be made after the indicated submission deadlines.

Submissions on any topic relevant to trials are welcome. This includes but is not limited to: study design, trial conduct methodology, data management, research ethics, information technology, data quality and safety monitoring, study coordination and management, education and mentoring, policy, regulation, statistical analysis, and reporting of results. We particularly welcome submissions covering the following topics: note that submissions do not need to cover all the items listed in that topic’s bullet point below (e.g. a clinical trial conduct workshop could cover recruitment but not start-up):

1. Adaptive designs
2. Pragmatic trials
3. Designing informative feasibility studies and pilot trials
4. Core outcome set development and selection of patient-centered outcomes
5. Clinical trial conduct: effective trial recruitment and retention, study start up and close-out in multi-center trials
6. The working of Data Monitoring Committees and preparing reports for DMCs
7. Health economics and cost effectiveness analysis
8. Data sharing: Preparing, submitting, and accessing trial data from data sharing platforms