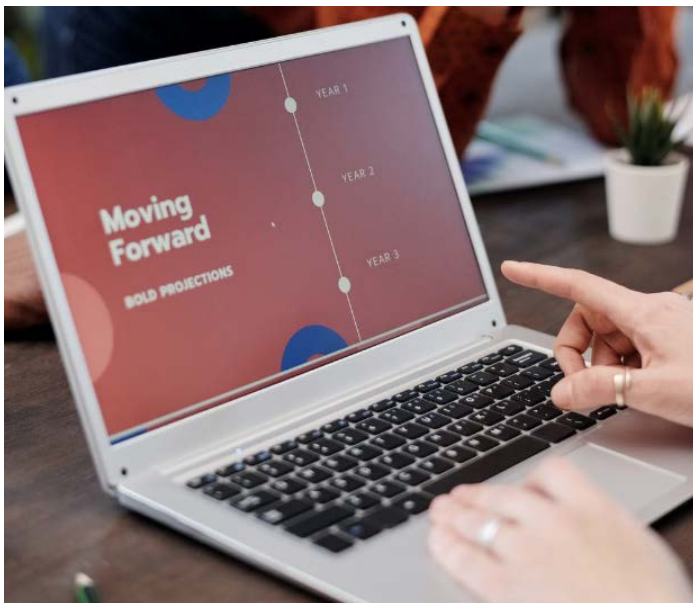




## A+ Science Newsletter

*"The way to get started is to quit talking and begin doing."*

Walt Disney



Teamwork and strong open communication are critical for project success. Formal and informal activities and trainings will play a part in bonding the team members and are particularly valuable and especially when the team operates from remote or different locations.

If you choose A+ Science as your CRO partner, you will receive a complimentary Good Clinical Practice (GCP) or Good

Pharmacovigilance Practice (GVP) webinar where your project team will receive a basic training together with the team at A+ Science allocated for your project. This complimentary training will be led by A+ Science Quality Assurance Consultant, Björn Edwall who has more than 30 years of experience in leading

positions within the pharma industry at global companies including Novartis and AstraZeneca. It is a perfect starting point for the collaboration between our companies and provides an opportunity for our respective project teams to get to know each other before project start. We realise that there is such a need and it will only be for the benefit of everyone. The training webinar can provide the project team members with learning opportunities that are effective and enjoyable as well as achieve development goals. What could be better than this?

Contact us at [info@a-plusscience.com](mailto:info@a-plusscience.com) to hear more about our offer.

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Are you aware of all the services A+ Science provides?

- Our services include clinical trial management, we can conduct the entire clinical study, from start to finish or parts of it, depending on our customers' needs.
- We also provide full pharmacovigilance services, pre-and post-marketing, including 24/7 availability of QPPV, Deputy QPPV and Drug Safety Physician, safety database license including MedDRA license, ICSR processing (pre-and post-marketing) including medical review and submission to Eudravigilance, Monitoring of new legislation, literature search and review (global and local, ICSR collection from EVWEB, entry and maintenance into the XEVMPD, PSMF maintenance, signal management, writing of SOPs, PSURs, DSURs and RMPs.
- We can empower your team by outsourcing personnel (short- and long-term solutions).
- Since 2004, we offer services within Continuing Medical Education (CME) in collaboration with University Hospital of Umeå and the Sahlgrenska Academy, University of Gothenburg.

Contact us at [info@a-plusscience.com](mailto:info@a-plusscience.com) to find out more about how our team at A+ Science can help you reach your goals and objectives.

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