Biopharmaceutical companies are committed to enhancing appropriate transparency of their research in a responsible manner. Clinical trial data transparency has multiple elements, all of which are aimed at improving access to clinical trial information in order to inform medical decision-making, advance scientific discovery and accelerate development of new treatments to benefit patients. They include the following:

**What is Clinical Trial Data Transparency?**

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- **Data Sharing**
  Clinical trials generate incredible amounts of data and responsibly sharing clinical trial data with qualified researchers has the great potential to advance medical research and benefit patients by accelerating the development of new medicines and improving patient care.

- **Registration and Results Reporting**
  Certain clinical trials are required to be registered and have their results reported on registries such as ClinicalTrials.gov. America's biopharmaceutical industry remains committed to enhancing public health and advancing medical research by sponsoring and conducting clinical research that fully complies with all legal and regulatory requirements.

- **Voluntary Disclosures**
  PhRMA’s member companies provide clinical trial information above and beyond legal and regulatory requirements, including publishing trial results in scientific literature and posting the results of clinical trials on company websites.

- **Lay/Plain Language Summaries**
  In order to make complicated clinical trial data more accessible to the public, PhRMA supports working with regulators to adopt mechanisms for the preparation and distribution of lay summaries to research participants.

Learn more at PhRMA.org/Clinical-Trials