A master protocol is designed to allow drug developers to test multiple interventions in one or more patient types or diseases within the same overall trial structure.

On April 15, MDA’s 2019 Clinical & Scientific Conference kicked off with a keynote address and panel discussion, where experts weighed in on the number of new drugs being submitted to the FDA for review and the best strategies for bringing them to market.

Keynote: From Discovery to Delivery
MDA was honored to have Janet Woodcock, M.D., director of the Center for Drug Evaluation and Research (CDER) at the U.S. Food and Drug Administration (FDA), present the keynote address.

Dr. Woodcock spoke about the current explosion of scientific information that will lead to new treatments for neuromuscular disease (NMD). According to Dr. Woodcock, new scientific discoveries are “pouring out of the lab,” providing hope for people with previously untreatable diseases.

But she also noted the challenges of developing drugs for neuromuscular diseases.

Clinical development for neuromuscular disease drugs is a long, expensive process, which can be tragic for patients.
Master protocols are a way to overcome some of the challenges so researchers can answer more questions in less time.

A master protocol is designed to allow drug developers to test multiple interventions in one or more patient types or diseases within the same overall trial structure. As a collection of trials, a master protocol shares key design components and operational aspects to achieve better coordination than can be achieved in single trials designed and conducted independently.

FDA Panel:
Seeking Change for the Better
Following the keynote, a group of experts took the stage for an FDA panel discussion. Joining Dr. Woodcock were Peter Marks, M.D., Ph.D., director, Center for Biologics Evaluation and Research (CBER) at the FDA; John Day, M.D., Ph.D., professor of Neurology at Stanford University; and Frank Sasinowski, M.P.H., J.D., adjunct professor of Neurology at the University of Rochester Medical Center.

The panel’s discussion stemmed from Dr. Woodcock’s keynote remarks on prescription, generic and over-the-counter drug regulation.

The number of Investigational New Drug (IND) applications to the FDA was increasing noticeably, said Dr. Marks. He cited a prediction from the Massachusetts Institute of Technology stating that by 2030, 40 to 60 new gene therapies will launch, and more than 500,000 individuals will be treated with gene therapies in the United States. He added that as of April 1, of the 97 requests the FDA received for Regenerative Medicine Advanced Therapy (RMAT) designation, 33 products were granted RMAT designation. Twenty of these were given Orphan Product designation (otherwise known as Orphan Drug designation). Most of these products are cellular therapy or cell-based gene therapy products.

Dr. Day was enthusiastic about gene therapies. “It’s exciting for medicine,” he said, “because these platforms have now been proven to be effective, and we can be excited about where this is going.”

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— JOHN DAY, M.D., PH.D.

Sasinowski applauded the FDA for not only listening to patients but acting on what patients tell them, primarily during and after FDA-supported patient-focused drug development meetings. By empowering patients, the FDA tries to understand and learn from those patients’ experiences.

The panel also gave advice for drug developers: Enter conversations with the FDA in early development stages. The panelists referenced times in which they’d had discussions with developers early in their process on broadening the agenda so results would be more applicable to more populations. In the end, the panel stressed the importance of continued conversations and sharing in advancement of research as a whole.

A key takeaway from the panel was that when it comes to neuromuscular diseases there is no limit to what we can achieve. It will, however, take a patient-centered approach, communication and cross-collaboration to best reach patients and to arrive at treatments and cures.