## FDA Issues Final Rule on Regulation of Laboratory Developed Tests

On April 29, 2024, the U.S Food and Drug Administration (FDA) announced its **final rule** on Laboratory Developed Tests (LDTs). This final ruling amends the FDA's regulations to make explicit that *in vitro* diagnostic products (IVDs), including those manufactured by laboratories, are devices under the Federal Food, Drug, and Cosmetic Act (FD&C Act). Alongside the amendment, FDA issued its policy to phase in regulatory requirements for certain LDTs over the course of four years.

The FDA will host a webinar to provide an overview of the final rule on May 14, 2024. A link to register can be found <a href="here">here</a>. The final rule is expected to have profound effects on many LDT developers. Goodwin's <a href="Life Sciences Regulatory & Compliance Team">Life Sciences Regulatory & Compliance Team</a> are ready to work with clients to navigate the challenges that the final rule may pose. Our breakdown and analysis of the rule will be upcoming on <a href="Goodwin's LDT Resource page">Goodwin's LDT Resource page</a>.