<u>How the Trump Administration Could</u> <u>Reshape Regulation in the Life Sciences</u> <u>Sector</u>



Based on recent policy signals and statements from incoming administration officials, a picture of potential regulatory and policy changes that could affect biotech, pharmaceutical, and medical device companies in coming months and years is emerging.

Anticipated changes span multiple regulatory fronts: a revamped approach to antitrust review at the Federal Trade Commission (FTC), continued momentum on biosecurity measures, and a fundamental rethinking of agency regulation to streamline "red tape" and accelerate patient access to innovative treatments. The Trump administration's stated focus on "making America healthy again" suggests a broader transformation in how healthcare is delivered and regulated, with emphasis on nutrition, prevention, longevity, enhanced physician autonomy, and a more holistic approach to health to reduce the burdens of chronic disease.

While some changes may create opportunities for innovation and growth, others could pose compliance and operational challenges. Understanding these emerging dynamics will be crucial for industry stakeholders as they position themselves for success under the new administration.

The following six sections are based on discussions from a regulatory panel held on January 15 at the <u>Goodwin + KPMG 6th Annual Symposium</u>, which was held during the 2025 JPM Healthcare Conference.

Read the full insight **<u>here</u>**.