Moving from the Informed Consent to Approved Labeling: Preparing for Risks in Product Marketing & Use Webinar Recording



On February 3, 2021 Goodwin FDA Regulatory partner, <u>Julie Tibbets</u>, Products Litigation + Counseling partner, <u>Nilda Isidro</u>, and Risk Management & Insurance counsel <u>Brian</u> <u>Mukherjee</u> discussed what drug and biologic companies with late-stage product candidates can do to best position their products to mitigate the risks that come with transitioning from clinical trials to marketing and sales.

Our speakers – leaders in life sciences regulatory compliance, product litigation preparedness, risk management, and insurance – highlighted best practices surrounding pharmaceutical promotion, preparing for risks inherent in the marketing and sale of prescription drugs, and the ways in which insurance can help mitigate those risks. This webinar identified key takeaways for companies that are nearing FDA approval and are poised to launch their commercial products.

Click **here** to view the slides and webinar recording.