

[Mark Your Calendars: This Halloween, Don't Miss FDA's LDT Webinar](#)



The U.S. Food and Drug Administration (FDA) has announced an upcoming [webinar](#) on its [proposed rule](#) on the regulation of laboratory developed tests (LDTs).

The webinar is scheduled for **October 31, 2023 from 1:00 - 2:00 PM ET** and will include an overview of the proposed rule, a description of the proposed phaseout of FDA's general enforcement discretion approach to LDTs, and a question and answer session. Stakeholders must submit questions by **October 23, 2023** to be considered for the discussion.

For our detailed analysis of the 83-page proposed rule, please see our two-part Insight series: [Part I: Underpinnings of FDA's Proposed Rule](#) and [Part II: FDA's Proposed Phaseout Policy - Key Considerations & Open Questions](#).

If you have questions on the proposed rule or its potential impact, contact the authors or a member of the [Goodwin Life Sciences Regulatory & Compliance](#) team.