

## Janice M. Hogan

Partner

Philadelphia

Washington, D.C.

### Biography

Janice Hogan has been involved in medical technology for over 25 years. From her engineering training at the Massachusetts Institute of Technology to work in the pharmaceutical industry, to her current practice representing medical device companies before the U.S. Food and Drug Administration (FDA), Janice has focused her career on the intersection of technology, regulation, and health care.

Widely recognized as a leader in FDA regulation of devices, Janice leverages her technical background to help companies with cutting-edge technologies navigate and optimize the FDA approval process.

Janice focuses on FDA regulation of high-tech products in women's health, diagnostics, neurology, cardiovascular, and orthopedics. She has assisted companies to obtain "first-of-a-kind" FDA approvals, providing guidance on regulatory strategy, clinical study design, advisory panel proceedings, and tools to expedite product approval.

Janice brings her lifelong passion for science and innovation to bear in her client advocacy. Through her extensive experience representing companies in a wide range of FDA interactions, Janice has been at the forefront of several of the most innovative medical device approvals, including the first successful FDA/Centers for Medicare and Medicaid Services parallel review project, as well as first-in-class



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### Languages

English

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### Practices

Medical Device and Technology  
Regulatory

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### Industries

Life Sciences and Health Care

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### Areas of focus

Advisory Panel Preparation

Combination Products

In Vitro Diagnostics

Medical Devices

approvals for devices used in treatment of breast cancer, diabetes management, obesity, spinal surgery, and neurology, as well as a variety of drug/device combination products. Janice also has substantial experience using newer FDA approval mechanisms such as the de novo process to reduce review time and bring products to market earlier, accelerating patient access.

## Awards and rankings

- Healthcare: Life Sciences, *Legal 500 US*, 2014-2015, 2017-2020
- Most Highly Regarded Firm for Life Sciences 2018, *Who's Who Legal*, 2018
- Life Sciences Star, *LMG Life Sciences*, 2018
- Who's Who Legal Life Sciences: Regulatory Lawyers, *Who's Who Legal*, 2008-2018
- Regulatory: Medical Devices, *PLC Life Sciences Cross-border Handbook*, 2011-2012

## Latest thinking and events

- Press Releases
  - Hogan Lovells advises CartiHeal in securing FDA premarket approval for knee implant
- News
  - COVID-19 Report for Life Sciences and Health Care Companies
- News
  - COVID-19 Report for Life Sciences and Health Care Companies (Jan - Feb 2022)
- Press Releases
  - Hogan Lovells advises Soliton on its US\$550 million acquisition by Allergan Aesthetics
- News
  - FDA warns over use of robotically-assisted surgical devices
- Press Releases
  - Hogan Lovells advises Soliton in US\$550 million acquisition by AbbVie's Allergan Aesthetics

Premarket Review

Cell, Tissue, and Gene Therapies

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## Education and admissions

### Education

J.D., Georgetown University Law Center, magna cum laude, Order of the Coif, 1995

B.S. Mechanical Engineering and Literature, Massachusetts Institute of Technology, 1988

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### Memberships

Member, American Bar Association

Member, Maryland State Bar Association

Member, National Health Lawyers Association

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### Bar admissions and qualifications

Pennsylvania

District of Columbia

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