

# **Premarket Approval (PMA) Submissions Workshop**

## Virtual Event March 1-3, 2022

\*Schedule Reflected in Eastern Time

#### March 1, 2022

11:00 am - 11:05 am Welcome and Introductions

11:05 am - 12:00 pm Beginning at the Beginning

When is a De Novo or PMA required

PMA: what to expect

What are the standards of evidence

What are the standards of review

o Will submission go to panel

o How much will it cost

How long will it take to get approval

12:00 pm – 12:45 pm Development of a PMA Submission Strategy

Product definition

Development of testing requirements and strategy

Desired patient population

Desired claims

Early interactions with FDA

Planning for product iterations

12:45 pm - 1:00 pm Q&A

1:00 pm - 1:15 pm Break

1:15 pm – 1:55 pm Mechanics of PMA Quality System Submission Development and Review

- Defining data requirements
- Required elements
- Presentation of information with clarity
- · Expectations during review
- Best practices
- Manufacturing & Quality Systems
- Case for Quality

#### **Important Notice**

The information provided in this course represents the personal opinions of the instructors and does not necessarily represent the opinions of AdvaMed staff. Companies relying on the information do so at their own risk and assume the risk of any subsequent liability that results from relying on the information. The information does not constitute legal advice.



1:55 pm - 2:40 pm

#### **During Submission Review**

- Interactions with the FDA
- When/How to expect questions
- Types of letters
- Timelines
- Day 100 meetings
- · Labeling review

2:40 pm - 2:55 pm

Q&A

2:55 pm - 3:10 pm

**Break** 

3:10 pm - 3:55 pm

### **Conditions of Approval Studies**

- Criteria and objectives
- Early collaboration with FDA
- Reaching agreement
- Reporting outcomes
- 522 Studies

3:55 pm - 4:40 pm

#### **Preparation for Advisory Panels**

- When?
- Who are the panel members?
- Why have a panel meeting?
- Preparation for a panel meeting
- What to expect before, during, and after
- Best practices

4:40 pm - 4:55 pm

Q&A

#### March 2, 2022

11:00 am - 11:45 am

#### **Inspection Activity**

- Pre-approval inspections
- How to prepare for an inspection

11:45 am - 12:45 pm

#### **Dealing with the Unexpected**

- Clinical outcomes
- Animal test results
- Adverse panel recommendation

12:45 pm - 1:00 pm

Q&A

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1:00 pm – 1:15 pm Break

1:15 pm – 2:00 pm The Care and Feeding of Approved PMAs

- Periodic ("Annual") ReportsSupplemental Submissions
- 30-day notices

2:00 pm - 2:30 pm CDRH Ombudsman's Office

- Roles & Responsibilities
- The Appeals Process

2:30 pm- 2:45 pm Q&A

2:45 pm Closing Remarks and Adjourn

#### March 3, 2022

12:00 pm – 1:15 pm Applied Learning and Discussion

- Hypothetical Case Study Review
- Facilitated Breakout Group Deep Dive
- Key Takeaways
- Final Program Q&A

#### **Important Notice**