# Challenges and Achievement: Japanese Regulatory View

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## Agenda

- 1. Background: What is burdens for development of pediatric medical devices?
- 2. Measures to meet unmet medical needs in Japan
- 3. Significance of HBD-for-children activity
- 4. Conclusion



#### What is the burdens for development of pediatric medical devices?

We have done a questionnaires about development of pediatric medical devices to industries in the Japan and U.S. through HBD for children activities.





"Small market" is majority opinion.

Variety of opinions are expressed.

- Development cost
- Application and approval
- Clinical trial
- Designing of device



### Measures to meet unmet medical needs in Japan

2006

 Program for requesting medical devices with high clinical needs started in 2006

2013

- Application with a clinical evaluation report started in 2008 (Notification by MHLW, No. 0804001)
- Support program for medical device used for treatment of orphan disease started in 2013
- New Ministerial Ordinance for Enforcement of the Act on Securing Quality, Efficacy and Safety of Pharmaceuticals, Medical Devices, Regenerative and Cellular Therapy Products, Gene Therapy Products, and Cosmetics in 2014
- Scientific advisory board about pediatric medical devices was held in 2014-2015
  - SAKIGAKE Designation System started in 2015
  - Fast-break scheme for innovative medical devices started in 2017
  - Subsidization program for application of pediatric medical device started in 2019

2019



#### Fast-break scheme for innovative medical devices

	Original	Fast-break scheme
Clinical data which is submitted when the device is applied	a prospective clinical trial under GCP or clinical evaluation report	Reliable clinical data (GCP trial is not always necessary)
Review term	12 months (if brand-new medical device)	
Post-marketing surveillance	If need	Necessary
Collaboration with academic society at the post-marketing phase	If need	Necessary
Increase facilities by stages for safety use of the medical device at the post-	If need	Necessary

The program was legalized in the Pharmaceutical and Medical Device Act last month!

Collaboration between regulatory, clinicians and manufactures is necessary!

marketing phase

## Subsidization program

Very new!

**◆**Target devices

A medical device for treatment of pediatric or congenital disease which was selected by a program for requesting medical devices with high clinical needs

**♦**Subsidization

MHLW subsidizes maximum 90% of a regulatory application fee of pediatric medical devices.

(The fund is limited.)

The program just started in September 2019!!



# Significance of the HBD-for-children activity

Scientific sessions related in the activity were held in following conferences!











Activity of HBD-for-children has been globally recognized.



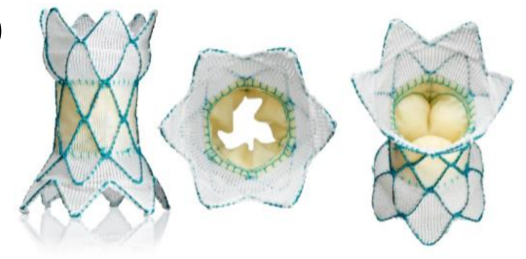
# Significance of the HBD-for-children activity

The collaboration among regulatory, industry, and academia in both Japan and U.S.A has made it easier to conduct global clinical trials.

First Japan-US global clinical trial of pediatric medical device started!

The Harmony™ TPV system (Medtronic, Inc)

Self-Expanding Stent Graft Porcine Pericardial Tissue Valve





# Significance of the HBD-for-children activity



Title: "Research project for promotion of early development of pediatric medical devices"

#### 1. Finding issues and solutions of clinical trials in the pediatric field

- Capturing the issues around Japan-US clinical trial of pediatric medical devices and proposing the solutions by the following actions
  - Questionnaire to pediatricians and industries
    Clarifying the burdens
  - Proposal for the solutions to the challenges
- Finding similarities and differences of GCP inspection processes between Japan and USA by observing the GCP inspection by FDA and PMDA each other

#### 2. Utilization of real world evidence

- Proposing requirements for utilization of real-world-data of pediatric medical devices by the following actions
  - Hearing survey from registries of Japan-US
  - Investigation past experiences of using registry data for regulatory use
  - Measuring for assuring minimum quality and quantity of registry data



## Goal of the research project

#### Making suggestions for effective development

- 1. Streamlining the process of clinical trial
  - Infrastructure development of clinical trial
  - Efficient GCP inspection for global clinical trial
- 2. Utilization of registry database
  - Use for post-marketing surveillance
  - Continual organizational operation of registry
- Proposal for consistent rational evaluation methods for pre- and post-marketing of pediatric medical devices



#### Conclusion

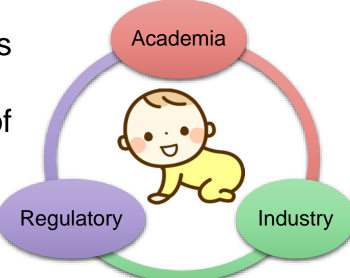
 Japanese regulatory carried out some enforcement of policies to support the development of pediatric medical devices.

 On the issue of funds, Japanese regulatory remits the 90% of fee for review and inspection.

 HBD-for-children activity contributes to the development of pediatric medical devices by supporting the Japan-US global clinical trial.

Research project related in the activity has started.

We will continue to seek a possibility of global clinical trials of pediatric medical devices and utilization of real-world evidence for an early approval of pediatric medical devices in the Japan and U.S..







# Thanks for your kind listening!

Please contact me or HBD secretariat, if you have any questions.

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Let's continue to think about highly feasible measures together by sharing information through industry-regulatory-academia and conducting frank discussions.