



IMDRF International Medical
Device Regulators Forum

Japan Update

- **New measures to improve access to innovative MDs/IVDs -**

September, 2015





Topics

1. New international regulatory harmonization strategies by MHLW and PMDA
2. Official participation in MDSAP Pilot
3. Implementation of Strategy of Sakigake
4. Clinical Innovation Network



1. Establishment of new international regulatory harmonization strategies by MHLW and PMDA

International Regulatory Harmonization Strategy by MHLW and PMDA International Strategic Plan 2015 by PMDA have been published on 26 June 2015.

Based on the *mutually complementary* strategies, the following measures will be taken:

A) Promotion of Regulatory Science

Guidelines related to medical device regulations in Japan will be prepared and internationally announced.

B) Establishment of Training Center for regulatory matters

PMDA will provide regulators outside Japan with training for capacity building.

C) Active commitment to IMDRF as well as advancement of bilateral collaboration

*IMDRF has been reaffirmed as one of the most important activities.*³



2. Official Participation in MDSAP

Japan made an announcement on the [official participation in MDSAP Pilot](#) on 23 June 2015.

Further information will be provided in a timely manner.

Shown at right is the press release on the official participation in MDSAP Pilot in Japan (written in Japanese).

You can find the announcement in English here;

<http://www.fda.gov/MedicalDevice/InternationalPrograms/MDSAPPilot/ucm452243.htm>



Press Release

平成 27 年 6 月 23 日

【照会先】

医薬食品局医療機器・再生医療等製品担当参事官室

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Medical Device Single Audit Program Pilot に正式参加します

～国際協力の下、医療機器の品質確保を推進～

医療機器の品質確保に関する国際協力活動として、2014年1月から米国、カナダ、オーストラリア及びブラジルにより「MDSAP (Medical Device Single Audit Program (医療機器単一調査プログラム)) Pilot」が試行的に運用されています。本日、米国ワシントンDCで行われるMDSAPフォーラムの場で、日本も正式メンバーとして本プログラムに参加することを表明します。今後とも本活動の下で、より一層の医療機器の品質確保を徹底してまいります。



3. Implementation of Strategy of Sakigake

An *innovative MD/IVD for patients in urgent need of innovative therapy* may be designated as a Sakigake Product if;

- 1) its premarket application will be filed in Japan firstly or simultaneously in some countries including Japan, *AND*
- 2) prominent effectiveness can be expected.

Once an MD/IVD is designated, its developer can enjoy such benefits as:

A) Prioritized Consultation by PMDA

C) Prioritized Review
(12 months → 6 months [MD])

B) Pre-application substantive review

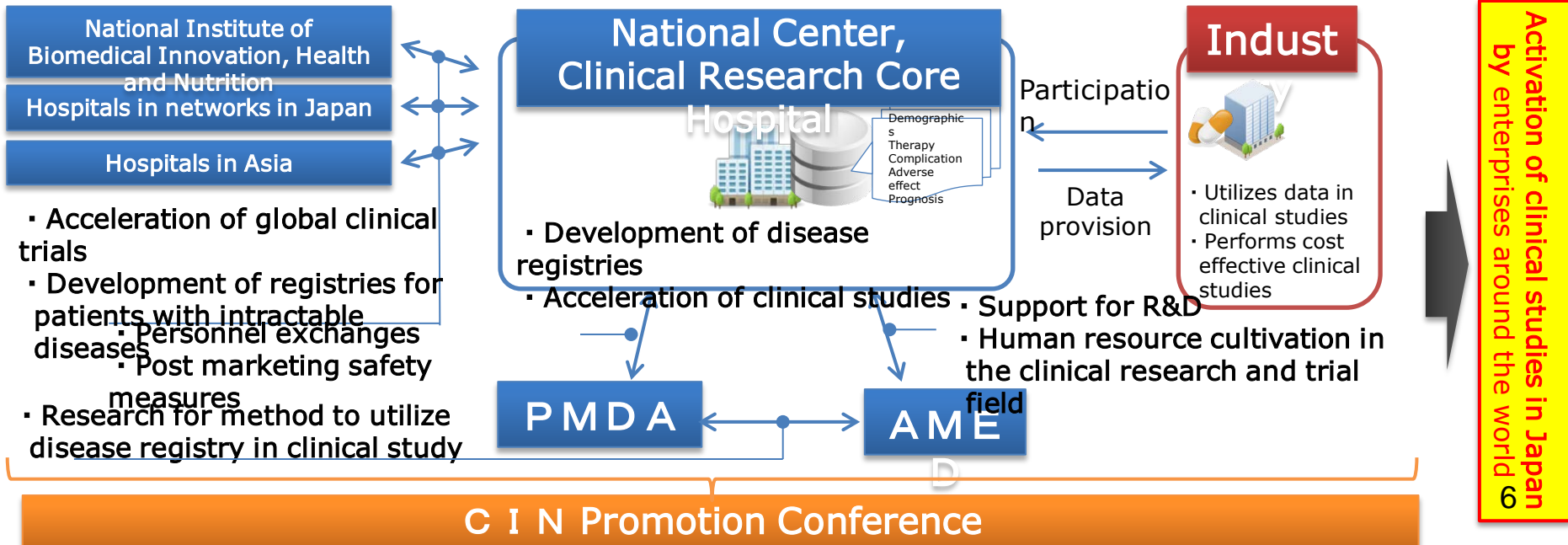
D) Review Concierge assigned
by PMDA



4. Clinical Innovation Network (CIN)

The clinical study infrastructure in Japan will be improved under the CIN project so that cost effective clinical studies can be performed with disease registries, based on Regulatory Science.

The improvement will accelerate clinical studies in Japan by enterprises around the world, which would result in the contribution to extended healthy life expectancy for people.





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Thank you



MHLW



PMDA