

Regulatory Update of Regenerative Medicine in Japan

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Outline

• The Act on the Safety of Regenerative Medicine

The Act on Pharmaceuticals and Medical Devices

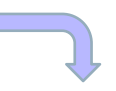


Dual Regulation of Regenerative Medicine in Japan

Technology & Product



Regenerative Medicine



Enacted on 25 November 2014

All medical **technologies** using processed cells which safety and efficacy have not yet been



Safety Act

The Act on the Safety of Regenerative Medicine

Medical Care or Academic Research Purpose Production and marketing of regenerative and cellular therapeutic **products** by firms





The Act on Pharmaceuticals and Medical Devices



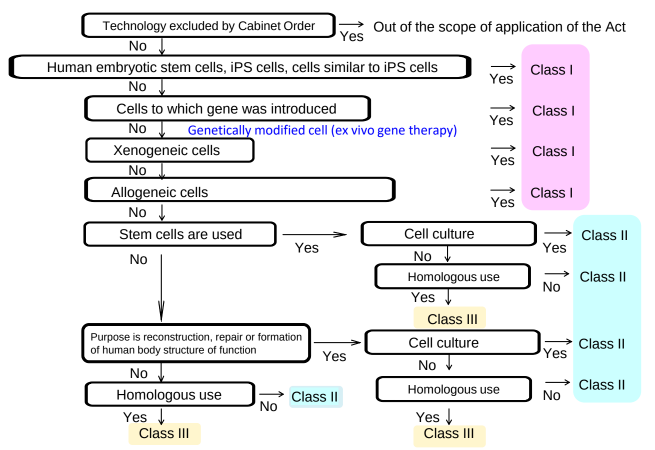
Commercial Product

Marketing Authorization Purpose



Risk Classification Regenerative Medical Technology





Medical Care	Clinical Research
7	16
1,571	43
3,898	42
	7 1,571

(As of 1 December 2023)

Current Activity

WG has been establish to review the Safety act, and studies are being conducted. *in vivo* gene therapy is currently out of scope. *in vivo* gene therapy is also under consideration for inclusion in the scope of Safety act.



Approved Plans (Class I, Clinical Research)



iPS cells

Health Condition(s) or Problem(s) Studied	Hospital/Clinic	Trial ID
Aplastic anemia with platelet transfusion refractoriness due to anti-platelet alloantibo	Kyoto University Hospital	jRCTa050190117
Bullous keratopathy	Keio University	jRCTa031210199
Damage of articular cartilage of the Knee	Kyoto University Hospital	jRCTa050190104
Limbal stem-cell deficiency	Osaka University Graduate School of Medicine	jRCTa050190084
Recurrent or advanced head and neck cancer	Chiba University Hospital	jRCTa030220741
Retinitis pigmentosa	Kobe City Eye Hospital	jRCTa050200027
RPE impaired disease	Kobe City Eye Hospital	jRCTa050210178 jRCTa050200122
Severe heart failure patients with NYHA class III or higher (HFrEF by dilated cardiomyopathy)	Keio University School of Medicine	jRCTa032200189
Spinal cord injury at subacute stage	Keio University School of Medicine	jRCTa031190228

Autologous cells to which gene was introduced

Health Condition(s) or Problem(s) Studied	Hospital/Clinic	Trial ID
CD19 positive acute lymphoblastic leukemia	Nagoya University Graduate School of Medicine	jRCTa040190099
Familial LCAT deficiency	Chiba university	jRCTa030190230

Allogopoio colle

Allogeneic cells		
Health Condition(s) or Problem(s) Studied	Hospital/Clinic	Trial ID
Bullous keratopathy	Kyoto Prefectural University of Medicine	jRCTa050190118
Cartilage defects associated with osteoarthritis of the knee	Tokai University	jRCTa030190242
Cerebral palsy	Kochi Medical School Hospital	jRCTa060200017 jRCTa060200018
Decompensated liver cirrhosis patients who will receive liver transplantation	Hiroshima University	jRCTa060190036 jRCTa030230316
EBV associated lymphoproliferative disorder after allogeneic stem cell transplantation	Nagoya University Graduate School of Medicine	jRCTa040190110
Hematological disorders applicable to cord blood transplantation	Aichi Medical University	jRCTa040190133
Idiopathic peripheral ulcer and cGVHD related corneal ulcer	Keio University	jRCTa030220167
Insulin-dependent diabetes mellitus	Shinshu University School of Medicine	jRCTa030190164
Persistent viral infection after hematopoietic cell transplantation	Tokyo Medical and Dental University	jRCTa030190229
Refractory chronic skin ulcer	Keio University School of Medicine	jRCTa030200053
Refractory cutaneous ulcers	Research Institute for Radiation Biology and Medicine(RIRBM), Hiroshima University	jRCTa060210018

Source: https://jrct.niph.go.jp/search

Search condition: Clinical Research for Regenerative Medicine (Class I)

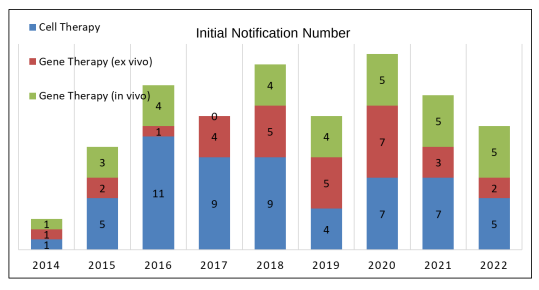


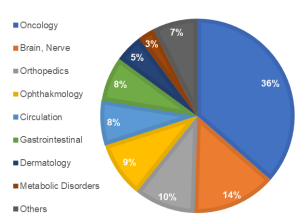
INDs Reviewed by PMDA

/	PMD	
\	Act	

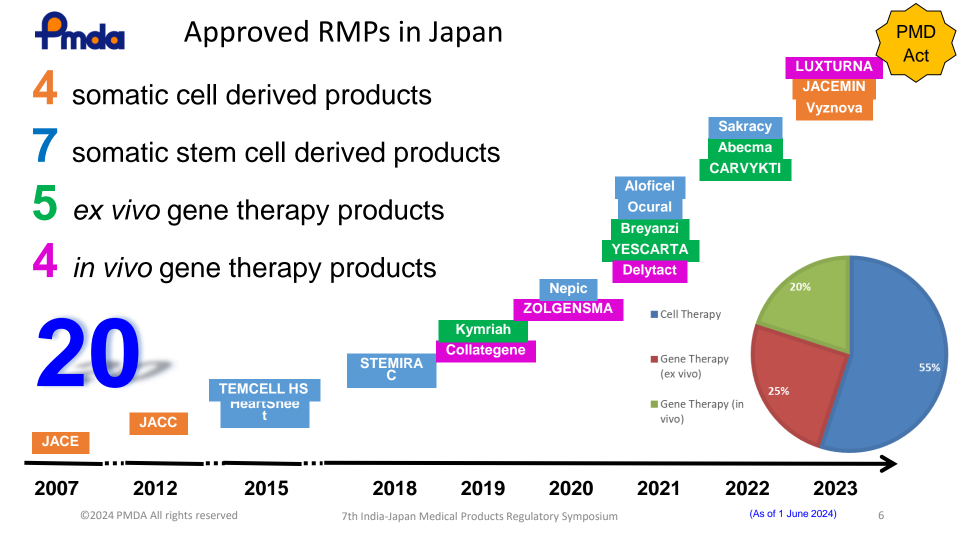
Notification	2014	2015	2016	2017	2018	2019	2020	2021	2022	Total
Initial	3 [1]	10 [2]	16 [7]	13 [8]	18 [8]	13 [7]	19 [9]	15 [7]	12 [3]	119 [51]
2 nd or later	1 [1]	3 [2]	5 [0]	14 [10]	17 [3]	16 [7]	22 [5]	18 [9]	25 [14]	121 [51]

Note: The table in brackets in parentheses indicate the number of notifications of "investigator-initiated clinical trials (IIT).





Area of disease





Accelerating Approval Pathways for RMPs in Japan



	Priority review (Orphan)	SAKIGAKE	Conditional & Time-limited Approval
Abecma	✓		
Breyanzi	✓		
CARVYKTI	✓		
Delytact	✓	✓	✓
Kymriah	✓		
YESCARTA	✓		
LUXTRNA	✓		
Nepic	✓		
Ocural	✓		
Sakracy	✓		
Vyznova	✓		
STEMIRAC		✓	✓
ZOLGENSMA	✓	✓	
Collategene			✓
HeartSheet			✓
JACE	✓ GCMN*, EB**		
JACEMIN			
Aloficel	✓		
JACC			
TEMCELL	✓		

Area of disease	# of products
Oncology	6
Ophthalmology	5
Brain, Nerve	2
Circulation	2
Dermatology	2
Others	3

The target review time Priority products: 9 month Regular products: 12 month

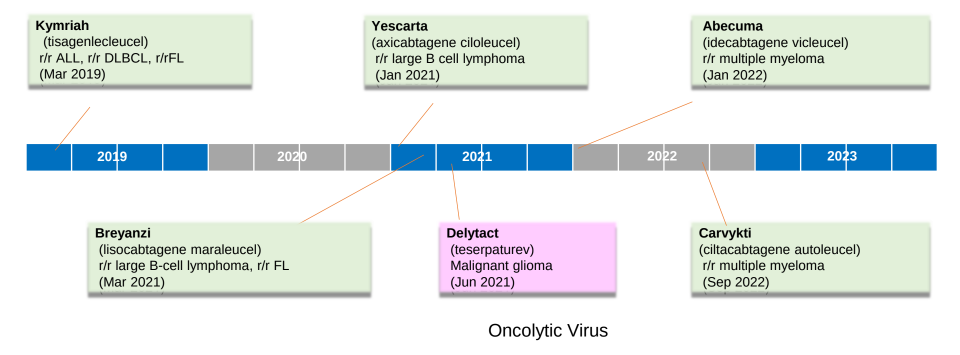
*GCMN: Giant congenital melanocytic nevi

**EB: Dystrophic epidermolysis bullosa



Oncology Area







Ophthalmology Area





https://www.jpte.co.jp/business/regenerative/

Nepic

(Human (autologous) corneal limbus-derived corneal epithelial cell sheet) (Mar 2020)

Epithelial cell sheet for limbal stem cell deficiency (LSCD), a rare and intractable corneal epithelial disease



Ocural

https://www.jpte.co.jp/business/regenerative (Human (autologous) oral mucosa-derived epithelial cell sheet) (Jun 2021)



Sakracy

http://hirosaki-li.co.jp/products_sakracy.html

(Human (autologous) oral mucosa-derived epithelial cell sheet using human amniotic membrane substrate)

Aketa et al., The Ocular Surface, 29, 220-225 (2023)

(Jan 2022)

2020 2021 2022 2023

Vyznova

(Human (allogenic) corneal endothelium-derived endothelial cell injection) (Mar 2023)

Endothelial cell injection for Bullous Keratopathy, a rare and intractable corneal endothelial disease

LUXTURNA

(voretigene neparvovec) Confirmed biallelic RPE65 Mutation-associated retinal dystrophy (Jun 2023)

Adeno-associated virus (AAV) vector



Summary of Approved RMPs in Japan

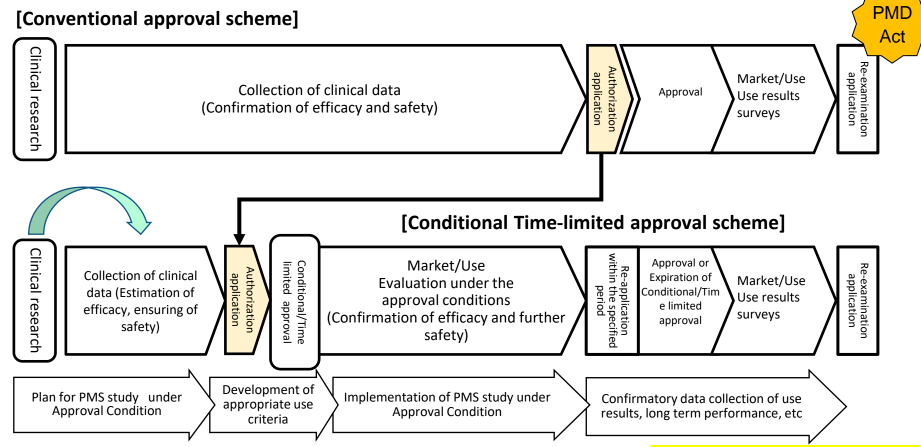


	Japan Origin (12)	Global (8)
Cell Therapy Products (16)		
Somatic cell derived products (9)	JACE (Skin) JACC (Knee) JACEMIN (Skin) Vyznova (Eye)	Kymriah YESCARTA Breyanzi Amecma CARVYKTI
Somatic stem cell derived products (7)	HeartSheet* TEMCELL HS STEMIRAC* Nepic (Eye) Ocural (Eye) Sakracy (Eye)	Aloficel
iPS cell derived products (0)		
Embryonic stem cell derived products (0)		
Gene Therapy Products (4)		
Plasmid vector products (1)	Collategene*	
Viral vector products (2)		ZOLGENSMA LUXTURNA AAV
Others (1)	Delytact*	

^{*}Conditional and Time limited Approval



Early Access Scheme; Conditional and Time-limited Approval





Outline of the Condition for Approval and Granted Time-period for PMS Study



Products	HeartSheet	Stemirac	Collategene	Delytact
	Treatment of patients with severe heart failure due to ischemic heart disease unresponsive to standard		The treatment of ulcers in patients with chronic arterial occlusion	Malignant glioma
Granted time- period	8 years (17/09/2023) (Extend on 20/11/2018 after hearing the opinion of the Pharmaceutical Affairs and Food Sanitation Council)	7 years (27/12/2025)	5 years (25/03/2024)	7 years (10/06/2028)
		Efficacy evaluation		
Primary endpoint	Time to cardiac death (at ≥ 2 years post transplantation)	Cohort I; Patients with AIS Grade A at 6 to 8 weeks (49 ± 7 days) after injury Percentage of patients achieving ≥ 2 grade improvement in AIS at 180 ± 30 days from 6 to 8 weeks (49 ± 7 days) after injury Cohort II; Patients with AIS Grade B or C at 6 to 8 weeks (49 ± 7 days) after injury Percentage of patients with AIS Grade B or C achieving ≥ 1 grade improvement in AIS at 180 ± 30 days from 6 to 8 weeks (49 ± 7 days) after injury	The proportion of patients with completely closed ulcer at 12 week later after injection	OS (from the day of diagnosis of malignant glioma to death [from any cause]): For each population of patients with primary glioblastoma and patients with recurrent glioblastoma, conduct a trend score matching so that the Delytact and control groups include the same number of patients (1:1), and perform a log-rank test with the two-sided significance level of 5% on OS in the sample population.
		Number of subject		
Product	60	Cohort I; 27 Cohort II; 63	120	Glioblastoma: 250 Grade III malignant glioma: 60 to 100
Control (External)	120	Cohort I; 54 Cohort II; 125	80	Glioblastoma: 500 Grade III malignant glioma: 120 to 200



Benefit and Risk Balance Assessment



- Discussion of acceptable level of clinical effectiveness vs.
 patient access to the new therapy.
- Weighing acceptable risk against expected benefit.



 Considerations for Conditional and Time-limited Approval of Cell and Gene Therapy Products, PSEHB/MDED Notification No. 0329-3, 2024.

https://www.pmda.go.jp/files/000267914.pdf

 Guidance on Conditional and Time-limited Approval and Protocol for Post Marketing Study of Stem Cell Therapy Products, PSEHB/MDED Notification No. 0329-4, 2024.

https://www.pmda.go.jp/files/000267915.pdf



Regulators' Action to Novel Technologies (1)





- Current Perspective on Evaluation of **Tumorigenicity** of **Cellular and** Tissue-based Products Derived from induced Pluripotent Stem Cells (iPSCs)* and iPSCs as Their Starting Materials (20 August 2013)
- Proposal on Basic Principle to Quality Assurance of Cell Therapy (CT) Products (14 June, 2015)

CAR-T for blodd cancers (2017-)

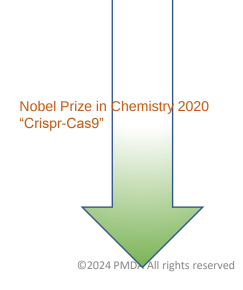
(PMDA Science Board has started the discussion on gene editing (2018))

- Points to Consider for Ensuring Quality and Safety of Gene Therapy Products (9 July, 2019)
- Report on Considerations on Quality and Safety of Gene Therapy **Products** Using Genome Editing Technology (7 February, 2020)

(Discussion on vectors with targeting function has begun(2023))

 Points to Consider for Development of in vivo Gene Therapy **Products** with Target Specific Viral/non-Viral Vector (To be publish in 2024)

https://www.pmda.go.jp/english/rs-sb-std/sb/subcommittees/0031.html





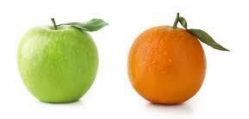
Regulators' Action to Novel Technologies (2)

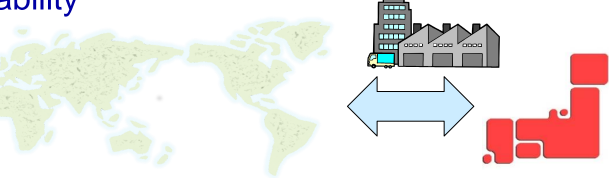


Comparability









 Comparability of Cell Therapy Products Subject to Changes in Their Manufacturing Process PSEHB/MDED Notification No. 0329-1, 2024.

https://www.pmda.go.jp/files/000267916.pdf



Summary

- Medical technologies using processed cells are active under the Safety Act.
- 20 RMPs, including 5 *ex vivo* and 4 *in vivo* gene therapy products, have been approved under the PMD Act.
- 4 of 20 RMPs have been approved through comprehensive framework for patient access (conditional and time-limited approval scheme).
 - Sponsors are subject to strict PMS study to prepare re-marketing authorization submission with in the granted time-period.
- According to R&D trends, the GLs related to RMPs development have been prepared.



Where to Find Information?

Regenerative Medical Products

1. Regulatory Framework

Regenerative medicine, which is expected to overcome intractable and serious diseases, is expected to play an important role in conventional medicine worldwide. The Japanese government must implement comprehensive policies to promote the development of regenerative medicine, inform the public, and increase public acceptance, and ensure that medical professionals and investigators cooperate with the policies. In this background, two regulatory frameworks for regenerative medicine, "The Act on the Safety of Regenerative Medicine 37" (ASRM) and the "Pharmaceuticals, Medical Devices, and Other Therapeutic Products Act 57" (PMD Act), came into effect in November 2014. The ASRM sets out legal regulations not only for research, but also for the daily medical practice of cell therapy, which had previously been under the jurisdiction of the Medical Practitioners' Act 57" and the Medical Care Act 57".

The PMD Act regulates the commercialization of regenerative medical products. Regenerative medical products in the PMD Act are defined as:

- a. Processed (more than minimal manipulation) live human/animal cells that are intended to be used for either
 - o reconstruction, repair, or formation of structures or functions of the human body



Review Reports: Regenerative Medical Products

Α

Brand Name	Non-proprietary Name	Approved In	English	Japanese
Abecma	idecabtagene vicleucel	January 2022	™	7
Alofisel	darvadstrocel	September 2021	72	1

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В

Brand Name	Non-proprietary Name	Approved In	English	Japanese
Breyanzi Initial Approval	lisocabtagene maraleucel	March 2021	Z	B
Breyanzi Partial Change Approval	lisocabtagene maraleucel	December 2022	™	7 2



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Recent Publication

- Maruyama Y, Noda S, Okudaira S, Sakurai A, Okura N, Honda F. Regulatory Aspects of Cell and Gene Therapy Products: The Japanese Perspective, Adv Exp Med Biol, 1430, 155-179 (2023) https://doi.org/10.1007/978-3-031-34567-8_9
- Maruyama Y, Sakurai A, Noda S, Fujiwara Y, Okura N, Takagi T, Asano J, Honda F. Regulatory Issues: PMDA Review of Sakigake Designation Products: Oncolytic virus therapy with Delytact Injection (teserpaturev) for malignant glioma, The Oncologist, 28(8) 664-670 (2023) https://doi.org/10.1093/oncolo/oyad041
- Aketa N, Kasai M, Noda S, Asano J, Kunieda A, Kawanishi S, Maruyama Y, Honda F. Insights Into the Clinical Development of Regenerative Medical Products Through a Comparison of Three Cell-based Products Recently Approved for Limbal Stem Cell Deficiency. The Ocular Surface, 29, 220-225 (2023) https://doi.org/10.1016/j.jtos.2023.05.008
- Sakurai A, Kanzaki S, Honda F. Japanese pharmaceutical regulations of engineered viral vectors for medical use compared with those in the US and EU. Clinical Pharmacology & Therapeutics (2023) https://doi.org/10.1002/cpt.2788
- Fujiwara Y, Maruyama Y, Honda F. Balancing safety and efficacy with early availability in the regulation of regenerative medicine product. Clin Pharmacol Ther, 109:1182-1185 (2021). https://doi.org/10.1002/cpt.2034



Thank you for your attention!

Please visit the PMDA website
http://www.pmda.go.jp
http://www.pmda.go.jp/english/index.html

