

FDA WEEKLY INTELLIGENCE

# FDA Intelligence

Coverage: 2026-08-06 – 2026-08-12 (drug approval date basis)

Published: August 16, 2026

Plan: Individual Plan

TABLE OF CONTENTS

|                                                                                                                                                |    |
|------------------------------------------------------------------------------------------------------------------------------------------------|----|
| This Week at FDA .....                                                                                                                         | 2  |
|  Asian Company Watch — FDA Activity This Week .....           | 3  |
|  Drug Approvals .....                                         | 7  |
|  Medical Devices .....                                       | 8  |
|  Warning Letters .....                                      | 9  |
|  Drug Recalls .....                                         | 10 |
|  FDA Guidance & Regulatory Actions .....                    | 12 |
|  Upcoming FDA Regulatory Event Calendar .....               | 13 |
| —  Weekly Outlook .....                                     | 13 |
| —  This Week's Spotlight (Editor's Picks) .....             | 14 |
| —  Changes Since Last Week .....                            | 15 |
| —  Upcoming FDA Schedule (Official, Not Predicted) .....    | 16 |
| —  PDUFA Target Action Dates Disclosed by Sponsors .....    | 17 |
| —  Annual Guidance Development Plans (CDER/CBER/CDRH) ..... | 21 |
|  Terms of Use & Disclaimer (Individual Plan) .....          | 23 |

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# FDA Weekly Report — Week 33, 2026

Coverage: 2026-08-06 – 2026-08-12 (drug approval date basis)

Published: August 16, 2026

*Note: the reporting period above is based on drug (NDA/BLA) approval dates. FDA typically publishes approvals 1-2 weeks after the decision date. Device data (510(k)/PMA) is published on an even longer lag, so the Medical Devices section covers 2026-07-25 to 2026-07-31.*

*Note: the issue number ("Week 33, 2026") is the ISO calendar week containing the END of the coverage period above, used only for stable issue numbering — it is not a claim that the coverage period spans that full Monday-Sunday week.*

## This Week at FDA

The FDA's drug approval activity was notably silent this week, with no new molecular entities or other novel drugs receiving marketing authorization.

0

NME Approvals

0

Total Drug Approvals

0

Efficacy Supplements

75

Device Clearances

1

PMA Originals

17

Asian Co. Events

## Asian Company Watch — FDA Activity This Week

Coverage: drugs 2026-08-06–2026-08-12, devices 2026-07-25–2026-07-31 (device data is published on a longer lag).

Asian companies with FDA approvals or clearances this week. The highest-priority items below include strategic commentary on what it means for the competitive landscape; the remaining approvals are listed for completeness (full detail for every item is available in the accompanying Excel data set). Country classification is based on ultimate corporate group origin, not necessarily the filing legal entity's place of incorporation (e.g. a US-registered subsidiary of a Japanese parent group is classified as Japanese).

### **AUROBINDO PHARMA LTD** India

#### TRIAMCINOLONE ACETONIDE

AUROBINDO PHARMA LTD — Generic approval (ANDA) for TRIAMCINOLONE ACETONIDE (TRIAMCINOLONE ACETONIDE).

**Intelligence:** This approval is part of Aurobindo's consistent strategy of bringing a broad portfolio of generic pharmaceuticals to the US market.

### **AUROBINDO PHARMA** India

#### BRIVARACETAM (BRIVARACETAM)

AUROBINDO PHARMA — Generic approval (ANDA) for BRIVARACETAM (BRIVARACETAM).

### **AUROBINDO PHARMA** India

#### VANCOMYCIN HYDROCHLORIDE

AUROBINDO PHARMA — Generic approval (ANDA) for VANCOMYCIN HYDROCHLORIDE (VANCOMYCIN HYDROCHLORIDE).

### **LUPIN** India

#### SODIUM ZIRCONIUM CYCLOSILICATE

LUPIN — Generic approval (ANDA) for SODIUM ZIRCONIUM CYCLOSILICATE (SODIUM ZIRCONIUM CYCLOSILICATE).

**DR REDDYS** India**LORATADINE**

DR REDDYS — Generic approval (ANDA) for LORATADINE (LORATADINE AND PSEUDOEPHEDRINE SULFATE).

**Samsung Medison Co., Ltd.** South Korea**V8/XV8/XH8/H8/cV8/V8 ONE/XV8 ONE/XH8...**

 3rd clearance in 2026 (73 total since 2011) — established filer

 Predicate: K250999 (Samsung Medison Co., Ltd., KR)

South Korea's Samsung Medison received 510(k) clearance for its V-series line of pulsed Doppler diagnostic ultrasound systems.

**Intelligence:** This clearance for a wide range of models suggests a strategy to offer a tiered portfolio of ultrasound systems to different segments of the US healthcare market.

**Shanghai United Imaging Healthcare Co., Ltd.** China**uMI Panvivo (uMI Panvivo M)**

 6th clearance in 2026 (84 total since 2018) — established filer

 Predicate: K241166 (Shanghai United Imaging Healthcare Co., Ltd., CN)

Shanghai United Imaging Healthcare Co., Ltd. received a 510(k) clearance for uMI Panvivo (uMI Panvivo M). FDA classifies this device as: System, Tomography, Computed, Emission.

**Nipro Medical Corporation** Japan**ELISIO™ -HX**

 3rd clearance in 2026 (35 total since 2005) — established filer

 Predicate: K260533 (Nipro Medical Corporation, US)

Japanese firm Nipro Medical Corporation received 510(k) clearance for its ELISIO-HX, a hemodialyzer with an expanded solute removal profile for dialysis treatment.

**Intelligence:** This clearance adds a new high-flux dialyzer to Nipro's US product line, potentially strengthening its offerings for the nephrology and dialysis market.

**NeuroLogica Corp., a Samsung Company** South Korea

## BodyTom Elite (NL4000)

 First FDA clearance — new US market entrant Predicate: K170238 (Neurologica Corporation, A Subsidiary of Samsung Electronics, US)

NeuroLogica Corp., a US-based Samsung company, received 510(k) clearance for its BodyTom Elite mobile computed tomography (CT) x-ray system.

**Intelligence:** This clearance for a portable CT scanner could enhance Samsung's portfolio in the point-of-care and intraoperative imaging segments of the US market.

**T&R BIOFAB CO., Ltd.** South Korea

## TnR Cranial Implant

 1st clearance in 2026 (2 total since 2023) — established filer Predicate: K051093 (Osteopore, Inc., US)

South Korean company T&R BIOFAB CO., Ltd. secured 510(k) clearance for its TnR Cranial Implant, a methyl methacrylate device for cranioplasty.

**Intelligence:** This clearance provides a US market entry point for T&R BIOFAB in the specialized field of neurosurgical cranial reconstruction implants.

**IMGFAX CO., LTD.** China

## Handheld Ultrasound Scanner

 First FDA clearance — new US market entrant Predicate: K241161 (Leltek, Inc., TW)

China-based IMGFAX CO., LTD. obtained 510(k) clearance for its handheld pulsed Doppler ultrasound scanner.

**Intelligence:** The clearance of a portable ultrasound device could position IMGFAX to compete in the growing US market for point-of-care and mobile diagnostic imaging solutions.

**Rayence Co., Ltd.** South Korea

## 0909FCE, 0909FCE-HS

 1st clearance in 2026 (39 total since 2011) — established filer Predicate: K240371 (Rayence Co., Ltd., KR)

South Korea's Rayence Co., Ltd. received 510(k) clearance for its solid-state flat-panel digital x-ray imagers.

**Intelligence:** As a key component manufacturer, this clearance for its digital detectors may support Rayence's business with medical and dental imaging system integrators in the US.

**Suzhou Haowei Medical Technology Co., Ltd.** China

## UniPearls® Embolic Microspheres

✓ 1st clearance in 2026 (2 total since 2023) — established filer

🔗 Predicate: K231554 (Suzhou Haowei Medical Technology Co., Ltd., CN)

Suzhou Haowei Medical Technology from China received 510(k) clearance for its UniPearls Embolic Microspheres, a vascular device used for embolization procedures.

**Intelligence:** This clearance marks an entry for Suzhou Haowei into the US interventional radiology market with a device used in procedures such as tumor treatment.

**Shanghai Hulu Devices Co., Ltd.** China

## Smart Wireless Stethoscope

✓ 1st clearance in 2026 (3 total since 2018) — established filer

🔗 Predicate: K220470 (Shanghai Hulu Devices Co., Ltd., CN)

Shanghai Hulu Devices Co., Ltd. obtained 510(k) clearance for its line of smart wireless electronic stethoscopes.

**Intelligence:** The introduction of multiple electronic stethoscope models could position the Chinese company to serve the US telehealth and digital health monitoring markets.

**Olympus Medical Systems Corporation** Japan

## EVIS EUS Ultrasound Gastrointestinal Videoscope

✓ 3rd clearance in 2026 (83 total since 2004) — established filer

🔗 Predicate: K251859 (Olympus Medical Systems Corporation, JP)

Olympus Medical Systems Corporation received a 510(k) clearance for EVIS EUS Ultrasound Gastrointestinal Videoscope (Olympus GF-UE190). FDA classifies this device as: Endoscopic Ultrasound System, Gastroenterology-Urology.

**Samsung Electronics Co., Ltd.** South Korea

## Hearing Aid Feature

✓ 2nd clearance in 2026 (42 total since 2013) — established filer

Samsung received 510(k) clearance for software that functions as an air-conduction hearing aid.

**Xiamen High Top Electronic Technology Co., Ltd.** **China**

**Air Compression Massager (HY-1165)**

 1st clearance in 2026 (2 total since 2022) — established filer

 Predicate: K182668 (Rapid Reboot Recovery Products, LLC, US)

Xiamen High Top received 510(k) clearance for a powered inflatable tube massager.

 **Drug Approvals**

0 new drug approvals this week (0 NMEs). 0 efficacy supplements. 9 generics (ANDA).

 Medical Devices

Coverage period for this section: **2026-07-25 to 2026-07-31** — FDA publishes device decisions later than drug approvals, so this differs from the report period (2026-08-06 to 2026-08-12) and is anchored to the latest available device data.

510(k) clearances: **74** (plus 1 separate De Novo authorization). PMA (premarket approval) totals **61**, broken down below — Original/Panel-Track represent new or significant changes.

PMA Original: **1** | Panel-Track: **0** | 30-Day Notice: **48** | Other: **12**

PMA Approvals

**PGDx elio tissue complete CDx**

Personal Genome Diagnostics, Inc.

Next generation sequencing oncology panel for detecting somatic or germline variants.

510(k) / De Novo Clearances (most recent 10)

| Device                                                                                                                        | Applicant                                       | Date       |
|-------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------|------------|
| <b>Air Compression Massager (HY-1165)</b><br>FDA classifies this device as: Massager, Powered Inflatable Tube.                | Xiamen High Top Electronic Technology Co., Ltd. | 2026-07-27 |
| <b>V8/XV8/XH8/H8/cV8/V8 ONE/XV8 ONE/XH8...</b><br>FDA classifies this device as: System, Imaging, Pulsed Doppler, Ultrasonic. | Samsung Medison Co., Ltd.                       | 2026-07-28 |
| <b>uMI Panvivo (uMI Panvivo M)</b><br>FDA classifies this device as: System, Tomography, Computed, Emission.                  | Shanghai United Imaging Healthcare Co., Ltd.    | 2026-07-28 |
| <b>ELISIO™-HX</b><br>A hemodialyzer with expanded solute removal profile for renal therapy.                                   | Nipro Medical Corporation                       | 2026-07-28 |
| <b>CaRi-Heart</b> <b>De Novo</b><br>Caristo Diagnostics, Ltd. received FDA clearance for this device.                         | Caristo Diagnostics, Ltd.                       | 2026-07-28 |
| <b>BodyTom Elite (NL4000)</b><br>FDA classifies this device as: System, X-Ray, Tomography, Computed.                          | NeuroLogica Corp., a Samsung Company            | 2026-07-28 |
| <b>TnR Cranial Implant</b><br>FDA classifies this device as: Methyl Methacrylate For Cranioplasty.                            | T&R BIOFAB CO., Ltd.                            | 2026-07-30 |

|                                                                                                                        |                                            |            |
|------------------------------------------------------------------------------------------------------------------------|--------------------------------------------|------------|
| <b>Handheld Ultrasound Scanner</b><br>FDA classifies this device as: System, Imaging, Pulsed Doppler, Ultrasonic.      | IMGFAX CO., LTD.                           | 2026-07-30 |
| <b>0909FCE, 0909FCE-HS</b><br>FDA classifies this device as: Solid State X-Ray Imager (Flat Panel/Digital Imager).     | Rayence Co., Ltd.                          | 2026-07-31 |
| <b>UniPearls® Embolic Microspheres</b><br>FDA classifies this device as: Device, Vascular, For Promoting Embolization. | Suzhou Haowei Medical Technology Co., Ltd. | 2026-07-31 |

\* Full 75-item 510(k)/De Novo dataset available in the Excel file attached to this issue.

 **Warning Letters**

Total this week: 0 | Significant (manufacturing/GMP, or not classifiable as unapproved/labeling-only): 0 | Unapproved/labeling ONLY (no manufacturing-quality finding; mostly supplement/personal-import cases): 0

## Drug Recalls

**Drug Recalls (Top Items)** (showing 10 rows covering 41 of 49 total recalls in the latest available recall data (2026-07-22 to 2026-08-05); recalls from the same company for the same reason are grouped into one row. Full list available in the accompanying Excel data file.)

Note: "Recalling Firm" is the firm that filed the recall with the FDA (openFDA's recalling\_firm field) — it is not necessarily the actual manufacturer, which may differ from the firm named in the Product column.

| Recalling Firm                        | Product                                                                                                                                                                                                                                                                                            | Reason                                                                                                                                                                                             | Class    |
|---------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------|
| JB Chemicals and Pharmaceuticals Ltd  | Cetirizine Hydrochloride Tablets USP 5 mg, 100-count bottle, Manufactured by: Unique Pharmaceuticals Labs, (A Div. of J.B. Chemicals & Pharmaceuticals, Ltd.), Mumbai 400 030, India. Distributed by: Rising Pharma Holdings, Inc., East Brunswick, NJ 08816, NDC 16571-401-10.                    | Cross Contamination with Other Products: Customer complaints for appearance of discolored tablets and red dots observed on Cetirizine Hydrochloride Tablets USP 5 mg. Determined to be Ranitidine. | Class I  |
| Rohto-Mentholatum (Vietnam) Co., Ltd. | Rohto Cooling Eye Drops Max Strength (naphazoline hydrochloride 0.03%, polysorbate 90, 0.2%), Sterile 0.4 FL OZ (13 mL), Distributed by: The Mentholatum Company, Orchard Park, NY 14127, Made in Vietnam, UPC 310742011012, NDC 10742-8158-1 (+7 similar recalls, same company & reason, grouped) | Lack of Assurance of Sterility                                                                                                                                                                     | Class II |
| Chiesi USA, Inc.                      | CLEVIPREX (clevidipine injectable emulsion) 50 mg/100 mL (0.5 mg/mL), 10 Single Use Vials, Rx Only, Manufactured for: Chiesi USA, Inc., Cary, NC 27518, by Fresenius Kabi, Graz, Austria, NDC 10122-611-10.                                                                                        | Lack of Assurance of Sterility                                                                                                                                                                     | Class II |
| CareFusion 213, LLC                   | BD Chloraprep Clear, (2% w/v chlorhexidine gluconate (CHG) and 70% v/v isopropyl alcohol), 60 x 1 mL applicators/carton, 0.03fl oz (1 mL) each, STERILE SOLUTION, CareFusion 123, LLC, El Paso, TX, subsidiary of Beckton, Dickson and Co. NDC 54365-400-31                                        | Lack of Assurance of Sterility: Affected product may exhibit an open or incomplete seal on the packaging of the applicator                                                                         | Class II |
| Glenmark Pharmaceuticals Inc., USA    | Azelaic Acid Gel, 15%, 50 gram tubes, For Topical Use only, Rx only, Distributed by: Glenmark Pharmaceuticals Inc., USA, Elmwood Park, NJ 07407, NDC 68462-626-52.                                                                                                                                 | CGMP Deviations: Product quality complaints concerning abnormal texture or consistency (described as grainy, gritty, or sandy) were received.                                                      | Class II |

|                           |                                                                                                                                                                                                                                                                                                                                                            |                                                                                                      |          |
|---------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------|----------|
| Major Pharmaceuticals     | Levothyroxine Sodium Tablets, USP, 75 mcg (0.075 mg), 10 Tablets (10 x 1) unit dose blisters per carton, Rx only, Packaged and Distributed by: MAJOR PHARMACEUTICALS, Indianapolis, IN 46268 USA. Distributed by Cardinal Health, Dublin, OH 43017. NDC Bag: 55154-3560-0; NDC Blister: 0904-6951-61 (+11 similar recalls, same company & reason, grouped) | Subpotent Drug                                                                                       | Class II |
| Bell Pharmaceuticals, Inc | QC Quality Choice, Carbamide Peroxide, 6.5%, 0.5 FL. OZ. (15 mL) a) Washer Bulb included (NDC 63868-026-15); b) 1 bottle (NDC 63868-027-16), Distributed by C.D.M.A., Inc., 43157 W. Nine Mile, Novi, MI 48376. (+10 similar recalls, same company & reason, grouped)                                                                                      | SubPotent Drug: low pH and significantly reduced assay (~0.6% vs expected ~6 7%).                    | Class II |
| Padagis US LLC            | Nystatin Cream USP, (100, 000 USP Nystatin Units), NET WT 15 g, Rx only, 15 g tube, Manufactured by Padagis, Yeruham, Israel, NDC 45802-059-35                                                                                                                                                                                                             | Subpotent Drug: Low out of specification assay results performed during long-term stability testing. | Class II |
| Fresenius Kabi USA, LLC   | Naropin (ropivacaine hydrochloride Injection, USP), 0.5%, 100 mg per 20 mL (5 mg per mL), Twenty-five, 20 mL Single-Dose Vials, Rx only, Fresenius Kabi, Lake Zurich, IL 60047 NDC Tray: 63323-286-23; NDC Vial: 63323-286-05 (+3 similar recalls, same company & reason, grouped)                                                                         | Presence of Particulate Matter: Hair was found in products                                           | Class II |
| MYLAN PHARMACEUTICALS INC | Mycophenolate Mofetil for Injection USP, 500 mg/Vial, Single Dose Vial, 4 vials per carton, Rx only, Manufactured for: Mylan Institutional LLC, Morgantown, WV, 26505 USA, Made in India, NDC 67457-386-00.                                                                                                                                                | Presence of precipitate                                                                              | Class II |

 **FDA Guidance & Regulatory Actions**

13 FDA regulatory documents this week.

[Determination That ANSAID \(Flurbiprofen\) Tablets, 50 Milligrams and 100 Milligrams, Were Not Withdrawn From Sale for Reasons of Safety or Effectiveness](#) (2026-08-14 • Notice)

This notice determines that ANSAID (Flurbiprofen) tablets were not withdrawn for safety or effectiveness reasons, affecting generic drug manufacturers seeking bioequivalent versions.

[Reauthorization of the Prescription Drug User Fee Act; Public Meeting; Request for Comments](#) (2026-08-14 • Notice)

This notice announces a public meeting and requests comments on the reauthorization of the Prescription Drug User Fee Act (PDUFA), affecting pharmaceutical companies developing prescription drugs.

[Determination That LASIX \(Furosemide\) Tablets, 20 Milligrams, 40 Milligrams, and 80 Milligrams, Were Not Withdrawn From Sale for Reasons of Safety or Effectiveness](#) (2026-08-14 • Notice)

This notice determines that LASIX (Furosemide) tablets were not withdrawn for safety or effectiveness reasons, affecting generic drug manufacturers seeking bioequivalent versions.

[Container Closure Systems for Human Drugs and Biological Products; Draft Guidance for Industry; Availability](#) (2026-08-14 • Notice)

This notice announces the availability of a draft guidance for industry on container closure systems for human drugs and biological products, affecting pharmaceutical and biologics manufacturers.

[Kremers Urban Pharmaceuticals, Inc.; Grant of Hearing Request Regarding a Proposal To Withdraw Approval of an Abbreviated New Drug Application for Extended-Release Methylphenidate Tablets](#) (2026-08-13 • Notice)

This notice grants a hearing request regarding a proposal to withdraw approval for an Abbreviated New Drug Application (ANDA) for extended-release methylphenidate tablets, affecting Kremers Urban Pharmaceuticals.

[Guide To Minimize Biological Hazards in Ready-to-Eat Fresh-Cut Produce; Guidance for Industry; Availability](#) (2026-08-12 • Rule)

This final rule announces the availability of guidance to minimize biological hazards in ready-to-eat fresh-cut produce, affecting producers and distributors of fresh-cut produce.

[Reauthorization of the Generic Drug User Fee Amendments; Public Meeting; Request for Comments](#) (2026-08-11 • Notice)

This notice announces a public meeting and requests comments on the reauthorization of the Generic Drug User Fee Amendments (GDUFA), affecting generic drug manufacturers.

[Substances Generally Recognized as Safe](#) (2026-08-11 • Proposed Rule)

This proposed rule outlines regulations for substances generally recognized as safe (GRAS), affecting food manufacturers and ingredient suppliers.

#### [Determination That OZOBAX \(Baclofen\) Oral Solution, 5 Milligrams/5 Milliliters Was Not Withdrawn From Sale for Reasons of Safety or Effectiveness](#) (2026-08-10 • Notice)

This notice determines that OZOBAX (Baclofen) oral solution was not withdrawn for safety or effectiveness reasons, affecting generic drug manufacturers seeking bioequivalent versions.

#### [Determination of Regulatory Review Period for Purposes of Patent Extension; BRINSUPRI](#) (2026-08-10 • Notice)

This notice establishes the regulatory review period FDA uses to calculate a potential patent-term extension for BRINSUPRI, to be decided by the USPTO. It affects the patent holder of BRINSUPRI.

#### [Determination of Regulatory Review Period for Purposes of Patent Extension; DAYVIGO](#) (2026-08-10 • Notice)

This notice establishes the regulatory review period FDA uses to calculate a potential patent-term extension for DAYVIGO, to be decided by the USPTO. It affects the patent holder of DAYVIGO.

#### [Molecular and Clinical Genetics Panel of the Medical Devices Advisory Committee; Notice of Meeting; Establishment of a Public Docket; Request for Comments-GRAIL, Inc. Galleri](#) (2026-08-10 • Notice)

This notice announces a public meeting of the Molecular and Clinical Genetics Panel regarding GRAIL, Inc.'s Galleri device, and requests comments. It affects medical device manufacturers, particularly in genetic diagnostics.

#### [Radiology Devices; Reclassification of Digital Breast Tomosynthesis System](#) (2026-08-10 • Proposed Rule)

This proposed rule reclassifies the Digital Breast Tomosynthesis System, affecting manufacturers and distributors of radiology devices, specifically breast imaging systems.



## Upcoming FDA Regulatory Event Calendar



### Weekly Outlook

This week's notable events include Deramiciocel (2026-08-22) and Zilganerssen (2026-09-22). In the Guidance Agenda, CDER's Generics category has the most items (24).

This text is generated mechanically from existing aggregate data (priority, countdown, category counts). It contains no forecasts or speculation.

★ This Week's Spotlight (Editor's Picks)

Selected mechanically by priority tier and proximity (days until event) — the notable items for this week.

| Date                 | Priority | Event         | Type  |
|----------------------|----------|---------------|-------|
| 2026-08-22<br>in 6d  | ★★★★★    | Deramiciocel  | PDUFA |
| 2026-09-22<br>in 37d | ★★★★★    | Zilganersen   | PDUFA |
| 2026-09-26<br>in 41d | ★★★★★    | zilurgisertib | PDUFA |
| 2026-09-30<br>in 45d | ★★★★★    | apitegromab   | PDUFA |
| 2026-08-25<br>in 9d  | ★★★★☆    | zanidatamab   | PDUFA |

**Changes Since Last Week**

| Category                              | Count |
|---------------------------------------|-------|
| New (PDUFA 33 / AdCom 2 / Guidance 0) | 35    |
| Postponed / rescheduled               | 0     |
| Removed / withdrawn                   | 27    |

**New this week (PDUFA / AdCom)** (showing top 10 of 35)

- [PDUFA] Deramiocele (2026-08-22)
- [PDUFA] zanidatamab (2026-08-25)
- [PDUFA] zanidatamab (2026-08-25)
- [PDUFA] Zilganerser (2026-09-22)
- [PDUFA] zilurgisertib (2026-09-26)
- [PDUFA] apitegromab (2026-09-30)
- [PDUFA] Bepirovirsen (2026-10-26)
- [PDUFA] INO-3107 (2026-10-30)
- [PDUFA] ivonescimab (2026-11-14)
- [PDUFA] BBP-418 (2026-11-27)

**Removed / withdrawn** (showing top 10 of 27)

- [AdCom] Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Food and Drug Administration Advisory Committees
- [PDUFA] Deramiocele
- [PDUFA] Deramiocele
- [PDUFA] zanidatamab
- [PDUFA] zanidatamab
- [PDUFA] Zilganerser
- [PDUFA] zilurgisertib
- [PDUFA] apitegromab
- [PDUFA] Bepirovirsen
- [PDUFA] ivonescimab



Upcoming FDA Schedule (Official, Not Predicted)

This section contains no forecasts. It lists only officially announced FDA events, each linked to its primary source.

| Date                 | Priority | Advisory Committee / Topic                                                                                                                                                            | Product / Company | Source                           |
|----------------------|----------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------|----------------------------------|
| 2026-09-23<br>in 38d | ★★★★☆    | Molecular and Clinical Genetics Panel of the Medical Devices Advisory Committee; Notice of Meeting; Establishment of a Public Docket; Request for Comments-GRAIL, Inc. Galleri        | —                 | <a href="#">Federal Register</a> |
| 2026-10-16<br>in 61d | ★★★★☆    | Gastroenterology-Urology Devices; Reclassification of Diagnostic Endoscopic Light Source Systems To Be Renamed Cystoscopic Systems Intended as an Aid for Detection of Bladder Cancer | —                 | <a href="#">Federal Register</a> |

**PDUFA Target Action Dates Disclosed by Sponsors**

This list is compiled from information disclosed by companies in SEC filings, official press releases, and investor materials. It is not a comprehensive official schedule published by FDA. Dates may change, be delayed, or be withdrawn. Always verify against the primary source linked below before making investment, regulatory, or business decisions.

**Confirmed (exact date disclosed)**

| Target Action Date   | Priority | Product       | Company                                           | Indication                                                                                                 | Source                                  |
|----------------------|----------|---------------|---------------------------------------------------|------------------------------------------------------------------------------------------------------------|-----------------------------------------|
| 2026-08-22<br>in 6d  | ★★★★★    | Deramiciel    | CAPRICOR THERAPEUTICS, INC.                       | cardiomyopathy in patients with Duchenne muscular dystrophy                                                | <a href="#">SEC 8-K</a>                 |
| 2026-08-25<br>in 9d  | ★★★★☆    | zanidatamab   | Jazz Pharmaceuticals plc                          | first-line (1L) gastroesophageal adenocarcinoma (GEA)                                                      | <a href="#">SEC 8-K (Press Release)</a> |
| 2026-08-25<br>in 9d  | ★★★★☆    | zanidatamab   | Zymeworks Inc.                                    | first-line (1L) HER2-positive (HER2+) locally advanced or metastatic gastroesophageal adenocarcinoma (GEA) | <a href="#">SEC 8-K (Press Release)</a> |
| 2026-09-22<br>in 37d | ★★★★★    | Zilganersen   | Ionis Pharmaceuticals, Inc.                       | Alexander disease (AxD)                                                                                    | <a href="#">SEC 8-K (Press Release)</a> |
| 2026-09-26<br>in 41d | ★★★★★    | zilurgisertib | Mirum Pharmaceuticals, Inc.                       | fibrodysplasia ossificans progressiva (FOP)                                                                | <a href="#">SEC 8-K (Press Release)</a> |
| 2026-09-30<br>in 45d | ★★★★★    | apitegromab   | Scholar Rock                                      | spinal muscular atrophy (SMA)                                                                              | <a href="#">SEC 8-K (Press Release)</a> |
| 2026-10-26<br>in 71d | ★★★★★    | Bepirovirsen  | IONIS PHARMACEUTICALS INC (IONS) (CIK 0000874015) | chronic hepatitis B (CHB)                                                                                  | <a href="#">SEC 8-K (Press Release)</a> |
| 2026-10-30<br>in 75d | ★★★★★    | INO-3107      | INOVIO PHARMACEUTICALS, INC.                      | Recurrent Respiratory Papillomatosis (RRP)                                                                 | <a href="#">SEC 8-K (Press Release)</a> |

|                       |       |                |                                                                 |                                                                                                                                                      |                                   |
|-----------------------|-------|----------------|-----------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------|
| 2026-11-14<br>in 90d  | ★★★★★ | ivonescimab    | Summit<br>Therapeutics Inc.                                     | EGFR-mutated,<br>locally advanced<br>or metastatic<br>non-squamous<br>NSCLC who were<br>previously<br>treated with a<br>third-generation<br>EGFR TKI | <a href="#">SEC 8-K (Press Re</a> |
| 2026-11-27<br>in 103d | ★★★★★ | BBP-418        | BridgeBio Pharma,<br>Inc.                                       | LGMD2I/R9                                                                                                                                            | <a href="#">SEC 8-K (Press Re</a> |
| 2026-11-27<br>in 103d | ★★★★★ | &mdash;        | BridgeBio<br>Disclosed in<br>BridgeBio Pharma,<br>Inc.'s filing | &mdash;                                                                                                                                              | <a href="#">SEC 8-K (Press Re</a> |
| 2026-11-30<br>in 106d | ★★★★★ | Povetacicept   | Vertex                                                          | immunoglobulin<br>A nephropathy<br>(IgAN)                                                                                                            | <a href="#">SEC 8-K (Press Re</a> |
| 2026-12-03<br>in 109d | ★★★★★ | zanzalintinib  | Exelixis, Inc.                                                  | metastatic<br>colorectal cancer                                                                                                                      | <a href="#">SEC 8-K (Press Re</a> |
| 2026-12-12<br>in 118d | ★★★★★ | Quimilza TM    | Vanda<br>Pharmaceuticals<br>Inc.                                | Generalized<br>Pustular Psoriasis<br>(GPP)                                                                                                           | <a href="#">SEC 8-K (Press Re</a> |
| 2026-12-27<br>in 133d | ★★★★★ | relutrigine    | Praxis Precision<br>Medicines, Inc.                             | SCN2A and<br>SCN8A<br>developmental<br>and epileptic<br>encephalopathies<br>(DEEs)                                                                   | <a href="#">SEC 8-K (Press Re</a> |
| 2026-12-30<br>in 136d | ★★★★★ | bezuclastinib  | Cogent Biosciences,<br>Inc.                                     | NonAdvSM                                                                                                                                             | <a href="#">SEC 8-K (Press Re</a> |
| 2027-01-06<br>in 143d | ★★★★☆ | ANKTIVA        | ImmunityBio, Inc.                                               | BCG-<br>unresponsive<br>NMIBC with<br>papillary disease<br>without CIS                                                                               | <a href="#">SEC 8-K (Press Re</a> |
| 2027-01-21<br>in 158d | ★★★★★ | z-rostudirsen  | Dyne Therapeutics,<br>Inc.                                      | Duchenne<br>muscular<br>dystrophy (DMD)<br>amenable to<br>exon 51 skipping                                                                           | <a href="#">SEC 8-K (Press Re</a> |
| 2027-01-29<br>in 166d | ★★★★★ | ulixacaltamide | Praxis Precision<br>Medicines, Inc.                             | Essential Tremor                                                                                                                                     | <a href="#">SEC 8-K (Press Re</a> |

|                       |       |                       |                                                              |                                                                                                                                                                                    |                                         |
|-----------------------|-------|-----------------------|--------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------|
| 2027-02-23<br>in 191d | ★★★★☆ | ZORYVE<br>cream 0.05% | Arcutis<br>Biotherapeutics,<br>Inc.                          | mild to moderate<br>atopic dermatitis<br>in infants aged 3<br>to 24 months                                                                                                         | <a href="#">SEC 8-K (Press Release)</a> |
| 2027-02-27<br>in 195d | ★★★★★ | zipalertinib          | Cullinan<br>Therapeutics, Inc.<br>(CGEM) (CIK<br>0001789972) | locally advanced<br>or metastatic<br>EGFR ex20ins<br>NSCLC whose<br>disease has<br>progressed on or<br>after platinum-<br>based<br>chemotherapy,<br>with or without<br>amivantamab | <a href="#">SEC 8-K (Press Release)</a> |
| 2027-02-28<br>in 196d | ★★★★☆ | VOXZOGO               | BioMarin<br>Pharmaceutical Inc.                              | achondroplasia                                                                                                                                                                     | <a href="#">SEC 8-K (Press Release)</a> |
| 2027-04-01<br>in 228d | ★★★★★ | Pitolisant GR         | Harmony<br>Biosciences<br>Holdings, Inc.                     | narcolepsy                                                                                                                                                                         | <a href="#">SEC 8-K (Press Release)</a> |
| 2027-04-28<br>in 255d | ★★★★★ | Varegacestat          | Immunome, Inc.                                               | desmoid tumors                                                                                                                                                                     | <a href="#">SEC 8-K (Press Release)</a> |
| 2027-05-01<br>in 258d | ★★★★★ | AXS-12                | Axsome<br>Therapeutics, Inc.                                 | cataplexy in<br>narcolepsy                                                                                                                                                         | <a href="#">SEC 8-K (Press Release)</a> |
| 2027-05-08<br>in 265d | ★★★★★ | encaleret             | BridgeBio Pharma,<br>Inc.                                    | ADH1                                                                                                                                                                               | <a href="#">SEC 8-K (Press Release)</a> |
| 2027-05-27<br>in 284d | ★★★★★ | obexelimab            | Zenas BioPharma,<br>Inc.                                     | Immunoglobulin<br>G4-Related<br>Disease (IgG4-<br>RD)                                                                                                                              | <a href="#">SEC 8-K (Press Release)</a> |

### Partially disclosed (quarter / month / half-year only)

| Expected Timing   | Priority | Product            | Company                               | Source                                  |
|-------------------|----------|--------------------|---------------------------------------|-----------------------------------------|
| August 2026       | ★★★★★    | rusfertide         | Protagonist<br>Therapeutics           | <a href="#">SEC 8-K (Press Release)</a> |
| September 2026    | ★★★★★    | Ameluz             | Biofrontera Inc.                      | <a href="#">SEC 8-K (Press Release)</a> |
| 2028 (timing TBD) | ★★★★★    | Pitolisant HD      | Harmony Biosciences<br>Holdings, Inc. | <a href="#">SEC 8-K (Press Release)</a> |
| 2028 (timing TBD) | ★★★★★    | WAKIX (pitolisant) | Harmony Biosciences<br>Holdings, Inc. | <a href="#">SEC 8-K (Press Release)</a> |

|                   |       |                                   |                                    |                                         |
|-------------------|-------|-----------------------------------|------------------------------------|-----------------------------------------|
| 2028 (timing TBD) | ★★★★★ | EPX-100 (clemizole hydrochloride) | Harmony Biosciences Holdings, Inc. | <a href="#">SEC 8-K (Press Release)</a> |
| 2028 (timing TBD) | ★★★★★ | Pitolisant                        | Harmony Biosciences                | <a href="#">SEC 8-K (Press Release)</a> |

## Annual Guidance Development Plans (CDER/CBER/CDRH)

Each center publishes an annual list of guidance topics it is considering. Individual items have no committed publication date, and FDA does not guarantee every listed topic will be issued.

### **CDER (CY2026)** — 85 items • [Official page](#)

By category: Generics (24), Pharmaceutical Quality/CMC (10), Clinical/Medical (9), Administrative/Procedural (8), Compounding (8), Biosimilars (6) and more

- Drug and Device Manufacture Communications with Payors, Formulary Committees, and Small Entities - Questions and Answers; Revised Draft
- Engaging with FDA to Discuss the Use of Digital Health Technologies in Clinical Investigations of Drug and Biological Products
- Exclusivity for First Interchangeable Biosimilar Biological Products
- Expedited Programs: Fast Track, Breakthrough Therapy, and Priority Review
- NDC Creation, Assignment, Listing and Appropriate Use for Human Drugs, Including Biological Products
- Priority Review Voucher Programs

+79 more (see official page for the full list)

### **CBER (CY2026)** — 35 items • [Official page](#)

By category: Therapeutic Products (19), Vaccines and Allergens (9), Blood and Blood Components (5), Other (2)

- Considerations for the Development of Blood Collection, Processing, and Storage Systems for the Manufacture of Blood Components Using the Buffy Coat Method; Guidance for Industry
- Collection of Platelets by Automated Methods; Draft Guidance for Industry
- Recommendations for Testing Blood Donations for Hepatitis B Virus; Guidance for Industry
- Revised Recommendations to Reduce the Risk of Transfusion-Transmitted Malaria; Guidance for Industry
- Recommendations for Testing Blood Donations for Human T-lymphotropic Viruses I and II; Draft Guidance for Industry
- Frequently Asked Questions - Cell and Gene Therapy Products; Guidance for Industry

+29 more (see official page for the full list)

### **CDRH (FY2026)** — 20 items • [Official page](#)

By category: |UC (9), |A (7), |B (3), ||A (1)

- Validation of Diagnostic Tests for Emerging Pathogens following a Declaration and Determination under Section 564
- Use of Real-World Evidence to Support Regulatory Decision-Making for Medical Devices
- Predetermined Change Control Plans for Medical Devices
- Incorporating Voluntary Patient Preference Information over the Total Product Life Cycle
- Enforcement Discretion Policy for Premarket and Other Requirements for NIOSH-Approved Air Purifying Respirators
- Menstrual Products - Labeling and Performance Testing Recommendations

+14 more (see official page for the full list)

**Priority (★) criteria**

- ★★★★★ Major approval decision / highest market impact
- ★★★★☆ AdCom tied to a specific product / initial BLA-NDA approval-related
- ★★★☆☆ Advisory Committee (general business, not tied to a specific product)
- ★★☆☆☆ Committee administration / procedural notice
- ★☆☆☆☆ Administrative/procedural public meeting

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