



**U.S. FOOD & DRUG  
ADMINISTRATION**

# FDA CDER NextGen Portal

## Regulatory Education for Industry (REdI) Annual Conference 2024: Innovation in Medical Product Development

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US FDA



## Disclaimer

The views and opinions presented here represent those of the speakers and should not be considered to represent advice or guidance on behalf of the U.S. Food and Drug Administration.

# Agenda

FDA CDER NextGen Portal Overview

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Before and After NextGen Portal

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NextGen Portal Products

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What is New ?

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User's Adoption

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# FDA CDER NextGen Portal



One stop shop for the purpose of non-eCTD Submission, Collaboration and Reporting. This platform continues to reduce regulatory overhead for sponsors, academia, research institutes, and small businesses.



CDER **NextGen** Portal

Welcome to

# CDER NextGen

Your direct line to the FDA

Learn More

## Sign In

Username

Password

Under 18 U.S.C. 1001, anyone who makes a materially false, fictitious, or fraudulent statement to the U.S. Government is subjected to criminal penalties.

☐ I have read and agree to the Terms and Conditions stated above and below.

Sign In

[Need help signing in?](#)

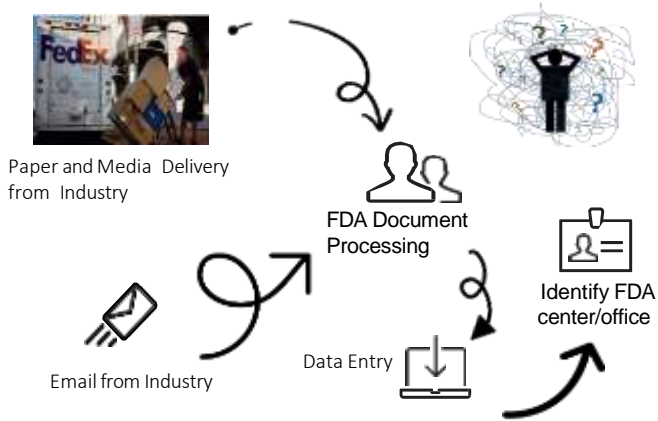


# Digital Transformation

In action to promote safe and effective human drug review and approval

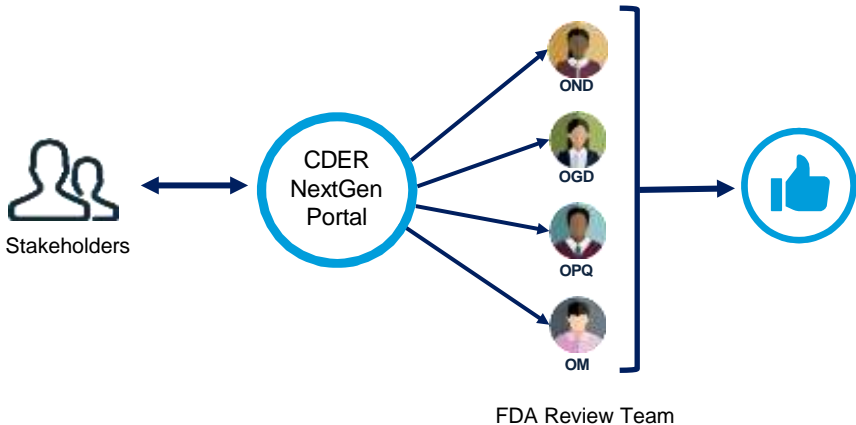


## Before NextGen Portal



- ⚠️ Inefficient paper and Media processing
- ⚠️ Manually intensive
- ⚠️ Time and resource consuming

## After NextGen Portal

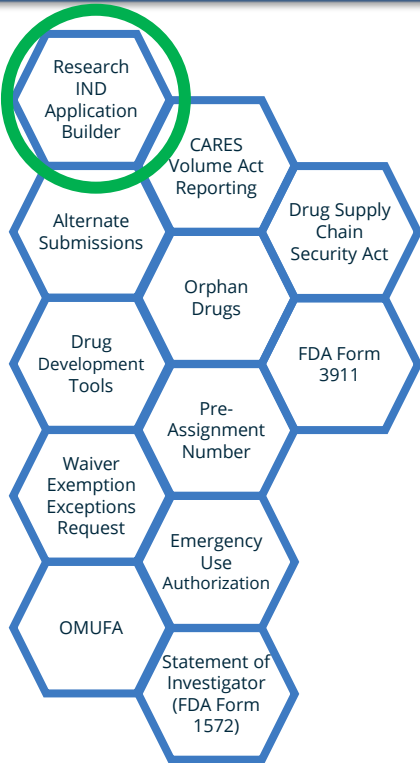


- ✅ Streamlined submissions with clean, complete, and validated data
- ✅ Maximized API led technology to improve efficiency
- ✅ Improved collaboration between the FDA and Stakeholders

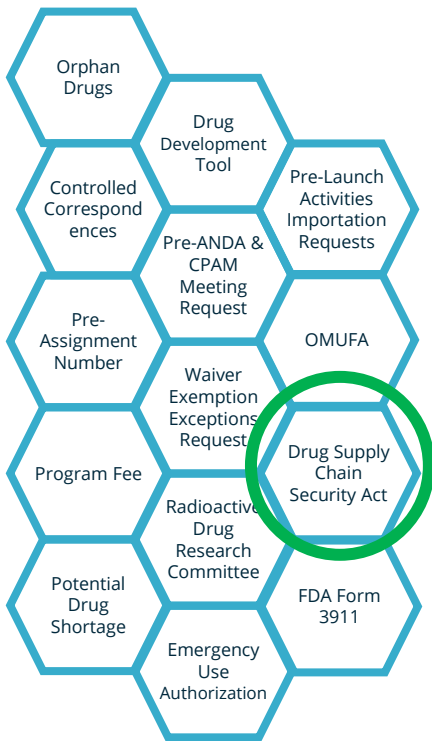
# FDA CDER NextGen Portal Products



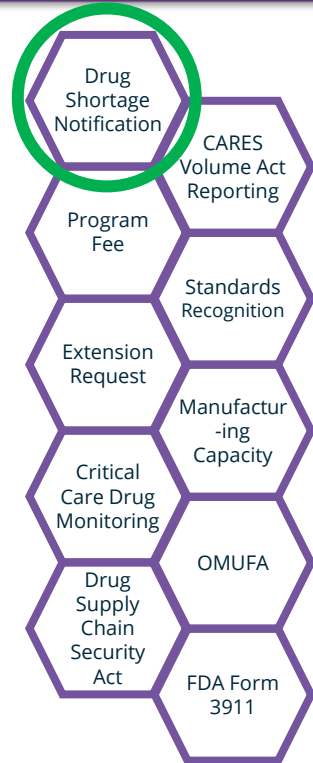
## Regulatory Submissions



## Streamlined Collaboration



## Congressional Reporting



# Application Submission Simplified

From days to minutes ....



## User has the following Information:

### Application / Submission Details

- **IND Number:** IND24840
- **IND Serial Number:** 0000

### Company and Contact Details

- **Company Name:** NIH
- **Company Address:** Bethesda, MD
- **Person Responsible:** Adam Kohl

### Product Details

- **Drug Name:** AIK12
- **UNII:** 36209ITL9D
- **Indication of Use:** SCTID  
404684003

### Study Details

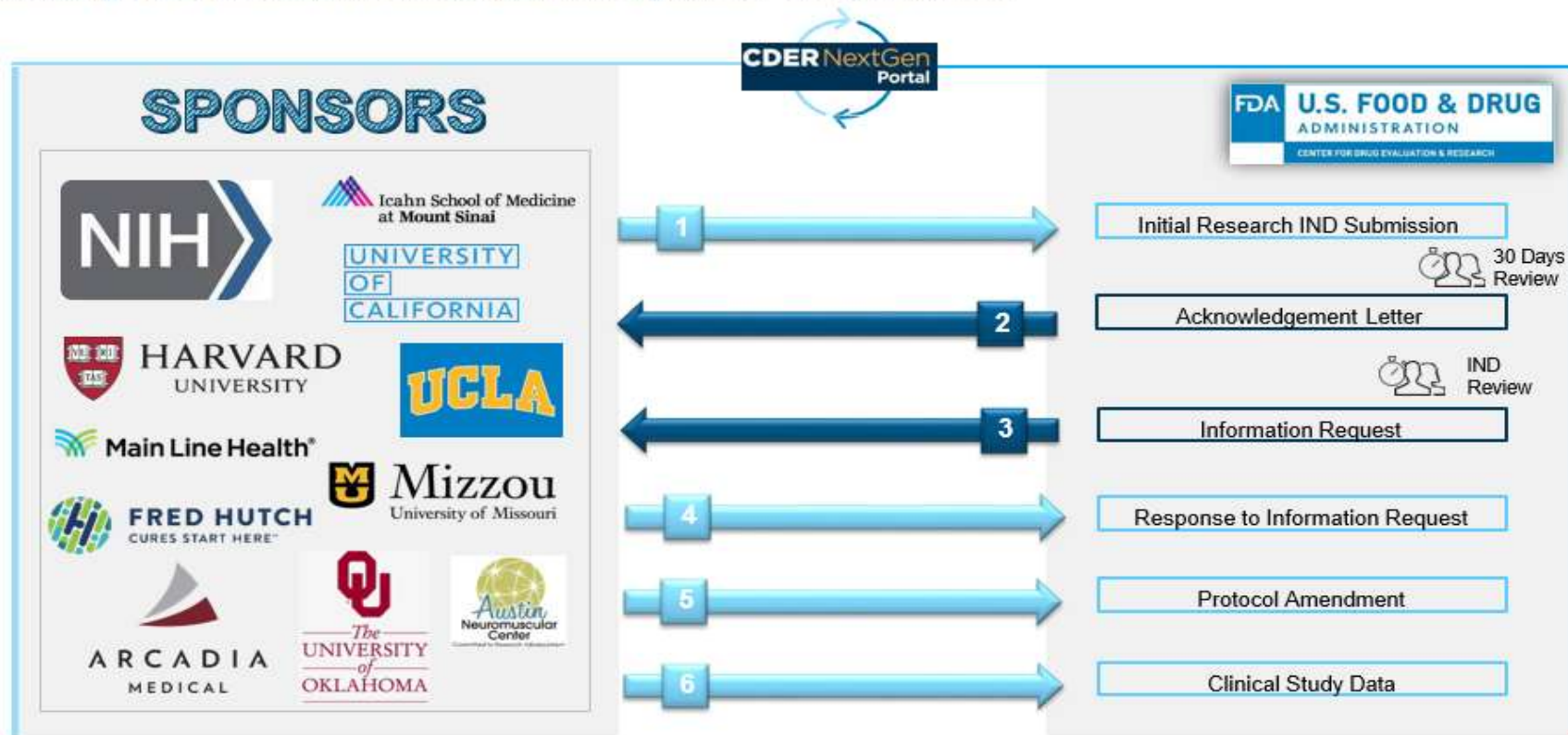
- **NCT Number:** 000032344

## Make a Research IND Submission in **less than 10 Steps**



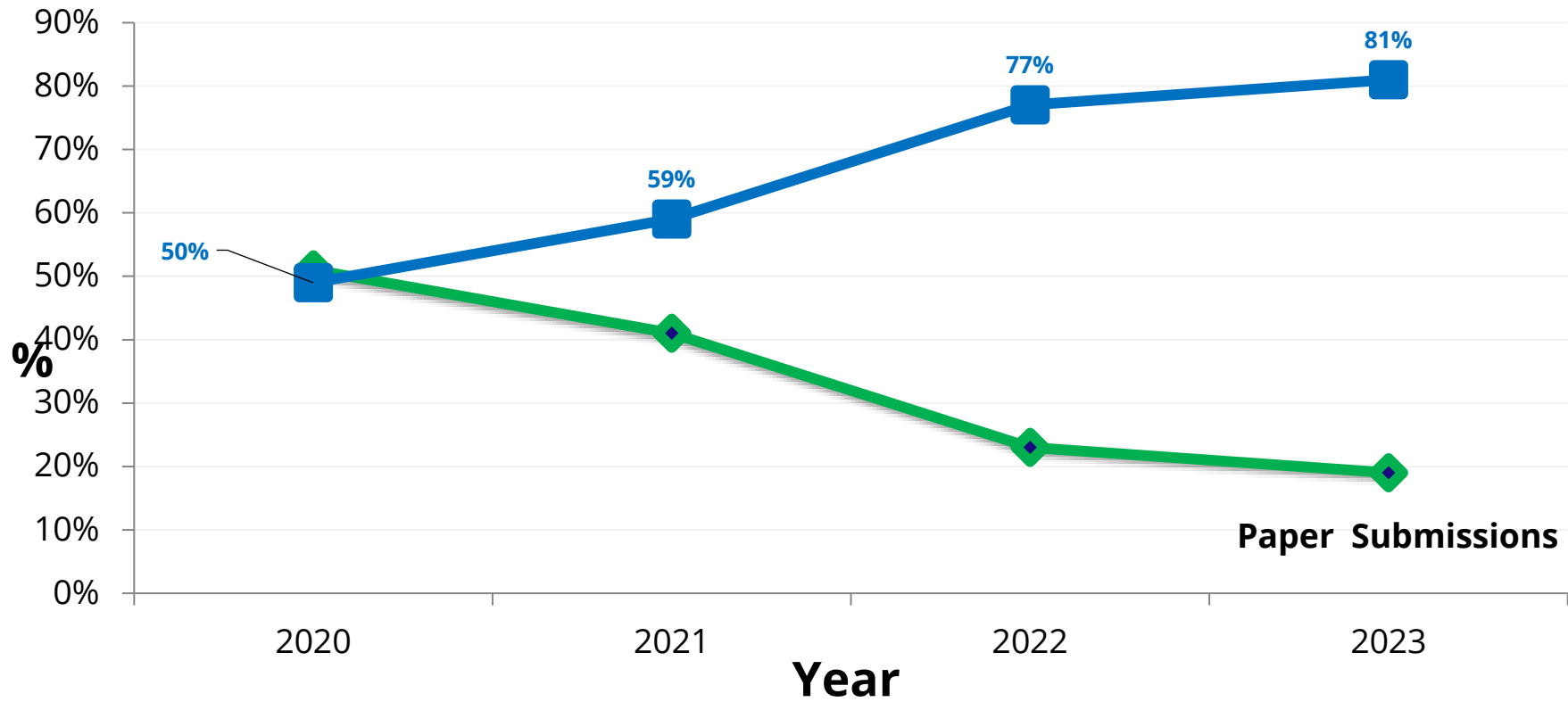
# Near Real-Time 2 Way Interactions

Enabling FDA CDER Reviewer and the Sponsor collaboration





# Research IND Submissions : CDER NextGen Portal Vs Paper



## What is New ?...

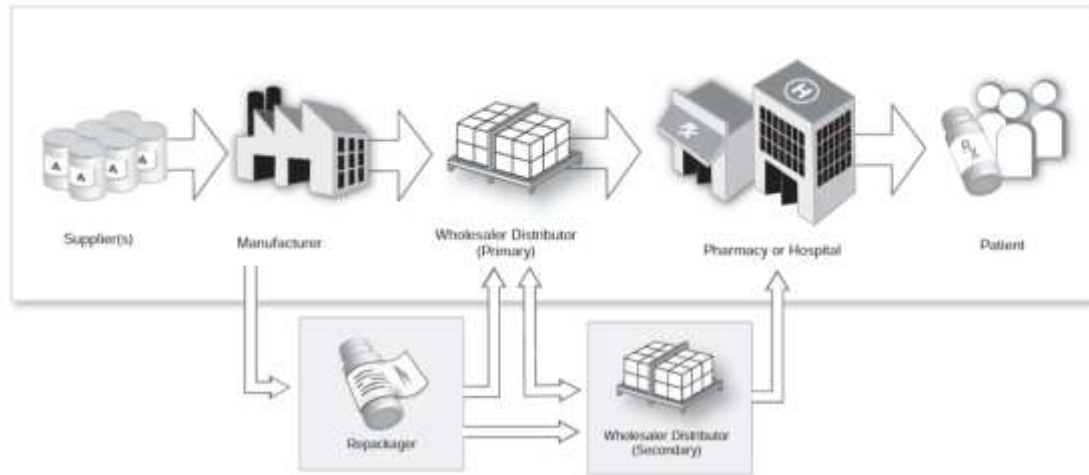
### Drug Supply Chain Security Act (DSCSA) Portal

Steps to achieve interoperable, electronic tracing of products at the package level to identify and trace certain prescription drugs as they are distributed in the United States. This will enhance FDA's ability to help protect consumers from exposure to drugs that may be counterfeit, stolen, contaminated, or otherwise harmful.



#### A Drug Supply Chain Example

From Supplier to Patient



Established near real time 2-way interactions for reporting and respond to inquires such as :

- Transaction Information (TI)
- Transaction Statement (TS)
- Information Request (IR)

# Drug Shortage Modernization Overview and Impact

The Drug Shortage Modernization solution is **critical to ensuring drugs are available for the American public**

## Overview

- Section 3112 of the CARES Act amends the Federal Food, Drug and Cosmetic Act Section 506 and requires FDA to prioritize and expedite reviews and inspections for submissions that could help mitigate or prevent shortages.
- Drug shortage Notification can be submitted in multiple ways such as:
  - Industry Drug Shortage Notification Portal - Safety and Innovation Act (FDASIA) mandate
  - Public Drug Shortage Portal – E.g patients, caregivers, and healthcare providers

## Impact



Reduce manual processes

### Improved Capability

Reduce time intensive manual processes and ensure information is in a centralized location



Enhanced data quality

### Validate Data

Validate data coming through the systems (e.g., NDC code lookup) and keep historical tracking of records in a centralized location

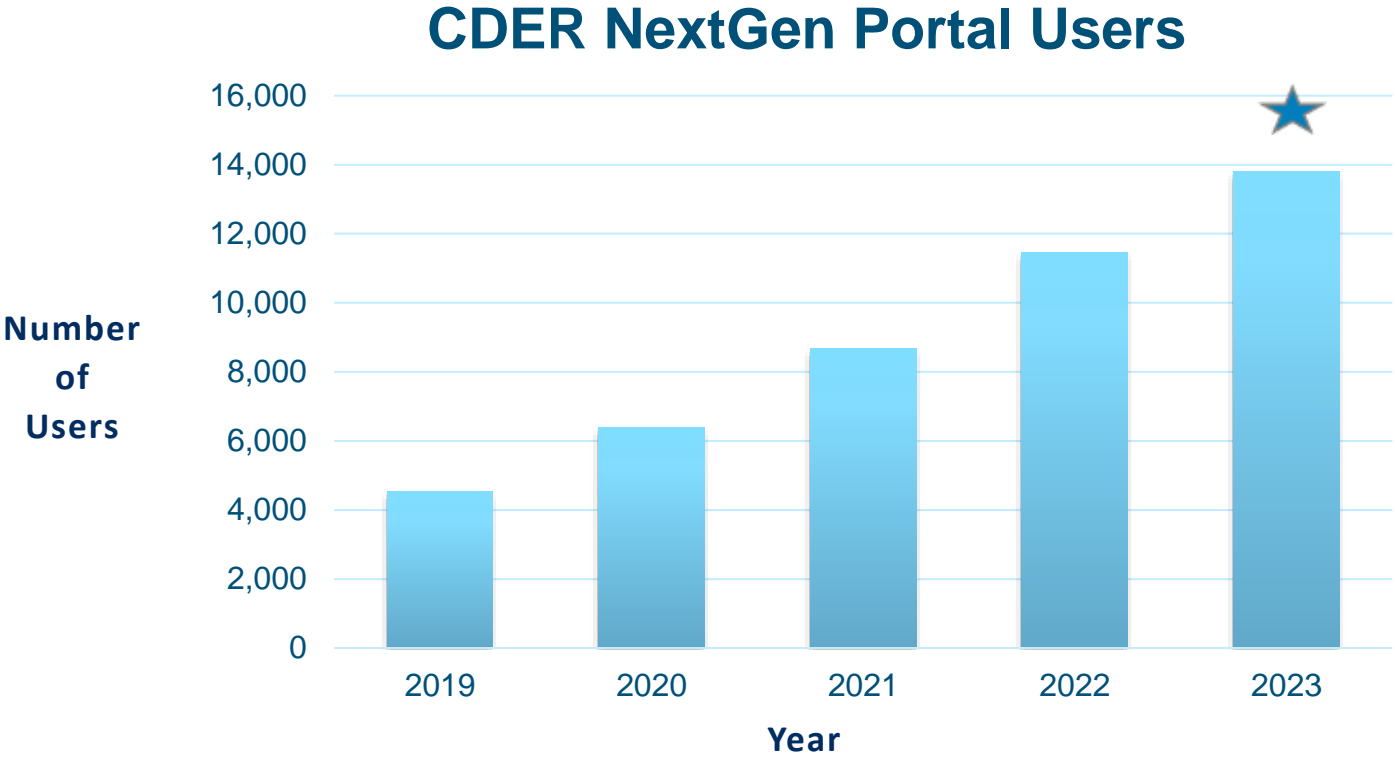


Business Transformation

### Improved Supply Chain

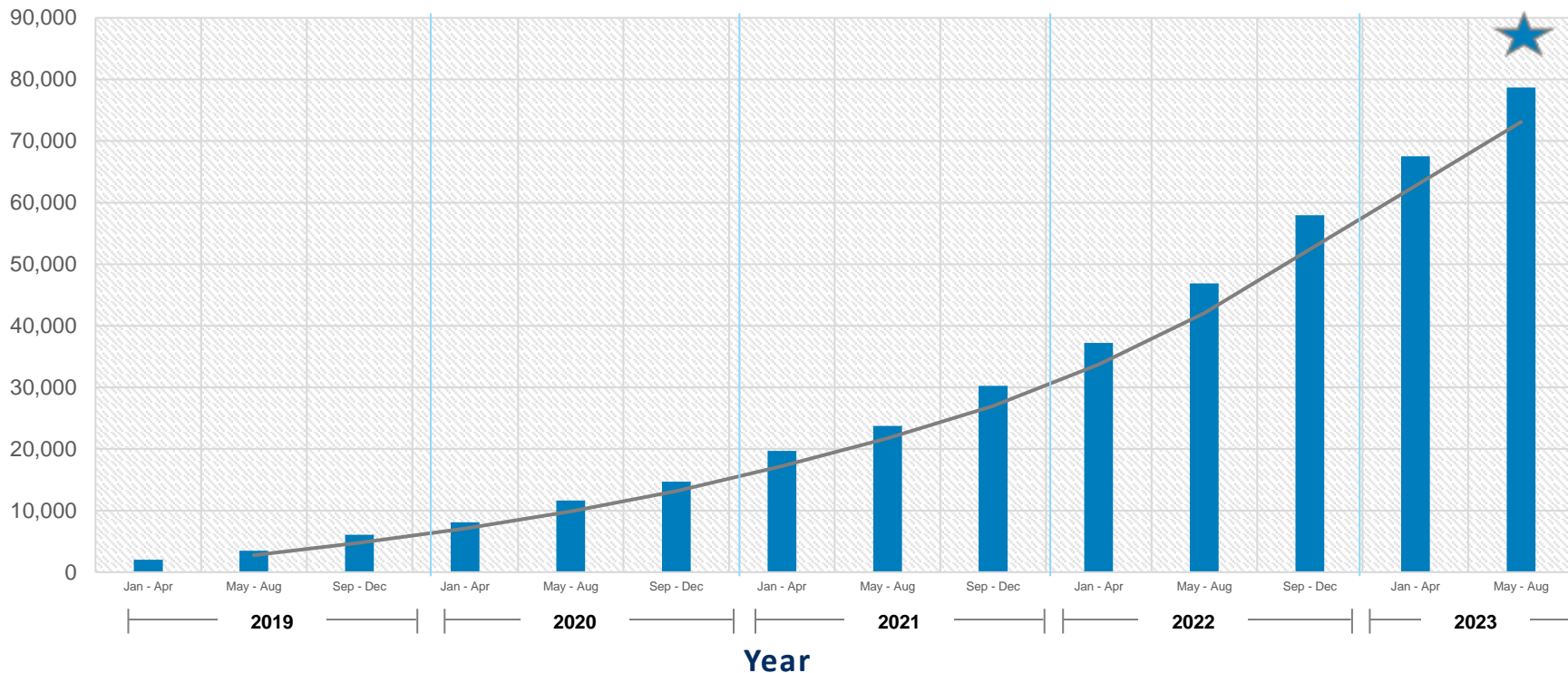
Accurate and recent data are posted on the drug shortage website.

# NextGen Portal Users adoption the last 5 years



# Number of Submissions

## CDER NextGen Portal Submissions



# Need Support ?

- The following support materials can help you get started

## Research IND Application Builder User Guides

<https://cdernextgenportal.fda.gov/s/indhelppcenterinfo>

## User Registration Guides

[https://edm.fda.gov/customThemeStatic/themes/customTheme/docs/CDERDirectNextGen\\_ReferenceGuide\\_MFA.pdf](https://edm.fda.gov/customThemeStatic/themes/customTheme/docs/CDERDirectNextGen_ReferenceGuide_MFA.pdf)

## General FAQs

[https://edm.fda.gov/customThemeStatic/themes/customTheme/docs/CDERDirectNextGen\\_ReferenceGuide\\_MFA.pdf](https://edm.fda.gov/customThemeStatic/themes/customTheme/docs/CDERDirectNextGen_ReferenceGuide_MFA.pdf)

## The Paperwork Reduction Act (PRA)

<https://pra.digital.gov/>

## Benefits of CDER NextGen

<https://www.fda.gov/media/136301/download>



**Contact the Platform Support Team at [edmsupport@fda.hhs.gov](mailto:edmsupport@fda.hhs.gov)**



**Thank You !!**