# How CI/CD Enhances the Development of R Packages in the Pharmaverse

PHUSE- US Connect March 7th, 2023 Ben Straub (GSK) & Dinakar Kulkarni (Roche)

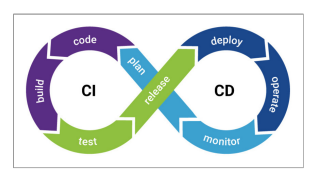




#### What is CI/CD?

- Continuous Integration (CI): Frequent merging of several small changes into a main branch
- Continuous Delivery (CD): Repeatable deployment process when deciding to deploy

CI/CD bridges the gaps between development and operation activities and teams by **enforcing automation** in building, testing and deployment of applications. CI/CD services compile the incremental code changes made by developers, then link and package them into software deliverables

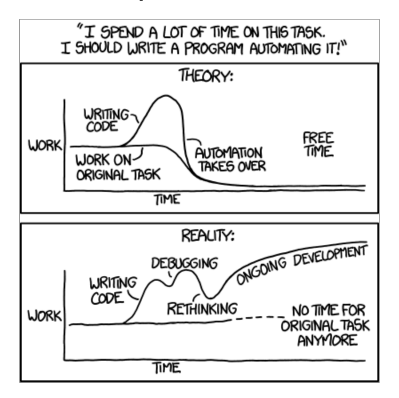








### Does it help?



...Yes! Yes, it does!!



### How does CI/CD help R packages?

- Catch issues (bugs) early on
- User base on multiple OSes and multiple R versions
- Faster turnaround on Code Review
- Multiple Contributors on your R Package
- Enforce style conventions and preferences
- Measure test coverage for new code
- Keep docs up-to-date
- And we can just keep going!

We covered a lot of custom CI/CD actions for R packages in the R/Pharma Workshop in 2022: <u>Intro to CI/CD for R</u> <u>Packages</u> <sup>¬</sup>





## **Ensuring technical innovation**





### **Two Case Studies**





<<u>CI-CD GitHub</u> >



## Case Study - admiral





#### **About admiral**

- Provide an open source, modularized toolbox that enables the pharmaceutical programming community to develop ADaM datasets in R.
- ADaM is one of the required standards for data submission to FDA (U.S.) and PMDA (Japan) for clinical trials
- Links
  - CDISC ☑
  - https://github.com/pharmaverse/admiral
- Issue 1: Checking ADaM Template code
- Issue 2: Common CI/CD workflows for the admiral family of packages





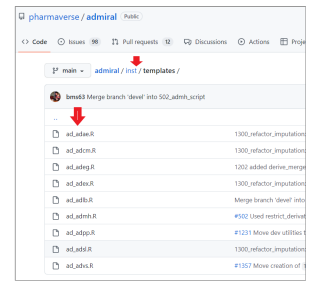


### Issue 1 - How to Check our Template Code

- Create a reference files to build common ADaM datasets that shows users how to implement our functions
- Way less text than a
   Vignette Code is ready
   to go and build a dataset
- Where we store this code is not checked by R-CMD
- How to ensure code stays up to date with deprecated functions or unforeseen bugs get in from functions working together?
- CI/CD for the win!





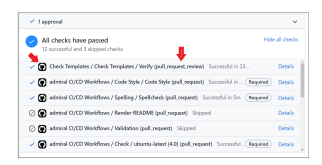






### Solution 1 - CI/CD for Templates

- Dedicated CI/CD workflow that executes the Template code
- Once a Code Review is completed the Check Template Workflow is executed
- If any errors or warnings are detected the CI/CD check fails and the contributor must fix the error or warning.
  - .github/workflows/check-templates.yml







## Issue 2 - admiral upstream and downstream dependencies

- As you can imagine there can be a lot of different types of ADaMs!
- Extension packages focus on specific disease areas like oncology
- The admiral family has a package for developers, template R package repo and dummy data
- Eek!! How to keep this all in line!





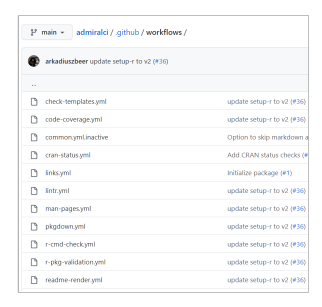
admiraldev [Public] Development Tools for the admiral family	
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## Solution 2 - Common CI/CD workflows for admiral upstream and downstream dependencies

- Using admiralci, we have a common set of CI/CD workflows
- Developers moving between packages are familiar with these workflows
- Common documentation between packages for CI/CD workflows - easy to maintain and provide to new contributors



### <CI-CD GitHub >



## Case Study - NEST





### **About NEST**

- A collection of R packages for creating TLGs/TFLs and exploratory clinical trials data visualization
- tern for creating TLGs
- teal for creating exploratory web applications for analyzing clinical trial data
- Links
  - <u>rtables</u> 🗹
  - NEST GitHub Organization 🗹







### Use Case 1 - Testing Packages as a Cohort

- An in-development package must be tested against the latest versions of upstream dependencies
- Monorepo emulation via a git branch naming strategy is achieved by using
  - the <u>staged.dependencies R package</u> ☑
  - and the **staged.dependencies GitHub Action** ☑
- Testing as a cohort can be done at any stage (eg. development, pre-release, release)





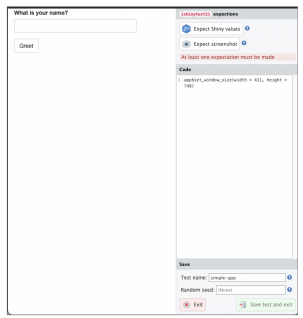


### Use Case 2 - Shiny App Testing & Deployment

- Analysts create Shiny web apps via the teal framework for analyzing data
- Apps deployed to an Posit Connect Server instance via a CD pipeline
  - With the help of the <u>rsconnect</u> 
     and <u>connectapi</u> 
     R packages









### Use Case 3 - Validating R Packages

- R packages are validated by an internal validation team that uses CI/CD pipelines to automatically
  - accept new package submissions via a form
  - running tests against the new package to ensure package integrity
  - enforcing criteria to ensure that the package meets regulatory requirements
- Also validated externally via an open source project called thevalidatoR







### **Additional Materials**

- Further Reading
  - GitHub Actions 🗹
  - GitLab Cl
- Advanced Examples
  - <u>r-lib/actions</u> 🗹
  - {admiralci} ☑
  - Docker 🗷
- Presentation built with **Quarto** ☑



