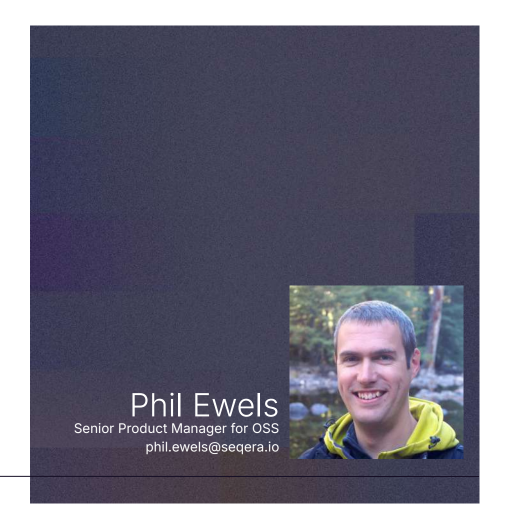


# Seqera AI: How we're using LLMs and Agents with Nextflow code

Creating systems that help scientists to build and run complex workflows



#### Outline

- **01.** Introduction
- **02.** Seqera Al Chat
- **03.** Tool integration
- **04.** Al agents
- **05.** Moving fast
- **06.** Looking ahead





### Introduction





#### Nextflow

Open-source orchestrator for writing & deploying workflows at scale.



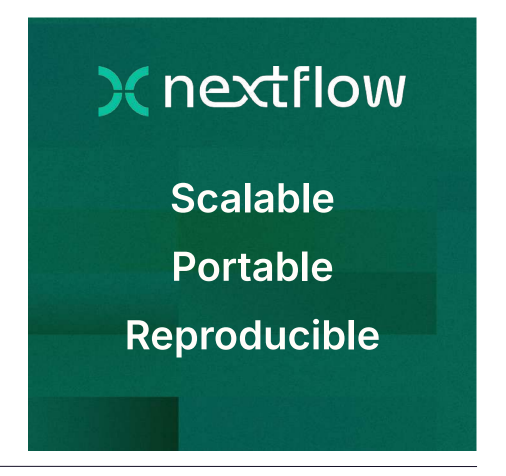




#### Nextflow

Open-source orchestrator for writing & deploying workflows at scale.

- Programming language for bioinformatics workflows
- Orchestrator for executing workflows
- Runs on everything, with data from anywhere
- Reproducibility built-in, native support for software packaging and code version control





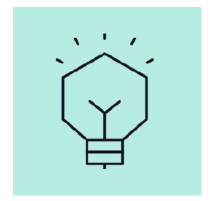
### Segera Al Chat

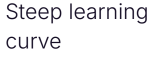
02



# What's the problem?

What do Nextflow users complain about the most?



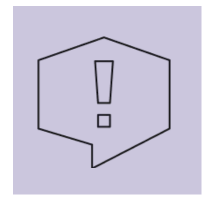


It's Groovy, baby



Syntax changed between versions

DSL1 → DSL2



Difficult / unusual errors

Generalists need not apply



#### Base models struggle with Nextflow

- Nextflow questions are a small subset of data science
- Answers are often very specific to Nextflow and not generally applicable

- Can't retrain the base model (too expensive)
- Can't include a long enough prompt (context window)

My Nextflow pipeline exited with an error in the FastQC process, saying exit code 127. What should I do?

Exit code 127 usually means a command not found error.

Check if it's installed:

fastqc --version

If it's not found, you need to install it, for example using conda:

conda install -c bioconda fastqc

Please don't tell people to do that.. 🤵





### Base models struggle with writing Nextflow\*

- Most public Nextflow code is old
- DSL1 → DSL2 syntax change in 2022+
- Hallucinates syntax from other (more common) languages
- Isn't as opinionated as we'd like it to be!

\* Though they're getting better all the time!

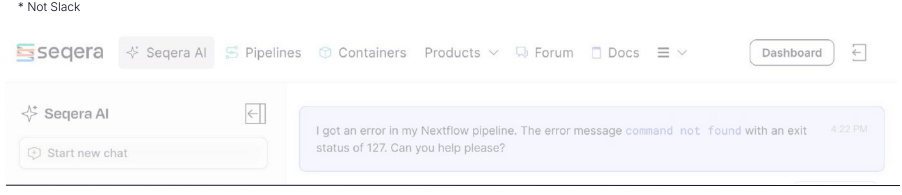


#### Segera Al

Bioinformatics AI that helps you get from 0 to 1 for all your omics

https://seqera.io/ask-ai/

- Latest Nextflow documentation is prioritised over everything else
- Also includes docs from nf-core, nf-test and other trusted sources\*
- Prioritises nf-core best practices and knows available pipelines



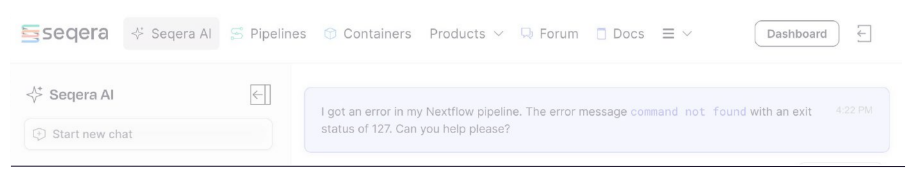


#### Seqera Al

Bioinformatics AI that helps you get from 0 to 1 for all your omics

https://seqera.io/ask-ai/

#### [demo]





### Tool integration

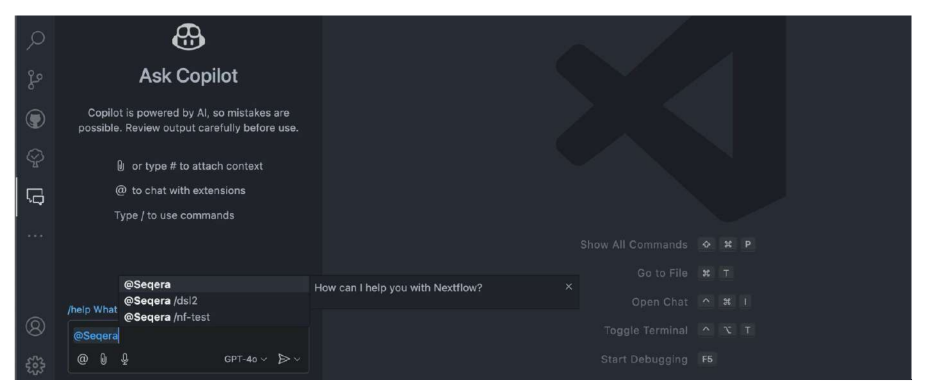
03



#### Segera AI in VS Code

Nextflow help right where you need it

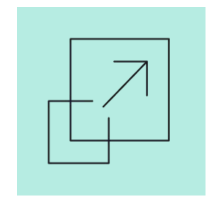
https://nextflow.io/vscode





## What's the problem?

What's most difficult about MultiQC reports?



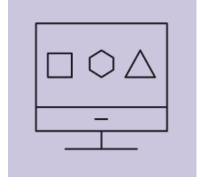
Large sample numbers

Big data is..big



Understanding results

Is this wiggle normal?



Seeing the big picture

Detecting patterns



#### Segera AI in MultiQC

Analysis help right where you need it

https://docs.seqera.io/multiqc/ai

- Help from AI, in the context of a report
- Generation at report time, or on demand
- Flexible, can be used with most providers





### Al agents

04



#### The problem with AI code generation

Al interns can create more work than they solve

- Al writes code, but doesn't understand it
- Looks impressive until you try to run it
- Now you're debugging an entire pipeline looking for the error, rather than iteratively writing + testing

ERROR ~ No signature of method:
groovyx.gpars.dataflow.DataflowBroadcast.into() is
applicable for argument types:
(Script\_85c4d82870d584dc\$\_runScript\_closure1) values:
[Script\_85c4d82870d584dc\$\_runScript\_closure1@53aa2fc9]
Possible solutions: any(), find(), bind(java.lang.Object),
any(groovy.lang.Closure), find(groovy.lang.Closure),
is(java.lang.Object)

-- Check script 'demo.nf' at line: 6 or see '.nextflow.log' file for more details



#### Testing Nextflow code isn't trivial

- Proper modularity and code structure
- Finding example data
- Using the nf-test framework

Al agents can help





### Al agents

#### Going beyond prompts and responses

Agency Level	Description	What that's called	Example Pattern
***	LLM output has no impact on program flow	Simple Processor	<pre>process_llm_output(llm_response)</pre>
***	LLM output determines an if/else switch	Router	<pre>if llm_decision(): path_a() else: path_b()</pre>
***	LLM output determines function execution	Tool Call	<pre>run_function(llm_chosen_tool, llm_chosen_args)</pre>
***	LLM output controls iteration and program continuation	Multi-step Agent	<pre>while llm_should_continue(): execute_next_step()</pre>
***	One agentic workflow can start another agentic workflow	Multi-Agent	<pre>if llm_trigger(): execute_agent()</pre>

Source: https://huggingface.co/docs/smolagents/en/conceptual\_guides/intro\_agents



### Moving fast

06



#### Keeping pace with progress

How Seqera Al adapts to a rapidly changing landscape

Claude 4 release timeline (yesterday)

- 7:01 PM Claude 4 announced on X/Twitter
- 7:16 PM Started testing internally
- 9:06 PM Testing finished, good results
- 10:18 PM Sonnet 4 live in production

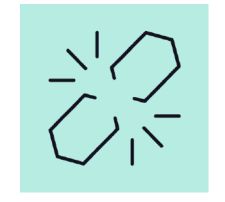


Sasha Red with Sonnet 4 in prod. nice work team; ভেলাল-the pipelines demo should be much more snappy now

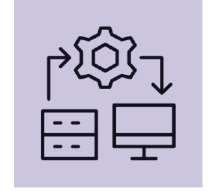


### But how can we move so fast?

Updating with confidence







Decouple

Tools like Langchain decouple application logic from LLM providers

Evaluate

Testing rigs give objective metrics about how well a given setup works

Deploy

Robust and automated deployment infrastructure allows fast deployment and rollback (if needed)



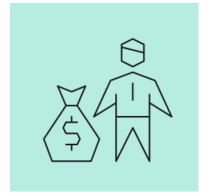
### Looking ahead



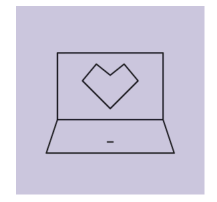


#### Helpful LLMs

Approaches with Segera Al







It's impossible to win against the big model providers, better to use them: standardise and be flexible.

#### Use the big models RAG and prompts

No-one else can do RAG (retrieval augmented generation) quite like you..

Fine-tuned LLMs can provide additional value over major providers.

#### Integration is king

Bringing LLMs out of a chat window and into applications helps to make them part of people's workflows.



#### What's next for Seqera Al

- Configure and launch pipelines
- Help when things go wrong
- Reduce the learning curve
- More integrations for Al agents
- Do all this in a way that is open, trusted and transparent

#### nf-core/rnaseq: 11 sections with 113 configurable fields

- Input/Output Options: 4
- Reference Genome Options: 21
- Read Trimming Options: 4
- Read Filtering Options: 5
- UMI Options: 9
- Alignment Options: 16
- Optional Outputs: 10
- Quality Control: 6
- Process Skipping Options: 20
- Institutional Config Options: 6
- Generic Options: 12



#### Find out more



https://seqera.io/podcasts/









https://summit.nextflow.io/2 025/boston/







### Phil Ewels phil.ewels@segera.io

### Thank you

https://seqera.io/ask-ai/

https://docs.seqera.io/multiqc/ai https://nextflow.io/vscode



nextflow.io



https://nf-co.re



segera.io/multiqc