**Geniac: Automatic Configuration GENerator and Installer for nextflow pipelines [version 2; peer review: 2 approved]**

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**Abstract**

With the advent of high-throughput biotechnological platforms and their ever-growing capacity, life science has turned into a digitized, computational and data-intensive discipline. As a consequence, standard analysis with a bioinformatics pipeline in the context of routine production has become a challenge such that the data can be processed in real-time and delivered to the end-users as fast as possible. The usage of workflow management systems along with packaging systems and containerization technologies offer an opportunity to tackle this challenge. While very powerful, they can be used and combined in many multiple ways which may differ from one developer to another. Therefore, promoting the homogeneity of the workflow implementation requires guidelines and protocols which detail how the source code of the bioinformatics pipeline should be written and organized to ensure its usability, maintainability, interoperability, sustainability, portability, reproducibility, scalability and efficiency. Capitalizing on Nextflow, Conda, Docker, Singularity and the nf-core initiative, we propose a set of best practices along the development life cycle of the bioinformatics pipeline and deployment for production operations which target different expert communities including i) the bioinformaticians and statisticians ii) the software engineers and iii) the data managers and core facility engineers. We implemented Geniac (Automatic Configuration GENerator and Installer for nextflow pipelines) which consists of a toolbox with three components: i) a technical documentation available at https://geniac.readthedocs.io to detail coding guidelines for the
bioinformatics pipeline with Nextflow, ii) a command line interface with a linter to check that the code respects the guidelines, and iii) an add-on to generate configuration files, build the containers and deploy the pipeline. The Geniac toolbox aims at the harmonization of development practices across developers and automation of the generation of configuration files and containers by parsing the source code of the Nextflow pipeline.

**Keywords**
workflow management systems, containerization, reproducibility, high-performance computing, bioinformatics pipelines
Introduction

With the advent of high-throughput biotechnological platforms and their ever-growing capacity, life science has turned into a digitized, computational and data-intensive discipline. While genomics was the major driving force for high-throughput sequencing, other fields including proteomics, imaging and microscopy now contribute to this data explosion (Goh & Wong, 2020). Therefore, the bottleneck shifted from data generation to the ability to process and analyze data with efficient algorithms.

The data analysis actually encompasses two levels, corresponding to different time scales. The first level consists of the iterative research development of innovative and state-of-the-art analysis methods to address novel scientific questions. This level usually takes several months or years. In contrast, the second level consists of routine production analysis using mature and validated methods to apply standard analysis (including quality control) to the new samples processed by these high-throughput biotechnological platforms (Reiter et al., 2021). The production analysis must be performed within a short-delay (few hours up to few days) in order to process in real-time the huge data flow produced by these platforms, such that the relevant biological information can be extracted and provided to the end-users as fast as possible for their downstream analysis. This time constraint is even more important when the data are used in healthcare to support therapeutic decisions: in this case, quick parallel algorithms are required to process high-throughput sequencing data (Jarlier et al., 2020). Both levels feed each other in a virtuous circle.

Whatever the level, the analysis is generally a complex workflow (or pipeline) which involves many steps with different informatics languages and software, whether developed in-house or by third-parties. Coordinating the execution and allowing the scalability of these different steps to benefit from high-performance computing infrastructures to speed-up computation have been simplified with the development of workflow management systems (Leipzig, 2017; Strozzi et al., 2019) in combination with several technologies (Gruening et al., 2018; Jarlier et al., 2021) including package management systems with Conda or Bioconda (Grüning et al., 2018), and containers with Docker (Merkel, 2014) or Singularity (Kurtzer et al., 2017). Importantly, the use of workflow management systems also promotes reusable, reproducible and shareable analysis according to the FAIR principles (Wilkinson et al., 2016) which not only apply to data but also to workflows (Goble et al., 2020). Among the workflow management systems, Nextflow (Di Tommaso et al., 2017) became very popular and offers many interesting features (Jackson et al., 2021). It uses a dataflow programming model which implicitly defines the dependencies between the processes via their outputs and inputs such that different processes can be run in parallel and/or wait for each other when needed.

Whenever run for production analysis, the workflow has no choice but to succeed such that the results can be delivered in time to the end-users. If the workflow fails because of hardware issues or edge/corner cases due to the data themselves, then debugging, fixing, re-installing and resuming the workflow must be performed as fast as possible. This obviously requires harmonized guidelines and protocols across the different developers involved in the development life cycle of the pipeline. This way, it is easier to maintain many different pipelines on one hand and it simplifies the skill transfer from one person to another on the other hand. We developed the biogitflow protocols which cover the code versioning using git and GitLab (Kamoun et al., 2020). However, these protocols do not address how to write the code itself. While very valuable guidelines have been proposed to avoid pitfalls in the implementation of bioinformatics pipelines using the technological stack with workflows management systems, package management systems and containers (Gruening et al., 2018; Reiter et al., 2021; Jarlier et al., 2021), they remain very generic. Therefore, software engineering best practices and technical protocols for coding are necessary in order to promote software quality in bioinformatics to ensure the usability, maintainability, interoperability, sustainability, portability, reproducibility, scalability and efficiency of the workflows, as highlighted by several authors (da Veiga Leprevost et al., 2014; Georgeson et al., 2019; Lawlor & Walsh, 2015).
We acknowledge the nf-core initiative (Ewels et al., 2020) which plays a major role in promoting the software engineering best practices to implement bioinformatics pipelines with Nextflow including code template, linter, code reviewing and continuous integration for better quality. Our Bioinformatics Core Facility decided to capitalize on both Nextflow and the nf-core initiative to provide the end-users with bioinformatics pipelines in the context of production analysis. In this article, we describe additional guidelines on how to write the Nextflow code. We propose a set of best practices along the development life cycle of the pipeline and deployment for production operations which address different expert communities: i) the bioinformaticians and statisticians who prototype the pipeline with state-of-the-art methods in order to extract most of the hidden value from the data and provide the end-users with summary reports, ii) the software engineers who optimize the pipeline to reduce the amount of required informatics resources, to shorten the time to result delivery, iii) the data managers and core facility engineers who deploy and operate the pipeline for daily production analysis for the end-users. The guidelines were motivated by: i) allowing the different expert communities to still work with their preferred work habits, ii) reducing the overall development cycle from the prototyping stage to deployment in a production environment, iii) providing portable pipelines with containers (Docker and Singularity), and iv) automating (whenever possible) the building of containers from the source code of the pipeline. Therefore, we implemented Geniac (Automatic Configuration GENerator and Installer for nextflow pipelines) (Hupé et al., 2022) to address these challenges. Geniac may be used alone for any pipeline built with Nextflow or as an extension of the nf-core guidelines to reduce the code complexity during the prototyping stage and standardize the deployment process.

Geniac actually consists of a toolbox with three components: a technical documentation available at https://geniac.readthedocs.io, a command line interface (CLI) with a linter to check that the code respects the guidelines, and an add-on to generate configuration files, build the containers and deploy the pipeline. We introduce here the main features of the Geniac toolbox.

**Methods**

**General principle**

The ultimate goal of the proposed best practices of coding with Nextflow was to ensure the portability and reproducibility of the pipeline with containerization techniques (such as Docker and Singularity). Therefore, the rationale behind Geniac was motivated by the automation of the construction of the containers without explicitly writing the Dockerfile or the Singularity Definition File recipes (whenever possible). To do so, the pipeline source code is parsed by Geniac. The portability of the pipeline with containers can be achieved in two ways: either a single container including all the tools required by the pipeline is provided, or several containers (one container for each tool to be as modular as possible) are provided. We decided to retain this second way, as this one container - one tool strategy was strongly recommended by several experts (Gruening et al., 2018) who stated that Each container should encapsulate only one piece of software that performs a unique task with a well-defined goal (e.g. sequence aligner, mass spectra identification). Moreover, the one container - one tool strategy has the following advantages:

- most of the pipelines share tools in common meaning that, once a container is built for one tool, it can be easily reused in other pipelines,

- sometimes, different tools that might be incompatible with each other are required, making the usage of a single container including all the tools for the pipeline impossible,

- building a container with all the tools can be very long and possibly tedious. Each time the pipeline changes, the single container has to be rebuilt. With the one container - one tool strategy, only the container in which the tool has changed has to be updated (which is faster),

- the building of the containers can be parallelized to speed-up the deployment of containers.

Generally, the building of the containers occurs in the very late stage of the development cycle of the pipeline, once it has been validated by the end-users and is ready to be deployed in production. Therefore, the guidelines also took into consideration that the pipeline could simply be run during its very early stage of prototyping by bioinformaticians and statisticians, without systematically building the containers whenever a new version of the tool is tested or a new tool is added. This challenge can be easily tackled with the Conda packaging system (Gruening et al., 2018) which offers great flexibility to bioinformaticians and statisticians in order to install and test new tools (when available in the different Conda channels, which is the case for most of the bioinformatics tools). For this reason, the Geniac toolbox allows for the possibility of running the pipeline with different Nextflow profiles (a profile is a set of configuration attributes
which can be activated/chosen when launching a pipeline execution by using the `-profile` command line option with Nextflow) including:

- **singularity** to use the Singularity containers.
- **docker** to use the Docker containers.
- **conda** to use a single Conda environment on which all the tools are available.
- **multiconda** to use a dedicated Conda environment for each tool.
- **path** to define the PATH environment variable for which all the tools are assumed to be installed in the same directory.
- **multipath** to define a dedicated PATH environment variable for which each tool is installed individually.

In addition to these profiles which set where the tools are accessible, we defined the **cluster** profile which can be combined with the previous profiles so that Nextflow launches the computation on a high-performance computing cluster.

As the **Geniac** toolbox automatically generates the configuration files used by these different profiles from the source code, it ensures that they are consistent with each other. Moreover, it saves time for the developer who does not have to write them one by one. All these different profiles are made available with the **Geniac** toolbox so that it covers all of the preferred work habits of a developer. However, we strongly encourage the use of **multiconda** during prototyping and **singularity** for production. Of note, both **conda** and **path** profiles may not work if the pipeline needs tools that are not compatible with each other.

The **one container - one tool** strategy has to be translated in terms of coding best practices with Nextflow so that the source code can be automatically parsed to generate these Nextflow configuration files and build the containers. The coding relies mainly on the use of the **label** directive for each process which uses only one tool and one container at a time, along with the **withLabel** process selector defined in each configuration file for the different profiles which can be run with the `-profile` Nextflow option. Figure 1 illustrates this principle on two tools and three Nextflow processes.

**Availability of the tool**
When implementing a new tool in a pipeline, the main question to answer is **Where is the tool available?** Depending on the answer, the developer has to follow specific guidelines which are fully detailed in the

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**Figure 1.** **one container - one tool** strategy and coding of the Nextflow pipeline: each tool that is used by the pipeline has its own container. In the different configuration files which are specific to a Nextflow profile, each tool is declared with the **withLabel** process selector. To use the appropriate tool in a Nextflow process, the use of the **label** directive with the name of the tool makes the link. The same tool can be used in different processes when different tasks have to be performed at different stages or with different options.
Geniac documentation. Whenever possible, it is recommended to use the tool from the Conda packaging system (Gruening et al., 2018). However, not all tools are available in Conda. Therefore, we guide the developer through a series of chronological questions which redirect the developer to the appropriate section whenever an answer is Yes. The questions are the following: i) Is it just a standard Unix command?, ii) Is it available in Conda?, iii) Is it available only as a binary or as an executable script?, iv) Is the source code available?, v) Have you still not answered yes? Briefly, the developer has to perform some actions such as modify the conf/geniac.config file, copy the source code of the tool in the modules/fromSource folder, write some recipes in the folder recipes, copy some files in the bin or recipes/dependencies folders, depending on the answer to the questions. In most of the cases, the tool should be available in Conda which is the most comfortable situation as it allows the automation of the creation of the containers and the configuration files for Nextflow profiles. However, the toolbox is able to tackle any other situations, although this requires more configuration on the developer’s side. Figure 2 describes the overall workflow with Geniac.

Implementation

It is important to note that the Geniac toolbox (Hupé et al., 2022a) relies on the structure of the Geniac Template (Servant & Hupé, 2022) while the latter could work without Geniac (provided that the missing configuration files for the different Nextflow profiles are added manually). It also means that the Geniac documentation explains how the Geniac Template works with the Geniac toolbox. Therefore, the source code must be organized as shown in Figure 3A with some mandatory files from the Geniac Template in blue and the Geniac toolbox itself in the eponymous folder in green. Based on this organization, the automatic generation of the Nextflow configuration files, container recipes and images are performed by some Nextflow scripts (geniac/install/singularity.nf and geniac/install/docker.nf) which are invoked by Cmake scripts. The file nextflow.config includes all the configuration files that are automatically generated by Geniac to define the different Nextflow profiles. As it will be illustrated in the Use Cases section, the file conf/geniac.config is essential as it allows registration of the tools available from Conda. The main Cmake script is the file geniac/CMakeLists.txt which invokes a series of functions from the geniac/cmake folder. The command line interface (CLI) corresponds to a standalone python package within the geniac/src folder. It includes a linter which offers the possibility for the developer to check the code’s compliance with the guidelines. The linter includes the following tests: i) existence of mandatory or optional files and folders in the Nextflow project tree structure, ii) configuration and consistency of labels with process directives in the Nextflow pipeline, iii) validation of the geniac configuration file, iv) availability of tools and their dependencies. Similarly to the nf-core linter, the output can be exploited manually or automatically within a continuous integration pipeline since any critical error reported by the linter changes the exit code from 0 to 1. The CLI also provides other commands for users not familiar with cmake or make commands to initiate a working directory, install and test the pipeline.

Figure 2. General principle of the Geniac guidelines and toolbox: 1/ a new tool is added in the pipeline according to the guidelines in the Geniac documentation depending on where the tool is available, 2/ the toolbox parses the structure of the source repository and the content of the conf/geniac.config file in order to automatically generate all the configuration files which define the Nextflow profiles, 3/ during the parsing, it also generates the container recipes (here the Singularity Definition Files) which are used to 4/ build the containers (here the Singularity images).
In order to ensure the reproducibility with Singularity, the `geniac/install/singularity.nf` script creates the `singularity.config` file such that Singularity is executed with specific options. Indeed, the default behavior of Singularity makes it possible to access the user’s `$HOME` directory from the host on which the process is executed. Some programming languages such as Python or R preemptively load libraries from the user `$HOME` if they exist rather than the libraries from the canonical installation. To tackle this major issue, the `singularity.config` file i) sets the option “-B "$PWD":/tmp --containall” which is passed on the Singularity command line when invoked by Nextflow and ii) sets “singularity.autoMounts = false” such that the developer explicitly decides during the deployment what folders have to be available within the container. The `--containall` sets some directories as empty (including `$HOME` and `/tmp`) thus avoiding possible libraries installed in the user’s HOME to be loaded by these programming languages. The `-B` option allowing the binding of the `/tmp` (inside the container) to the work directory of the Nextflow process (i.e. in the folder like `work/2d/8202294` created by Nextflow during the execution) is also necessary. Indeed, the default overlay size of `/tmp` inside the Singularity image is few MB that makes the writing of temporary files required by many tools impossible due to space limitation. We preferred to bind `/tmp` to the work directory of the Nextflow process instead of on the actual `/tmp` of the host since it might use some RAM (tmpfs) which is not reported to the job scheduler (such as SLURM or PBS). Our configuration isolates the `$HOME` from the host whatever the programming language used inside a Nextflow process which makes this solution universal. However, this solves this issue only for the `singularity` profile. In the case of the conda or miconda profiles, we addressed this issue by adding ad-hoc environment variables using the beforeScript directive `beforeScript = "export R_LIBS_USER="-"; export R_PROFILE_USER="-"; export PYTHONNOUSERSITE=1"` in the files.

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**Figure 3. Organization of the different directories.** A) source code directory: in blue are listed the mandatory files which could be retrieved from the Geniac Template, in green appears the folder of the Geniac toolbox itself which contains the files from the Geniac repository. Examples of source code are available on the GitHub repositories for both the Geniac Demo and the Geniac Demo DSL2 (in red is highlighted what is specific to a Nextflow pipeline which is implemented with DSL2). B) build directory: once the pipeline has been configured with `cmake` and built with `make`, a `workDir` folder appears with all the files which are automatically generated by Geniac. C) install directory: during the installation with `make install` several folders are created such as the `pipeline` folder which contains the Nextflow code. The files located within the same block in the build directory are copied in the folder of the same color in the install directory.
which are automatically generated by Geniac. If any other programming language suffers from the same behavior, 
*ad-hoc* solutions will have to be added. Note that the default options when using Docker containers within 
Nextflow avoid access to $HOME, this profile is therefore not concerned by the reproducibility issue.

**Operation**
The Geniac toolbox is intended to run on a Linux distribution with Nextflow (>= 21.10.6), git (>= 2.0), Cmake
(>= 3.0), Make (>= 4.1) and Conda (>= 4.10.1). To use the containers, either Apptainer/Singularity (>= 3.8.5) or Docker (>= 18.0) are needed. The Geniac command line interface requires python (>= 3.10).

**Use cases**
In this section, several use cases are described. They are available in the bash script data/useCases.bash
from the Geniac repository so that the reader can reproduce them step-by-step. The use cases depend on
each other meaning that they must be sequentially performed from the beginning. They also require the
installation of Nextflow (>= 20.01.0), git (>= 2.0), Cmake (>= 3.0), Make (>= 4.1), Conda (>= 4.10.1) and
Singularity. The conducting line of this section is the deployment of a pipeline to be run in routine
*production* with Singularity on a high-performance computing cluster having SLURM as a job-scheduler. A
comprehensive overview of the Geniac functionalities is available in the Geniac documentation.

Create the geniac conda environment
In order to run the different use cases, create the geniac conda environment which provides all the dependencies
needed:

```bash
export GENIAC_CONDA="https://raw.githubusercontent.com/bioinfo-pf-curie/geniac/release/environment.yml"

wget ${GENIAC_CONDA}
conda env create -f environment.yml
conda activate geniac
```

Add a Nextflow process for a tool available in Conda
We explain here how to add a new process in the pipeline when the tool is available in Conda. First, the
source code from the Geniac Demo ([Hupé et al., 2022b](#)) repository can be downloaded as follows:

```bash
export WORK_DIR="${HOME}/tmp/myPipeline"
export SRC_DIR="${WORK_DIR}/src"
export INSTALL_DIR="${WORK_DIR}/install"
export BUILD_DIR="${WORK_DIR}/build"
export GIT_URL="https://github.com/bioinfo-pf-curie/geniac-demo.git"

mkdir -p ${INSTALL_DIR} ${BUILD_DIR}
# clone the repository
# the option --recursive is needed if you use geniac as a submodule
git clone --recursive ${GIT_URL} ${SRC_DIR}
```

Let’s assume that the tool `multiqc` ([Ewels et al., 2016](#)) must be added in the pipeline. The section
params.geniac.tools of the file `${SRC_DIR}/conf/geniac.config` must contain the following line:

```plaintext
multiqc = "conda-forge::lzstring=1.0.4=py_1001
             conda-forge::matplotlib-base=3.1.1=py37h250f245_2
             conda-forge::spectra=0.0.11=py_1 bioconda::multiqc=1.8=py_2"
```

The syntax follows the pattern from the conda package naming convention, so that the version is
explicit as recommended in some guidelines ([Gruening et al., 2018](#)).

```plaintext
toolName = "condaChannelName::softName=version=buildString"
```

If other Conda dependencies are required, they are added according to the same convention as shown for `multiqc`. In order to use the tool in a Nextflow process, use the `label` directive using the exact same
Add a Nextflow process for a tool available from source code

As not all the tools are available in Conda, we explain here how to add a new process in the pipeline when the tool is available from source code. It offers a great flexibility as the software developer can control the source code and tune the installation process. However, this obviously requires more configuration. In particular, the software developer has to be fluent with Cmake.

Let’s assume that the tool helloWorld must be added in the pipeline. Its source code must be copied in the folder `${SRC_DIR}/modules/fromSource/helloWorld`. In order to use the tool in a Nextflow process, use the label directive using the exact same name as given to this folder (i.e. helloWorld). Then, the Cmake template script `{SRC_DIR}/geniac/data/modules/fromSource/CMakeLists.txt` is copied into `{SRC_DIR}/modules/fromSource/CmakeLists.txt` and the following Cmake directive is added:

```cmake
ExternalProject_Add(
    helloWorld
    SOURCE_DIR ${CMAKE_CURRENT_SOURCE_DIR}/helloWorld
    CMAKE_ARGS
    -DCMAKE_INSTALL_PREFIX=${CMAKE_BINARY_DIR}/externalProject/bin)
```

Lastly, the `${SRC_DIR}/modules/fromSource/helloWorld/CmakeLists.txt` Cmake script must be written to detail how the tool must be compiled and installed, for example:

```cmake
project(helloWorld LANGUAGES CXX)
project (helloWorld)
add_executable(helloWorld main.cpp)
install(TARGETS helloWorld DESTINATION ${CMAKE_INSTALL_PREFIX})
```

Check the code with the linter

The Geniac command line interface is available in the conda environment which has been created and activated at the beginning of the Use cases section.

Run the linter on your repository:

```bash
geniac lint ${SRC_DIR}
```

Build the Singularity containers

It consists of two main steps. First, the pipeline is configured with the required cmake options. The options with the pattern `-DCMAKE_<UPPER_CASE>=<value>` correspond to the variables available in the cmake language (CMAKE_INSTALL_PREFIX defines where the analysis pipeline will be deployed). The options with the pattern `-Dap_<lower_case>=<value>` correspond to the ad-hoc variables defined by the Geniac toolbox (see the Geniac documentation for the list of available options). In this use case, the option `-Dap_install_singularity_images=ON` tells Geniac to build and install the Singularity images and `-Dap_nf_executor=slurm` to set SLURM as the executor (i.e. the job-scheduler) in the cluster profile used by Nextflow. Secondly, it starts the build of the pipeline with the make command. These two steps work as follows:
### configure the pipeline

```bash
cd ${BUILD_DIR}

### cmake

cmake ${SRC_DIR}/geniac -DCMAKE_INSTALL_PREFIX=${INSTALL_DIR}
  -Dap_install_singularity_images=ON -Dap_nf_executor=slurm

### with sudo, the singularity and nextflow commands must be
### in the secure_path option declared in the file /etc/sudoers

### build the files needed by the pipeline

sudo make

### change file owner/group to the current user

sudo chown -R $((id -gn):$(id -gn)) ${BUILD_DIR}
```

During the build process, Genia c generates the different configuration files, the Singularity recipes and images in the `${BUILD_DIR}/workDir` folder as shown in Figure 3B. In particular, the `${BUILD_DIR}/workDir/results/conf/singularity.config` file needed to run the singularity profile is created at this stage. Note that these files are not present in the `${SRC_DIR}` as they are automatically generated. The root credentials are needed to build the Singularity images thus requiring the use of the `sudo` commands.

### Deploy the pipeline

The deployment of the pipeline is performed with the `make` command:

```
make install
```

After the deployment, the pipeline is available in the `${INSTALL_DIR}/pipeline` folder in which the `main.nf` Nextflow file is available (see Figure 3C). Other folders are also created which contain the Singularity images, a folder (or a symlink depending on the option used during the configuration) for the genome annotations, and folders for the path and multipath profiles. In particular, the `${BUILD_DIR}/workDir/results/conf/singularity.config` file created in the previous stage is deployed in the `${INSTALL_DIR}/conf/singularity.config` file needed to run the singularity profile.

### Test and run the pipeline

As testing is generally a challenging part of the development and maintenance of the pipeline, we strongly recommend to create a `conf/test.config` profile along with a minimal dataset in the `test/data` folder such that it can be processed quickly to check that new developments did not introduce error. In this section, we show how to run the pipeline using this test profile.

The pipeline can be run with the `singularity` profile locally on the computer as follows:

```
cd ${INSTALL_DIR}/pipeline
nextflow -c conf/test.config run main.nf -profile singularity
```

The pipeline can be run with the `singularity` and `cluster` profiles on a high-performance computing cluster with the SLURM job-scheduler as follows:

```
nextflow -c conf/test.config run main.nf -profile singularity,cluster
```

Of note, running the pipeline with the `test` profile can be directly done from the `${BUILD_DIR}` directory using the commands `make test_singularity` or `make test_singularity_cluster` respectively.

### Geniac command line interface

For users not familiar with `cmake` or `make` commands we propose a command line interface. The use case can be performed with the following commands:

```
```
```
export WORK_DIR="${HOME}/tmp/myPipeline_CLI"
export INSTALL_DIR="${WORK_DIR}/install"
geniac init -w ${WORK_DIR} ${GIT_URL}
cd ${WORK_DIR}
geniac lint
/geniac install . ${INSTALL_DIR} -m singularity
sudo chown -R $((id -gn)):$(id -gn) build
geniac test singularity
/geniac test singularity --check-cluster
```

Compatibility with Nextflow DSL2

Geniac is fully compatible with Nextflow DSL2 provided that the process, workflows and subworkflows are organized in the `${SRC_DIR}/nf-modules` folder as shown in red in Figure 3A. The Geniac Demo DSL2 (La Rosa et al., 2022) repository can be used instead of the Geniac Demo. To do so just set, at the very beginning of the Use cases section, the value of GIT_URL to:

```
export GIT_URL="https://github.com/bioinfo-pf-curie/geniac-demo-dsl2.git"
```

Conclusions

Geniac consists of a toolbox with three components: a technical documentation available at https://geniac.readthedocs.io, a command line interface with a linter to check that the code respects the guidelines, and an add-on to generate configuration files, build the container and deploy the pipeline.

Compared to nf-core, Geniac builds upon their best-practises by adding additional guidelines, but is more flexible and can be easily adjusted to any pipeline template. More precisely, the only requirement in Geniac is the use of the label directive for each process which should use only one tool and one container at a time. The use of labels enables the one container - one tool strategy which was motivated by the development of Geniac. To do so, the specific `conf/geniac.config` file is added in the pipeline repository along with a specific profiles section (which differs from nf-core) in the nextflow.config to use Geniac.

The Geniac Demo and Geniac Demo DSL2 pipelines are available so that the reader can practice the Geniac guidelines. The Geniac Template makes it possible to start a new pipeline from scratch. The Geniac toolbox is fully compatible with Nextflow DSL1 and DSL2. While it supports the automatic generation of Docker and Singularity containers, it is straightforward to add any other containerization framework such as podman or shifter. The development of the Geniac toolbox was strongly motivated by the context of production analysis. However, we strongly encourage the use of Geniac in the context of the research development analysis. Our Bioinformatics Core facility uses the Geniac toolbox for a ten of pipelines deployed for production analysis covering a large variety of sequencing applications with both short-read and long-read technologies: it includes quality controls, analysis of ChIP-seq, ATAC-seq, RNA-seq, Whole Exome-seq, Whole Genome-seq and CRISPR data.

Data availability

All data underlying the results are available as part of the article and no additional source data are required.

Software availability

- Source code for geniac available from: https://github.com/bioinfo-pf-curie/geniac
- Archived source code at time of publication: https://doi.org/10.5281/zenodo.6039812 (Hupé et al., 2022a)
- Source code for geniac-demo available from: https://github.com/bioinfo-pf-curie/geniac-demo
- Archived source code at time of publication: https://doi.org/10.5281/zenodo.6040173 (Hupé et al., 2022b)
- Source code for geniac-demo-dsl2 available from: https://github.com/bioinfo-pf-curie/geniac-demo-dsl2
- Archived source code at time of publication: https://doi.org/10.5281/zenodo.6040166 (La Rosa et al., 2022)
- Source code for geniac-template available from: https://github.com/bioinfo-pf-curie/geniac-template
- Archived source code at time of publication: https://doi.org/10.5281/zenodo.6040058 (Servant & Hupé, 2022)
- All documentation is available at: https://geniac.readthedocs.io
- License: CeCILL Version 2.1

Author contributions
J.R. and P.H. conceived the coding best practices and automation principles proposed by Geniac. P.H. developed Geniac and Geniac Demo. P.L.R. developed Geniac Demo DSL2. N.S. and P.H. developed the Geniac Demo command line interface. F.J. tested the different demo pipelines. P.H. wrote the manuscript and the documentation. P.H. supervised the study.

Acknowledgements
We are grateful to our colleagues from the Bioinformatics Core Facility for their feedback on the Geniac toolbox and from the ICT department.

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Version 2

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I thank the authors for the work they did to address the points I raised and to make the manuscript clearer.

I have no further comments to make.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: I work in the field of evolutionary bioinformatics where I develop tools and workflows. I am interested in the reproducibility of bioinformatics analyses, and I use Nextflow and container technologies on a daily basis.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Reviewer Report 31 August 2021

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Frédéric Lemoine

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This manuscript describes Geniac, a toolbox aiming at increasing the reproducibility and reusability of Nextflow workflows. To do so, Geniac has the following features:

1. It defines a set of good practices for the development of Nextflow workflows, via a detailed documentation.

2. It provides a linter to check that the best coding practices are respected.

3. It provides a "add-on" that automatically builds dependencies, generates the singularity (or docker) containers needed to run the workflow, checks the configuration files, and installs the workflow in the specified directory ("deploy").

Geniac tackles an important issue with bioinformatic analyses: Pipelines are rarely homogeneous in terms of description and implementation, which leads to reproducibility, maintenance, and reusability difficulties. In terms of goals, Geniac present some similarities with nf-core, which aims at curating and harmonizing pipeline implementations. The solution proposed by Geniac is to define a set of homogeneous practices for workflow development (e.g. how to describe process dependencies, how to name variables, channels and files, and provide tools to check workflows, build containers and install the workflow). The price to pay for more reproducible and homogeneous workflows is a more complex technological stack (with CMake specifically) and workflow architecture. Technically, Geniac is based on the following technologies: Nextflow, conda, Docker (or Singularity) and CMake. Although technical, the manuscript explains clearly the philosophy, the goals, the usage and the usefulness of Geniac. The tools are usable in the provided workflow example, I was able to run the geniac-demo and the linter works well and is easy to use.

I believe that Geniac is a useful tool that answers important questions. However, I think it would be easier to understand and use with a few clarifications in my opinion:

- I had some initial difficulties to run the geniac-demo workflow. I think the documentation (on readthedoc and on the git repositories) could be clearer about how to run the workflow in different situations, at least with docker or singularity. For example, I initially had an error when the install and the build directories were included in the geniac-demo directory. Before understanding the general philosophy of Geniac, I also had some difficulties to understand how to generate the singularity.config file needed to generate singularity images and run the workflow. A more visible step by step concrete quick-start example could be very useful.

- Following the previous point, I was a bit confused by the README in the geniac-demo and geniac-template repositories, where I was looking for information on how to run the example. The documentation could be better adapted to the workflows. For example the geniac.md file refers to nf-CRISPR repository.

- Testing the workflow is also generally a difficult part of the development and maintenance
of workflows. It is not mentioned clearly in the introduction or in the discussion, though I think that tests can be performed using test profiles in Geniac. I think it would be great to give a little bit more insights on what tests consist of in geniac workflows.

- The abstract states that workflow systems + containers increase usage complexity. I am not sure that Geniac tackles this problem, since it adds a new layer of complexity in the top of Nextflow, Docker, etc (which is unavoidable). If I understand correctly, Geniac is mainly solving the problem of workflow homogeneity in terms of structure, description, and documentation, and thus make the workflow more reproducible, maintainable and reusable.

- A more detailed description of the specificities of Geniac compared to nf-core may be a great addition to the manuscript.

- Are there already any public real-world Geniac compatible workflows in production? If so, it could be interesting to mention a few of them.

A few typos/suggestions:

- Abstract+Introduction: "which address" => "which target"?

- Introduction: "the workflow has no choice but deliver": Isn't it more the facility rather than the workflow that has no choice?

- Introduction: "being less invasive to the different expert communities": This point is not very clear to me. Would it be possible to rephrase it?

- Methods: "to be as granular as possible", maybe clearer with "to obtain the finest level of granularity"?

- Methods: "The building of the containers can be parallelized to reduce the time in which...". It is not very clear to me. Would something like "to speed up the deployment of new containers" make sense?

- The DOI link in the ref Servant & Hupé 2021 does not work (there is an additional space in the URL).

- Methods: "Other folders are also created which contains"=>"contain".

Is the rationale for developing the new software tool clearly explained?
Yes

Is the description of the software tool technically sound?
Partly

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?
Partly
Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?  
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** I work in the field of evolutionary bioinformatics where I develop tools and workflows. I am interested in the reproducibility of bioinformatics analyses, and I use Nextflow and container technologies on a daily basis.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 11 Feb 2022

Philippe HUPE

First, we would like to thank the reviewer. We are very grateful for his time, contribution and very valuable comments which significantly helped to improve the article and the documentation of the Geniac. You will find below a detailed answer to the different issues that we have addressed in the revised manuscript. Best regards, Philippe Hupé. (Reviewer comments in italics)

I had some initial difficulties to run the geniac-demo workflow. I think the documentation (on readthedoc and on the git repositories) could be clearer about how to run the workflow in different situations, at least with docker or singularity. For example, I initially had an error when the install and the build directories were included in the geniac-demo directory. Before understanding the general philosophy of Geniac, I also had some difficulties to understand how to generate the singularity.config file needed to generate singularity images and run the workflow. A more visible step-by-step concrete quick-start example could be very useful. Following the previous point, I was a bit confused by the README in the geniac-demo and geniac-template repositories, where I was looking for information on how to run the example. The documentation could be better adapted to the workflows. For example the geniac.md file refers to nf-CRISPR repository.  

Reply: We are sorry that the geniac-demo was not relevant as it was simply a copy/paste from the geniac-template. Therefore, the geniac-demo documentation has been modified. As suggested by the reviewer1, we included in the README a "Quick start" section such that the reader can test and run the pipeline either with the multiconda or the singularity profile using the new command line interface.

Testing the workflow is also generally a difficult part of the development and maintenance of workflows. It is not mentioned clearly in the introduction or in the discussion, though I think that tests can be performed using test profiles in Geniac. I think it would be great to give a little bit more insights on what tests consist of in geniac workflows.  

Reply: Testing is indeed a major component of software development. Therefore, the section "Run the pipeline" has been renamed "Test and run the pipeline". We highlighted
the necessity to include a test profile and test data. We have also indicated that geniac integrates several Make commands to facilitate the test of the pipeline.

The abstract states that workflow systems + containers increase usage complexity. I am not sure that Geniac tackles this problem, since it adds a new layer of complexity in the top of Nextflow, Docker, etc (which is unavoidable). If I understand correctly, Geniac is mainly solving the problem of workflow homogeneity in terms of structure, description, and documentation, and thus make the workflow more reproducible, maintainable and reusable.

Reply: The abstract has been modified as suggested to mention that "promoting the homogeneity of the workflow implementation requires guidelines and protocols" such as Geniac intends.

A more detailed description of the specificities of Geniac compared to nf-core may be a great addition to the manuscript.

Reply: We added a new paragraph in the Conclusion to explain how Geniac compares to nf-core.

Are there already any public real-world Geniac compatible workflows in production? If so, it could be interesting to mention a few of them. Indeed, geniac is used in our Bioinformatics Core Facility.

Reply: This has now been added in the Conclusion.

A few typos/suggestions: > * Abstract+Introduction: "which address" => "which target"?

Reply: This has been corrected.

Introduction: "the workflow has no choice but deliver": Isn't it more the facility rather than the workflow that has no choice?

Reply: This is right, the sentence has been rephrased.

Introduction: “being less invasive to the different expert communities”: This point is not very clear to me. Would it be possible to rephrase it?

Reply: This has been rephrased to explain that geniac wants to preserved the preferred work habits of the developers.

Methods: "to be as granular as possible", maybe clearer with "to obtain the finest level of granularity"?

Reply: We have replaced granular by modular since the idea of "one container - one tool" strategy is that each container can be seen as a module which performs a given task.

Methods: "The building of the containers can be parallelized to reduce the time in which...". It is not very clear to me. Would something like "to speed up the deployment of new containers" make sense?

Reply: Indeed, the sentence has been rephrased as suggested.

The DOI link in the ref Servant & Hupé 2021 does not work (there is an additional space in the URL).

Reply: This has been corrected.
Toni Hermoso Pulido

1 Centre for Genomic Regulation, The Barcelona Institute of Science and Technology, Barcelona, Spain
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In this article, the Geniac toolbox is presented. The software consists of several tools to address some common difficulties of deploying and adapting Nextflow pipelines to fit into different computing environments. Its major emphasis is ensuring both code reproducibility and production-level reliability. Technically speaking, it depends on Nextflow itself and CMake automation and packaging tool. A Python-based code analysis (linter) tool is also provided for checking pipeline setup in advance before the actual build and deployment process takes place.

Provided software for the pipelines relies primarily on Conda package manager, which Geniac facilitates to adapt to different approaches (e.g., single or multi), and converts to handy container images such as Singularity-based ones.

Experienced practitioners familiar with Nextflow, Conda and container technologies can easily follow the instructions and customize the provided examples (which are based on both legacy and DSL2 Nextflow syntax). Maybe the final result and associated folders of the pipeline preparation (build, install), as depicted in Figure 3, could be more clearly stated.

Despite it being mentioned in the article and in the toolbox documentation as an actual possibility, a more explicit example case of a software step that does not rely on Conda could be a nice addition.

As a user experience suggestion for future releases, it could be interesting that the geniac executable could be already provided straight from the Python Package Index (PyPI). It would also help new adopters, specially if not so familiar with CMake, if geniac could wrap some of the execution calls that are included in the useCases.bash script.
Is the rationale for developing the new software tool clearly explained? 
Yes

Is the description of the software tool technically sound? 
Yes

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others? 
Yes

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool? 
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** I work in Bioinformatics. I have experience developing and deploying Nextflow pipelines, and I regularly take care of infrastructure and devops-related tasks.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Author Response 11 Feb 2022

Philippe HUPE

First, we would like to thank the reviewer. We are very grateful for his time, contribution and very valuable comments which significantly helped to improve the article and the documentation of the Geniac. You will find below a detailed answer to the different issues that we have addressed in the revised manuscript. Best regards, Philippe Hupé. (Reviewer comments in italics)

In this article, the Geniac toolbox is presented. The software consists of several tools to address some common difficulties of deploying and adapting Nextflow pipelines to fit into different computing environments. Its major emphasis is ensuring both code reproducibility and production-level reliability. Technically speaking, it depends on Nextflow itself and CMake automation and packaging tool. A Python-based code analysis (linter) tool is also provided for checking pipeline setup in advance before the actual build and deployment process takes place. Provided software for the pipelines relies primarily on Conda package manager, which Geniac facilitates to adapt to different approaches (e.g., single or multi), and converts to handy container images such as Singularity-based ones. Experienced practitioners familiar with Nextflow, Conda and container technologies can easily follow the instructions and customize the provided examples (which are based on both legacy and DSL2 Nextflow syntax). Maybe the final result and associated folders of the pipeline preparation (build, install), as depicted in Figure 3, could be more clearly stated.

**Reply:** The Figure 3B and 3C have been modified to highlight where the files automatically generated in the build directory are copied in the install directory.
Despite it being mentioned in the article and in the toolbox documentation as an actual possibility, a more explicit example case of a software step that does not rely on Conda could be a nice addition.

Reply: We added a new section in the manuscript to describe how to "Add a Nextflow process for a tool available from source code".

As a user experience suggestion for future releases, it could be interesting that the geniac executable could be already provided straight from the Python Package Index (PyPI).

Reply: The Geniac command line interface is now available on PyPI at https://pypi.org/project/geniac/.

It would also help new adopters, specially if not so familiar with CMake, if geniac could wrap some of the execution calls that are included in the useCases.bash script.

Reply: New options have been added to the Geniac command line interface to facilitate the use of geniac for developers not familiar with CMake commands. The new section "Geniac command line interface" has been added in the manuscript to illustrate what command lines can be used for the proposed use case. The documentation has been updated with these new functionalities.

Competing Interests: No competing interests were disclosed.