SOFTWARE TOOL ARTICLE

**chaste codegen: automatic CellML to C++ code generation with fixes for singularities and automatically generated Jacobians [version 1; peer review: 2 approved, 1 approved with reservations]**

Maurice Hendrix\(^1,2\), Michael Clerx\(^1\), Asif U Tamuri\(^3\), Sarah M Keating\(^3\), Ross H Johnstone\(^4\), Jonathan Cooper\(^3\), Gary R Mirams\(^1\)

\(^1\)Centre for Mathematical Medicine & Biology, University of Nottingham, Nottingham, UK
\(^2\)Digital Research Service, Information Sciences, University of Nottingham, Nottingham, NG8 1BB, UK
\(^3\)Centre for Advanced Research Computing, University College London, London, WC1E 6BT, UK
\(^4\)Computational Biology & Healthcare Informatics, Department of Computer Science, University of Oxford, Oxford, OX1 3QD, UK

**Abstract**

Hundreds of different mathematical models have been proposed for describing electrophysiology of various cell types. These models are quite complex (nonlinear systems of typically tens of ODEs and sometimes hundreds of parameters) and software packages such as the Cancer, Heart and Soft Tissue Environment (Chaste) C++ library have been designed to run simulations with these models in isolation or coupled to form a tissue simulation. The complexity of many of these models makes sharing and translating them to new simulation environments difficult. CellML is an XML format that offers a solution to this problem and has been widely-adopted. This paper specifically describes the capabilities of chaste_codegen, a Python-based CellML to C++ converter based on the new cellmlmanip Python library for reading and manipulating CellML models. While chaste_codegen is a Python 3 redevelopment of a previous Python 2 tool (called PyCML) it has some additional new features that this paper describes. Most notably, chaste_codegen has the ability to generate analytic Jacobians without the use of proprietary software, and also to find singularities occurring in equations and automatically generate and apply linear approximations to prevent numerical problems at these points.

**Keywords**

CellML, cardiac electrophysiology, code generation, C++, jacobian, singularity, GHK equation

**Open Peer Review**

Approval Status

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1. Axel Loewe\(^1\), Karlsruhe Institute of Technology (KIT), Karlsruhe, Germany
2. David Nickerson\(^1\), University of Auckland, Auckland, New Zealand
3. Rahuman S Malik-Sheriff\(^1\), Wellcome Genome Campus, Hinxton, UK

Any reports and responses or comments on the article can be found at the end of the article.
Corresponding author: Maurice Hendrix (maurice.hendrix@nottingham.ac.uk)

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**Introduction**

Within the area of electrophysiology, there are hundreds of models describing biological behaviour. Many of these models are complex systems of tens of ordinary differential equations (ODEs) with hundreds of parameters, making translation into different simulation software time-consuming and prone to transcription errors. This also makes sharing models between different tools and application areas difficult. CellML addresses this problem by offering a way to describe mathematical models in an XML-based format, independent of the choice of programming language or tools used to simulate or analyse the models. It was originally created with the Physiome Project in mind, and a large repository of well over a hundred CellML electrophysiology models is available on the Physiome Model Repository (PMR, https://models.physiomeproject.org/cellml). CellML sees continued development by the research user base and several tools are available to support modelling using CellML models.

CellML models can be imported into various simulation tools such as the Cancer Heart and Soft Tissue Environment (Chaste), OpenCOR, Myokit, and model comparison tools such as the Cardiac Electrophysiology Web Lab. This paper describes the development of chaste_codegen — a Python 3 CellML code generator which is now used by Chaste. Its functionality is largely inspired by PyCML, a Python 2 implementation of CellML model import into Chaste. However chaste_codegen is an all-new implementation which includes a number of new features previously unavailable to the community, as described below.

**Methods**

**Implementation**

chaste_codegen builds on the cellmlmanip library and Jinja2 templates to generate Chaste C++ code from CellML files. It is available as a standalone command-line tool for Python 3 on Windows Linux and Mac, and has been integrated into the latest release of Chaste.

CellML is a model definition language and as such it describes the model and its equations, but does not describe how the the equations should be solved or how experiments are run. cellmlmanip is a new Python 3 library for parsing CellML into ordered equations and metadata. It is a flexible component that can read CellML and enable it to be used for a variety of purposes, such as translating CellML models into other formats, or into code for various simulation packages. Separation of the parser and simulator has the advantage of creating a resource for the CellML community, which will ultimately be much easier to maintain than writing a bespoke CellML parser for each application. Key features of cellmlmanip are: using SymPy to represent mathematics (with the full manipulation capabilities thereof available), tracking physical units and performing conversions as needed, and managing and querying RDF metadata annotations on models. Cellmlmanip currently supports CellML version no 1.0, but it could easily be adapted to support CellML version no 2.0. However there are currently no plans to do this, because of the ongoing development of libcellml. libcellml is a more general propose CellMI 2.0 library should offer the ability to not only read and manipulate CellMI 2.0 models but also write adjusted models back again.

SymPy is a python library for symbolic mathematics that offers a number of convenient features for chaste_codegen. Most notably, it provides us with the ability to calculate Jacobians algebraically, to recognise patterns, rewrite equations, and to extract common terms in a set of equations. It also comes with a convenient printing mechanism, separating the mathematics from their representation.

Jinja2 is a templating language for Python, modelled after Django’s templates. Using Jinja2 allows us to separate the logic from the code output, which allows generating code for a number of different solvers, as described in detail in Cooper et al. Jinja2 templates should allow easy adaptation to export code in other programming languages. chaste_codegen and its stack of dependencies are all free and open source.

**Analytic Jacobians**

Within Chaste one of the main solver types available is CVODE, which performs well for the stiff systems that feature in many electrophysiology models. CVODE can be sped up if the user provides a method to return the Jacobian matrix for the ODE system (with entries defined as the partial derivative of each ODE’s right-hand-side with respect to each state variable). If this is missing CVODE derives an approximation for it based on finite differences. PyCML required a Jacobian matrix to be pre-computed by the proprietary Maple software. In chaste_codegen we have integrated calculation of a Jacobian matrix at runtime, by making use of the SymPy library.
We contrasted previously existing code, using externally created Jacobians, with the newly generated models with automatically generated Jacobians. The validity of the generated Jacobians was assessed by comparing numerical results of runs of a number of different models.

Singularities
Many electrophysiology models use a formulation for ion currents based on the Goldman-Hodgkin-Katz (GHK) flux equation\textsuperscript{15}, or feature equations with similar structure. Unfortunately such equations can introduce \textit{singularities} into a model. Here, we define singularities as situations where an equation tends to 0/0 close to a particular value of a model variable (usually membrane voltage, $V$).

In general terms, these GHK-style equations take the form:

$$C(V) = f(V) \frac{(V - v_0)}{e^{B(V-v_0)} - 1},$$

where $B$ and $v_0$ are constants, and $f(V)$ is any function of $V$ (that may also be simply a constant) and $v_0$ is the voltage at which we hit the singularity. As per Johnstone\textsuperscript{16}, we can simplify this notation by defining

$$A(V) = \frac{f(V)}{B},$$

and use the substitution

$$U = B \times (V - v_0),$$

to leave Equation (1) with a nondimensional fraction term that encapsulates the singularity,

$$C(V) = A(V) \frac{U}{e^U - 1}.$$  

For convenience we also define

$$g(U) = \frac{U}{e^U - 1}.$$  

We can now see that at as $U \to 0$ there is a singularity as both the top and bottom of the fraction in Equation (5) tend to zero ($e^U \to 1$, so $e^U - 1$ tends to zero).

It is important to point out that mathematically the value of the equation remains well defined; no physics in the model is breaking down as we get close to 0/0. That is, there is a finite value to which the expression evaluates which we will tend towards as we get closer to the singularity (a \textit{limit}), and this can be found analytically using approaches such as L’Hôpital’s rule, as we will discuss later. But when evaluating such equations numerically, it is possible to reach voltages close to (or at) the singularity where the numerical evaluation is not just inaccurate but often tends to $\pm \infty$.

As an example, Figure 1 shows the background calcium current $I_{Ca_{ab}}$ in the Davies \textit{et al.}\textsuperscript{17} model of a canine cardiomyocyte, as a function of trans-membrane voltage ($V$). The figure shows unstable, asymptotically-increasing oscillatory behaviour close to $V = 0$.

By plotting such graphs for a number of cases, Johnstone\textsuperscript{16} found that the range in which instability occurs numerically is within approximately $10^{-7}$ of $U = 0$ when using double precision (64 bits to represent a floating point number in computer memory).

Problems with these singularities are most apparent when running voltage-clamp experiments, when the voltage $V$ is clamped exactly at a singularity voltage $v_0$. Given the infinite number of choices for voltage clamps and parameters within the models, one might expect this to be unusual. However, this situation is very common. Rounded numbers appear as the clamp voltage in experiments, where steps to a handful of voltages are chosen manually by an experimenter; and also in the model equations themselves. This unfortunate collision is most commonly encountered when the model equations feature $v_0 = 0$ mV, as in the standard form of the GHK flux equation itself (Equation (11) in Goldman\textsuperscript{15}). But the situation also occurs in the more general form of
Equation (1) because parameterisation has commonly been done ‘by hand’ in ion channel modelling, and so the \((V – v_0)\) terms often feature round numbers for \(v_0\). As an example, equations of the form of Equation (1) with both \((V + 10)\) and \((V + 25)\) appeared in the original Hodgkin and Huxley\(^{18}\) model, meaning that even this fundamental model requires modification to do voltage clamps to –10 or –25 mV (N.B. the voltage in the original paper is defined relative to resting potential rather than extracellular potential, and some CellML implementations have updated these numbers to the modern convention using a similar process to that described in Brown\(^{19}\)).

Apart from voltage clamp experiments and singularities at ‘round numbers’, any singularity within normal physiological voltage ranges will be crossed during the simulation of an action potential. Perhaps frustratingly, the more accurately you solve the ODEs, the more likely you are to hit these singularities. This is because a simple fixed-timestep solver is very unlikely to hit the narrow region of voltage which causes a problem, but using an adaptive timestep solver with automatic error correction (like CVODE\(^{20}\)) can detect large gradients or sudden changes, introduce more timesteps to refine the solution around these points, ‘home in’ on the singularities and crash. This is definitely not to say that the adaptive solvers are unsuitable for these models, quite the opposite — electrophysiology models frequently have stiff systems with ‘fast sodium current’ upstrokes and other slower processes, and adaptive ODE solvers can provide huge improvements in speed and accuracy. So rather, enabling use of adaptive solvers is an added incentive to avoid the numerical problems associated with singularities.

**Approaches to fix singularities**

It is worth noting that the use of computational optimisations such as using the function ‘expm1’ to represent \(e^U – 1\) do help by increasing precision for small values of \(U\), but they do not alleviate the problem entirely.

There are a number of ways the singularity can be removed. When inspecting existing models, we found many CellML files where L’Hôpital’s rule had already been applied manually when numerical problems had been encountered. In the notation we introduced above it is simple to see how L’Hôpital’s rule applies

\[
\lim_{U \to 0} g(U) = \lim_{U \to 0} \frac{U}{e^U - 1} = \lim_{U \to 0} \frac{dU}{dU} e^U = \lim_{U \to 0} \frac{1}{e^U} = \frac{1}{1} = 1. 
\]

\[\text{Equation (6)}\]
So the ‘fixes’ in most CellML files apply this constant limit value, \( g(U) = 1 \), across a region close to the singularity (with a ‘piecewise’ statement in CellML). Substituting the L’Hôpital limit into Equation (4), this fix is applied as

\[
C(V) = \begin{cases} 
  A(V), & |U| < \epsilon, \\
  A(V) \frac{U}{e^U - 1}, & \text{otherwise.} 
\end{cases}
\]

(7)

Where \( \epsilon \) is a small range, which varies depending on the CellML file but translates to a small region of voltage. In our notation this is equivalent to \(|V - V_0| < \epsilon B|\).

A Taylor expansion around the singularity gives \( g(U) \approx 1 - \frac{U}{2} + \frac{U^2}{12} + \ldots \) and so as a refinement of Equation (7), Johnstone\(^{16}\) suggested using the first two terms of the Taylor expansion rather than just the first term, giving a more accurate approximation close to the singularity:

\[
C(V) = \begin{cases} 
  A(V) \left(1 - \frac{U}{2}\right), & |U| < \epsilon, \\
  A(V) \frac{U}{e^U - 1}, & \text{otherwise.} 
\end{cases}
\]

(8)

In \texttt{chaste_codegen}, we take this approach with a small modification. Rather than using the above expression we decided to simply ‘draw a line’ between the values of \( C(V) \) evaluated using Equation (1) at the \( \epsilon \) bounds of our region, and interpolate from this. This was not actually any simpler to implement, as we still identify \( U \) in order to pick bounds over which to apply the linear approximation. But a benefit of this approach over using Equation (8) is that we get at least \( C^6 \) continuity as we leave the region, even if some curvature in \( C(V) \) is apparent, whereas there can still be (typically very small) discontinuities in \( C(V) \) as we transition between the cases in Equation (8). Note that any discontinuities in Equation (8) are, by definition, much smaller than the discontinuities that appear using Equation (7), as we can see in Figure 1. This continuity can be important to avoid problems in numerical solutions of ODEs, but in practice Equation (8) and the approach we have taken are almost indistinguishable given the size of \( \epsilon \).

The algorithm we implemented within \texttt{cellmlmanip} has two parts. To identify whether an equation has a singularity and apply a fix, the algorithm is:

1. Recursively check within each equation for singularities, term by term.
2. Skip (sub-)equations that are piecewise statements, we assume that if these have a singularity it has a manual fix applied.
3. Find \( U \) using SymPy’s pattern matching capabilities.
4. Solve for \( U = 0 \) to find the singularity point in terms of model variables (usually Voltage, \( V \)).
5. Introduce a piecewise statement to replace the original expression within \(-10^{-7} \leq U \leq +10^{-7}\) (the fact this is now a nondimensional range means that the same fixed range appears to work well for all singularities we have found; it is translated back into voltage ranges of different widths at code generation time).
6. Within this range we use linear interpolation between values evaluated using the original expression at the boundaries (\( U = \pm 10^{-8} \)).
7. We note that \( \frac{U}{-1 + e^U} \) leads to a similar situation, as do \( \frac{e^U - 1}{U} \) and \( \frac{-1 + e^U}{U} \). Therefore these cases are treated similarly.

To ensure we fix the appropriate equation, the algorithm is:

1. A graph is constructed for dependencies of all equations in the model. We can then order them from ODEs at the top level to state variables at the bottom level.
2. All equations in the model are rewritten in terms of state variables, by substituting them into intermediate variables, so that (for instance) \( V \) appears explicitly in all equations that have any dependence on it.
3. We start at the bottom level of the graph, look for singularities and if none are found using the procedure above we progress to the next level of the graph. If a singularity is found we introduce a fix at that node of the graph.

Testing
To test the process we searched a set of CellML models (all those annotated for use with Chaste and the WebLab and available at [https://github.com/chaste/cellml](https://github.com/chaste/cellml)) for any piecewise elements. We then manually identified the subset of these being used to fix singularities. We removed these fixes from the CellML files then verified that our code would indeed find and fix these singularities automatically. The result of this exercise is shown in Table 1. The table shows the number of previously hard-coded fixes and the number of

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</table>
extra fixes detected for the range of CellML files. In short, all 106 singularities with previously hard-coded fixes were identified, and the code found 263 new singularities that had never been ‘fixed’ within the CellML files. An overview of all 369 fixes, each with a similar plot to Figure 1, is available in the ‘assets’ branch of the chaste_codegen GitHub page.

There are a number of advantages in removing hard-coded singularity fixes from the CellML files: we can make more accurate fixes around singularities (as shown in Figure 1); others could choose how to treat the singularities and adapt our code to use other methods if they wish; and finally the CellML model is simplified to represent the equations and not how they should be solved. A disadvantage in removing hard-coded singularity fixes from the main CellML repository (PMR) is that all translation code would need to adopt an approach similar to the one we have outlined to avoid hitting singularities, whereas currently around 80 to 90 hard-coded fixes will appear in any generated code without code generation tools needing to do any special treatment. Note that the figure of 106 singularities includes approximately 20 hard-coded fixes we applied to our subset of CellML models using Equation (8) in previous work, which do not feature in the main CellML repository (PMR). Dealing with singularities at code generation time is more future-proof for when people add new models, rather than hard-coding the fixes we applied here into the CellML files in the Physiome Model Repository (PMR) and then having to periodically repeat this exercise for any new models that have been added. On balance, we think that this automated approach at code generation time is the preferable route and hope it has been described so that others can reproduce it in their own code generation software, or re-use components of our open source implementation. In the remainder of this article we discuss the practicalities of using chaste_codegen.

<table>
<thead>
<tr>
<th>CellML file</th>
<th># previously hard-coded fixes (all auto-fixed when removed)</th>
<th># extra detected (all auto-fixed)</th>
<th>total (all auto-fixed)</th>
</tr>
</thead>
<tbody>
<tr>
<td>sachse model 2007</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>sakmann model 2000 epi</td>
<td>4</td>
<td>6</td>
<td>10</td>
</tr>
<tr>
<td>shannon wang puglisi weber bers 2004 model updated</td>
<td>0</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>stewart zhang model 2008</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>tentusscher model 2004 endo</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>tentusscher model 2004 epi</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>tentusscher model 2004 M</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>tentusscher model 2006 endo</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>tentusscher model 2006 epi</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>tentusscher model 2006 M</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Tomek model 13 endo</td>
<td>0</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Tomek model 13 epi</td>
<td>0</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Trovato2020</td>
<td>0</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>viswanathan model 1999 epi</td>
<td>2</td>
<td>7</td>
<td>9</td>
</tr>
<tr>
<td>wang model 2008</td>
<td>0</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>winslow model 1999</td>
<td>1</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>zhang SAN model 2000 0D capable</td>
<td>0</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>106</strong></td>
<td><strong>263</strong></td>
<td><strong>369</strong></td>
</tr>
</tbody>
</table>
Operation
There are two main ways to use chaste_codegen. It can be used integrated as part of the Chaste build process, from the 2021.1 release onward. It can also be used as a standalone command line tool or Python library.

The minimal system requirements for chaste_codegen as a standalone command-line tool are:

- python3 (3.5+), tested on Windows 10, Ubuntu Linux 18.04 and MacOS.
- python3 pip (usually bundled with a python installation).
- python3 venv (or other python virtual environment) is recommended, to ensure the right versions of dependencies are available. However chaste_codegen will still work without a virtual environment. Python3_venv is required for use within Chaste and is usually bundled with a python installation.

For use integrated into the Chaste build process, follow the regular guides on installing and building Chaste. As part of the Chaste installation process chaste_codegen will be installed in a virtual environment and all CellML files in the source will be converted using the appropriate settings.

To install as a stand-alone command line tool, run pip install chaste_codegen. You may want to create a python virtual environment (venv) first. The basic usage is: chaste_codegen cellml_file where cellml_file is the CellML file to be converted. To get a detailed overview of the various options run the command chaste_codegen -h.

Use cases
In this section we will briefly show a number of common conversions in action. For a more detailed guide see the ‘Code generation from CellML’ section in the Chaste guides. Due to the size of both the import and generated code, however, we will mention only key snippets and refer to the full files available in the assets branch of the chaste_codegen GitHub page.

In the following examples a CellML 1.0 file for the Hodgkin-Huxley model will be converted into code for a number of different ODE solvers. The CellML file consists of a number of components (membrane, sodium channel, sodium channel m gate, sodium channel h gate, potassium channel, potassium channel n gate and leakage current) and within these components variables are defined. The model also includes and links between components and unit definitions.

CellML files may include metadata through the use of RDF, the Resource Description Framework. chaste_codegen makes use of these annotations when generating C++ source code for Chaste, some of which are optional. In particular, chaste_codegen needs to know which variables represent voltage and stimulus_current, in order to link the models into the mono/bi-domain equations.

Below an example of a variable called V tagged as voltage is shown.

<variable name="V" units =" millivolt “ initial_value ="−80” public_interface="out”

    cmeta : id="membrane_voltage”>

    <rdf : RDF xmlns : rdf ="http://www.w3.org/1999/02/22-rdf-syntax-ns#”

        xmlns : bqbiol="http://biomodels.net/biology-qualifiers/”>

    <rdf : Description rdf : about="#membrane_voltage”>

        <bqbiol : is

            rdf : resource="https://chaste.comlab.ox.ac.uk/cellml/ns/oxford-metadata#membrane_voltage”/>

    </rdf : Description>

</rdf : RDF>

</variable>
‘Plain’ C++ code
The following command generates what we call ‘Plain’ C++ code. This code is used for solvers that only require the right hand side of the ODE, such as Forward Euler and Runge-Kutta solvers. This kind of code generation does not require any specific flags.

    chaste_codegen ModelName.cellml

This generates ModelName.cpp and ModelName.hpp. The code generates a class called CellModelNameFromCellML which inherits from AbstractCardiacCell. It contains the following key methods:
- a constructor and destructor
- UseCellMLDefaultStimulus calculates a stimulus based on parameters (amplitude, duration, start- and end-time) set in the model. These are identified using metadata as described above.
- GetIIonic calculates total ionic current at the present time (for use in tissue simulations).
- EvaluateYDerivatives calculates the derivatives of the the state values when provided with their current values, defining the ODEs of the model.
- ComputeDerivedQuantities gives a way to calculate the value of quantities that are derived directly from state variables, for example currents such as “the fast sodium current”.
- OdeSystemInformation::Initialise gives a way to retrieve information about the model such as name, free variable (usually time), state variable, modifiable parameters and named derived quantities.

CVODE
The following command generates code for the CVODE solver which has its own vector class.

    chaste_codegen –cvode –use-analytic-jacobian ModelName.cellml

This generates .cpp and .hpp files with the same name as before. The generated class now inherits from AbstractCvodeCell, containing the same methods but with SUNDIALS’ vector class for use directly with CVODE. It also has a method EvaluateAnalyticJacobian in which the analytic Jacobian is defined, to be used by CVODE.

Backward Euler
The following command generates Backward Euler code.

    chaste_codegen –backward-euler ModelName.cellml

This generates .cpp and .hpp files with the same name as before. The generated class inherits from AbstractBackwardEulerCardiacCell. The class does not have EvaluateYDerivatives, but instead it has:
- UpdateTransmembranePotential where the voltage is updated based on the ODE for voltage.
- ComputeOneStepExceptVoltage where the other state variables are updated. The variables are described by linear equations use a backward Euler fashion e.g. $n = (n + (\alpha * mDt)) / (1.0 - ((-\alpha - \beta) * mDt))$

Rush-Larsen
The following command generates code in which some state variables are updated using analytic solutions using the Rush-Larsen scheme\(^4\).

    chaste_codegen –rush-larsen ModelName.cellml

This generates .cpp and .hpp files with the same name as before. The generated class now inherits from AbstractRushLarsenCardiacCell. The concrete class does not contain a method EvaluateYDerivatives, but instead it has methods:
- EvaluateEquations where the voltage and non-linear state-variables are updated using the Forward Euler method. For the linear equation state variables variables are stored to capture the analytic solution.
• ComputeOneStepExceptVoltage where the linear state-variables are updated using their analytic solutions.

Summary
This paper has introduced the software tool chaste_codegen, designed to translate electrophysiology models from the CellML XML format into C++ code for use by the Chaste simulation package. We have shown how the tool can be used, highlighted the main different types of code it can generate for different solvers and shown a number of advanced features that chaste_codegen implements over a previous tool called PyCML. The most notable are the ability to generate analytic Jacobians and to evaluate and fix singularities in equations. We have shown that the singularity analysis works as expected with an analysis of a large number of popular models, by identifying all previously-identified singularities and finding over three times as many in total.

Contributing to development
chaste_codegen and the cellmlmanip libraries are open-source and publicly available and we welcome contributions: from questions about the current functionality to suggestions for improvements and source code contributions. It should also be relatively simple to extend the use of templates to generate code for other simulation packages in C++, Python, or other languages. Contributions are made in the first instance using GitHub issues. In order to contribute, users create a new issue in either GitHub repository or comment on an existing issue. Contributions in the form of source code can be made by issuing a pull request on either repository (ideally a new GitHub issue should be created, which can then be linked to the pull-request). The pull request will trigger an automated test suite and a number of other checks for things such as code formatting and test coverage.

Software availability
Software available from: https://pypi.org/project/chaste-codegen/
Source code available from: https://github.com/ModellingWebLab/chaste-codege
Archived source code as at time of publication: https://doi.org/10.5281/zenodo.5527756
BSD 3-Clause License

Author contributions
MH led the development of chaste_codegen, contributed to cellmlmanip development and co-wrote this paper. MC contributed to cellmlmanip development and provided code reviews and suggestions for chaste_codegen development. AUT and SMK contributed to cellmlmanip development. RHJ identified the singularities issue and provided analysis around how this could be solved. JC led the development of cellmlmanip and provided code reviews and suggestions to chaste_codegen. GRM worked on the singularity fixing algorithm, supervised the work and co-wrote this paper.

References
10. On the application of partial evaluation to the optimisation of cardiac electrophysiological simulations, PEPM ’06, New York, NY USA, Association for Computing Machinery. 2006. Publisher Full Text

11. Jinja Documentation (3.0.x). Reference Source


This article describes the chaste_codegen tool developed to convert a CellML model to C++ code. chaste_codegen is a Python3 implementation of the previously used Python 2 based PyCML tool. The authors describe in detail two new additional features implemented in chaste_codegen (1) Jacobian matrix generation (2) the ability to automatically detect and fix singularities in the equation with linear approximation. And authors clearly demonstrate the validity of their solution to detect and fix singularities with several examples. chaste_codegen along with the new features is a quite useful tool.

The article is well written, however, it could benefit the readers further if the following minor suggestions are considered.

1. It would be useful to provide a short summary of the major functionalities of chaste_codegen in the introduction. Although one would expect them to be similar to PyCML, it would useful to briefly describe them for the benefit of the new users.

2. Also, it would be useful to discuss the performance of chaste_codegen when compared against PyCML. For example, it will be useful to briefly mention how well does the computational time compares in general. Were the functionalities from PyCML comparable to those in chaste_codegen excluding the new features (automated Jacobians and singularity fixing)?

3. The article describes in detail the rationale and solution to fix singularities in the equations (where the value of variable such as Voltage tends to become 0/0). The chaste_codegen has been already integrated into chaste. Authors could discuss how this tool or approach could be used beyond chaste and provide a few suggestions.

4. I tried to run and test the standalone chaste_codegen python tool. Although I tried only for
an hour or so, I couldn't get it working. I managed to get it installed in a virtual environment as suggested, but couldn't export the cpp file from the CellML model. It will be useful to provide some sort of “test_codegen” command to run sample tests and report errors to help to troubleshoot.

5. A concern on the tool is that it is based on cellmlmanip which supports only CellML 1.0 and the author described there are no plans to support CellML 2.0. Would this mean the chaste_codegen will be outdated when CellML 2.0 is adapted widely? It would be useful to discuss this in the article, as a part of the limitation if so. Also, would chaste_codegen throw a clear warning if CellML 2.0 is used?

6. I would suggest authors, to thoroughly check all the equations (1 - 8) and ensure all components are properly defined. For example, \( V_0 \) is defined as the voltage at which we hit the singularity, but there is no definition for \( B \) in equation 1. Although, it might be trivial to those working in electrophysiology modeling and references are provided, it will be useful to describe functions such as \( C(V) \), \( A(V) \) in the text to help general readers.

7. Validity of the singularity detection by chaste_codegen was shown through the comparison of the number of singularities detected and the hardcoded fixes in previous CellML models (Table 1). Similarly, the authors described that the validity of the generated Jacobians was assessed by comparing numerical results of runs of a number of different models without providing further details. If not in detail, at least a few sentences on how many models were used and so on would be beneficial.

8. The authors have discussed the advantages of the tool and the newly developed features, it would be useful to also discuss limitations if any.

9. It will be also useful if the authors discuss any comparable existing approaches and implementation for automatic singularities detection and fixing.

**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?**
Yes

**Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?**
Yes

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

**Reviewer Expertise:** I am a mathematical modeler and one of the editorial board members of the community standard, SBML. I am familiar with XML-based formats to describe models and the libraries developed to handle them. I also lead the BioModels, a repository of mathematical models of biological systems.

**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.**

Reviewer Report 01 March 2022

https://doi.org/10.21956/wellcomeopenres.19013.r48112

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David Nickerson
Auckland Bioengineering Institute, University of Auckland, Auckland, New Zealand

This article very nicely lays out the rationale for the chaste_codegen tool, particularly in the evolution from the previous PyCML software used in Chaste. While clearly the emphasis is on producing code suitable for use in Chaste, the authors do suggest how the tool could be used independently. Some introduction or pointers as to how readers might be able to edit/alter the Jinja2 templates to produce non-Chaste code would be helpful.

While the performance and correctness of the analytic Jacobians generated by chaste_codegen using SymPy is very briefly mentioned, seeing some of those checks would I think be interesting to readers.

A large portion of this manuscript is describing the use of cellmlmanip to detect and fix singularities. While this is a valuable contribution the authors preface this by suggesting the replacement of cellmlmanip with the libCellML when looking at handling more recent versions of the CellML specification. Perhaps a bit more clarity regarding the future use of cellmlmanip and chaste_codegen regarding the singularity detection and fixing would be helpful.

I needed the assistance of google to learn what expm1 is, so perhaps other readers might appreciate a pointer.

Step 2 in the algorithm to identify singularities in equations seems a bit too general. While historically most cardiac electrophysiological models have been encoded in CellML in a similar manner for which this assumption holds true, it is not likely to always be true. Particularly if readers were looking to build on chaste_codegen to work with models from other domains.
The discussion regarding the removal of hard-coded singularity fixes from the original models available in the main CellML repository is great to see. And something that the authors should be encouraged to discuss with the broader CellML community.

(Minor comment: `chaste_codegen -h` suggests that the cellml_file could be a URI, but pointing to, for example, https://raw.githubusercontent.com/Chaste/cellml/master/cellml/hodgkin_huxley_squid_axon_model_1952_modified.cellml fails with a file not found error.)

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?  
Yes

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?  
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Standards development, cardiac electrophysiology modelling, software development.

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.

Reviewer Report 08 February 2022

https://doi.org/10.21956/wellcomeopenres.19013.r48330

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Axel Loewe
Institute of Biomedical Engineering (IBT), Karlsruhe Institute of Technology (KIT), Karlsruhe, Germany

Hendrix et al. present *chaste_codegen*, a Python 3 reimplementation of the *PyCML* software package which adds new functionality, mostly through integration of the *cellmlmanip* library. Flexible code generation for CellML models is a relevant need in the cardiac modeling community and the tool is well presented in general.

**A few remarks:**
- It seems that most of the new functionality (fixes for singularities) comes directly through *cellmlmanip*: https://github.com/ModellingWebLab/cellmlmanip/. As such, it might be helpful to explain the relation between cellmlmanip and chaste_codegen early in the manuscript. In the current form it seems a bit odd that half of the 12 text pages of the manuscript deal not with chaste_codegen but with one of its dependencies.
- It might be helpful for the reader to point out the main differences between chaste_codegen and other tools providing similar functionality like for example Myokit (partly from the same authors)
- In general, the tool could be valuable for developers (and potentially also users) of other cardiac electrophysiology simulators. To increase the value beyond the Chaste user community, it would be helpful to outline how the Jinja2 templates would need to be adapted or extended. For example Introduction in the (second paragraph, first sentence) and Methods (fourth paragraph, second-last sentence) sections.
- Why are specific operating systems mentioned? I would have assumed that chaste_codegen runs on all platforms that provide a Python interpreter.

**Typos and other trivia:**
- Title: consider adding the dash in "chaste_codegen"
- Abstract: consider "new cellmlmanip Python library" -> "cellmlmanip Python library"
- Introduction:
  - consider "hundreds of models" -> "hundreds of mathematical models"
  - The first sentence of the second paragraph is a bit confusing. Why is chaste_codegen needed if CellML models can be imported in Chaste already? Or is this only the case with PyCML/chaste_codegen?
- Methods:
  - Please double check appropriate references to software mentioned in the manuscript (cellmlmanip, SymPy etc.)
  - CellMI -> CellML
  - Cellmlmanip is typed with capital c and in normal font in the second paragraph
- RDF should be spelled out when it is first mentioned
- Second paragraph, last sentence: is an "and" missing here?
- Often, the Goldman–Hodgkin–Katz voltage equation is referred to as "the GHK equation". Here, you refer to the GHK flux equation. Mentioning this explicitly can help to avoid confusion.
- p.5: some readers might not be familiar with "expm1", please provide some more background
- p.6, point 5: hyphenation in nondimensional
- Table 1: model naming (w, w/o "model" etc.) and capitalization appear arbitrary
- Operation:
  - "can be used integrated" -> "can be used"?
- p.12: "pull-request" -> "pull request" for the sake of consistency

**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?**
Yes

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Yes

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

**Reviewer Expertise:** Computational cardiac modeling

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.