BIODEG

Biodegradation and Corrosion Simulation using Finite Element

User Manual

Version 0.8

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KU LEUVEN

BioDeg website

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1 Introduction

BIODEG is an open source software written in FreeFEM (a domain-specific language for finite element programming), C++, and Python for modeling the degradation of metallic biomaterials and simulating the biodegradation behavior of medical devices, implants and scaffolds in corrosion experiments. It can handle any geometry of desire and supports parallel computing to simulate large scale models.

1.1 Authors

BIODEG is developed by the Biomechanics Research group at KU Leuven and University of Liege. The code is currently maintained by its principal developer, who manage the development of the mathematical models and the core functionalities.

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1.3 Referencing BioDeg

Please refer to BIODEG repository, section "Publications and referencing" to properly cite the use of BIODEG in your scientific work.

2 Useful background information

You may refer to the following articles for a background of the methods and algorithms implemented in BioDeg.

In addition, below are some useful references on finite element method and some online resources that provide a background of finite elements and their application to the solution of partial differential equations.

3 Installation

Installing BIODEG is a straightforward procedure. You need to install a couple of prerequisites and download and run BIODEG; that's all you need to do. But for advanced users, it might be necessary or more interesting to build everything from scratch to have more control on customizing features and improving performance. As a result, we have provided 2 sets of instructions, one for easy installation using the compiled binaries and one for building things from the source codes. The installation instructions are provided for Linux and Windows operating systems, but the procedure should be very similar for macOS.

It is possible to use BIODEG without the user interface (UI) if this scenario is required by the user (like for running it on a super-computer). The core of BIODEG is written in FreeFEM, so in this case, all you need to do is to installing/building FreeFEM and the required libraries, and then, cloning the BIODEG core repository and running the code according the provided instruction in the README file of the repository.

The BIODEG UI contains all the bundles for pre-processing and post-processing simulation input/results. These features are being hosted on their own repositories (), but with obtaining the user interface, you can have them all together. If you choose to use BIODEG without the user interface and still want to use the provided script for pre/post-processing, you may need to obtain them separately.

For building the source codes, we assume standard tools and libraries like CMake, compilers (for C, C++ and Fortran), and MPI libraries are pre-installed on your machine. If you are going to build BioDeg and required dependencies on a super-computer or cluster, you should notice that most high-performance computers would have the latest version of these compilers and libraries in the default environment.

3.1 Easy installation

The simplest way to install and run BIODEG is via the pre-built binaries you can download from GitHub. The same principle applies to prerequisites, which is in this case FreeFEM only. So, following these steps will install BIODEG on your machine:

- 1. Download FreeFEM installer for the platform you use and install it. You can find the .exe installer for Windows and the .deb installer for Linux (Ubuntu) in the Release page of FreeFEM repository. You will find these files under the Assets section of the latest (or any other version). Execute the download file and follow the installation procedure appearing on your screen.
- 2. Download BIODEG tarballs for your preferred platform (Windows or Linux) from the Release page of BIODEG repository. This is indeed the BIODEG UI bundle that contains the BIODEG core, the user

interface, the pre-processor, and the post-processor.

3. Extract the downloaded tarball (zip) file and execute runBioDeg.cmd in Windows or runBioDeg.sh in Linux. By doing this you see BIODEG interface showing up on the screen.

3.2 Advanced installation for improved performance/flexibility

Building BIODEG and required libraries from source code will increase the performance since the program and the libraries will get optimized for the platform in which they are going to run. Moreover, this enables users to customize the software in the way they want. Additionally, this is an inevitable aspect if you are going to use BIODEG for development purposes or you want to contribute to it.

3.2.1 Compiling and installing external libraries

3.2.1.1 PETSc and Qt

BIODEG uses PETSc for parallel computing. You may choose to build a customized version of PETSc or use the version that comes with FreeFEM. The version that is bundled with FreeFEM has the following build configuration:

```
--with-debugging=0 COPTFLAGS="-03 -mtune=native" CXXOPTFLAGS="-03 -mtune=native"
FOPTFLAGS="-03 -mtune=native" --with-cxx-dialect=C++11 --with-ssl=0 --with-x=0
--with-fortran-bindings=0 --with-cc=/usr/ --with-scalar-type=complex
--with-blaslapack-include= --with-blaslapack-lib="-llapack -lblas"
--with-scalapack --with-metis --with-ptscotch --with-suitesparse --with-suitesparse-lib=
"-Wl, -lumfpack -lklu -lcholmod -lbtf -lccolamd -lcolamd -lcamd -lamd -lsuitesparseconfig"
--with-mumps --with-parmetis --with-tetgen --download-slepc --download-hpddm PETSC_ARCH=fc
```

You may need to build your own version if this configuration is not suitable for you. You can find the instruction for building custom version of PETSc here.

BIODEG UI is developed using Qt so it should be installed on your system if you want to compile BIODEG UI. You can find the installation instruction for various platforms here.

3.2.1.2 FreeFEM

The full build documentation of FreeFEM is available here, but the following steps is what you need to do to build it on any platform. By default, FreeFEM downloads and builds PETSc during the build process.

- 1. Install required prerequisites
 - \$ sudo apt-get install cpp freeglut3-dev g++ gcc gfortran m4 make patch pkg-config wget python unzip liblapack-dev libhdf5-dev libgsl-dev autoconf automake autotools-dev bison flex gdb git cmake
 - \$ sudo apt-get install mpich
- 2. Make a new directory for FreeFEM and navigate to it

- \$ cd
- \$ mkdir FreeFEM
- \$ cd FreeFEM/
- 3. Clone the source code repository and navigate to the downloaded directory

```
$ git clone https://github.com/FreeFem/FreeFem-sources.git
$ cd FreeFem-sources/
```

- 4. Generate the configure scripts
 - \$ autoreconf -i
- 5. Run the configure script to specify build options, including the location to install the program

```
$ ./configure --enable-download --enable-optim
--prefix=/home/<your_profile>/FreeFEM/freefem-install
```

- 6. Download the source code of 3rd-party libraries
 - \$./3rdparty/getall -a
- 7. Build PETSc and all the 3rd-party libraries
 - \$ cd 3rdparty/ff-petsc/
 \$ make petsc-slepc
- 8. Navigate back and reconfigure the build
 - \$ cd -
 - \$./reconfigure
- 9. Build the source code of FreeFEM using 4 parallel processes (or nay other number you like
 - make -j4
- 10. Check the build by running some examples
 - \$ make -j2 check
- 11. Install the built binaries to the specified directory
 - \$ make install
- 12. Navigate to the installation location and run FreeFEM

```
$ cd ../freefem-install/
$ cd bin/
$ ./FreeFem++
```

13. Navigate to the home directory and add FreeFEM to the PATH variable in the .bashrc' file

```
$ cd
$ nano .bashrc
```

and add export PATH= $PATH:/home/<your_profile>/FreeFEM/freefem-install/bin to the end of the file and save (press <math>Ctrl+X$ and then Y).

After doing this, you should be abale to run FreeFEM. Start a new terminal and run FreeFem++ and FreeFem++-mpi. See ing no error in the output means that you have successfully installed it.

3.2.2 Building and installing BioDeg

Since BIODEG UI is developed using Qt, compiling the source files is quite straightforward. Upon installing Qt on your machine, clone the BIODEG UI repository and follow one of the following scenarios to build it.

3.2.2.1 Build BioDeg UI using Qt Creator IDE

This is the simplest technique to build the program, and it has a similar procedure for all the supported platforms. Qt Creator is the default IDE for Qt development, so it is automatically installed along with Qt. Simply open the Qt project file (CMakeLists.txt) in Qt Creator (by executing qtcreator CMakeLists.txt or selecting File->Open Project from the IDE) and build the project (by pressing Shift+B).

3.2.2.2 Build BioDeg UI using Qt tools

Building the source files using CMake is also quite simple. Navigate to the source files directory (the cloned repository) and run the following commands (this assumes that you have already added Qt bin directory to the 'PATH' variable so that the CMake script can find Qt libraries and binaries):

```
$ mkdir build
$ cd build
$ cmake ..
```

In Windows, you may need to call the correct build system installed along with Qt. For example, by assuming that you have installed the MinGW integration for Qt, the make command should be written as mingw32-make. Moreover, in this case, you need to call CMake with a suitable generator, so the cmake command should be replaced by something like cmake -G "MinGW Makefiles" (don't forget to insert the double dots).

After doing this, you can find the BioDeg UI executable in the source directory and run it by executing ./BioDeg in Linux or .\BioDeg.exe in Windows.

4 Running BioDeg

After installing/compiling BIODEG as described in Section 3, we are ready to run it. There are 2 ways to run BIODEG simulations: 1) using the UI to configure and execute BIODEG, or 2) running BIODEG directly from command line and provide simulation parameters via command line arguments. Moreover, in a hybrid approach, the UI can be used to configure and generate the command for method #2, which can be useful in which you want to configure the simulation only and run it later in another environment like on a super-computer or cluster.

The UI can be run simply by double clicking on the BioDeg-UI.exe in Windows or by executing ./BioDeg-UI in Linux. For running BioDeg directly, one need to execute the following command:

\$ mpirun -n N FreeFem++-mpi BioDeg-core/src/main.edp <command line args>

in which N defines the number of MPI processes to be used. The full list of command line arguments can be found in Section Index.

In order to clarify and demonstrate the procedure of performing simulations using BioDeg, 2 step-by-step examples are provided in Section 4.1, showing how to configure and run simulations with and without the UI. The mesh files needed to run these examples can be found in the demo directory.

4.1 Configuring the simulation

4.1.1 Example 1 - degradation of a simple screw

Let us consider the first example given in the demo directory, where we simulate the biodegradation of a small screw. The size of the screw was chosen to be very small intentionally to make the simulation shorter such that the user can see the effect of the degradation much faster. The input mesh file is named screw.mesh. For this example, we use the BioDeg-UI interface to perform the simulation.

Let's conduct the first simulation with most of the parameters left with their default values. Run the user interface, and mark Geometry & Mesh Import external mesh. Then click the browse button in front of the "File" box (Geometry & Mesh Import mesh File), navigate to the demo directory, and select the screw.mesh file. The path should be inserted in the "File" box. Selecting appropriate label numbers of the external mesh is very crucial in BIODEG, so you should always check the labels before importing the mesh. One of the best tools to do this is GMSH, in which you can view the labels of mesh files by opening the mesh and navigating to Tools Visibility. Doing this on the screw.mesh shows that the label number of the scaffold is 2 while the medium has a label number of 1 (Fig. 1). So, we need to switch the default labels by selecting 2 for the scaffold (Geometry & Mesh Import mesh Scaffold label) and 1 for the medium (Geometry & Mesh Import mesh Medium label).

We need to enable the VTK output if we want to see the graphical output of the simulations. Navigate to Output Output options Write VTK output and mark it. You should also specify the output directory by clicking on the browse button in front of the "Output directory" (Output Output options) Output directory) and select a directory of desire.

That was all we needed to do to setup a simulation in BIODEG, and you can start the simulation by pressing the Run simulation button located in the middle of the main window. There are lot more parameters we can deal with (see Section Index), but the input and output are the most essential ones.

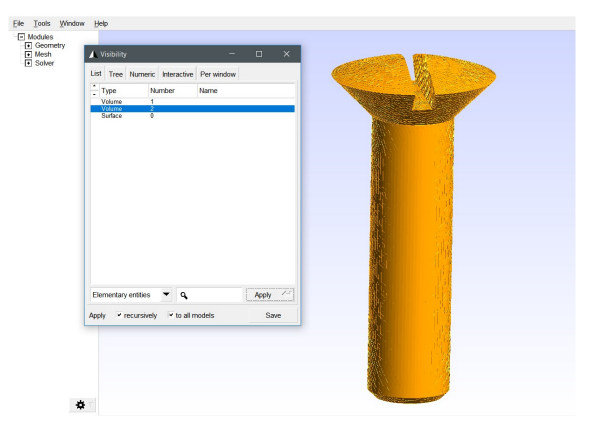


Figure 1: Using GMSH to read the labels of the mesh

Although you can start the simulation of this example right now, there is a couple of more things you may want to change:

- In BioDeg, simulations are by default carried out in parallel using domain decomposition, meaning that the simulation is distributed among available computing nodes. If you are running BioDeg on your local machine, you may need to adjust the parallel computing settings. Navigate to Solver Parallel computing Enable parallel computing and disable it if you don't want to parallelize the simulation. If you want to continue with parallelization enabled (default behavior), you may need to adjust the number of parallel processes to match the number of free CPU cores you have on your machine. You can change it in Solver Parallel computing CPU/MPI cores.
- The default degradation rate is quite fast, so you may want to decrease it by reducing the diffusion coefficient of the metallic ions (please refer to "Theory Guide" if you want to know more about diffusion controls the rate of degradation). The default diffusion rate is the value we have estimated for saline solutions, which leads to a high rate of corrosion. You can apply this it by changing the value in Material & BCs Reaction-diffusion properties Metal ion diffusion coefficient and reduce it to something like 0.0005, which is its order when it comes to buffered solutions and simulated body fluids.
- The results write interval, implying how frequently you want to store the results, affects the resource consumption (which is storage in this case) and the quality of the graphical postprocessing. So, you

should configure this carefully to keep the balance of the quality and resource consumption. The default save interval is 0.25 hours of simulation time, but you can change it in Output Output options Save results every ... hours. For this simulation, since the screw geometry is small and degrades very fast, you can reduce this to 0.1 to be able to see the degradation steps better.

• Final simulation time does not affect the simulation progress, but it is always a good practice to adjust it, enabling us to track the progress of the simulation better and avoid wasting resources (both computing power and storage). The default simulation time is 21 hours, but you can reduce it in Solver Time control Final simulation time (hour). For this simulation, you can reduce it to 2.

After running the simulation (by pressing the Run simulation) button), you can view the progress of the simulation on the UI, showing you how many steps have been taken and how much material degradation has happened. The UI also shows you the details info of the size of the problem, including the degrees of freedom (DOF) of each equation and the number of elements, as well as the number of DOFs for each sub-domain after mesh partitioning (for parallel computing).

Running this simulation leads to the results demonstrated in Fig 2, showing how the screw degrades. For more information on how to postprocess the results, please refer to the postporcessing section.

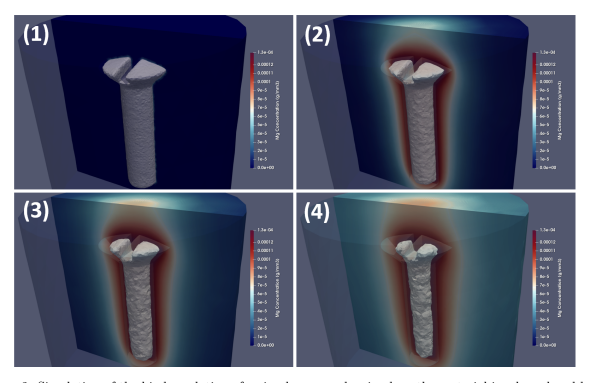


Figure 2: Simulation of the biodegradation of a simple screw, showing how the material is released and how the implant degrades.

4.1.2 Example 2

5 Future extensions to BioDeg

The future versions of BioDeg will focus on implementing the following methodologies/features.

- Extending the core models to capture more complex chemistry of biodegradation by considering more reactions occurring in buffered solutions.
- Adding support for more base materials like Fe and Zn.
- Considering the effect of alloying elements and complex compositions.
- Adding more post-processing features to BioDeg UI.
- Adding basic visualization to BIODEG UI using ParaView Glance.
- Improving the performance of fluid flow solver by employing a gradient-based solver.
- Considering GPU support by enabling GPU computing in recent versions of PETSc.

6 Finding answers to more questions

If you have questions that go beyond this manual, there are a number of resources:

- For questions/suggestions about BIODEG installation, bugs, or similar stuff please use the BIODEG issue tracker.
- BIODEG is primarily based on the FreeFEM. If you have particular questions about FreeFEM, contact the community at https://community.freefem.org/.
- If you have specific questions about BIODEG that are not suitable for public and archived mailing lists, you can contact the primary developer and mentor:
 - Mojtaba Barzegari: mojtaba.barzegari@kuleuven.be.
 - Liesbet Geris: liesbet.geris@kuleuven.be (Mentor).

A Run-time input parameters

The underlying description of the input parameters also includes a "Standard/Advanced" label, which signifies whether an input parameter is a standard one or an advanced level parameter. The default values of the "Advanced" parameters are good enough for almost all cases. However, in some cases user may need to use "Advanced" labeled parameters. For user convenience, all input parameters are also indexed at the end of this manual in Section Index.

A.1 Geometry and mesh parameters

• Parameter name: import_mesh

Default: true (1)

Description: [Standard] Boolean parameter specifying whether an external mesh file is imported or a container box as well as a cubic scaffold would be created on the fly for simulation.

Possible values: A boolean value (1 or 0)

• Parameter name: mesh_file

Default: Should be provided

Description: [Standard] Path to the input mesh file (in MEDIT .mesh format), which can be either absolute or relative. Is relevant only if parameter import_mesh is set to TRUE.

Possible values: Any string value

• Parameter name: label_scaffold

Default: 1

Description: [Standard] The label of the (volume) region supposed to be scaffold in the input mesh (can be viewed in programs like GMSH before importing into BioDeg).

Possible values: Any positive integer value

• Parameter name: label_medium

Default: 2

Description: [Standard] The label of the (volume) region supposed to be the medium (electrolyte) in the input mesh.

Possible values: Any positive integer value

• Parameter name: label_wall

Default: 3

Description: [Advanced] The label of the surface in the input mesh to be assigned as wall (no slip boundary condition) in the fluid flow simulations.

Possible values: Any positive integer value

• Parameter name: label inlet

Default: 4

Description: [Advanced] The label of the surface in the input mesh to be assigned as flow inlet (constant velocity boundary condition) in the fluid flow simulations.

Possible values: Any positive integer value

• Parameter name: label_outlet

Default: 5

Description: [Advanced] The label of the surface in the input mesh to be assigned as flow outlet (zero pressure boundary condition) in the fluid flow simulations.

Possible values: Any positive integer value

• Parameter name: box_length

Default: 20.0

Description: [Standard] In case of import_mesh being FALSE, specifies the length of the container box (for the electrolyte) in mm.

Possible values: Any positive floating point number

• Parameter name: cube_size_x

Default: 13.0

Description: [Standard] In case of import_mesh being FALSE, specifies the length of the scaffold cuboid along the x axis in mm.

Possible values: Any positive floating point number

• Parameter name: cube_size_y

Default: 13.0

Description: [Standard] In case of import_mesh being FALSE, specifies the length of the scaffold cuboid along the y axis in mm.

Possible values: Any positive floating point number

• Parameter name: cube_size_z

Default: 4.0

Description: [Standard] In case of import_mesh being FALSE, specifies the length of the scaffold cuboid along the z axis in mm.

Possible values: Any positive floating point number

• Parameter name: mesh_size

Default: 32

Description: [Standard] Number of elements on each edge of the container box, so a higher number means a finer mesh. The mesh size of the cuboid will be adjusted accordingly or can be adaptively refined by setting parameter refine_initial_mesh to TRUE.

Possible values: Any positive integer number

• Parameter name: refine_initial_mesh

Default: false (0)

Description: [Advanced] A boolean parameter specifying if the mesh (no matter if imported or generated) should be adaptively refined on the metal-medium interface (corrosion surface). This affects the beginning of the simulation only (on the initial mesh).

Possible values: A boolean value (1 or 0)

• Parameter name: mshmet error

Default: 0.01

Description: [Advanced] Since the open source tool mshmet is used for creating a metric for refining the mesh on the level set signed distance function, a tolerance should be specified for it. A lower value results to a finer mesh.

Possible values: Any floating point number

• Parameter name: mesh_size_min

Default: 0.04

Description: [Advanced] Specifies the smallest element size to be passed to the tetgen mesh generator for refining the initial mesh.

Possible values: Any floating point number

• Parameter name: mesh_size_max

Default: 0.8

Description: [Advanced] Specifies the largest element size to be passed to the tetgen mesh generator for refining the initial mesh.

Possible values: Any floating point number

A.2 Materials and boundary conditions parameters

• Parameter name: material_density

Default: 1.735e-3

Description: [Standard] The density of the metallic material. The default value is the density of magnesium.

Possible values: Any floating point number

• Parameter name: film_density

Default: 2.3446e-3

Description: [Standard] The density of the protective film that forms on the corrosion surface. The default value is the density of magnesium hydroxide.

Possible values: Any floating point number

• Parameter name: material_satur

Default: 0.134e-3

Description: [Advanced] The saturation concentration at which the metallic material (here, the ions) saturates through the medium. The default value is defined for magnesium ions.

Possible values: Any floating point number

• Parameter name: material_eps

Default: 0.55

Description: [Advanced] The porosity of the formed protective layer in the range [0, 1].

Possible values: Any floating point number between 0 and 1

• Parameter name: material_tau

Default: 1.0

Description: [Advanced] The tortuosity of the formed protective layer.

Possible values: Any floating point number

• Parameter name: d_mg

Default: 0.05

Description: [Standard]

Possible values: The diffusion coefficient of the metallic ions transport. This parameter is one of the most effective ones on the rate of degradation.

• Parameter name: d_cl

Default: 0.05

Description: [Standard] The diffusion coefficient of the chloride ions transport

Possible values: Any floating point number

• Parameter name: d_oh

Default: 25.2

Description: [Standard] The diffusion coefficient of the hydroxide ions transport

Possible values: Any floating point number

• Parameter name: k1

Default: 7.0

Description: [Standard] The reaction rate at which the chemical reaction of the protective film formation occurs.

Possible values: Any floating point number

• Parameter name: k2

Default: 1e15

Description: [Standard] The reaction rate at which the chemical reaction of the protective film disso-

lution occurs.

Possible values: Any floating point number

• Parameter name: fluid_nu

Default: 0.85

Description: [Advanced] The effective viscosity of the fluid used to simulate hydrodynamics condition.

Possible values: Any floating point number

• Parameter name: fluid_in_x

Default: 0.1

Description: [Advanced] The X component of the fluid velocity defined on inlet (see label_inlet) as the boundary condition for the fluid flow.

Possible values: Any floating point number

• Parameter name: fluid_in_y

Default: 0

Description: [Advanced] The Y component of the fluid velocity defined on inlet (see label_inlet) as the boundary condition for the fluid flow.

Possible values: Any floating point number

• Parameter name: fluid_in_z

Default: 0

Description: [Advanced] The Z component of the fluid velocity defined on inlet (see label_inlet) as the boundary condition for the fluid flow.

Possible values: Any floating point number

• Parameter name: initial_cl

Default: 5.175e-6

Description: [Standard] Initial concentration of chloride ions in the medium.

Possible values: Any floating point number

• Parameter name: initial_oh

Default: 1e-7

Description: [Standard] Initial concentration of hydroxide ions in the medium, used for computing pH.

The default value indicates a pH of 7.

Possible values: Any floating point number

A.3 Solver parameters

• Parameter name: solve_mg

Default: true (1)

Description: [Standard] Boolean parameter indicating whether the equation for material dissolution and ions release should be solved or not. This equation is the most essential equation and in most use-cases should be solved.

Possible values: Any boolean value (1 or 0)

• Parameter name: solve film

Default: true (1)

Description: [Standard] Boolean parameter indicating whether the equation for the protective film formation should be solved or not.

Possible values: Any boolean value (1 or 0)

• Parameter name: solve cl

Default: true (1)

Description: [Standard] Boolean parameter indicating whether the equation for the transport of chloride ions should be solved or not.

Possible values: Any boolean value (1 or 0)

• Parameter name: solve_oh

Default: true (1)

Description: [Standard] Boolean parameter indicating whether the equation for the transport of hydroxide ions should be solved or not. This equation is essential for calculating the pH changes, if desired.

Possible values: Any boolean value (1 or 0)

• Parameter name: solve_ls

Default: true (1)

Description: [Standard] Boolean parameter indicating whether the level set surface tracking equation should be solved or not. Surface tracking is essential for computing the mass loss.

Possible values: Any boolean value (1 or 0)

• Parameter name: solve_fluid

Default: false (0)

Description: [Standard] Boolean parameter indicating whether the fluid flow equation should be solved or not.

Possible values: Any boolean value (1 or 0)

• Parameter name: solve_full_ns

Default: true (1)

Description: [Advanced] Boolean parameter specifying which fluid flow equation to solve: the full transient Navier-Stokes equations or a steady-state Stokes equation. A true value (1) results in BioDeg solving the former equation.

Possible values: Any boolean value (1 or 0)

• Parameter name: write_fluid_output

Default: true (1)

Description: [Advanced] Boolean parameter specifying whether the fluid flow quantities (velocity field components and pressure) should be saved in the output VTK file or not. Requires write_vtk be TRUE.

Possible values: Any boolean value (1 or 0)

• Parameter name: solve_fluid_each

Default: 10

Description: [Advanced] Determines the number of time steps to skip before solving the specified fluid flow equation. For example, if the value is set to 10 (default value), the fluid equation gets solved in time steps 1, 11, 21,

Possible values: Any positive integer number

• Parameter name: time_step

Default: 0.025

Description: [Advanced] The time step value of the simulations.

Possible values: Any floating point number

• Parameter name: final_time

Default: 21.0

Description: [Standard] The final simulation time, meaning the duration of interest for the biodegradation model.

Possible values: Any floating point number

• Parameter name: do_redistance

Default: true (1)

Description: [Advanced] Indicates whether the redistancing of the level-est distance function should be done or not. Please refer to the theory guides to see how this affects the simulation.

Possible values: Any boolean value (1 or 0)

• Parameter name: redistance_time

Default: 1.0

Description: [Advanced] In case do_redistance being true, this parameter indicates the interval between each level set function re-initialization.

Possible values: Any floating point number

A.4 Output parameters

• Parameter name: text_output_file

Default: "output/result.txt"

Description: [Standard] Path to the text file in which text output, like the mass loss and evolved hydrogen production, are written over time. The path can be relative or absolute.

Possible values: Any string value referring to a valid path

• Parameter name: write_vtk

Default: true (1)

Description: [Standard] Indicates whether VTK output (in the VTU format) should be written or not. This is required for further post-processing of the results using ParaView.

Possible values: Any boolean value (1 or 0)

• Parameter name: vtk_output_name

Default: "output/output"

Description: [Standard] The naming scheme for the VTK output. This is mainly for the PVD file, and the final name of the rest of the files will be determined by the number of employed MPI computing nodes and the time step. The number of VTU files saved per time step equals to the number of employed MPI nodes.

Possible values: Any string value referring to a valid path

• Parameter name: save each

Default: 0.25

Description: [Standard] The interval of saving results, text and VTK (if selected), to disk.

Possible values: Any floating point number

• Parameter name: save_last_state

Default: true (1)

Description: [Standard] Indicates whether the last state of the system should be saved or not. The last state will be always saved on a global (non-partitioned) mesh, meaning that it will be a single VTU file, in contrast to normal save in which the number of written files equals to the number of computing nodes.

Possible values: Any boolean value (1 or 0)

• Parameter name: output_per_area

Default: false (0)

Description: [Advanced] Indicated whether the side hydrogen evolution results should be computed per unit area of exposed surface.

Possible values: Any boolean value (1 or 0)

• Parameter name: save_multiplier

Default: 1.0

Description: [Advanced] In case of symmetrical conditions, this parameter can be used to multiply the obtained quantitative results to have easier comparison with experimental results. A value of 1.0 indicates no multiplication.

Possible values: Any floating point number

• Parameter name: export_scaffold

Default: false (0)

Description: [Advanced] Indicates if the degrading material should be exported as a single entity for further analysis like in a structural mechanics simulation. It works based on Mmg level set meshing.

Possible values: Any boolean value (1 or 0)

• Parameter name: export_scaffold_each

Default: 1.0

Description: [Advanced] In case export_scaffold being true, this parameter determines the interval of saving the material mesh to the disk.

Possible values: Any floating point number

• Parameter name: export_scaffold_volume

Default: true (1)

Description: [Advanced] In case export_scaffold being true, this parameter indicates whether a volume mesh should be saved as output or not.

Possible values: Any boolean value (1 or 0)

• Parameter name: export_scaffold_surface

Default: true (1)

Description: [Advanced] In case export_scaffold being true, this parameter indicates whether a surface mesh should be saved as output or not.

Possible values: Any boolean value (1 or 0)

• Parameter name: save_initial_mesh

Default: false (0)

Description: [Advanced] Determines whether the initial computational mesh should be saved for further debugging or not. Can be used to investigate the mesh refinement if it is asked by setting refine_initial_mesh to true.

Possible values: Any boolean value (1 or 0)

• Parameter name: save_initial_partitioned_mesh

Default: false (0)

Description: [Advanced] Determines whether the partitioned computational mesh should be saved for further debugging or not. Can be used to debug and view the output of the mesh partitioning process before going for the actual simulation.

Possible values: Any boolean value (1 or 0)

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