

New functionalities of the 3.0 version of TFEL, MFront and MTest

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Abstract — MFront is a tool which allows easy implementation of arbitrary complex mechanical behaviours in an efficient way. Those implementations are portable between various finite element solvers and solvers based on FFT. MFront is part of the open-source TFEL project which also provide an useful point-wise solver called MTest.

The purpose of this paper is to present the 3.0 version of TFEL, MFront and MTest.

1 Overview of TFEL, MFront and MTest

The TFEL project is an open-source collaborative development of the French Alternative Energies and Atomic Energy Commission (CEA) and Électricité de France (EDF) in the framework of the PLEIADES platform (see (1)). TFEL provides mathematical libraries which are the basis of the MFront code generator and the MTest solver (see (2, 3)).

MFront translates a set of closely related domain specific languages into plain C++ on top of the TFEL library. Those languages are meant to be easy to use and learn by researchers and engineers and cover three kinds of material knowledge:

- Material properties (for instance Young modulus, thermal conductivity, etc.)
- Mechanical behaviours.
- Simple point-wise models, such as material swelling used in fuel performance codes.

Authors of MFront paid particular attention to the robustness, reliability and numerical efficiency of the generated code, in particular for mechanical behaviours: various benchmarks show that MFront implementations are competitive with native implementations available in the Cast3M (4), Code_Aster (5), Europlexus (6), Abaqus/Standard, Abaqus/Explicit (7, 8) and Cyrano3 (9) solvers.

Portability is also a very important issue: a behaviour written in MFront shall be usable in a solvers for which an interface exists. For example, for finite strain analysis, each solver has chosen a specific definition of the consistent tangent operator. MFront will convert the operator provided by the user, who is free to choose the most convenient operator to ease the implementation effort, into the one expected by the solver. The conversion paths currently available are depicted on Figure 1.

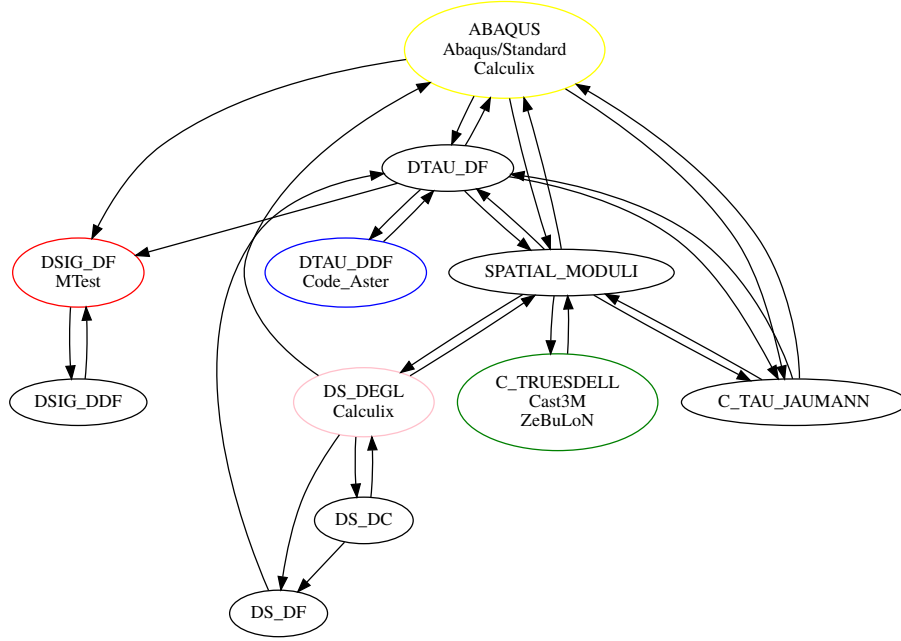


Figure 1: “Relation between tangent operators”

MTest is a tool created to perform unit-testings on the mechanical behaviours generated by **MFront**.

1.1 How to get **TFEL** and **MFront**

TFEL is distributed with the **Code_Aster** finite element solver (see (5)), the **Salome-Meca** platform (see also (5)) and the **Material Ageing Plateform** (MAP), which is available for all members of the Material Ageing Institute (MAI) (see (10, 11)).

Sources of **TFEL** can be downloaded from its website (see (12)). Binaries for the **Windows** operating system which are built upon versions of the **Cast3M** finite element solver are also available.

2 Highlights

From a user point of view, **TFEL 3.0** brings many game-changing features which are described hereafter:

- New mechanical behaviour interfaces to several finite element solvers (**Europlexus**, **Abaqus/Standard**, **Abaqus/Explicit**, **CalculiX**)
- The support of behaviours bricks, described below.
- The ability to simulate pipes using a rigorous finite strain framework in **MTest**
- The introduction of various accelerations algorithms used for solving the mechanical equilibrium when the consistent tangent operator is not available (see (13)).
- The development of a stable API (application programming interface) in **C++** and **python** which allow building external tools upon **MFront** and **MTest**. This point is illustrated by the development by EDF MMC in the **Material Ageing Plateform** of a new identification tool which is particularly interesting.

Many improvements for mechanical behaviours have also been made, which won't be detailed in this paper:

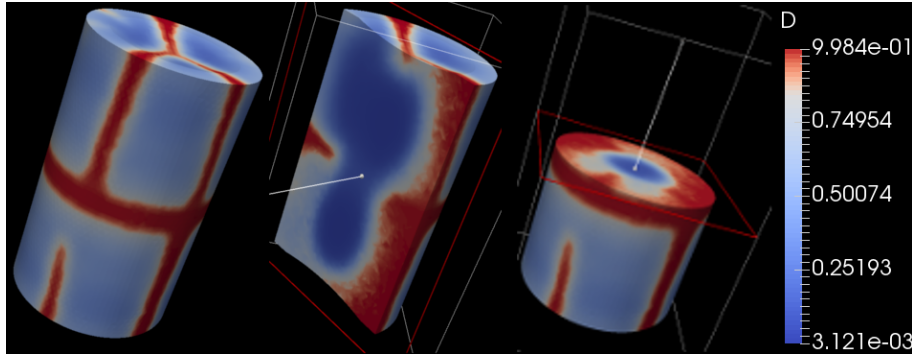


Figure 2: Phase field modelling of the nuclear fuel pellet fragmentation during the reactor start-up (see (15) for details)

- Definition and automatic computation of elastic material properties from external **MFront** files.
- Support for the automatic computation of the thermal expansion and swelling in the mechanical behaviour.
- Better support for orthotropic behaviours with the notion of orthotropic convention which affects various keywords (`@ComputeStiffnessTensor`, `@ComputeThermalExpansion`, `@Swelling`, `@AxialGrowth`, `@HillTensor` etc..).
- An initial support of non local mechanical behaviours. See for example Figure 2.
- Time steps management from the mechanical behaviour.
- Consistent tangent operator support in the **Cast3M** interface.
- Easier computations of isotropic functions of symmetric tensors and their derivatives.
- New material properties interfaces for various programming languages (`fortran03`, `java`, `GNU/Octave`).

2.1 New interfaces

2.1.1 Europlexus

Europlexus (EPX) is a simulation software dedicated to the analysis of fast transient phenomena involving structures and fluids in interaction. The program is co-owned by the French Alternative Energies and Atomic Energy Commission (CEA) and the Joint Research Centre of the European Commission (EC/JRC). Its development is carried out through a Consortium involving the co-owners and so-called major partners who are granted a complete access to the source code and development tools (see (6)).

The ability to call external libraries generated by **MFront** has been introduced in the current development version of **EPX**. When loading a **MFront** behaviour, **Europlexus** automatically retrieves various helpful information about the mechanical behaviour:

- Number and name of material properties
- Number, names and type of internal state variables
- Number and names of external variables
- Symmetry of the behaviour (isotropy, entropy)
- etc...

This information allows to check that the material declaration is consistent and reduces the inputs provided by the user to the bare minimal. A similar technique is used in **Code_Aster** and **MTest** when calling **MFront** generated behaviours.

2.1.1.1 Supported features

The following features are currently available:

- Support for finite strain behaviours
- Support for isotropic and orthotropic behaviours
- Automatic generation of an input file example to help the user in assigning a **MFront** behaviour to a material.

2.1.1.2 Finite strain strategies

Small strain behaviours can be embedded through finite strain strategies: currently, **MFront** provides (see (16, 17)):

- the **FiniteRotationSmallStrain** strategy: based on the Green-Lagrange strain measure.
- the **MieheApellLambrechtLogarithmicStrain** strategy based on the lagrangian Hencky strain measure.

In both cases, the strain measure is computed before calling the behaviour and the stress tensor resulting from the behaviour integration is interpreted as the dual of this strain measure. This stress tensor is then converted into the Cauchy stress tensor for computing the nodal forces.

2.1.1.3 Performances

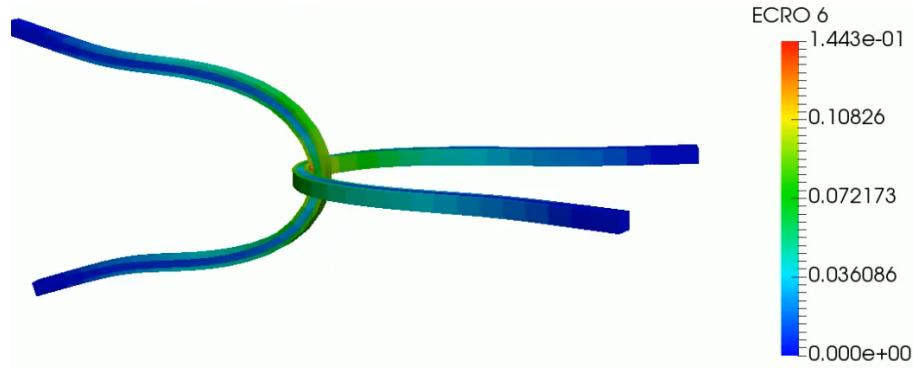


Figure 3: Impact of two bars

Figure 3 shows a simple test simulating the impact of two bars.

Table 1: Example of comparison of the computation times for the test depicted on Figure 3 for various behaviours describing isotropic plasticity with linear isotropic hardening.

Behaviour	Computational times
native	174s
MFront FiniteRotationSmallStrain	232s
MFront ImplicitSimoMieheElastoPlasticity	250s
MFront MieheApellLambrechtLogarithmicStrain	528s

This tests show that, for simple behaviours describing isotropic plasticity, **MFront** behaviours seems a bit less efficient than Fortran built-in implementations, as reported on Table 1. However, it shall be stated that the behaviours tested are not strictly equivalent (native behaviours are based on objective stress rates while **MFront** ones are hyperelastoplastic). **MFront** computational times can be improved by keeping some intermediate results for one step to the next to the expense of memory usage.

This interface is fully described here: <http://tfel.sourceforge.net/epx.html>

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2.1.2 Abaqus/Standard, Abaqus/Explicit and CalculiX

The Abaqus Unified FEA product suite provides:

- an implicit finite element solver, called **Abaqus/Standard**, which can be extended using the **umat** subroutine.
- an explicit finite element solver, called **Abaqus/Explicit**, which can be extended using the **vumat** subroutine.

MFront provides an interface for these two solvers and strives to allow the user restart a computation while switching from one solver to the other: a feature which is not easy to implement as these solvers use different conventions and, by default, different kinematics assumptions.

The **Abaqus/Standard** interface can also be used to extend the **CalculiX** finite element solver (see (18)).

2.1.2.1 Calling MFront behaviours from Abaqus/Standard or Abaqus/Explicit

Calling a behaviour from **Abaqus/Standard** or **Abaqus/Explicit** is a two steps process:

- Compiling a **MFront** library before launching either **Abaqus/Standard** or **Abaqus/Explicit**. This allows the creation of a set of libraries that can be shared between co-workers (for example, by copying them in shared folders).
- Launching **Abaqus/Standard** or **Abaqus/Explicit** with a general purpose **umat** or **vumat** subroutine which handles the loading of the **MFront** library and the call of the appropriate behaviour. The material name as declared in the input file must contains the name of the library and the name of the behaviour and the modelling hypothesis.

2.1.2.2 Features supported

For every behaviour, example of input file are automatically generated.

The **MFront** interface for the **Abaqus/Standard** solver supports orthotropic and isotropic, small and finite strain behaviours. For isotropic behaviours written in the small strain framework, internal state variables are automatically and properly rotated in finite strain analyses.

Concerning **Abaqus/Explicit**, we support orthotropic and isotropic finite strain behaviours.

In both cases, we offer three finite strain strategies to use small strain behaviours implementations in finite strain:

- The **Native** one which uses **Abaqus** native strategy to handle behaviour written in rate-form.
- The **FiniteRotationSmallStrain** strategy (see (16)).
- The **MieheApellLambrechtLogarithmicStrain** strategy (see (17)).

2.1.2.3 Performances

In **Abaqus/Standard**, our tests show that the performances of **MFront** generated behaviours are on par with native behaviours (even slightly better in some cases): those results were quite surprising because we expected the manual handling of the external libraries at the Gauss point level to be a performance bottleneck.

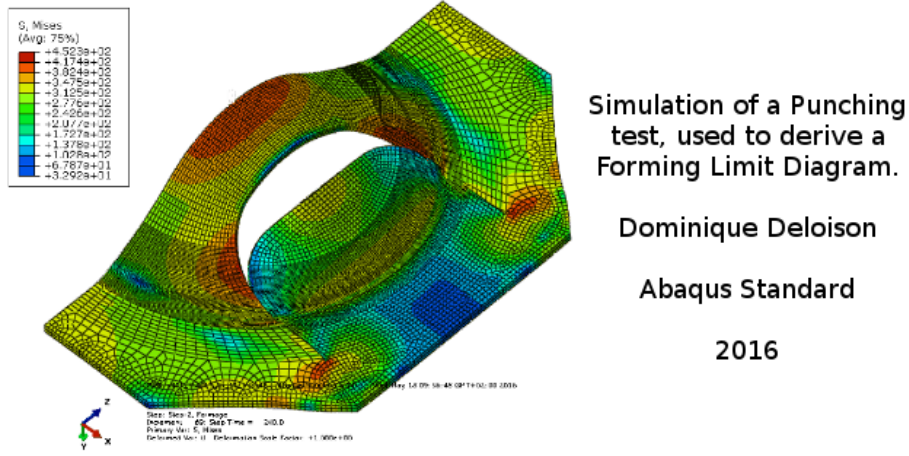


Figure 4: Simulation of a Punching test

Table 2: Comparison of computational times for a simple isotropic plasticity behaviour with linear kinematic hardening applied to the punching test depicted on Figure 4.

Implementation	Computational times
Native	638s
VUMAT (Fortran)	762s
MFront	788s

In **Abaqus/Explicit**, we found that our implementations are about 20% slower than the native ones on the simulation of the punching test depicted on Figure 4 for an isotropic plastic behaviour with linear isotropic hardening. However, our implementations are nearly equivalent to **VUMAT** implementations described in (19) (see Table 2). This difference can have several explanations:

- the **VUMAT** implementation is based on an update of the stresses whereas the **MFront** implementation keeps the elastic strain as an internal state variable.
- the **VUMAT** implementation is compiled using the **INTEL** compiler and the **MFront** implementation is compiled with **gcc**.

2.1.2.4 Additional documentations

A dedicated page has been created describing the current features of these interfaces, the potential pitfalls, and the choices that were made to support some behaviours, for example orthotropic finite strain behaviours: <http://tfel.sourceforge.net/abaqus.html>

A specific talk of the Second **MFront** User Day has been dedicated to those interfaces (see (8)).

2.2 Behaviour bricks

Behaviour bricks allow the use of predefined parts of a behaviour.

For example, the **StandardElasticity** brick handles:

- The computation of the stress according to the Hooke law, taking into account the evolution of the elastic material properties if necessary.
- The computation of an elastic prediction of the stresses, taking into account the modelling hypothesis.
- The computation of the tangent operator.
- The computation of a minimal prediction operator.
- The computation of the stored and dissipated energy.
- The additional equations associated with the plane stress and generalised plane stress modelling hypotheses support.

```

5  @Epsilon 1.e-16;
6
7  @ElasticMaterialProperties {150e9,0.3};
8
9  @StateVariable strain p;
10 p.setGlossaryName("EquivalentViscoplasticStrain");
11
12 @Parameter A = 8.e-67;
13 A.setEntryName("NortonCoefficient");
14 @Parameter E = 8.2;
15 E.setEntryName("NortonExponent");
16
17 @Integrator{
18     const auto seq = sigmaeq(sig);
19     if(seq>1.e-8*young){
20         const auto n = 3*deviator(sig)/(2*seq);
21         const auto tmp = A*pow(seq,E-1);
22         const auto df_dseq = E*tmp;
23         feel += dp*n;
24         fp -= tmp*seq*dt;
25         // jacobian
26         dfeel_ddeel += (2*mu*theta*dp/seq)*(Stensor4::M()-(n^n));
27         dfeel_ddp = n;
28         dfp_ddeel = -2*mu*theta*df_dseq*dt*n;
29     }
30 } // end of @Integrator

```

The following bricks are available:

- **StandardElasticity** which have been shortly described earlier.
- **FiniteStrainSingleCrystal** which can be used to hide many details about the classical $F_e F_p$ formalism used in the implementations of finite strain single crystal behaviours (see (20)).
- **DDIF2** which describes a damage model widely used in CEA nuclear fuel behaviours (see (21, 22)).

2.3 Pipe Modelling in MTest

Most mechanical behaviours available for the cladding are identified on tests where a pipe is submitted to an internal pressure loading.

MTest have been extended to describe various tests on pipes with various axial or radial loadings using a mono-dimensional finite strain framework described in depth in (23):

- imposed inner pressure or external pressure
- imposed external radius evolution rate, controlled through the internal pressure (which becomes one of the unknowns)
- end-cap effect
- imposed axial strain
- imposed axial force

This extension allows a consistent identification of the mechanical behaviour at finite strain.

Table 3: Example of comparison of the computation times for a simple test where a pipe with a Norton behaviour is submitted to internal pressure. The mesh size has been chosen large enough so that **MTest** computational times become significant

Solver	Computational times
MTest (C++)	0.024s
MTest (python)	0.071s
Castem 2015 PASAPAS	5.805s

Being very specialized, this extension leads to computational times significantly lower than with general purpose finite element solver (see Table 3).

2.4 New acceleration algorithms in MTest

The **Cast3M** finite element solver does not use by default a standard Newton-Raphson algorithm which would require the computation of the consistent tangent operator, but relies on a quasi-Newton algorithm using the elastic stiffness. The displacement is corrected by this fixed point iteration:

$$\Delta \mathbf{U}^{n+1} = \Delta \mathbf{U}^n - \mathbb{K}_{\text{el}}^{-1} \cdot \mathbb{R}(\Delta \mathbf{U}^n)$$

The convergence of this algorithm is greatly improved by the use of an acceleration algorithm closed to the one introduced by Anderson (see (24)). There is no point in discussing whether this strategy is better than the standard Newton-Raphson algorithm, as the answer is very sensitive to the case treated. However, we can outline that the elastic stiffness is only decomposed once when using the **Cast3M** strategy.

In **MTest**, systems solved are so small and the situation treated so simple that the full Newton-Raphson algorithm is always the better option.

However, when the consistent tangent operator is not available, the **Cast3M** strategy can be used in **MTest**. As described by I. Ramière (see (13)), various acceleration algorithms were introduced and tested in **MTest**.

Those algorithms were improved by É. Castelier to accelerate the equilibrium iterations of the **TMFFT** solver (see (14)) for systems with several millions of degrees of freedom (see Figure 5). They are available in **MTest** with the name **UAnderson** and **FAnderson**. The implementations of those algorithms, which require a special care to get accurate and reliable results, are available as autonomous classes which can be used outside **MTest**.

2.5 A stable API to build external tools: application to the mechanical behaviours identification developed in the MAP platform

With version 3.0.x, **MFront** and **MTest** now provide stable API in C++ and Python upon which external tools can be built.

This is illustrated by the recent development by EDF of an identification tool, called `c_solver_constitutive_law_0d`, in the Material Ageing Platform (MAP) based on:

- **MFront** to generate the mechanical behaviour. The **MFront python** module is used to extract information about the behaviour, such as the parameters that has to be identified.
- **MTest** to perform simulations on simple mechanical tests such as tensile tests or tests on pipe as described earlier. More complex simulations requires a fully-fledge finite element solver such as **Code_Aster** or **Cast3M**. **MTest** is used through its **python** module.

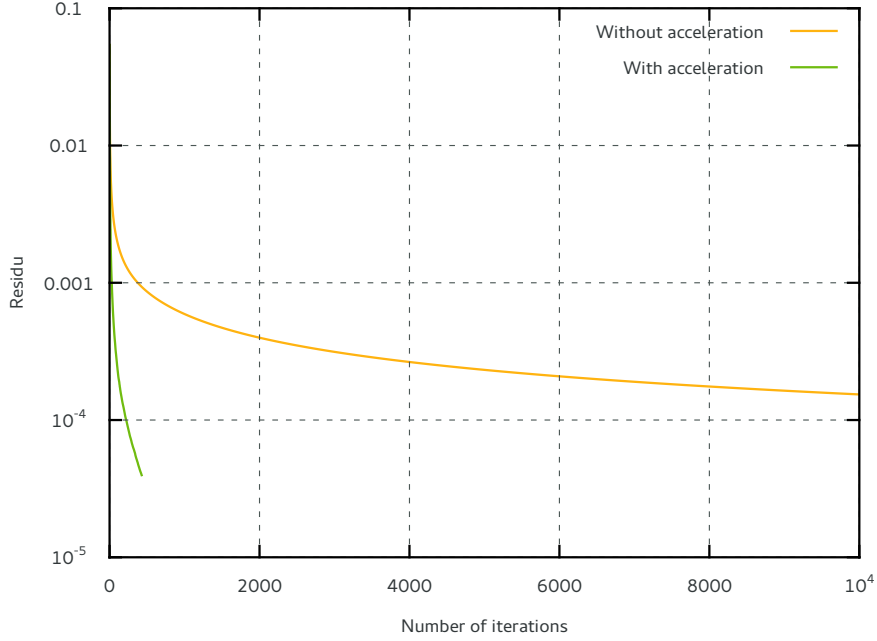


Figure 5: Convergence of the TMFFT solver with and without acceleration (see (14) for details)

- **ADAO** for parameters optimisation. **ADAO** is a specific software dedicated to data assimilation and distributed in the platform **Salome** (see (25)).

This identification tool, co-developed by two departments of EDF R&D (SINETICS and MMC), is a central part of the material knowledge management of the MAP platform and thus addresses the following issues:

- Ease of use by engineers without advanced programming skills.
- Provide a robust and reliable identification of mechanical behaviours.
- Guarantee a consistent approach of the identification process and the behaviour usage in engineering studies. In particular, its is the same implementation that will be used in the identification process and the engineering simulations.
- Maintainability and tracability. EDF wants to guarantee that the identification process of each behaviour used in engineering studies can be replayed for years.
- Integration in the EDF material knowledge database (called **CADEEX**) which stores every information from the material specification, the measures, the thermal treatments, the experimental data and the identification process and finally the mechanical behaviour implementation. This integration is currently under heavy developments.

Though relatively young - it has been developed for one year now -, this tool gives quite promising results, as discussed during the MFront users day (see (26)).

This ambitious project highlights the need for high quality coding standards for **TFEL**, **MFront** and **MTest**. This point is discussed in depth in the following section.

3 An industrial strength software

The version 3.0.x is based on the C++11 standard. This implied a major code refactoring. In particular, the expression template engine was greatly simplified and is now much more reliable and maintainable.

3.1 Systems supported

Version 3.0.x of **TFEL/MFront** will be available on the following systems:

- **LiNuX**
- **Windows**. The port to **Visual Studio 2015** is the first step toward an industrial strength support of this platform. However, due to the very nature of this platform, a direct use of **MFront** is discouraged. One shall consider creating a material knowledge management project based on **cmake** to build material librairies.
- **FreeBSD** and **TrueOS**.

Various Unix-based systems, including **Mac Os X**, have been tested at various stages of the development of this version and shall work out of the box.

3.2 Compiler support

Version 3.0.x were tested using the following compilers:

- **gcc**: versions 4.7, 4.8, 4.9, 5.1, 5.2, 6.1, 6.2
- **clang**: versions 3.5, 3.6, 3.7, 3.8, 3.9
- **intel**. The only supported version is the 2016 version. Intel compilers 15 are known not to work due to a bug in the EDG front-end that can't parse a syntax mandatory for the expression template engine. The same bug affects the **Blitz++** library (see <http://dsec.pku.edu.cn/~mendl/blitz/manual/blitz11.html>)
- **Visual Studio** The only supported version is the 2015 version. Previous versions does not provide a suitable **C++11** support.

3.3 Documentation

A vast amount of documentation has been written for **TFEL** and **MFront**, mostly in French: (see <http://tfel.sourceforge.net/publications.html>).

With version 3.0, we followed the example of the **CMake** software and introduced the ability to query documentation from the command line for **MFront** and **MTest** which now provide the following options:

- **--help-keywords** which displays the help associated with all keywords.
- **--help-keywords-list** which displays the list of available keywords.
- **--help-keyword** which displays the help associated to a specific keyword.

The documentations of the keywords are now written in English and displayed using **pandoc** markdown language.

3.4 Code quality

Code quality is an important matter in the development of **TFEL** and **MFront**. As an example, one may consider the number of code defects measured by the **Coverity** analysis tool: this indicator shows that the code is nearly as good as great open-source projects such as **Python** and much lower that what is considered as well developed industrial projects.

Many static analysis tools (**Coverty**, **PVS-Studio**, **cppcheck**, **clang-tidy**) were used to improve the overall quality of the code.

3.4.1 A myriad of tests

TFEL and MFront use a test-driven development scheme: each new functionality has at least one associated test. The current version is delivered with more than 3300 tests, each of them containing various units-tests.

4 Conclusions

This paper have highlighted the improvements made in the version 3.0 of TFEL, MFront and MTest. Mechanical behaviours can be written even more easily than in previous versions and performances are competitive with built-in behaviour implementation of most mechanical solvers. Being mostly developed for implicit solvers, we were pleased to see that performances obtained in explicit solvers such as Europlexus and Abaqus/Explicit are quite decent.

The MFront users' community is steadily increasing outside the nuclear industry and the french mechanical community: its use now encompasses a wide range of materials and applications. Creation of new interfaces is relatively easy, extensive testing is time consuming. Every new user is thus welcomed, even with a solver which is not currently supported yet, and everybody's contribution in the improvement of TFEL and MFront is much appreciated.

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