

# tmech a C++ library for the numerical study of the physics of continuous materials using higher-order tensors

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## Summary

The objective of the tmech framework is to propose high-level semantics, inspired by different linear algebra packages, allowing fast software prototyping in a low-level compiled language numerical study of the physics of continuous materials using higher-order tensors.

The tmech framework is provided as an open source software under the BSD-3-Clause License, compiled and validated with clang and gcc

## Statement of need

For the numerical study of the physics of continuous materials using higher-order tensors. Partial Differential Equations (PDEs) describing natural phenomena are modelled using tensors of different order. Two commonly studied problems are heat transfer, which include temperature and heat flux (rank-0 and rank-1 tensor, respectively), and continuum mechanics, which include stress and strain (rank-2 tensors) and the so-called tangent stiffness (rank-4 tensor).

## Overview and features

Non-indexed lower case light face Latin letters (e.g. f and h) are used for scalars

## Tensor operations involving overloaded C++ operator functions

- Addition of tensors of same rank and dimension
- Subtraction of tensors of same rank and dimension
- Scalar update of tensors

## General inner product

Hence if one of the original tensors is of rank-m and the other is of rank-n, the inner product will be of rank-(m + n - 2). Controlled by template parameters SeqLHS and SeqRHS. Bases contained in sequence SeqLHS are contracted with bases contained in sequence SeqRHS. tmech provides some wrapper functions for the most common inner products  $a : B$ ,  $A : b$ ,  $A : B$  and  $A :: B$ , see documentation for more details.

```
using SeqL = tmech::sequence<3,4>;
using SeqR = tmech::sequence<1,2>;
//Double contraction of two 4th order tensors
tmech::tensor<double, 3, 4> A, B, C;
C = tmech::inner_product<SeqL, SeqR>(A,B);
//Double contraction of a 4th and a 2th order tensor
```

```
tmech::tensor<double, 3, 2> a, c;
c = tmech::inner_product<SeqL, SeqR>(C,a);
```

## General outer product

On multiplying each component of a tensor of rank  $r$  by each component of a tensor of rank  $k$ , both of dimension  $m$ , a tensor of rank  $(r + k)$  with  $mr+k$  components is obtained. The outer product of a tensor of type  $(m, n)$  by a tensor of type  $(p, q)$  results in a tensor of type  $(m + p, n + q)$  the outer product of two tensors their outer product is a tensor. The outer product of tensors is also referred to as their tensor product, controlled by template parameters SeqLHS and SeqRHS. Bases contained in sequence SeqLHS are used for ordered element access in A. Bases contained in SeqRHS are used for ordered element access in B expression. tmech provides some wrapper functions for the most common outer products  $\otimes$ ,  $\otimes$  and  $\otimes$ , see documentation for more details. An example code snippet, where two second-order tensor are multiplied to form a fourth-order tensor is given as

```
using SeqL = tmech::sequence<1,2>;
using SeqR = tmech::sequence<3,4>;
...
tmech::tensor<double, 3, 2> a, b;
tmech::tensor<double, 3, 4> C;
C = tmech::outer_product<SeqL, SeqR>(a,b);
```

where bases 1,2 of the new tensor C are given by a and bases 3,4 are given by b.

## General basis rearrangement

Controlled by template parameter Sequence, which contains the new order of bases. tmech provides some wrapper functions for the most common basis rearrangement like transposition of a second-order tensor and a major-transposition of a fourth-order tensor. An example code snippet

```
//Basis 1,2,3,4 is swaped to 3,4,1,2.
tmech::tensor<double, 3, 4> A, B;
A.randn();
B = tmech::basis_change<tmech::sequence<3,4,1,2>>(A);
B = tmech::basis_change<tmech::sequence<3,4,1,2>>(A+2*A);
```

## Decompositions

### Eigendecomposition

Eigendecomposition of a positive semi-definite symmetric second-order tensor

$$Y = \sum_{i=1}^m \lambda_i \mathbf{e}_i \otimes \mathbf{e}_i = \sum_{i=1}^m \lambda_i \mathbf{E}_i, \quad (1)$$

where  $m$  is the number of non repeated eigenvalues,  $\lambda_i$  are the corresponding eigenvalues,  $\mathbf{e}_i$  are eigenvectors and  $\mathbf{E}_i$  are eigenbasen.

```
tmech::tensor<double, 3, 2> A, A_inv;
A = tmech::sym(tmech::randn<double,3,2>());
auto A_eig = tmech::eigen_decomposition(A);
const auto [eigenvalues, eigenbasis]{A_eig.decompose_eigenbasis()};
const auto idx{A_eig.non_repeated_eigenvalues_index()};
for(auto idx : A_eig.non_repeated_eigenvalues_index()){
    A_inv += (1.0/eigenvalues[idx])*eigenbasis[idx];
}
```

```
}
std::cout<<std::boolalpha<<tmech::almost_equal(tmech::inv(A), A_inv, 5e-7)<<std::endl;
```

## 51 Polar decomposition

52 Polar decomposition of a positive semi-definite symmetric second-order tensor

$$\mathbf{F} = \mathbf{R}\mathbf{U} = \mathbf{V}\mathbf{R} \quad (2)$$

53 where  $\mathbf{R}$  is an orthogonal tensor also known as the rotation tensor,  $\mathbf{U}$  and  $\mathbf{V}$  are symmetric  
54 tensors called the right and the left stretch tensor, respectively. This function provides two  
55 different methods to determine  $\mathbf{U}$ ,  $\mathbf{V}$  and  $\mathbf{R}$ . The first method uses spectral decomposition

$$\mathbf{U} = \sqrt{\mathbf{F}^T \mathbf{F}}, \quad \mathbf{R} = \mathbf{F}\mathbf{U}^{-1}, \quad \mathbf{V} = \mathbf{R}\mathbf{U}\mathbf{R}^T, \quad (3)$$

```
//use the spectral decomposition
```

```
tmech::tensor<double, 3, 2> F;
F.randn();
auto F_polar = tmech::polar_decomposition(F);
F = F_polar.R()*F_polar.U();
F = F_polar.V()*F_polar.R();
```

56 The second one is based on a Newton iteration

$$\mathbf{R}_{k+1} = \frac{1}{2} (\mathbf{R}_k + \mathbf{R}_k^{-T}), \quad \text{with } \mathbf{R}_0 = \mathbf{F} \quad (4)$$

```
//use the newton iteration
```

```
tmech::tensor<double, 3, 2> F;
F.randn();
auto F_polar = tmech::polar_decomposition(F, // tensor to decompose
true, //use newton iteration
5e-7, // tolerance
5 // number of max steps
);
F = F_polar.R()*F_polar.U();
F = F_polar.V()*F_polar.R();
```

57 The following derivatives are also important

$$\frac{\partial \mathbf{U}}{\partial \mathbf{F}}, \quad \frac{\partial \mathbf{V}}{\partial \mathbf{F}}, \quad \frac{\partial \mathbf{R}}{\partial \mathbf{F}} \quad (5)$$

58 explicit results are given here

```
//use the spectral decomposition
```

```
tmech::tensor<double, 3, 2> F;
F.randn();
auto F_polar = tmech::polar_decomposition(F);
auto dR = F_polar.R().derivative();
auto dU = F_polar.U().derivative();
auto dV = F_polar.V().derivative();
```

## 59 Numerical differentiation

60 Numerical differentiation based on central difference scheme  $f'(x) \approx \frac{f(x+h)-f(x-h)}{2h}$

## 61 Non-symmetric tensor functions

```
tmech::tensor<double,3,2> X;
X.randn();
auto func = [&](auto const& F){return 1.5*(tmech::trace(tmech::trans(F)*F) - 3);};
auto dFunc = tmech::num_diff_central(func, X);
auto dFunc_ = [&](auto const& F){return tmech::num_diff_central(func, F);};
auto ddFunc = tmech::num_diff_central<tmech::sequence<1,2,3,4>>(func, X);
```

## 62 Symmetric tensor functions

```
using Sym2x2 = std::tuple<tmech::sequence<1,2>,tmech::sequence<2,1>>;
//symmetry of the result
using Sym4x4 = std::tuple<
    tmech::sequence<1,2,3,4>,
    tmech::sequence<2,1,3,4>,
    tmech::sequence<1,2,4,3>,
    tmech::sequence<2,1,4,3>>;
tmech::tensor<double,3,2> X;
X = tmech::sym(tmech::rand<double,3,2>());
//first deriv
auto func = [&](auto const& F){return 1.5*(tmech::trace(tmech::trans(F)*F) - 3);};
auto dfunc_res = tmech::num_diff_sym_central<Sym2x2>(func, X);
//second deriv
auto dfunc = [&](auto const& F){return tmech::num_diff_sym_central<Sym2x2>(func, F);};
auto ddfunc_res = tmech::num_diff_sym_central<Sym2x2, Sym4x4>(dfunc, X);
```

## 63 Compile-time differentiation

64 As an illustrative example of using tmech, we here give the mathematical formulation of an  
 65 energy potential and stress in large deformation continuum mechanics and its implementation  
 66 as a function in Julia. For a deformation gradient  $F = I + \text{grad } u$ , where  $u$  is the displacement  
 67 from the reference to the current configuration, the right Cauchy-Green deformation tensor is  
 68 defined by  $C = F^T F$ . The Second Piola-Kirchhoff stress tensor  $S$  is derived from the Helmholtz  
 69 free energy  $\psi$  by the relation

$$S = 2 \frac{\partial \psi}{\partial C} \quad (6)$$

70 The strain energy density function for an incompressible Mooney–Rivlin material is

$$\psi = C_{10}(I_1 - 3) + C_{01}(I_2 - 3), \quad \text{where } I_1 = \text{trace} C, \quad I_2 = \frac{1}{2}(I_1^2 - \text{trace}(C^2)) \quad (7)$$

71 where is the isochoric part

## 72 User material in Abaqus

73 Abaqus is a commercial software and often used in research (Abaqus, 2014)

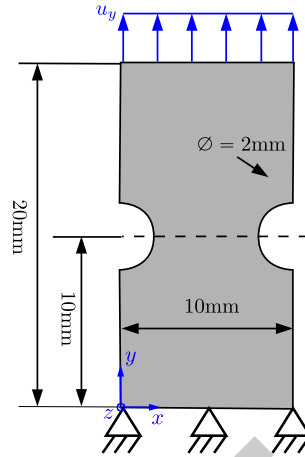


Figure 1: Caption for example figure.

$$\varepsilon = \varepsilon^e + \varepsilon^p, \quad \sigma = \lambda \text{trace}(\varepsilon^e) I + 2\mu \varepsilon^e, \quad f[\sigma, q] = \|\eta\| - \sqrt{\frac{2}{3}} K[\alpha], \quad (8)$$

$$\dot{\varepsilon}^p = \gamma \frac{\eta}{\eta}, \quad \dot{\beta} = \gamma \frac{2}{3} H[\alpha] \frac{\eta}{\eta}, \quad \dot{\alpha} = \gamma \sqrt{\frac{2}{3}} \quad (9)$$

```
#include <tmech/tmech.h>
using tensor2 = tmech::tensor<double,3,2>;
using tensor4 = tmech::tensor<double,3,4>;

extern "C" void umat_(double *stress, double *statev, double *ddsdde, double *sse,
double *spd, double *scd, double *rpl, double *ddsddt, double *drplde,
double *drpldt, double *stran, double *dstran, double *time, double *dtime,
double *temp, double *dtemp, double *predef, double *dpred, char *cmname,
int *ndi, int *nshr, int *ntens, int *nstatv, double *props, int *nprops,
double *coords, double *drot, double *pnewdt, double *celent, double *dfgrd0,
double *dfgrd1, int *noel, int *npt, int *layer, int *kspt,
int *kstep, int *kinc, short cmname_len){
//parameters
const double E      = props[0]; //210000
const double nue     = props[1]; //0.3
const double m       = props[2];
const double k       = props[3];
const double sig_y   = props[4];

//Lamé parameters
const double mu      = E/(2*(1+nue));
const double lambda  = E*nue/((1+nue)*(1-2*nue));

//second order tensor
const tmech::eye<double,3,2> I;
//fourth order symmetric identity tensor
const auto IIsym = 0.5*(tmech::otimesu(I,I) + tmech::otimesl(I,I));
//fourth order identity tensor
const auto II = tmech::otimes(I,I);
```

```
//voigt to tensor currently abq_std only Dim==3 supported
tmech::adaptor<double,3,2, tmech::abq_std<3,true> strain(stran);
tmech::adaptor<double,3,2, tmech::abq_std<3,true> dstrain(dstran);
tmech::adaptor<double,3,2, tmech::abq_std<3,true> epspl(statev);
tmech::adaptor<double,3,2, tmech::abq_std<3> sig(stress);
tmech::adaptor<double,3,4, tmech::abq_std<3> C(ddsde);

//accumulated plastic strain last step
const double epl_n{*(statev+6)};
//updated strain
const tensor2 eps = strain + dstrain;
//trial stress
const tensor2 sig_tr = lambda*tmech::trace(eps-epspl)*I + 2*mu*(eps-epspl);
//deviatoric stress
const auto sig_dev = tmech::dev(sig_tr);
//equivalent stress
const auto sig_eq = std::sqrt(tmech::dcontract(sig_dev,sig_dev));
//yield function
auto f = [&](double const& epl){return sig_eq - sig_y - k*std::pow(epl,m)};
if(f(epl_n) > 0){ //plastic
    //yield normal
    const tensor2 N = 3*sig_dev/(2*sig_eq);
    //nonlinear function
    auto f_non = [&](double const& depl){
        const auto f{sig_eq - 3*G*depl - k*std::pow(epl_n+epl,m)};
        const auto df{3*G - k*m*std::pow(epl_n+epl,m-1)};
        return std::make_pair(std::make_tuple(std::make_tuple(df)), std::make_tuple(f));
    };
    const auto [iter, norm, dlambd]{tmech::general_newton_raphson_iterate(f_non, 0,
}
}
}
```

74 : Abaqus example linear elasticity.

75 Compile to obtain the . With the following commands for a Linux based operating system  
76 the object file is generated

```
gcc -c -fPIC -O3 -march=native -std=c++17 -I/path_to_tmech -o object_file_output.o input
```

77 The compile flags -O3 and -march=native are used to enable optimization and auto vectorization  
78 of the code, respectively. After the object file is generated, a Abaqus job can be submitted by  
79 the following commands

```
abaqus job=input_file_name user=object_file_output
```

80 where input\_file\_name is the name of the .inp input file and object\_file\_output is the name  
81 of the generated object file.

## 82 References

83 Abaqus. (2014). *Abaqus/standard user's manual, version 6.14*. Dassault Systèmes Simulia  
84 Corporation.