# **Documentation of the pw85 library** *Release 2.0*

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# **CONTENTS**



#### **Abstract**

This library implements the "contact function" defined by Perram and Wertheim (J. Comp. Phys. 58(3), 409–416, [DOI:10.1016/0021-9991\(85\)90171-8\)](https://doi.org/10.1016/0021-9991(85)90171-8) for two ellipsoids. Given two ellipsoids, this function returns the *square* of the common factor by which both ellipsoids must be scaled (their centers being fixed) in order to be tangentially in contact.

This library is released under a BSD 3-Clause License.

#### **OVERVIEW**

## <span id="page-6-1"></span><span id="page-6-0"></span>**1.1 Introduction**

It is quite common in materials science to reason on assemblies of ellipsoids as model materials. Although simplified upscaling mean-field/effective-field theories exist for such microstructures, they often fail to capture the finest details of the microstructure, such as orientation correlations between anisotropic inclusions, or particle-size distributions. In order to account for such microstructural details, one must resort to so-called *full-field* numerical simulations (using dedicated tools such as [Damask](https://damask.mpie.de/) or [Janus](https://github.com/sbrisard/janus), for example).

Full-field simulations require *realizations* of the microstructure. For composites made of ellipsoidal inclusions embedded in a (homogeneous) matrix, this requires to be able to generate assemblies of (non-overlapping) ellipsoids. The basic ingredient of such microstructure simulations is of course the overlap test of two inclusions.

Checking for the overlap (or the absence of it) of two ellipsoids is not as trivial as checking for the overlap of two spheres. Several criteria can be found in the literature [\[VB72](#page-38-1)];[[PW85\]](#page-38-2);[[WWK01\]](#page-38-3);[[CYP07\]](#page-38-4);[[ABH18\]](#page-38-5). We propose an implementation of the *contact function* of Perram and Wertheim [\[PW85\]](#page-38-2).

The present chapter is organised as follows. We first give a brief description of the contact function. Then, we discuss two essential features of this function: robustness with respect to floating-point errors and suitability for application to Monte-Carlo simulations. Finally, we give a brief description of the pw85 library.

# <span id="page-6-2"></span>**1.2 The contact function of Perram and Wertheim [PW85]**

The origin being fixed, points are represented by the 3×1 column-vector of their coordinates in a global cartesian frame. For i = 1, 2,  $E_i \subset \mathbb{R}^3$  denotes the following ellipsoid:

(1)  $E_i = \{m \in \mathbb{R}^3: (m-c_i)^{\top} \cdot Q_i^{-1} \cdot (m-c_i) \leq 1\},$ 

where  $c_i \in \mathbb{R}^3$  is the center of E<sub>i</sub>, and Q<sub>i</sub> is a positive definite matrix. Perram and Wertheim define the following function:

(2)  $f(\lambda; r_{12}, Q_1, Q_2) = \lambda(1-\lambda)r_{12}r_{12}r_{12}$ ,

where  $0 \le \lambda \le 1$  is a scalar,  $Q = (1-\lambda)Q_1 + \lambda Q_2$ , and  $r_{12} = c_2 - c_1$  denotes the center-to-center radius-vector. The *contact function*  $\mu^2$  (E<sub>1</sub>, E<sub>2</sub>) of the two ellipsoids is defined as the unique maximum of f over (0, 1):

(3)  $\mu^2 = \max\{f(\lambda; r_{12}, Q_1, Q_2), 0 \le \lambda \le 1\}.$ 

It turns out that the contact function has a simple geometric interpretation. Indeed, μ is the quantity by which each of the two ellipsoids  $E_1$  and  $E_2$  must be scaled to bring them in contact. Therefore, an overlap test could be defined as follows

- $\mu^2(E_1, E_2)$  < 1: the two ellipsoids overlap,
- $\mu^2(E_1, E_2) > 1$ : the two ellipsoids do not overlap,
- $\mu^2$ (E<sub>1</sub>, E<sub>2</sub>) = 1: the two ellipsoids are tangent.

Despite its apparent complexity, this overlap test has two nice features that are discussed below.

## <span id="page-7-0"></span>**1.3 Features of the overlap test**

#### **1.3.1 Robustness with respect to floating-point errors**

All overlap tests amount to checking for the sign of a real quantity  $\Phi(E_1, E_2)$  that depends on the two ellipsoids E<sub>1</sub> and E<sub>2</sub>. The ellipsoids do not overlap when  $\Phi(E_1, E_2) < 0$ ; they do overlap when  $\Phi(E_1, E_2) > 0$ . Finally, we usually have  $\Phi(E_1, E_2) = 0$  when  $E_1$  and  $E_2$  are in tangent contact (but it is important to note that, depending on the overlap criterion, the converse is not necessarily true).

In a finite precision setting, we are bound to make wrong decisions about pairs of ellipsoids that are such that Φ is small. Indeed, let us consider a pair of ellipsoids (E<sub>1</sub>, E<sub>2</sub>) for which the true value of  $\Phi$ ,  $\Phi$ <sup>e</sup> (E<sub>1</sub>, E<sub>2</sub>), is close to the machine epsilon. Then, the numerical estimate of  $\Phi$ ,  $\Phi$ <sup>e</sup> (E<sub>1</sub>, E<sub>2</sub>), is also (hopefully) a very small value. However, whether  $\Phi_{\alpha}$  (E<sub>1</sub>, E<sub>2</sub>) is the same sign as  $\Phi_{\alpha}$  (E<sub>1</sub>, E<sub>2</sub>) (and therefore delivers the correct answer regarding overlap) is uncertain, owing to accumulation of round-off errors. Such misclassifications are acceptable provided that they occur for ellipsoids that are close (nearly in tangent contact). The overlap criterion will be deemed robust if it is such that  $\Phi(E_1, E_2)$  is small for nearly tangent ellipsoids only. This is obviously true of the overlap test based on the contact function of Perram and Wertheim. Note that some of the overlap tests that can be found in the literature do not exhibit such robustness.

#### **1.3.2 Application to Monte-Carlo simulations**

Generating compact assemblies of hard particles is a notoriously difficult task. Event-driven simulations [\[DTS05\]](#page-38-6); [\[DTS05a\]](#page-38-7) are often used, but require a lot of book-keeping. A comparatively simpler approach [\[BL13\]](#page-38-8) is similar to atomistic simulations with a non-physical energy. More precisely, starting from an initial configuration where the n ellipsoids  $E_1$ , …,  $E_n$  do overlap, a simulated annealing strategy is adopted to minimize the quantity  $U(E_1, ..., E_n)$ defined as follows:

```
(4) U(E_1, ..., E_n) = \sum u(E_i, E_j),
                      1 \leq i \leq j \leq n
```
where u(E<sub>1</sub>, E<sub>2</sub>) denotes an *ad-hoc* pair-wise (non-physical) potential, that should vanish when the two ellipsoids do not overlap, and be "more positive when the overlap is greater" (this sentense being deliberately kept vague). A possible choice for u is the following:

(5)  $u(E_1, E_2) = max\{0, \mu^{-1}(E_1, E_2)\}.$ 

Monte-Carlo simulations using previous implementations of the contact function of Perram and Wertheim and the above definition of the energy of the system were successfully used to produce extremely compact assemblies of ellipsoids [\[BL13](#page-38-8)].

# <span id="page-8-0"></span>**1.4 Implementation**

pw85 is a C library that implements the contact function of Perram and Wertheim. It is released under a BSD-3 license, and is available at <https://github.com/sbrisard/pw85>. It is fully documented at <https://sbrisard.github.io/pw85>.

The core library depends on The [boost::mathGNU](https://www.boost.org/doc/libs/1_76_0/libs/math/doc/html/math_toolkit/brent_minima.html) (for its implementation of the Brent algorithm).

The API is extremely simple; in particular it defines no custom objects: parameters of all functions are either simple types (size t, double) or arrays. Note that all arrays must be pre-allocated and are modified in-place. This minimizes the risk of creating memory leaks when implementing wrappers for higher-level (garbage-collected) languages.

A Python wrapper (based on [pybind11\)](https://pybind11.readthedocs.io/en/stable/) is also provided. It has the following (fairly standard) dependencies: [NumPy,](https://numpy.org/) [pytest](https://pytest.org/) and [h5py](https://www.h5py.org/).

Note that when developing the library, several strategies have been tested for the evaluation of the function f defined above, and its optimization. Evaluation of f relies on a Cholesky decomposition of Q; we tested the accuracy of this implementation over a comprehensive set of large-precision reference values that are available on Zenodo([https:](https://doi.org/10.5281/zenodo.3323683) [//doi.org/10.5281/zenodo.3323683](https://doi.org/10.5281/zenodo.3323683)). Optimization of f first starts with a few iterations of Brent's robust algorithm. Then, the estimate of the minimizer is refined through a few Newton–Raphson iterations.

# <span id="page-8-1"></span>**1.5 Extensions**

Several improvements/extensions are planned for this library:

- 1. Provide a 2D implementation of the contact function.
- 2. Allow for early stop of the iterations. If, during the iterations, a value of  $\lambda$  is found such that  $f > 1$ , then  $\mu^2$ must be greater than 1, and the ellipsoids certainly do not overlap, which might be sufficient if the user is not interested in the exact value of the contact function.
- 3. Return error codes when necessary. Note that this would be an extra safety net, as the optimization procedure is extremely robust. Indeed, it never failed for the thousands of test cases considered (the function to optimize has the required convexity over  $(0, 1)$ .

This project welcomes contributions. We definitely need help for the following points:

- 1. Define a "Code of conduct".
- 2. Improve the Python wrapper (see Issue XXX).
- 3. …

# <span id="page-8-2"></span>**1.6 Acknowledgements**

The author would like to thank Prof. Chloé Arson (GeorgiaTech Institute of Technology, School of Civil and Environmental Engineering) for stimulating exchanges and research ideas that motivated the exhumation of this project (which has long been a defunct Java library).

The author would also like to thank Xianda Shen (GeorgiaTech Institute of Technology, School of Civil and Environmental Engineering) for testing on fruity operating systems the installation procedure of this and related libraries. His dedication led him to valiantly fight long battles with setuptools and brew.

**TWO**

#### **INSTALLATION**

<span id="page-10-0"></span>First of all, clone the repository

\$ git clone https://github.com/sbrisard/pw85

## <span id="page-10-1"></span>**2.1 Installing the C++ library**

pw85 is a header-only library: there is no installation procedure *per se* and you can drop the header wherever you like (as long as it is located in a pw85 subdirectory). To use pw85 in a C++ project, you must include the header

*#include <pw85/pw85.hpp>*

and inform the compiler of its location.

**Note:** pw85 depends on [Boost::Math](https://www.boost.org/doc/libs/1_75_0/libs/math/) (for the implementation of the Brent algorithm). You must pass the relevant options to the compiler. Typically, these would be -I options. The C++ tutorials provides a *[CMake example](#page-15-0)*.

To run the tests or build the documentation properly, you need to first build the python bindings (see *[below](#page-10-2)*). To further test your installation, build the example in the *[C++ tutorial](#page-15-0)*.

<span id="page-10-2"></span>

# **2.2 Installing the Python bindings**

The Python bindings are built with [pybind11](https://pybind11.readthedocs.io/), which must be installed.

To install the pw85 module, cd into the python subdirectory and run the setup.py script as follows.

First, build the extension:

\$ python setup.py build\_ext -Ipath/to/boost/math

When the extension is built, installation is down as usual:

\$ python setup.py install --user

or (if you intend to edit the project):

\$ python setup.py develop --user

To run the tests with [Pytest](https://docs.pytest.org/):

\$ python -m pytest tests

(beware, these tests take some time!).

**Note:** Upon first execution, the test script will attempt to retrieve some precomputed reference data. In case of failure (e.g. if you sit behind a firewall), this reference file can be downloaded manually at this address: [https://zenodo.org/](https://zenodo.org/record/3323683/files/pw85_ref_data-20190712.h5) [record/3323683/files/pw85\\_ref\\_data-20190712.h5](https://zenodo.org/record/3323683/files/pw85_ref_data-20190712.h5)

The file should be placed in the data/ subdirectory, at the root of the project, and should be renamed pw85\_ref\_data. h5:



#### <span id="page-11-0"></span>**2.3 Building the documentation**

**Note:** For the documentation to build properly, the python module must be installed, as it is imported to retrieve the project metadata.

The documentation of pw85 requires [Sphinx](https://www.sphinx-doc.org/). The C++ API docs are built with [Doxygen](https://www.doxygen.nl/) and the [Breathe](https://breathe.readthedocs.io/) extension to [Sphinx](https://www.sphinx-doc.org/).

To build the HTML version of the docs in the docs subdirectory:

```
$ cd docs
$ sphinx-build -b html . ../docs
```
To build the LaTeX version of the docs:

\$ cd docs \$ make latex

#### **THREE**

# **TUTORIAL**

<span id="page-12-0"></span>In this tutorial, we consider two ellipsoids, and check wether or not they overlap.

Ellipsoid Eı is an [oblate spheroid](https://en.wikipedia.org/wiki/Spheroid) centered at point  $x_1 = (-0.5, 0.4, -0.7)$ , with equatorial radius  $a_1 = 10$ , polar radius  $c_1 = 0.1$  and polar axis  $(0, 0, 1)$ .

Ellipsoid E<sub>2</sub> is a [prolate spheroid](https://en.wikipedia.org/wiki/Spheroid) centered at point (0.2, -0.3, 0.4), with equatorial radius  $a_1 = 0.5$ , polar radius  $c_1 = 5$  and polar axis (1, 0, 0).

To carry out the overlap check, we must first create the representation of ellipsoids  $E_i$  as quadratic forms  $Q_i$  (see *[Mathematical representation of ellipsoids](#page-18-1)*). Convenience functions are provided to compute the matrix representation of a *spheroid*.

**Note:** In principle, the contact function implemented in PW85 applies to *any* ellipsoids (with unequal axes). However, at the time of writing this tutorial (2019-01-01), convenience functions to compute the matrix representation of a general ellipsoid is not yet implemented. Users must compute the matrices themselves.

We first check for the overlap of  $E_1$  and  $E_2$  using the Python wrapper of pw85. We will then illustrate the C API.

# <span id="page-12-1"></span>**3.1 Python tutorial**

The Python module relies on [NumPy](www.numpy.org) for passing arrays to the underlying C library. We therefore import both modules:

**>>> import numpy as np >>> import pw85**

and define the parameters of the simulation:

```
>>> x1 = np.array([-0.5, 0.4, -0.7])
\Rightarrow n1 = np.array([0., 0., 1.])
>>> a1, c1 = 10, 0.1
\Rightarrow \times 2 = np.array([0.2, -0.3, 0.4])>>> n2 = np.array([1., 0., 0.])
>>> a2, c2 = 0.5, 5.
>>> r12 = x2-x1
```
where  $r₁₂$  is the vector that joins the center of the first ellipsoid,  $x₁$ , to the center of the second ellipsoid,  $x₂$ .

We use the function pw85.spheroid() to create the matrix representations  $q_1$  and  $q_2$  of the two ellipsoids. Note that these arrays must be *preallocated*:

```
\Rightarrow q1 = np.empty((6, ), dtype=np.float64)
>>> pw85.spheroid(a1, c1, n1, q1)
>>> q1
array([ 1.e+02, -0.e+00, -0.e+00, 1.e+02, -0.e+00, 1.e-02])
\Rightarrow q2 = np.empty like(q1)
>>> pw85.spheroid(a2, c2, n2, q2)
>>> q2
array([25., 0., 0., 0.25, 0., 0.25])
```
We can now compute the value of the contact function — see the documentation of  $pw85$ . contact\_function():

```
\Rightarrow out = np.empty((2, ), dtype=np.float64)
>>> pw85.contact_function(r12, q1, q2, out)
>>> mu2, lambda_ = out
>>> print('μ² = {}'.format(mu2))
\Rightarrow print('\lambda = \{\}'.format(lambda))
\mu^2 = 3.362706040638343
\lambda = 0.1668589553405904
```
We find that  $\mu^2 > 1$ , hence  $\mu > 1$ . In other words, both ellipsoids must be *swollen* in order to bring them in contact: the ellipsoids do not overlap!

#### **3.1.1 Checking the output**

The output of this simulation can readily be checked. First, we can check that  $q_1$  and  $q_2$  indeed represent the ellipsoids E<sub>1</sub> and E<sub>2</sub>. To do so, we first construct the symmetric matrices  $Q_1$  and  $Q_2$  from their upper triangular part

```
>>> Q1 = np.zeros((3, 3), dtype=np.float64)
>>> i, j = np.triu_indices_from(Q1)
>>> Q1[i, j] = q1
>>> Q1[j, i] = q1
>>> Q1
array([[ 1.e+02, -0.e+00, -0.e+00],
       [-0.e+00, 1.e+02, -0.e+00],
       [-0.e+00, -0.e+00, 1.e-02]])
```

```
>>> Q2 = np.zeros_like(Q1)
>>> Q2[i, j] = q2
>>> Q2[j, i] = q2
>>> Q2
array([[25. , 0. , 0. ],
      [ 0. , 0.25, 0. ],
      [0. , 0. , 0.25]
```
We can now check these matrices for some remarkable points, first for ellipsoid  $E_1$ 

```
\Rightarrow Q1 inv = np.linalg.inv(Q1)
\Rightarrow f1 = lambda x: Q1_inv.dot(x).dot(x)
>>> f1((a1, 0., 0.))
1.0
>>> f1((-a1, 0., 0.))
1.0
>>> f1((0., a1, 0.))
1.0
>>> f1((0., -a1, 0.))
```
(continued from previous page)

```
1.0
>>> f1((0., 0., c1))
0.9999999999994884
>>> f1((0., 0., -c1))
0.9999999999994884
```
then for ellipsoid  $E_2$ 

```
>>> Q2_inv = np.linalg.inv(Q2)
\Rightarrow f2 = lambda x: Q2_inv.dot(x).dot(x)
>>> f2((c2, 0., 0.))
1.0
>>> f2((-c2, 0., 0.))
1.0
>>> f2((0., a2, 0.))
1.0
>>> f2((0., -a2, 0.))
1.0
>>> f2((0., 0., a2))
1.0
>>> f2((0., 0., -a2))
1.0
```
Note that in the above tests, we have omitted the centers of ellipsoids  $E_1$  et  $E_2$  (both ellipsoids were translated to the origin).

We will now verify the corectness of the value found for the scaling factor  $\mu$ . To do so, we will find the coordinates of the contact point of the two scaled ellipsoids, and check that the normals to the two ellipsoids at that point coincide.

Although we use formulæ from the *[Theory](#page-18-0)* section to find the coordinates of the contact point, x₀, it is not essential for the time being to fully understand this derivation. What really matters is to check that the resulting point  $x<sub>0</sub>$  is indeed the contact point of the two scaled ellipsoids; how the coordinates of this point were found is irrelevant.

From the value of λ returned by the function pw85.contact\_function(), we compute Q defined by Eq. *[\(10\)](#page-19-1)* in section *[Theory](#page-18-0)*, as well as  $x = Q^{-1} \cdot r_{12}$ 

```
>>> Q = (1-lambda_)*Q1+lambda_*Q2
\Rightarrow x = np. linalg. solve(Q, r12)
```
From which we find x₀, using either Eq. *[\(9a\)](#page-19-2)* or Eq. *[\(9b\)](#page-19-2)* (and we can check that both give the same result)

```
\Rightarrow \times 0a = x1+(1-1)ambda_\Rightarrownp.dot(Q1, x)
>>> x0a
array([ 0.16662271, -0.29964969, -0.51687799])
\Rightarrow \times 0b = \times2-lambda *np.dot(Q2, x)
>>> x0b
array([ 0.16662271, -0.29964969, -0.51687799])
```
We can now check that  $x<sub>θ</sub>$  belongs to the two scaled ellipsoids, that we first define, overriding the matrices of the unscaled ellipsoids, that are no longer needed. We observe that if ellispoid  $E_i$  is scaled by  $\mu$ , then its matrix representation Q<sub>i</sub> is scaled by  $\mu^2$ , and its inverse Q<sub>i</sub><sup>-1</sup> is scaled by  $\mu^{-2}$ .

**>>>** x0 = x0a **>>>** Q1 \*= mu2 **>>>** Q2 \*= mu2 **>>>** Q1\_inv /= mu2 **>>>** Q2\_inv /= mu2

```
\Rightarrow x = x0-x1>>> Q1_inv.dot(x).dot(x)
1.0000000000058238
```

```
\Rightarrow x = x0-x2>>> Q2_inv.dot(x).dot(x)
0.9999999999988334
```
Therefore  $x \circ$  indeed belongs to both ellipsoids. We now compute the normal  $n \circ$  to ellipsoid E<sub>i</sub> at point  $x \circ$ . Since ellipsoid E<sub>i</sub> is defined by the level-set:  $(x-x_i)^{-1} \cdot (x-x_i) = 1$ , the normal to E<sub>i</sub> is given by Q<sub>i</sub><sup>-1</sup> ·  $(x-x_i)$ (which is then suitably normalized)

```
\Rightarrow n1 = Q1 inv.dot(x0-x1)
>>> n1 /= np.linalg.norm(n1)
>>> n1
array([ 3.64031943e-04, -3.82067448e-04, 9.99999861e-01])
```

```
\Rightarrow n2 = Q2 inv.dot(x0-x2)
>>> n2 /= np.linalg.norm(n2)
>>> n2
array([-3.64031943e-04, 3.82067448e-04, -9.99999861e-01])
```
We find that  $n_1 = -n_2$ . Therefore, E<sub>1</sub> and E<sub>2</sub> are in external contact. QED

Follow this link to download the above Python script.

#### <span id="page-15-0"></span>**3.2 C++ tutorial**

The Python interface to PW85 has been kept close to the undelying C++ API. The following C++ program (download source file) defines the two ellipsoids, then computes  $\mu^2$  and  $\lambda$ :

```
#include <iostream>
#include <array>
#include "pw85/pw85.hpp"
using Vec = std::array<double, pw85::dim>;
using SymMat = std::array<double, pw85::sym>;
int main() {
 Vec x1{-0.5, 0.4, -0.7};
 Vec n1{0., 0., 1.};
 double a1 = 10.;
 double c1 = 0.1;
 Vec x2{0.2, -0.3, 0.4};
 Vec n2{1., 0., 0.};
 double a2 = 0.5;
 double c2 = 5.;
 Vec r12;
  for (int i = 0; i < pw85::dim; i++) r12[i] = x2[i] - x1[i];
  SymMat q1, q2;
```
(continued from previous page)

```
pw85::spheroid(a1, c1, n1.data(), q1.data());
pw85::spheroid(a2, c2, n2.data(), q2.data());
std::array<double, 2> out;
pw85::contact_function(r12.data(), q1.data(), q2.data(), out.data());
\mathsf{std}:\mathsf{cout}\ \texttt{<<}\ \texttt{"mu} \texttt{?} =\ \texttt{"<<}\ \texttt{out}[\texttt{\theta}] \ \texttt{<<}\ \texttt{std}:\texttt{endl};std::cout << "lambda = " << out[1] << std::endl;
```
A CMakeLists.txt file is provided for the compilation of the tutorial using [CMake](https://cmake.org/). You can reuse it in one of your own projects (download):

```
cmake_minimum_required(VERSION 3.13)
project("tutorial" LANGUAGES CXX)
set(CMAKE_CXX_STANDARD 17)
add_executable(${PROJECT_NAME} ${PROJECT_NAME}.cpp)
find_library(MATH_LIBRARY m)
if (MATH_LIBRARY)
    target link libraries(${PROJECT_NAME} INTERFACE ${MATH_LIBRARY})
endif()
find_package(Boost REQUIRED)
target_link_libraries(${PROJECT_NAME} PRIVATE Boost::headers)
target_include_directories(${PROJECT_NAME} PRIVATE "../../include")
```
cd into the cpp\_tutorial subdirectory. The provided example program should be compiled and linked against pw85:

```
$ mkdir build
$ cd build
$ cmake ..
$ cmake --build . --config Release
```
An executable called tutorial should be present in the build/Release subdirectory. On execution, it prints the following lines to stdout:

```
mu^2 = 3.36271lambda = 0.166859
```
}

**FOUR**

### **THEORY**

<span id="page-18-0"></span>This chapter provides the theoretical background to the Perram–Wertheim algorithm[[PW85\]](#page-38-2). We use matrices rather than tensors: a point/vector is therefore defined through the 3×1 column-vector of its coordinates. Likewise, a secondrank tensor is represented by its 3×3 matrix.

Only the global, cartesian frame is considered here, and there is no ambiguity about the basis to which these column vectors and square matrices refer.

## <span id="page-18-1"></span>**4.1 Mathematical representation of ellipsoids**

Ellipsoids are defined from their center c and a positive-definite quadratic form Q as the set of points m such that:

```
(1) (m-c)^T \cdot Q^{-1} \cdot (m-c) \leq 1.
```
Q is a symmetric, positive-definite matrix:

```
(2) Q = \sum a_i^2 V_i \cdot V_i^T,
                    \mathbf i
```
where  $a_1$ ,  $a_2$ ,  $a_3$  are the lengths of the principal semi-axes and  $v_1$ ,  $v_2$ ,  $v_3$  their directions (unit vectors).

In the PW85 library, Q is represented as a double[6] array q which stores the upper triangular part of Q in row-major order:



## <span id="page-18-2"></span>**4.2 The contact function of two ellipsoids**

Let E<sub>1</sub> and E<sub>2</sub> be two ellipsoids, defined by their centers  $c_1$  and  $c_2$  and quadratic forms  $Q_1$  and  $Q_2$ , respectively.

<span id="page-18-3"></span>For  $0 \leq \lambda \leq 1$  and a point x, we introduce the function:

```
(4) F(x, \lambda) = \lambda (x-c_1)^{\top} \cdot Q_1^{-1} \cdot (x-c_1) + (1-\lambda) (x-c_2)^{\top} \cdot Q_2^{-1} \cdot (x-c_2).
```
Forfixed  $λ$ ,  $F(x, λ)$  has a unique minimum [[PW85\]](#page-38-2)  $f(λ)$ , and we define:

(5)  $f(\lambda) = min\{F(x, \lambda), x \in \mathbb{R}^3\}, \theta \le \lambda \le 1.$ 

<span id="page-19-7"></span>Now, the function f has a unique maximum over [0, 1], and the "contact function"  $F(r_1_2, 0_1, 0_2)$  of ellipsoids  $E_1$ and E<sub>2</sub> is defined as:

(6)  $F(r_{12}, Q_1, Q_2) = max\{f(\lambda), 0 \le \lambda \le 1\},$ 

where  $r_{12} = c_2 - c_1$ . It can be shown that

- if  $F(r_1, q_1, q_2)$  < 1 then  $E_1$  and  $E_2$  overlap,
- if  $F(r_1, q_1, q_2) = 1$  then  $E_1$  and  $E_2$  are externally tangent,
- if  $F(r_1, q_1, q_2) > 1$  then  $E_1$  and  $E_2$  do not overlap.

The contact function therefore provides a criterion to check overlap of two ellipsoids. The PW85 library computes this value.

#### <span id="page-19-0"></span>**4.3 Geometric interpretation**

<span id="page-19-3"></span>The scalar  $\lambda$  being fixed, we introduce the minimizer  $x \circ (\lambda)$  of  $F(x, \lambda)$ . The stationarity of F w.r.t to x reads:

```
(7) \nabla F(x_0(\lambda), \lambda) = 0,
```
<span id="page-19-5"></span>which leads to:

$$
(8) \qquad \lambda Q_1^{-1} \cdot [x_0(\lambda) - c_1] + (1-\lambda)Q_2^{-1} \cdot [x_0(\lambda) - c_2] = 0,
$$

<span id="page-19-2"></span>and can be rearranged:

 $(9a)$   $x_{0}(\lambda)-c_{1} = (1-\lambda)Q_{1} \cdot Q^{-1} \cdot r_{12}$ , (9b)  $x \circ (\lambda) - c_2 = -\lambda Q_2 \cdot Q^{-1} \cdot r_{12}$ ,

<span id="page-19-1"></span>with:

(10)  $Q = (1-\lambda)Q_1 + \lambda Q_2$ .

It results from the above that:

(11) 
$$
f(\lambda) = F(x_0(\lambda), \lambda) = \lambda(1-\lambda)r_{12} \cdot r_1 \cdot r_{12}
$$
.

<span id="page-19-6"></span>Maximization of  $f$  with respect to  $\lambda$  now delivers the stationarity condition:

(12) 
$$
\theta = f'(\lambda) = \nabla F(x_0(\lambda), \lambda) \cdot x_0'(\lambda) + \frac{\partial F}{\partial \lambda}.
$$

<span id="page-19-4"></span>Using Eqs. [\(4\)](#page-18-3) and [\(7\)](#page-19-3), it is found that f is minimum for  $\lambda = \lambda_0$  such that:

(13) 
$$
[x\circ(\lambda\circ)-c_1]^{\top}\cdot Q_1^{-1}\cdot [x\circ(\lambda\circ)-c_1] = [x\circ(\lambda\circ)-c_2]^{\top}\cdot Q_2^{-1}\cdot [x\circ(\lambda\circ)-c_2].
$$

Let  $\mu^2$  be this common value. It trivially results from Eqs. [\(4\)](#page-18-3) and [\(13\)](#page-19-4) that  $\mu^2 = F(x \circ (\lambda \circ)$ ,  $\lambda \circ)$ . In other words,  $\mu^2$ is the value of the contact function.

We are now in a position to give a geometric interpretation of μ. It results from Eq. *[\(13\)](#page-19-4)* and the definition of μ that:

$$
(14a) \qquad [x_{\theta}(\lambda_{\theta})-c_1]^{\top} \cdot (\mu^2 Q_1)^{-1} \cdot [x_{\theta}(\lambda_{\theta})-c_1] = 1,
$$

and:

(14b)  $[x_{\theta}(\lambda_{\theta})-c_2]^{\top} \cdot (\mu^2 Q_2)^{-1} \cdot [x_{\theta}(\lambda_{\theta})-c_2] = 1.$ 

The above equations mean that  $x \circ (\lambda \circ)$  belongs to both ellipsoids centered at c<sub>i</sub> and defined by the symmetric, positivedefinite quadratic form  $\mu^2Q_j$  (j = 1, 2). These two ellipsoids are nothing but the initial ellipsoids E<sub>1</sub> and E<sub>2</sub>, scaled by the *same* factor μ.

Furthermore, Eq. [\(8\)](#page-19-5) applies for  $\lambda = \lambda_0$ . Therefore, the normals to the scaled ellipsoids coincide at  $x_0(\lambda_0)$ : the two scaled ellipsoids are externally tangent.

To sum up, μ is the common factor by wich ellipsoids E₁ and E₂ must be scaled in order for them to be externally tangent at point  $x \circ (\lambda \circ)$ .

## **IMPLEMENTATION OF THE FUNCTION F**

<span id="page-22-2"></span><span id="page-22-0"></span>In this chapter, we explain how the contact function is computed. From Eq. *[\(12\)](#page-19-6)* in chapter *[Theory](#page-18-0)*, the value of the contact function is found from the solution  $\lambda$  to equation  $f'(\lambda) = 0$ , where it is recalled that f is defined as follows:



with:

(2)  $Q = (1-\lambda)Q_1 + \lambda Q_2$ .

In the present chapter, we discuss two implementations for the evaluation of f. The *[first implementation](#page-22-1)* uses the Cholesky decomposition of Q. The *[second implementation](#page-23-0)* uses a representation of f as a quotient of two polynomials (rational fraction).

#### <span id="page-22-1"></span>**5.1 Implementation #1: using Cholesky decompositions**

Since Q is a symmetric, positive definite matrix, we can compute its [Cholesky decomposition,](https://en.wikipedia.org/wiki/Cholesky_decomposition) which reads as follows:

 $(3)$  Q = L⋅L<sup>T</sup>,

where L is a lower-triangular matrix. Using this decomposition, it is straightforward to compute  $s = Q^{-1} \cdot r$  (where we write r as a shorthand for  $r₁₂$ ), so that:

(4)  $f(\lambda) = \lambda(1-\lambda)r^{\top} \cdot s$ .

In order to solve  $f'(\lambda) = 0$  numerically, we use a [Newton–Raphson](https://en.wikipedia.org/wiki/Newton%27s_method) procedure, which requires the first and second derivatives of f. It is readily found that:

(5)  $s' = -Q^{-1} \cdot Q' \cdot Q^{-1} \cdot r = -Q^{-1} \cdot u$  and  $r^T \cdot s' = -r^T \cdot Q^{-1} \cdot u = -s^T \cdot u$ ,

with  $u = Q_{12} \cdot s$  and  $Q_{12} = Q_2 - Q_1$ . Therefore:

```
(6) f'(\lambda) = (1-2\lambda)r^{\top} \cdot s - \lambda(1-\lambda)s^{\top} \cdot u.
```
Similarly, introducing  $v = Q^{-1} \cdot u$ :

(7)  $S^T \cdot U' = S^T \cdot Q_{12} \cdot S' = -S^T \cdot Q_{12} \cdot Q^{-1} \cdot U = -U^T \cdot V,$ 

and:

(8)  $u^T \cdot s' = -u^T \cdot Q^{-1} \cdot u = -u^T \cdot v$ .

Therefore:

(9)  $f''(\lambda) = -2r^{\dagger} \cdot s - 2(1-2\lambda)s^{\dagger} \cdot u + 2\lambda(1-\lambda)u^{\dagger} \cdot v.$ 

## <span id="page-23-0"></span>**5.2 Implementation #2: using rational functions**

<span id="page-23-1"></span>We observe that  $f(\lambda)$  is a rational function [see Eq.  $(1)$ ], and we write:



with:



where adj(Q) denotes the adjugate matrix of Q (transpose of its cofactor matrix), see e.g [Wikipedia](https://en.wikipedia.org/wiki/Adjugate_matrix).

The coefficients  $a_{\perp}$  and  $b_{\perp}$  are found from the evaluation of  $a(\lambda)$  and  $b(\lambda)$  for specific values of  $\lambda$ :



This requires the implementation of the determinant and the adjugate matrix of a  $3\times3$ , symmetric matrix, see pw85\_\_det\_sym() and pw85\_\_xT\_adjA\_x().

Evaluating the derivative of  $f$  with respect to  $\lambda$  is fairly easy. The following [Sympy](https://www.sympy.org) script will do the job:

```
import sympy
from sympy import Equality, numer, pprint, Symbol
if name == '_main ':
   sympy.init_printing(use_latex=False, use_unicode=True)
```
(continued from previous page)

```
λ = Symbol('λ')
a = sum(sympy.Symbol('a{}'.format(i))*\lambda^{**}i for i in range(3))
b = sum(sympy.Symbol('b{}'.format(i))*\lambda**i for i in range(4))
f = \lambda * (1-\lambda) * a/bf_prime = f.diff(\lambda).ratsimp()
c = number(f_prime)c_dict = c.collect(λ, evaluate=False)
for i in range(sympy.degree(c, gen=λ)+1):
    pprint(Equality(Symbol('c{}'.format(i)), c_dict[λ**i]))
```
It is readily found that:

c(λ) (13)  $f'(\lambda) = \cdot$ b( $\lambda$ )<sup>2</sup>

where  $c(\lambda)$  is a sixth-order polynomial in  $\lambda$ :

(14)  $C(\lambda) = C_0 + C_1\lambda + C_2\lambda^2 + C_3\lambda^3 + C_4\lambda^4 + C_5\lambda^5 + C_6\lambda^6$ ,

with:

```
(15a) C<sub>0</sub> = a<sub>0</sub>b<sub>0</sub>,
(15b) C_1 = 2(a_1 - a_0)b_0,
(15c) c_2 = -a_0(b_1+b_2) + 3b_0(a_2-a_1) + a_1b_1,
(15d) c_3 = 2[b_1(a_2-a_1) - a_0b_3] - 4a_2b_0,
(15e) c_4 = (a_0 - a_1)b_3 + (a_2 - a_1)b_2 - 3a_2b_1(15f) C_5 = -2a_2b_2,
(15g) C_6 = -a_2b_3
```
Solving  $f'(\lambda) = 0$  for  $\lambda$  is therefore equivalent to finding the unique root of c in the interval  $0 \le \lambda \le 1$ . For the sake of robustness, the [bisection method](https://en.wikipedia.org/wiki/Bisection_method) has been implemented (more efficient methods will be implemented in future versions).

Once  $\lambda$  is found, μ is computed from  $\mu^2 = f(\lambda)$  using Eq. *[\(10\)](#page-23-1)*.

## <span id="page-24-0"></span>**5.3 Comparison of the two implementations**

High precision reference data was generated using the [mpmath](http://mpmath.org/) library. The reference dataset is fully described and freely downloadable from the [Zenodo platform](http://about.zenodo.org/) [\(DOI:10.5281/zenodo.3323683](https://doi.org/10.5281/zenodo.3323683)). Accuracy of both implementations is then evaluated through the following script (download source file):

```
#include <glib.h>
#include <hdf5.h>
#include <hdf5_hl.h>
#include <pw85.h>
#include <pw85_legacy.h>
void read_dataset_double(hid_t const hid, char const *dset_name, size_t *size,
                         double **buffer) {
  int ndims;
 H5LTget_dataset_ndims(hid, dset_name, &ndims);
  hsize t *dim = g new(hsize t, ndims);
 H5LTget dataset info(hid, dset name, dim, NULL, NULL);
```
(continued from previous page)

```
*size = 1;for (size_t i = 0; i < ndims; i++) {
   *size *= dim[i];
  }
  *buffer = g_new(double, *size);
 H5LTread_dataset_double(hid, dset_name, *buffer);
  g_free(dim);
}
void update_histogram(double act, double exp, size_t num_bins, size_t *hist) {
 double const err = fabs((act - exp) / exp);
  int prec;
 if (err == 0.0) {
   prec = num bins - 1;} else {
   prec = (int)(floor(-log10(err)));
   if (prec \leq 0) {
      prec = 0;}
   if (prec >= num bins) {
      prec = num bins - 1;}
  }
  ++hist[prec];
}
int main() {
 hid_t const hid = H5Fopen(PW85_REF_DATA_PATH, H5F_ACC_RDONLY, H5P_DEFAULT);
 size_t num_directions;
  double *directions;
  read_dataset_double(hid, "/directions", &num_directions, &directions);
  num directions /= PW85 DIM;
  size_t num_lambdas;
  double *lambdas;
  read_dataset_double(hid, "/lambdas", &num_lambdas, &lambdas);
  size_t num_radii;
  double *radii;
  read_dataset_double(hid, "/radii", &num_radii, &radii);
  size_t num_spheroids;
  double *spheroids;
  read_dataset_double(hid, "/spheroids", &num_spheroids, &spheroids);
  num spheroids /= PW85 SYM;
  size t num expecteds;
  double *expecteds;
  read_dataset_double(hid, "/F", &num_expecteds, &expecteds);
  double *exp = expecteds;
  double params[2 * PW85_SYM + PW85_DIM];
  size t num bins = 16;
  size_t hist1[num_bins];
  size_t hist2[num_bins];
```

```
(continued from previous page)
```

```
for (size_t i = 0; i < num_bins; i++) {
  hist1[i] = 0;
  hist2[i] = 0;
}
for (size_t i1 = 0; i1 < num_spheroids; i1++)memory(params + PW85_DIM, spheroids + PW85_SYM * i1,PW85_SYM * sizeof(double));
  for (size_t i2 = 0; i2 < num spheroids; i2++) {
    memcpy(params + PW85_DIM + PW85_SYM, spheroids + PW85_SYM * i2,
           PW85_SYM * sizeof(double));
    for (size t i = 0; i < num directions; i++) {
      memcpy(params, directions + PW85_DIM * i, PW85_DIM * sizeof(double));
      for (size t j = 0; j < num lambdas; j++, exp++) {
        double const act1 = -pw85_f_neg(lambdas[j], params);
        update_histogram(act1, *exp, num_bins, hist1);
        double out[2];
        pw85_legacy_f2(lambdas[j], params, params + PW85_DIM,
                       params + PW85_DIM + PW85_SYM, out);
        double const act2 = out[0];
        update_histogram(act2, *exp, num_bins, hist2);
      }
    }
  }
}
FILE *f = fopen(HISTOGRAM_PATH, "w");
for (size t i = 0; i < num bins; i++) {
  fprintf(f, "%d,%g,%g\n", (int)i,
          100. * ((double)hist1[i]) / ((double)num_expecteds),
          100. * ((double)hist2[i]) / ((double)num_expecteds));
}
fclose(f);
g_free(spheroids);
g_free(radii);
g_free(lambdas);
g_free(directions);
H5Fclose(hid);
return 0;
```
Note: To compute this program, you might need to pass the options -Dpw85 include=…, -Dpw85 lib=… and -Dpw85\_data=… to meson.

**Note:** The provided script refers to an old implementation of pw85.

We get the histograms shown in [Fig. 5.1](#page-27-0). These histograms show that *[Implementation #1](#page-22-1)* is more accurate than *[Imple](#page-23-0)[mentation #2](#page-23-0)*. The former will therefore be selected as default.

}

<span id="page-27-0"></span>

Fig. 5.1: Accuracy of the two implementations.

#### **OPTIMIZATION OF THE FUNCTION F**

<span id="page-28-0"></span>It was shown in chapter *[Theory](#page-18-0)* [see Eq. [\(6\)](#page-19-7)] that the contact function was defined as the maximum for  $0 \le \lambda \le 1$  of the function f discussed in chapter *[Implementation of the function f](#page-22-0)*.

Given that the first and second derivatives of f can be computed explicitely (see section *[Implementation #1: using](#page-22-1) [Cholesky decompositions](#page-22-1)*in chapter *[Implementation of the function f](#page-22-0)*) it would be tempting to use the Newton–Raphson method to solve f'(λ) iteratively. However, our experiments show that this method performs very poorly in the present case, because the variations of f can be quite sharp in the neighborhood of  $\lambda = 0$  or  $\lambda = 1$ . To carry out the otpimization of f, we therefore proceed in two steps.

In the first step, we use a robust optimization algorithm. We selected here [Brent's method,](https://en.wikipedia.org/wiki/Brent%27s_method) as implemented in the [Boost::Math library.](https://www.boost.org/doc/libs/1_75_0/libs/math/) However, this method delivers a relatively low accuracy of the maxmimizer and the maximum.

Therefore, in the second step, we use a few Newton–Raphson iterations to refine the previously obtained estimates of the minimizer and minimum of f. In the remainder of this chapter, we describe how these Newton–Raphson iterations are performed.

<span id="page-28-1"></span>Our starting point is Eqs. *[\(9\)](#page-19-2)* and *[\(13\)](#page-19-4)* in chapter *[Theory](#page-18-0)*, from which it results that for a given value of λ we can define two values of μ²: one is provided by Eq. *[\(9a\)](#page-19-2)*, the other one is given by Eq. *[\(9b\)](#page-19-2)* (both in chapter *[Theory](#page-18-0)*):

(1a)  $\mu_1^2 = [x \circ (\lambda_0) - c_1]^T \cdot Q_1^{-1} \cdot [x \circ (\lambda_0) - c_1] = (1 - \lambda)^2 S^T \cdot Q_1 \cdot S,$ (1b)  $\mu_2^2 = [x_\theta(\lambda_\theta) - c_2]^\top \cdot Q_2^{-1} \cdot [x_\theta(\lambda_\theta) - c_2] = \lambda^2 s^\top \cdot Q_2 \cdot s,$ 

<span id="page-28-2"></span>where we have introduced  $s = Q^{-1} \cdot r_{12}$ . We further define the matrix  $Q_{12} = Q_2 - Q_1$ , so that:

(2) 
$$
Q_1 = Q - \lambda Q_{12}
$$
 and  $Q_2 = Q + (1-\lambda)Q_{12}$ .

<span id="page-28-5"></span>Combining Eqs. [\(1\)](#page-28-1) and [\(2\)](#page-28-2) and recalling that  $Q \cdot s = r$  then delivers the following expressions:

```
(3a) \mu_1^2 = (1-\lambda)^2 r^{\dagger} \cdot s - \lambda (1-\lambda)^2 s^{\dagger} \cdot u,
(3b) \mu_2^2 = \lambda^2 r^\top \cdot s + \lambda^2 (1-\lambda) s^\top \cdot u,
```
<span id="page-28-3"></span>where we have introduced  $u = 0<sub>12</sub> · s$ .

The above expressions seem to behave slightly better from a numerical point of view. Our problem is now to find  $\lambda$ such that  $\mu_1^2 = \mu_2^2$ . We therefore define the following residual:

(4) 
$$
g(\lambda) = \mu_2^2 - \mu_1^2 = (2\lambda - 1)r^T \cdot s + \lambda(1 - \lambda)s^T \cdot u
$$
,

<span id="page-28-4"></span>and we need to find  $\lambda$  such that  $g(\lambda) = 0$ . In order to implement Newton–Raphson iterations, we need the expression of the derivative of the residual. Using results that are presented in section *[Implementation #1: using Cholesky](#page-22-1) [decompositions](#page-22-1)*, we readily find that:

(5) 
$$
g'(\lambda) = 2r^{\dagger} \cdot s + 2(1-2\lambda)s^{\dagger} \cdot u - 2\lambda(1-\lambda)u^{\dagger} \cdot v
$$
.

Eqs. [\(4\)](#page-28-3) and [\(5\)](#page-28-4) are then used for the final, refinement step of determination of  $\lambda$ .

**SEVEN**

# <span id="page-30-0"></span>**TESTING THE IMPLEMENTATION OF THE CONTACT FUNCTION**

This chapter describes how our implementation of the contact function is tested. The source of the unit tests can be found in the file src/test pw85.c. Note that the tests described here are repeated over a large set of tests case, including very flat and very slender sheroids, for various relative orientations and center-to-center distances.

In the present chapter, we assume that the two ellipsoids (their matrices  $Q_1$  and  $Q_2$  are given), as well as their centerto-center radius vector  $r₁₂$ . Then, a call to  $pw85$ : : contact\_function() delivers an estimate of  $\lambda$  and  $\mu²$ .

We first assert that μ₁² and μ₂² as defined by Eq. *[\(3\)](#page-28-5)* in chapter *[Optimization of the function f](#page-28-0)* are close to the value returned by  $pw85::contact\_function()$ . For all the cases considered here, this is true up to a relative error of  $10^{-1}$ <sup>o</sup>.

We also check that  $f'(\lambda) = 0$ , up to an absolute error of  $Δλf''(λ)$  where  $Δλ$  is the absolute tolerance on  $λ$  for the stopping criterion of the Brent iterations, as defined by the constant *[pw85::lambda\\_atol](#page-34-1)*.

**EIGHT**

# **THE C++ API**

<span id="page-32-0"></span>**Note:** functions whose name is prefixed with an underscore should be considered as "private": these functions are exposed for testing purposes. They should not be used, since they are susceptible of incompatible changes (or even removal) in future versions.

## <span id="page-32-1"></span>**8.1 Representation of vectors and matrices**

An ellipsoid is defined from its center c (a  $3\times1$ , column-vector) and quadratic form Q (a  $3\times3$ , symmetric, positive definite matrix) as the set of points m such that:

 $(m-c)^T \cdot Q^{-1} \cdot (m-c) \leq 1.$ 

In this module, objects referred to as "vectors" are double[3] arrays of coordinates. In other words, the representation of the vector  $x$  is the double [3] array  $x$  such that:

 $\lceil x[0] \rceil$  $x = | x[1] |$ .  $\lfloor x[2] \rfloor$ 

Objects referred to as "symmetric matrices" (or "quadratic forms") are of type double[6]. Such arrays list in rowmajor order the coefficients of the triangular upper part. In other words, the representation of a the symmetric matrix A is the double[6] array a such that:



## <span id="page-32-2"></span>**8.2 API**

namespace **pw85**

#### **Functions**

<span id="page-33-0"></span>void **cholesky** decomp (double const \*a, double \*l)

Compute the Cholesky decomposition of a symmetric, positive matrix.

Let A be a symmetric, positive matrix, defined by the double [6] array a. This function computes the lower-triangular matrix L, defined by the double[6] array l, such that  $L^\top \cdot L = A$ .

The array l must be pre-allocated; it is modified by this function. Note that storage of the coefficients of L is as follows



void **cholesky** solve(const double \*l, const double \*b, double \*x)

Compute the solution to a previously Cholesky decomposed linear system.

Let L be a lower-triangular matrix, defined by the double[6] array l (see *[pw85::\\_cholesky\\_decomp\(\)](#page-33-0)* for ordering of the coefficients). This function solves (by substitution) the linear system  $L^\top \cdot L \cdot x = b$ , where the vectors x and b are specified through their double[3] array of coordinates; x is modified by this function.

void **spheroid**(double a, double c, const double \*n, double \*q) Compute the quadratic form associated to a spheroid.

The spheroid is defined by its equatorial radius a, its polar radius c and the direction of its axis of revolution, n (unit-vector, double[3] array).

q is the representation of a symmetric matrix as a double[6] array. It is modified in-place.

double **f\_neg**(double lambda, const double \*r12, const double \*q1, const double \*q2) Return the value of the opposite of the function f defined as (see *[Theory](#page-18-0)*).

$$
f(\lambda) = \lambda \left(1 - \lambda\right) r_{12}^{\mathsf{T}} \cdot Q^{-1} \cdot r_{12},
$$

with

$$
Q = (1 - \lambda)Q_1 + \lambda Q_2,
$$

where ellipsoids 1 and 2 are defined as the sets of points *m* (column-vector) such that

$$
(m-c_i)\cdot Q_i^{-1}\cdot (m-c_i)\leq 1.
$$

In the above inequality,  $c_i$  is the center;  $r_{12} = c_2 - c_1$  is the center-to-center radius-vector, represented by the double<sup>[3]</sup> array r12. The symmetric, positive-definite matrices  $Q_1$  and  $Q_2$  are specified through the double[6] arrays q1 and q2.

The value of  $\lambda$  is specified through the parameter lambda.

This function returns the value of  $-f(\lambda)$  (the "minus" sign comes from the fact that we seek the maximum of *f*, or the minimum of −*f*).

This implementation uses *[Cholesky decompositions\\*](#page-22-1)*.

void **residual** (double lambda, const double \*r12, const double \*q1, const double \*q2, double \*out) Compute the residual  $g(\lambda) = \mu_2^2 - \mu_1^2$ .

See *[Optimization of the function f](#page-28-0)* for the definition of *g*. The value of  $\lambda$  is specified through the parameter lambda. See *[contact\\_function\(\)](#page-34-2)* for the definition of the parameters r12, q1 and q2.

The preallocated double<sup>[3]</sup> array out is updated as follows: out<sup>[0]</sup> =  $f(\lambda)$ , out<sup>[1]</sup> =  $g(\lambda)$  and out<sup>[2]</sup> =  $g'(\lambda)$ .

This function is used in function *[pw85::contact\\_function\(\)](#page-34-0)* for the final Newton–Raphson refinement step.

<span id="page-34-2"></span><span id="page-34-0"></span>int **contact\_function**(const double \*r12, const double \*q1, const double \*q2, double \*out) Compute the value of the contact function of two ellipsoids.

The center-to-center radius-vector  $r_{12}$  is specified by the double[3] array r12. The symmetric, positivedefinite matrices *Q*<sup>1</sup> and *Q*<sup>2</sup> that define the two ellipsoids are specified through the double[6] arrays q1 and q2.

This function computes the value of  $\mu^2$ , defined as

$$
\mu^{2} = \max_{0 \leq \lambda \leq 1} \left\{ \lambda (1 - \lambda) r_{12}^{T} \cdot \left[ (1 - \lambda) Q_{1} + \lambda Q_{2} \right]^{-1} \cdot r_{12} \right\},\
$$

and the maximizer *λ*, see *[Theory](#page-18-0)* . Both values are stored in the preallocated double[2] array out: out[0]  $=\mu^2$  and out [1]  $=\lambda$ .

 $\mu$  is the common factor by which the two ellipsoids must be scaled (their centers being fixed) in order to be tangentially in contact.

This function returns 0.

**Todo:** This function should return an error code.

#### **Variables**

```
constexpr size t \dim = 3The dimension of the physical space (3).
```
constexpr size\_t  $sym = 6$ The dimension of the space of symmetric matrices (6).

<span id="page-34-1"></span>constexpr double **lambda\_atol** = 1e-6

The absolute tolerance for the stopping criterion of Brent's method (in function *[contact\\_function\(\)](#page-34-2)*).

constexpr size\_t **max\_iter** = 25

The maximum number of iterations of Brent's method (in function *[contact\\_function\(\)](#page-34-2)*).

constexpr size\_t **nr\_iter** = 3

The total number of iterations of the Newton–Raphson refinement phase (in function *[contact\\_function\(\)](#page-34-2)*).

namespace **metadata**

#### **Variables**

constexpr std::string\_view **author** = {"S. Brisard"}

constexpr std::string\_view **description** = {"Implementation of the \"contact function\" defined by Perram and Wertheim (J. Comp. Phys. 58(3), 409-416, DOI:10.1016/0021-9991(85)90171-8) for two ellipsoids."}

constexpr std::string\_view **author\_email** = {"sebastien.brisard@univ-eiffel.fr"}

constexpr std::string\_view **license** = {"BSD 3-Clause License"}

constexpr std::string\_view **name** = {"pw85"}

constexpr std::string\_view **url** = {"https://github.com/sbrisard/pw85"}

constexpr std::string\_view **version** = {"2.0"}

constexpr std::string\_view **year** = {"2021"}

# **THE PYTHON API**

<span id="page-36-1"></span><span id="page-36-0"></span>pypw85.**\_cholesky\_decomp**(*a: numpy.ndarray[numpy.float64]*, *l: numpy.ndarray[numpy.float64]*) *→* None Compute the Cholesky decomposition of a symmetric, positive matrix.

Let A be a symmetric, positive matrix, defined by the double[6] array a. This function computes the lowertriangular matrix L, defined by the double[6] array l, such that  $L^{\top} \cdot L = A$ .

The array l must be pre-allocated; it is modified by this function. Note that storage of the coefficients of L is as follows:

 $l[0]$  0 0  $]$  $L = | U[1] U[3] 0 |.$  $[l[2] l[4] l[5] ]$ 

This function is exposed for testing purposes only.

pypw85.**\_cholesky\_solve**(*l: numpy.ndarray[numpy.float64]*, *b: numpy.ndarray[numpy.float64]*, *x:*

*numpy.ndarray[numpy.float64]*) *→* None

Compute the solution to a previously Cholesky decomposed linear system.

Let L be a lower-triangular matrix, defined by the double<sup>[6]</sup> array l (see *cholesky decomp()* for ordering of the coefficients). This function solves (by substitution) the linear system L<sup>T</sup>⋅L⋅x = b, where the vectors x and b are specified through their double[3] array of coordinates; x is modified by this function.

This function is exposed for testing purposes only.

pypw85.**contact\_function**(*r12: numpy.ndarray[numpy.float64]*, *q1: numpy.ndarray[numpy.float64]*, *q2: numpy.ndarray[numpy.float64]*, *out: numpy.ndarray[numpy.float64]*) *→* int

Compute the value of the contact function of two ellipsoids.

See *[f\\_neg\(\)](#page-36-2)* for the meaning of the parameters *r12*, *q1* and *q2*.

This function computes the value of  $\mu^2$ , defined as:

 $\mu^2$  = max{ $\lambda(1-\lambda)r_{12}$ <sup>T</sup> · [(1- $\lambda)$ Q<sub>1</sub> +  $\lambda$ Q<sub>2</sub>]<sup>-1</sup> · r<sub>12</sub>,  $0 \le \lambda \le 1$ }

and the maximizer λ, see *[Theory](#page-18-0)*. Both values are stored in the preallocated double[2] array out: out[0] =  $μ<sup>2</sup>$  and out [1] =  $λ$ .

μ is the common factor by which the two ellipsoids must be scaled (their centers being fixed) in order to be tangentially in contact.

This function returns 0.

<span id="page-36-2"></span>**Todo:** This function should return an error code.

pypw85.**f\_neg**(*lambda: float, r12: numpy.ndarray[numpy.float64], q1: numpy.ndarray[numpy.float64], q2: numpy.ndarray[numpy.float64]*) *→* float

Return the value of the opposite of the function f defined as (see *[Theory](#page-18-0)*):

 $f(\lambda) = \lambda(1-\lambda)r_{12}$ <sup>T</sup> $\cdot$ Q<sup>-1</sup> $\cdot$ r<sub>12</sub>,

with:

 $Q = (1-\lambda)Q_1 + \lambda Q_2$ ,

where ellipsoids 1 and 2 are defined as the sets of points m (column-vector) such that:

 $(m-c_i)^\top \cdot Q_i^{-1} \cdot (m-c_i) \leq 1.$ 

In the above inequality,  $c_1$  is the center;  $r_1$ ,  $c_2$  = c<sub>2</sub> - c<sub>1</sub> is the center-to-center radius-vector, represented by the double[3] array r12. The symmetric, positive-definite matrices  $Q_1$  and  $Q_2$  are specified through the double[6] arrays q1 and q2.

The value of  $\lambda$  is specified through the parameter lambda.

This function returns the value of  $-f(\lambda)$  (the "minus" sign comes from the fact that we seek the maximum of f, or the minimum of (-f).

This implementation uses *[Cholesky decompositions](#page-22-1)*.

pypw85.**spheroid**(*a: float*, *c: float*, *n: numpy.ndarray[numpy.float64]*, *q: numpy.ndarray[numpy.float64]*) *→* None Compute the quadratic form associated to a spheroid.

The spheroid is defined by its equatorial radius a, its polar radius c and the direction of its axis of revolution, n (unit-vector, double[3] array).

q is the representation of a symmetric matrix as a double[6] array. It is modified in-place.

#### **BIBLIOGRAPHY**

- <span id="page-38-5"></span><span id="page-38-0"></span>[ABH18] Anoukou, K., Brenner, R., Hong, F., Pellerin, M., & Danas, K. (2018). Random distribution of polydisperse ellipsoidal inclusions and homogenization estimates for porous elastic materials. Computers & Structures, 210, 87–101. <https://doi.org/10.1016/j.compstruc.2018.08.006>
- <span id="page-38-8"></span>[BL13] Brisard, S., & Levitz, P. (2013). Small-angle scattering of dense, polydisperse granular porous media: Computation free of size effects. Physical Review E, 87(1), 013305. [https://doi.org/10.1103/PhysRevE.](https://doi.org/10.1103/PhysRevE.87.013305) [87.013305](https://doi.org/10.1103/PhysRevE.87.013305)
- <span id="page-38-4"></span>[CYP07] Chen, X.-D., Yong, J.-H., Paul, J.-C., & Sun, J. (2007). Intersection Testing between an Ellipsoid and an Algebraic Surface. 2007 10th IEEE International Conference on Computer-Aided Design and Computer Graphics, 43–46. <https://doi.org/10.1109/CADCG.2007.4407853>
- <span id="page-38-6"></span>[DTS05] Donev, A., Torquato, S., & Stillinger, F. H. (2005). Neighbor list collision-driven molecular dynamics simulation for nonspherical hard particles. I. Algorithmic details. Journal of Computational Physics, 202(2), 737–764. <https://doi.org/10.1016/j.jcp.2004.08.014>
- <span id="page-38-7"></span>[DTS05a] Donev, A., Torquato, S., & Stillinger, F. H. (2005). Neighbor list collision-driven molecular dynamics simulation for nonspherical hard particles.: II. Applications to ellipses and ellipsoids. Journal of Computational Physics, 202(2), 765–793. <https://doi.org/10.1016/j.jcp.2004.08.025>
- <span id="page-38-2"></span>[PW85] Perram, J. W., & Wertheim, M. S. (1985). Statistical mechanics of hard ellipsoids. I. Overlap algorithm and the contact function. *Journal of Computational Physics*, 58(3), 409–416. [https://doi.org/10.1016/](https://doi.org/10.1016/0021-9991(85)90171-8) [0021-9991\(85\)90171-8](https://doi.org/10.1016/0021-9991(85)90171-8)
- <span id="page-38-1"></span>[VB72] Vieillard‐Baron, J. (1972). Phase Transitions of the Classical Hard‐Ellipse System. The Journal of Chemical Physics, 56(10), 4729–4744. <https://doi.org/doi:10.1063/1.1676946>
- <span id="page-38-3"></span>[WWK01] Wang, W., Wang, J., & Kim, M.-S. (2001). An algebraic condition for the separation of two ellipsoids. Computer Aided Geometric Design, 18(6), 531–539. [https://doi.org/10.1016/S0167-8396\(01\)00049-8](https://doi.org/10.1016/S0167-8396(01)00049-8)

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