BornAgain - simulating and fitting X-ray and neutron small angle scattering at grazing incidence.

User Guide
version 0.1

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August 28, 2013
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Introduction

BornAgain is a software to simulate and fit small-angle scattering at grazing incidence (GISAS). It supports analysis of both X-ray (GISAXS) and neutron (GISANS) data. The name of the software, BornAgain, indicates the central role of the distorted-wave Born approximation (DWBA) in the physical description of the scattering process. The software provides a generic framework for modeling multilayer samples with smooth or rough interfaces and with various types of embedded nanoparticles. In this way, it reproduces and enhances the functionality of the present reference software, IsGISAXS by R. Lazzari [1], and lays a solid base for future extensions in response to specific user needs.

To meet the growing demand for GISAS simulation of more complex structured materials, BornAgain has extended the IsGISAXS program’s functionality by removing the restrictions on the number of layers and particles, by providing diffuse reflection from rough layer interfaces and by adding particles with inner structures.

BornAgain is a platform independent software, with active support for Linux, MacOS and Microsoft Windows (planned for October, 2013). It is a free and open source software provided under terms of GNU General Public License (GPL). The authors will be grateful for all kind of feedback: criticism, praise, bug reports, feature requests or contributed modules. When BornAgain is used in preparing scientific papers, please cite this manual as follows:

C. Durniak, G. Pospelov, W. Van Herck, J. Wuttke,
BornAgain - simulating and fitting X-ray and neutron small angle scattering at grazing incidence, http://apps.jcns.fz-juelich.de/BornAgain

This user guide starts with a brief description of the steps necessary for compiling the source code and running the simulation in Section 1.1. More detailed overview of software architecture and installation procedure are given in Section 1.2 and Section 1.3. General methodology of simulation with BornAgain and detailed usage examples are given in Chapter 2.

Icons used in this manual:

manda: this sign highlights further remarks.

manda: this sign highlights essential points.
Chapter 1

Installation

1.1 Quick start

This section shortly describes how to build BornAgain from source and run the first simulation. More details about software architecture and installation procedure are given in Section 1.2 and Section 1.3.

Step I: installing third party software

- compilers: clang versions $\geq 3.1$ or GCC versions $\geq 4.2$
- cmake ($\geq 2.8$)
- boost library ($\geq 1.48$)
- GNU scientific library ($\geq 1.15$)
- fftw3 library ($\geq 3.3.1$)
- python-2.7, python-devel, python-numpy-devel

Step II: getting the source

```
git clone git://apps.jcns.fz-juelich.de/BornAgain.git
```

Step III: building the source

```
mkdir <build_dir>; cd <build_dir>;
cmake <source_dir> -DCMAKE_INSTALL_PREFIX=<install_dir>
make
clean
make install
```

Step IV: running example

```
<install_dir>/Examples/python/ex001_CylindersAndPrisms
python CylindersAndPrisms.py
```
1.2 Software architecture

Figure 1.1: Structure of BornAgain libraries.
1.3 Installation

This section describes how to build and install BornAgain libraries from the source. At the moment we support building on x86/x86_64 Linux and Mac OS X operating systems. Support for Windows systems is planned in next releases. There are three major steps to building BornAgain:

1. Acquire required third-party libraries.
2. Get BornAgain source code.
3. Use cmake to build and install software.

The remainder of this section explains each step in detail.

1.3.1 Third-party software.

To successfully build BornAgain a number of prerequisite packages must be installed.

- compilers: clang versions \( \geq 3.1 \) or GCC versions \( \geq 4.2 \)
- cmake \((\geq 2.8)\)
- boost library \((\geq 1.48)\)
- GNU scientific library \((\geq 1.15)\)
- fftw3 library \((\geq 3.3)\)
- python \((\geq 2.7, < 3.0), \) python-devel, python-numpy-devel

Other packages are optional

- ROOT framework (adds several additional fitting algorithms to BornAgain)
- python-matplotlib (allows to run usage examples with graphics)

All required packages can be easily installed on most Linux distributions using the system’s package manager. Below we give a few examples for several selected operation systems. Please note, that other distributions (Fedora, Mint, etc) may have different commands for invoking the package manager and slightly different names of packages (like “boost” instead of “libboost” etc). Besides that, the installation should be very similar.

**OpenSuse 12.3**

Adding “scientific” repository

```
sudo zypper ar http://download.opensuse.org/repositories/science/openSUSE_12.3 science
```

Installing required packages

```
sudo zypper install git-core cmake gsl-devel boost-devel fftw3-devel python-devel python-numpy-devel
```

Installing optional packages

```
sudo zypper install libroot-* root-plugin-* root-system-* root-ttf libeigen3-devel python-matplotlib
```
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**Ubuntu 12.10, 13.04**

Installing required packages

```
sudo apt-get install git cmake libgsl0-dev libboost-all-dev libfftw3-dev python-dev python-numpy
```

Installing optional packages

```
sudo apt-get install libroot-* root-plugin-* root-system-* ttf-root-installer libeigen3-dev python-matplotlib python-matplotlib-tk
```

**Mac OS X 10.8**

To simplify the installation of third party open-source software on a Mac OS X system we recommend the use of MacPorts package manager. The easiest way to install MacPorts is by downloading the dmg from [www.macports.org/install.php](http://www.macports.org/install.php) and running the system's installer. After the installation new command "port" will be available in terminal window of your Mac.

Installing required packages

```
sudo port -v selfupdate
sudo port install git-core cmake
sudo port install fftw-3 gsl
sudo port install boost -no_single -no_static +python27
```

Installing optional packages

```
sudo port install py27-matplotlib py27-numpy py27-scipy
sudo port install root +fftw3+python27
sudo port install eigen3
```

### 1.3.2 Getting source code

BornAgain source can be downloaded at [http://apps.jcns.fz-juelich.de/BornAgain](http://apps.jcns.fz-juelich.de/BornAgain) and unpacked with

```
tar xzf bornagain-<version>.tgz
```

Alternatively one can obtain BornAgain source from our public Git repository.

```
git clone git://apps.jcns.fz-juelich.de/BornAgain.git
```

**More about Git**

Our Git repository holds two main branches called “master” and “develop”. We consider “master” branch to be the main branch where the source code of HEAD always reflects latest stable release. `git clone` command shown above

1. gives you a source code snapshot corresponding to the latest stable release,

2. automatically sets up your local master branch to track our remote master branch, so you will be able to fetch changes from the remote branch at any time using “git pull” command.

Master branch is updating approximately once per month. The second branch, “develop” branch, is a snapshot of the current development. This is where any automatic nightly builds are built from. The develop branch is always expected to work, so to get the most recent features one can switch source code to it by
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cd BornAgain
git checkout develop
git pull

1.3.3 Building and installing the code

BornAgain should be build using CMake cross platform build system. Having third-party libraries installed on the system and BornAgain source code acquired as was explained in previous sections, type build commands

```
mkdir <build_dir>
cd <build_dir>
cmake <source_dir> -DCMAKE_INSTALL_PREFIX=<install_dir>
make
```

Here `<source_dir>` is the name of directory, where BornAgain source code has been copied, `<install_dir>` is the directory, where user wants the package to be installed, and `<build_dir>` is the directory where building will occur.

**About CMake**

Having dedicated directory `<build_dir>` for build process is recommended by CMake. That allows several builds with different compilers/options from the same source and keeps source directory clean from build remnants.

Compilation process invoked by the command “make” lasts about 10 min for an average laptop of 2012 edition. On multi-core machines the compilation time can be decreased by invoking command “make” with the parameter “make -j[N]”, where N is the number of cores.

Running functional tests is an optional but recommended step. Command “make check” will compile several additional tests and run them one by one. Every test contains the simulation of a typical GISAS geometry and the comparison on numerical level of simulation results with reference files. Having 100% tests passed ensures that your local installation is correct.

```
make check
...
100% tests passed, 0 tests failed out of 26
Total Test time (real) = 89.19 sec
[100%] Build target check
```

The last command “make install” copies compiled libraries and some usage examples into the installation directory.

```
make install
```

**Troubleshooting**

In the case of complex system setup, with variety of libraries of different versions scattered across multiple places (/opt/local, /usr etc.), you may want to help CMake to find libraries in proper place. In example below two system variables are defined to force CMake to prefer libraries found in /opt/local to other places.
export CMAKE_LIBRARY_PATH=/opt/local/lib:$CMAKE_LIBRARY_PATH
export CMAKE_INCLUDE_PATH=/opt/local/include:$CMAKE_INCLUDE_PATH

If compilation fails for some reason, please submit your bug report including compilation errors at http://apps.jcns.fz-juelich.de/redmine/projects/bornagain/issues

1.3.4 What is next?

In your installation directory you will find

```
./include - header files for compilation of your C++ program
./lib - libraries to import into python or link with your C++ program
./Examples - directory with examples
```

Run your first example and enjoy first BornAgain simulation plot.

```
cd <install_dir>/Examples/python/ex001_CylindersAndPrisms
python CylindersAndPrisms.py
```
Chapter 2

Examples

2.1 General methodology

A simulation of GISAXS using BornAgain platform can be decomposed into the following points:

- definition of the materials by specifying their names and their refractive indices,
- definition of particles: shapes, sizes, constituting materials, interference functions,
- definition of the layers: thicknesses, roughnesses, associations with the previously defined materials,
- inclusion of the particles in layers: density, positions, orientations,
- assembling the sample: generation of a multilayered system,
- specifying the input beam and the detector's characteristics,
- running the simulation,
- saving the data.

The sample is built from object oriented building blocks instead of loading data files.

2.2 Conventions

2.2.1 Geometry of the sample

The geometry used to describe the sample is shown in figure 2.1. The \( z \)-axis is perpendicular to the sample's surface and pointing upwards. The \( x \)-axis is perpendicular to the plane of the detector and the \( y \)-axis is along it. The input and the scattered output beams are each characterized by two angles \( \alpha_i, \phi_i \) and \( \alpha_f, \phi_f \) respectively. Our choice of orientation for the angles \( \alpha_i \) and \( \alpha_f \) is so that they are positive as shown in figure 2.1.

The layers are defined by their thicknesses (parallel to the \( z \)-direction), their possible roughnesses (equal to 0 by default) and the material they are made of. We do not define any dimensions in the \( x, y \) directions. And, except for roughness, the layer's vertical boundaries are plane and perpendicular to the \( z \)-axis. There is also no limitation to the number of layers that could be defined in BornAgain.
Note that the thickness of the top and bottom layer are not defined.

**Remark:** Order of the different steps for the simulation:

When assembling the sample, the layers are defined from top to bottom. So in most cases the first layer will be the air layer.

The particles are characterized by their form factors (i.e. the Fourier transform of the shape function - see the list of form factors implemented in BornAgain) and the composing material. The number of input parameters for the form factor depends on the particle symmetry; it ranges from one parameter for a sphere (its radius) to three for an ellipsoid (its three main axis lengths).

By placing the particles inside or on top of a layer, we impose their vertical positions, whose values corresponds to the bottoms of the particles. The in-plane distribution of particles is linked with the way the particles interfere with each other, which is therefore implemented when dealing with the interference function.

The complex refractive index associated with a layer or a particle is written as \( n = 1 - \delta + i\beta \), with \( \delta, \beta \in \mathbb{R} \). In our program, we input \( \delta \) and \( \beta \) directly.

![Figure 2.1: Representation of the scattering geometry.](image)

\( n_j \) is the refractive index of layer \( j \) and \( \alpha_i \) and \( \phi_i \) are the incident angle of the wave propagating. \( \alpha_f \) is the exit angle with respect to the sample's surface and \( \phi_f \) is the scattering angle with respect to the scattering plane.

The input beam is assumed to be monochromatic without any spatial divergence.

### 2.2.2 Units

By default the angles are expressed in radians and the lengths are given in nanometers. But it is possible to use other units by specifying them right after the value of the corresponding parameter like, for example, 20.0*micrometer.
2.2.3 Programs

The examples presented in the next paragraphs are written in Python. For tutorials about this programming language, the users are referred to [2].

2.3 Example 1: Two types of islands on top of substrate. No interference function

In this example, we simulate the scattering from a mixture of cylindrical and prismatic nanoparticles without any interference between them. These particles are placed in air, on top of a substrate. We are going to go through each step of the simulation. The Python script specific to each stage will be given at the beginning of the description. But for the sake of completeness the full code is given at the end of this section (Listing 2.1).

We start by importing different functions from external modules (line 1), for example NumPy, which is a fundamental package for scientific computing with Python [3]. In particular, line 3 imports the features of BornAgain software.

```python
import sys, os, numpy
from libBornAgainCore import *
```

**First step: Defining materials**

```python
def RunSimulation():
    # defining materials
    mAmbience = MaterialManager.getHomogeneousMaterial("Air", 0.0, 0.0)
    mSubstrate = MaterialManager.getHomogeneousMaterial("Substrate", 6e-6, 2e-8)
    mParticle = MaterialManager.getHomogeneousMaterial("Particle", 6e-4, 2e-8)
```

Line 4 marks the beginning of the function to define and run the simulation. Lines 6, 8 and 10 define different materials using function getHomogeneousMaterial from class MaterialManager. The general syntax is the following

```
<material_name> = MaterialManager.getHomogeneousMaterial("name", delta, beta)
```

where name is the name of the material associated with its complex refractive index n=1-delta +i beta. <material_name> is later used when referring to this particular material. The three defined materials in this example are Air with a refractive index of 1 (delta = beta =0), a Substrate associated with a complex refractive index equal to 1−6×10^{-6}+i2×10^{-8}, and the material of particles, whose refractive index is n= 1−6×10^{-4}+i2×10^{-8}. 

Second step: Defining the particles

```python
# collection of particles
cylinder_ff = FormFactorCylinder(5*nanometer, 5*nanometer)
cylinder = Particle(mParticle, cylinder_ff)
prism_ff = FormFactorPrism3(5*nanometer, 5*nanometer)
prism = Particle(mParticle, prism_ff)
```

We implement two different shapes of particles: cylinders and prisms (i.e. elongated particles with a constant equilateral triangular cross section). All particles implemented in BornAgain are defined by their form factors, their sizes and the material they are made of. Here, for the cylindrical particle, we input its radius and height. For the prism, the possible inputs are the length of one side of its equilateral triangular base and its height.

In order to define a particle, we proceed in two steps. For example for the cylindrical particle, we first specify the form factor of a cylinder with its radius and height, both equal to 5 nanometers in this particular case (see line 12). Then we associate this shape with the constituting material as in line 13.

The same procedure has been applied for the prism in lines 14 and 15 respectively.

Third step: Characterizing the layers and assembling the sample

Particle decoration

```python
particle_decoration = ParticleDecoration()
particle_decoration.addParticle(cylinder, 0.0, 0.5)
particle_decoration.addParticle(prism, 0.0, 0.5)
interference = InterferenceFunctionNone()
particle_decoration.addInterferenceFunction(interference)
```

The object which holds the information about the positions and densities of particles in our sample is called ParticleDecoration (line 16). We use the associated function addParticle for each particle shape (lines 17, 18). Its general syntax is addParticle(<particle_name>, depth, abundance)

where <particle_name> is the name used to define the particles (lines 13 and 15), depth (default value =0) is the vertical position, expressed in nanometers, of the particles in a given layer (the association with a particular layer will be done during the next step) and abundance is the proportion of this type of particles, normalized to the total number of particles. Here we have 50% of cylinders and 50% of prisms.

Remark: Depth of particles

The vertical positions of particles in a layer are given in relative coordinates. For the top layer, the bottom corresponds to depth=0 and negative values would correspond to particles floating above layer 1 since the vertical axis, shown in figure 2.1 is pointing upwards. But for all the other layers, it is the top of the layer which corresponds to depth=0.

Finally lines 19 and 20 specify that there is no coherent interference between the waves scattered by these particles. The intensity is calculated by the incoherent sum of the scattered waves: \( \langle |F_n|^2 \rangle \), where \( F_n \) is the form factor associated with the particle of type \( n \). The way these waves interfere imposes the horizontal distribution of the particles as the interference reflects the long or short-range...
order of the particles distribution (see Theory). On the contrary, the vertical position is imposed when we add the particles in a given layer by parameter depth, as shown in lines 17 and 18.

Multilayer

```python
# air layer with particles and substrate form multi layer
air_layer = Layer(mAmbience)
air_layer.setDecoration(particle_decoration)
substrate_layer = Layer(mSubstrate, 0)
multi_layer = MultiLayer()
multi_layer.addLayer(air_layer)
multi_layer.addLayer(substrate_layer)
```

We now have to configure our sample. For this first example, the particles, i.e. cylinders and prisms, are on top of a substrate in an air layer. The order in which we define these layers is important: we start from the top layer down to the bottom one.

Let us start with the air layer. It contains the particles. In line 22, we use the previously defined mAmbience (=“air” material) (line 6). The command written in line 23 shows that this layer is decorated by adding the particles using the function particle_decoration defined in lines 16-20. The substrate layer only contains the substrate material (line 24).

There are different possible syntaxes to define a layer. As shown in lines 22 and 24 we can use Layer(<material_name>, thickness) or Layer(<material_name>). The second case corresponds to the default value of the thickness, equal to 0. The thickness is expressed in nanometers.

Our two layers are now fully characterized. The sample is assembled using MultiLayer() constructor (line 25): we start with the air layer decorated with the particles (line 26), which is the layer at the top and end with the bottom layer, which is the substrate (line 27).

Fourth step: Characterizing the input beam and output detector and running the simulation

```python
# run simulation
simulation = Simulation()
simulation.setDetectorParameters(100, -1.0*degree, 1.0*degree,
                                 100, 0.0*degree, 2.0*degree, True)
simulation.setBeamParameters(1.0*angstrom, 0.2*degree, 0.0*degree)
simulation.setSample(multi_layer)
simulation.runSimulation()
```

The first stage is to define the Simulation() object (line 29). Then we define the detector (line 31) and beam parameters (line 32), which are associated with the sample previously defined (line 33). Finally we run the simulation (line 34). Those functions are part of the Simulation class. The different incident and exit angles are shown in figure 2.1.

The detector parameters are set using ranges of angles via the function:

```
setDetectorParameters(n_phi, phi_f_min, phi_f_max,
                     n_alpha, alpha_f_min, alpha_f_max, isgisaxs_style=false),
```
where \( n_{\phi}=100 \) is the number of iterations for \( \phi_f \), \( \phi_{f\min}=-1.0\text{degree} \) and \( \phi_{f\max}=1.0\text{degree} \) are the minimum and maximum values respectively of \( \phi_f \), \( n_{\alpha}=100 \) is the number of iterations for \( \alpha_f \), \( \alpha_{f\min}=0.0\text{degree} \) and \( \alpha_{f\max}=2.0\text{degree} \) are the minimum and maximum values respectively of \( \alpha_f \).

isgisaxs_style=True (default value = False) is a boolean used to characterise the structure of the output data. If isgisaxs_style=True, the output data is binned at constant values of the sine of the output angles, \( \alpha_f \) and \( \phi_f \), otherwise it is binned at constant values of these two angles.

For the beam the function to use is \texttt{setBeamParameters(lambda, alpha_i, phi_i)} where \( \lambda=1.0\text{angstrom} \) is the incident beam wavelength, \( \alpha_i=0.2\text{degree} \) is the incident grazing angle on the surface of the sample, \( \phi_i=0.0\text{degree} \) is the in-plane direction of the incident beam (measured with respect to the \( x \)-axis).

Remark: Note that, except for isgisaxs_style, there are no default values implemented for the parameters of the beam and detector.

Line 34 shows the command to run the simulation using the previously defined setup.

**Fifth step: Saving the data**

```python
# retrieving intensity data
return GetOutputData(simulation)
```

In line 36 we obtain the simulated intensity as a function of outgoing angles \( \alpha_f \) and \( \phi_f \) for further uses (plots, fits,...) as a NumPy array containing \( n_{\phi} \times n_{\alpha} \) datapoints. Some options are provided by BornAgain. For example, figure 2.2 shows the two-dimensional contourplot of the intensity as a function of \( \alpha_f \) and \( \phi_f \).
import sys, os, numpy

sys.path.append(os.path.abspath(os.path.join(os.path.split(__file__)[0], '..', '..', '..', 'lib')))

from libBornAgainCore import *

def RunSimulation():
    # defining materials
    mAmbience = MaterialManager.getHomogeneousMaterial("Air", 0.0, 0.0)
    mSubstrate = MaterialManager.getHomogeneousMaterial("Substrate", 6e-6, 2e-8)
    mParticle = MaterialManager.getHomogeneousMaterial("Particle", 6e-4, 2e-8)

    # collection of particles
    cylinder_ff = FormFactorCylinder(5*nanometer, 5*nanometer)
    cylinder = Particle(mParticle, cylinder_ff)
    prism_ff = FormFactorPrism3(5*nanometer, 5*nanometer)
    prism = Particle(mParticle, prism_ff)
    particle_decoration = ParticleDecoration()
particle_decoration.addParticle(cylinder, 0.0, 0.5)
particle_decoration.addParticle(prism, 0.0, 0.5)
interference = InterferenceFunctionNone()
particle_decoration.addInterferenceFunction(interference)
# air layer with particles and substrate form multi layer
air_layer = Layer(mAmbience)
air_layer.setDecoration(particle_decoration)
substrate_layer = Layer(mSubstrate, 0)
multi_layer = MultiLayer()
multi_layer.addLayer(air_layer)
multi_layer.addLayer(substrate_layer)

# build and run simulation
simulation = Simulation()
simulation.setDetectorParameters(100, -1.0*degree, 1.0*degree, 100, 0.0*degree, 2.0*degree, True)
simulation.setBeamParameters(1.0*angstrom, 0.2*degree, 0.0*degree)
simulation.setSample(multi_layer)
simulation.runSimulation()

# retrieving intensity data
return GetOutputData(simulation)

Listing 2.1: Python script of example 1

2.4 Example 2
Bibliography

