PDFmorph user guide

1.0.0 release June 2024

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Acknowledgments

The PDFmorph app was developed by the Billinge group at Columbia University. The examples in the tutorial part of the manual were made possible by data collected by Soham Banerjee at Brookhaven National Laboratory, Benjamin Frandsen during his PhD at Columbia University, and Long Yang from Tonji University.

The DiffPy development team is the set of all contributors to DiffPy projects. Each member of the development team maintains copyright on their individual contributions to the code base. For a detailed contribution history of PDFmorph, see the git logs at https://github.com/diffpy/diffpy.pdfmorph.

For more information about the application, please visit https://www.diffpy.org/diffpy.pdfmorph. Report any bugs to diffpy-users@googlegroups.com or submit an issue to https://github.com/diffpy/diffpy.pdfmorph/issues.

Preface

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Table of Contents

| 1 | Introduction | T |
|----------|----------------------------------|----------|
| 2 | Installation | 2 |
| | 2.1 Availability | 2 |
| | 2.2 Requirements | |
| | 2.3 Installation instructions | |
| 3 | Tutorials | 4 |
| | 3.1 Quick start tutorial | 4 |
| | 3.2 Extra tutorials | |
| | 3.2.1 Performing multiple morphs | 8 |
| | 3.2.2 Nanoparticle shape effect | 2 |
| | 3.2.2.1 Spherical shape | 3 |
| | 3.2.2.2 Spheroidal shape | 4 |
| 4 | Available morphs1 | 5 |
| | 4.1 Temperature-related morphs | 5 |
| | 4.1.1 Isotropic expansion | |
| | 4.1.2 Thermal broadening | 6 |
| | 4.2 Shape-related morphs | 7 |
| 5 | PDFmorph options 19 | 9 |
| Τı | ndex 2: | 3 |

1 Introduction

PDFmorph is a Python package that increases the insight researchers can obtain by comparing measured atomic pair distribution functions (PDFs) in a model-independent way. It was originally designed to help a researcher answer a question: "Has my material undergone a phase transition between these two measurements?"

A common approach to compare two PDFs is to plot the difference curve below the two PDFs. However, a significant signal can be seen in the difference curve due to benign effects such as thermal expansion (peak shifts), increased thermal motion (peak broadening), or a change in scale due to differences in the incident flux when computing the PDFs for example. Other commonly-used dissimilarity metrics such as the weighted Rietveld reliability factor R_W are also inflated by these effects. PDFmorph does its best to correct for these benign effects prior to computing the R_W and plotting the difference curve.

To use PDFmorph, supply one PDF as the "target" (suggested to the one collected under higher temperature) and a second PDF to be morphed. The user can choose from a list of morphs to apply, which includes "stretching" (scaling the r-axis, the abscissa, to simulate isotropic expansion/compression), "smearing" (peak broadening by convlution with a Gaussian), and "scaling" (scaling the ordinate axis). By default, PDFmorph will refine the intensity of the morphs to best match the morphed to the target PDF. All morphed PDFs generated by PDFmorph can easily be saved and exported (see the --save option in Chapter 5 [PDFmorph options], page 19).

PDFmorph includes other morphs including those that factor in nanoparticle shape effects. For a full list of available morphs, see Chapter 4 [Available morphs], page 15. For a quick tutorial on using PDFmorph, see Section 3.1 [Quick start tutorial], page 4. For a full list of command-line options, see Chapter 5 [PDFmorph options], page 19.

Finally, we note that though PDFmorph could be used on other spectra, it has not been extensively tested beyond the PDF.

2 Installation

2.1 Availability

PDFmorph is open source and distributed under the BSD license. It runs on Windows, Mac OS, Linux, and all major Unix systems. The source code is freely available at https://github.com/diffpy/diffpy.pdfmorph.

If you come across any bugs in the application, please open an issue (https://github.com/diffpy/diffpy.pdfmorph/issues) or email diffpy-dev@googlegroups.com.

2.2 Requirements

PDFmorph is currently run from the command-line, so it is recommended that users first familiarize themselves with their operating system's command-line interface.

PDFmorph currently requires Python 3.10 or higher to run. It also makes use of the following third party libraries:

- NumPy (https://numpy.org) Python's fast scientific computing library
- Matplotlib (https://matplotlib.org) Python plotting library
- SciPy (https://scipy.org) Python's technical computing library
- diffpy.utils (https://www.diffpy.org/diffpy.utils) DiffPy (https://www.diffpy.org) general utility functions

These dependencies will be installed automatically if you use the conda installation procedure described in the following section (Section 2.3 [Installation instructions], page 2).

2.3 Installation instructions

We recommend installing the package and its dependencies using conda. Miniconda, a minimal installer for conda, can be downloaded here (https://docs.anaconda.com/free/miniconda). Once Miniconda is installed, run conda on the command-line to test that it has been installed properly.

To create and activate a conda environment to use PDFmorph, run the following command.

```
conda create -n pdfmorph_env python=3 --yes
conda activate pdfmorph_env
```

A conda environment must be active to use the packages within. In future sessions, make sure to run conda activate pdfmorph_env to activate the environment before interacting with the PDFmorph program.

Once active, you can install PDFmorph into your environment from the conda-forge (https://conda-forge.org) channel of Anaconda packages by running

```
conda config --add channels conda-forge
conda install diffpy.pdfmorph
```

This installation only needs to be done once. The next time you activate the environment, these packages will already be installed. To see a list of all installed packages within an environment, run

```
conda list
```

When you are finished with the session, exit the environment by running conda deactivate pdfmorph_env

If you do not wish to use conda to install, PDFmorph is also available on the Python Package Index (https://pypi.org) or can be installed from source at https://github.com/diffpy/diffpy.pdfmorph. Note that these installation methods will require you to first install the dependencies listed in the previous section (Section 2.2 [Requirements], page 2).

3 Tutorials

This section includes a quick start tutorial to acquaint users with the core features of PDFmorph. Also included are a few extra tutorials covering some more specific PDFmorph functionalities.

Before you start this tutorial, make sure you have installed all necessary software and dependencies (Section 2.3 [Installation instructions], page 2). If you are looking for a full summary of PDFmorph command-line options, see Chapter 5 [PDFmorph options], page 19, or run pdfmorph --help in a PDFmorph-equipped conda environment.

3.1 Quick start tutorial

In this tutorial, we will demonstrate how to use PDFmorph to compare two PDFs measured from the same material at different temperatures. The morphs showcased include "stretch", "scale", and "smear". Throughout this tutorial, entries surrounded by angle brackets <> represent placeholder names. The user is free to use their own names in stead of those provided.

- 1. Open your terminal (Linux and MacOS) or Command Prompt (Windows).
- 2. If not already active, activate your conda environment with PDFmorph installed using conda activate <PDFmorph_env>
 - See Chapter 2 [Installation], page 2, for how to set up a conda environment with PDFmorph installed.
 - If you need to list your available conda environments, run conda env list
- 3. Create a directory to store the tutorial PDF files using

mkdir <PDFmorph_tutorial_directory>
and enter the directory with the cd command

cd <PDFmorph_tutorial_directory>

- 4. Download the quick start tutorial files tutorialData.zip (https://www.diffpy.org/diffpy.pdfmorph/tutorialfiles.html) into your current directory and extract the files. After this is done, type ls in the command-line to list your current directory contents and make sure there is a directory named tutorialData.
 - The files in this dataset were collected by Soham Banerjee at Brookhaven National Laboratory in Upton, New York. Note that these files have a .gr extension, indicating that they are measured PDFs. The .cgr file extension indicates a calculated PDF such as those generated by the PDFgui (https://www.diffpy.org/products/pdfgui.html) program.
 - Each file is PDF data collected on Iridium Telluride with 20% Rhodium Doping $IrRhTe_2$ at different temperatures with the first file (01) collected at 10K and the last (44) at 300K. The samples increase in temperature as their numbers increase. The "C" in their names indicates that they have undergone cooling.
- 5. Enter the tutorialData directory using the cd command and find the data labeled darkSub_rh20_C_##.gr within this directory.

```
(pdfmorph_env) ~ $cd PDFmorph_tutorial_directory/
(pdfmorph_env) PDFmorph_tutorial_directory $ts

tutorialData tutorialData 2.19

(pdfmorph_env) PDFmorph_tutorial_directory $cd tutorialData
(pdfmorph_env) tutorialData $ts

_MACOSX tutorialData $cd tutorialData $cd tutorialData
(pdfmorph_env) tutorialData $ts

darkSub_rh20_C.01.gr darkSub_rh20_C.29.gr darkSub_rh20_C.10.gr darkSub_rh20_C.25.gr

darkSub_rh20_C.02.gr darkSub_rh20_C.10.gr darkSub_rh20_C.19.gr darkSub_rh20_C.25.gr

darkSub_rh20_C.03.gr darkSub_rh20_C.10.gr darkSub_rh20_C.19.gr darkSub_rh20_C.25.gr

darkSub_rh20_C.03.gr darkSub_rh20_C.12.gr darkSub_rh20_C.21.9.gr darkSub_rh20_C.27.gr

darkSub_rh20_C.04.gr darkSub_rh20_C.12.gr darkSub_rh20_C.21.9.gr

darkSub_rh20_C.04.gr darkSub_rh20_C.13.gr

darkSub_rh20_C.04.gr darkSub_rh20_C.13.gr

darkSub_rh20_C.03.gr darkSub_rh20_C.13.gr

darkSub_rh20_C.03.gr darkSub_rh20_C.13.gr

darkSub_rh20_C.03.gr darkSub_rh20_C.13.gr

darkSub_rh20_C.23.gr darkSub_rh20_C.23.gr

darkSub_rh20_C.36.gr

darkSub_rh20_
```

Figure 3.1: Entering the tutorialData directory and searching for the PDF data (Linux Terminal Output).

6. Let us try try using the PDFmorph program to plot two PDFs against each other. Type the following command into the command-line

```
pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr
```

Underneath the two PDFs, we can see a substantial difference curve and large R_W value of 0.407

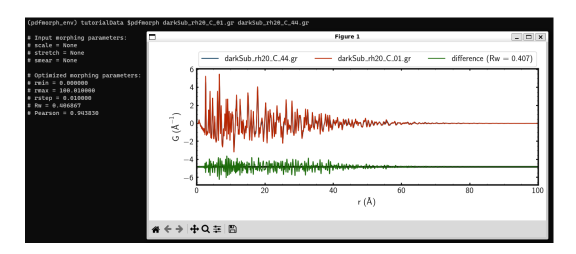


Figure 3.2: Using PDFmorph to compare two PDFs without morphing.

- 7. Now, we will start the morphing process.
 - Note that the morphs will apply only to the first PDF provided (which will be darkSub_rh20_C_01.gr in our case). We will henceforth refer to the first PDF as the "morphed" and the second (unmorphed) PDF as the "target". Our goal will be to apply morphs to see if we can reduce the difference between the morphed and target PDFs.
 - In this tutorial, we will apply the scale factor, Gaussian smear, and stretch morphs. To learn more about why we wish to apply these three morphs in particular, see Section 4.1 [Temperature-related morphs], page 15.
- 8. We will begin by applying a scaling factor of 2 by typing the command

pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=2 -a Now, the difference is much larger $R_W=1.457$. We should modify our initial value for the scaling factor until we see a reduction in R_W . We can try 0.9

pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=0.9 -a and see that the difference has dropped to $R_W=0.351$, lower than when comparing the unmorphed PDF. For PDFmorph to optimize the scale factor, drop the -a option

```
pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=0.9
```

Given a reasonable initial guess, PDFmorph will find the optimal value for each morph. In this example, PDFmorph should display scale=0.799025 meaning it has found this to be the optimal value for the scale factor. The difference $R_W=0.330$ has further decreased.

• The -a option prevents PDFmorph from refining the morph. It is the choice of the user whether or not to run values before removing -a when analyzing data with PDFmorph. By including it, you can attempt to manually move towards convergence before allowing the program to optimize by removing it; when including it, you may reach a highly optimized value on the first guess or diverge greatly.

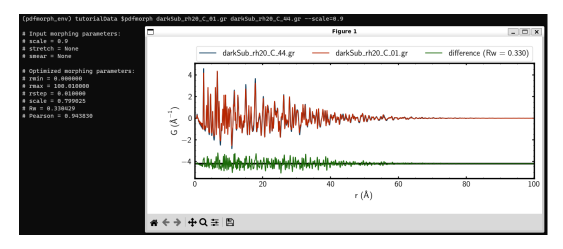


Figure 3.3: PDFmorph has found an optimal value (minimizes R_W) for the scale factor.

9. Now we will add on a Gaussian smear morph with an initial smear factor of 0.5. The absolute value of the smear factor is the standard deviation of the Gaussian that is convoluted with the morphed PDF.

```
pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=0.8 \
--smear=0.5 -a
```

• The \ in the above command is only used to break our long command into multiple lines for readability on this document. In your Terminal/Command Prompt, you can write the entire command in a single line.

We have tailored our initial scale factor to be close to the optimal value given by PDFmorph, but we should see that the $R_W = 0.897$ has increased substantially due to the smear factor. Remove the -a from above and run it again to refine the smear factor.

• Note that the warnings that the Terminal/Command Prompt displays are largely numerical in nature and do not indicate a physically irrelevant guess. These are somewhat superficial and in most cases can be ignored.

After refining, we should see that R_W has dropped back to 0.330, and the optimized smear factor is quite small (smear = 0.002940) indicating that the smear has hardly had an effect on our PDF. To see an effect, restrict the rmin and rmax values by typing

```
pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=0.8 \
--smear=0.5 --rmin=1.5 --rmax=30
```

The optimized smear factor is now a non-insignificant smear=-0.084138 and $R_W = 0.204$.

- We restricted the r (abscissa) values because some of the Gaussian smear effects are only visible in a fixed r range. We chose this r range by noting where most of our relevant data was that was not exponentially decayed by instrumental short-comings.
- Also note that a negative smear factor is equivalent to its absolute value. You can see that

```
pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=0.8 \ --smear=-0.84138 --rmin=1.5 --rmax=30 -a
```

and

```
pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=0.8 \
--smear=+0.84138 --rmin=1.5 --rmax=30 -a
```

give the exact same results.

10. Finally, we will examine the stretch morph. The r axis will be stretched by 1+s where s is the stretch factor.

```
pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=0.8 \
--smear=-0.84 --stretch=0.5 --rmin=1.5 --rmax=30 -a
```

Again, the R_W has increased. Before letting PDFmorph refine automatically, try seeing which direction (higher or lower) the initial stretch factor should go to decrease R_W . If you cannot, type

```
pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=0.8 \
--smear=-0.84 --stretch=0.005 --rmin=1.5 --rmax=30 -a
```

to observe a decreased difference and remove the -a to get the optimized stretch factor of stretch=0.001762 and smaller $R_W = 0.117$. We have reached the optimal fit for our PDF!

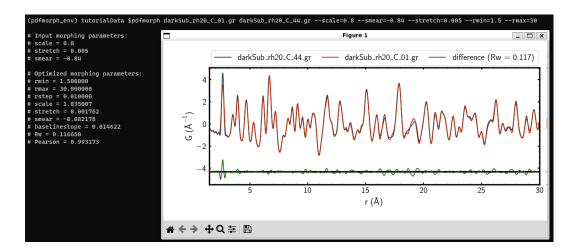


Figure 3.4: The optimal fit after applying the scale, smear, and stretch morphs.

11. To save your new morphed PDF as a file named morph.gr, use the save command

```
pdfmorph darkSub_rh20_C_01.gr darkSub_rh20_C_44.gr --scale=0.8 \
--smear=-0.84 --stretch=0.005 --rmin=1.5 --rmax=30 \
--save=morph.gr
```

Type 1s to confirm a new file named morph.gr has been saved.

12. Now, try it on your own! If you have personally collected or otherwise readily available PDF data, try this process to see if you can morph your PDFs to one another. Note that many of the parameters provided in this tutorial are unique to it, so be cautious about your choices and made sure that they remain physically relevant.

3.2 Extra tutorials

PDFmorph has some more functionalities not showcased in the basic workflow above (run pdfmorph ---help for an overview of all PDFmorph functionalities). Tutorials for some of these are included below. Additional files are used for these tutorials. Please download and extract the extra tutorial data additionalData.zip (https://www.diffpy.org/diffpy.pdfmorph/tutorialfiles.html) before proceeding.

3.2.1 Performing multiple morphs

It may be useful to morph a PDF against multiple targets: for example, you may want to morph a PDF against a sequence of PDFs measured at various temepratures to determine whether a phase change has occured. PDFmorph currently allows users to morph a PDF against all files in a selected directory and plot resulting R_W values from each morph. It is advised that the lowest temperature PDF be that morphed and the higher temperature PDFs act as targets as the smear morph is only able to account for increases in thermal motion.

1. Within additionalData, enter (using cd) the morphMultiple directory. Inside, you will find multiple PDFs of $SrFe_2As_2$ measured at various temperatures.

• These PDFs come from Atomic Pair Distribution Function Analysis: A Primer (https://github.com/Billingegroup/pdfttp_data) by Simon Billinge and Kirsten Jensen. Contributions to the dataset have been made by Benjamin Frandsen and Long Yang.

```
(pdfmorph_env) PDFmorph_tutorial_directory $cd additionalData
(pdfmorph_env) additionalData $ls
additionalData
(pdfmorph_env) additionalData $cd additionalData/
(pdfmorph_env) additionalData $ls
morphMultiple morphShape
(pdfmorph_env) additionalData $cd morphMultiple
(pdfmorph_env) morphMultiple $ls
SrFe2Asz_156K.gr SrFe2Asz_168K.gr SrFe2Asz_2186K.gr SrFe2Asz_204K.gr SrFe2Asz_222K.gr SrFe2Asz_246K.gr
SrFe2Asz_156K.gr SrFe2Asz_174K.gr SrFe2Asz_192K.gr SrFe2Asz_216K.gr SrFe2Asz_228K.gr SrFe2Asz_246K.gr
SrFe2Asz_162K.gr SrFe2Asz_186K.gr SrFe2Asz_198K.gr SrFe2Asz_216K.gr SrFe2Asz_234K.gr
(pdfmorph_env) morphMultiple $
```

Figure 3.5: The files within the morphMultiple directory.

2. Let us start by getting the R_W of SrFe2As2_150K.gr compared to the other, higher-temperature PDFs in the directory. Run

```
pdfmorph SrFe2As2_150K.gr . --multiple-targets
```

• The --multiple-targets tag tells PDFmorph to compare the morphed file SrFe2As2_150K.gr against all PDFs in a directory. The directory we have supplied is ., which is shorthand for the current working directory. In our case, this is the morphMultiple directory.

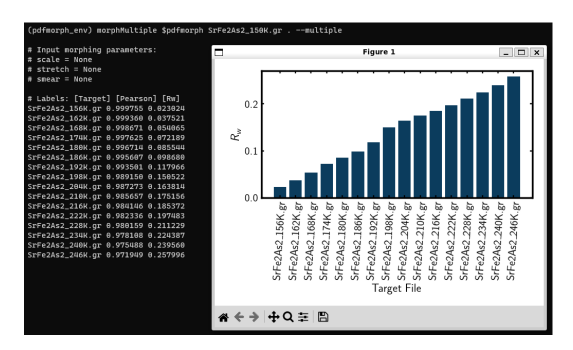


Figure 3.6: Bar chart of R_W values for each target file. Target files are listed in ASCII sort order.

3. After running this we get a chart of R_W values for the morphed file compared to each

target file. By default, a bar chart is generated. However, this plot can be difficult to interpret. We may instead wish to generate a line chart of R_W values against some numerical abscissa such as temperature. At the top of each .gr file in this directory is a list of parameters. These are of the form $\operatorname{parameter_name} = \operatorname{parameter_value}$ and are located above the r versus gr table. Included in these parameters is the temperature at which each PDF was measured at.

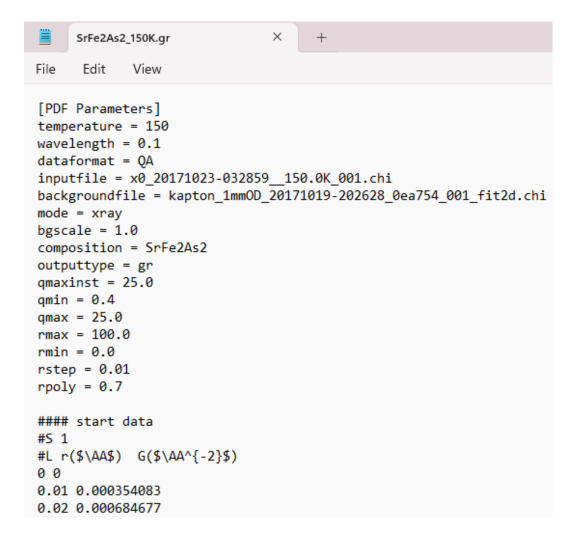


Figure 3.7: Parameters located at the top of the SrFe2As2_150K.gr file. Some parameters included are temperature, wavelength, and composition.

4. By running

pdfmorph SrFe2As2_150K.gr . --multiple-targets --sort-by=temperature we can sort the plotted R_W values by the temperature parameter included within each file.

- When the value of the parameter given to --sort-by is numerical, a line plot is generated.
- Otherwise, a bar chart with the parameter values sorted in ASCII sort order is

generated. You can observe this behavior by using inputfile as the --sort-by parameter instead of temperature.

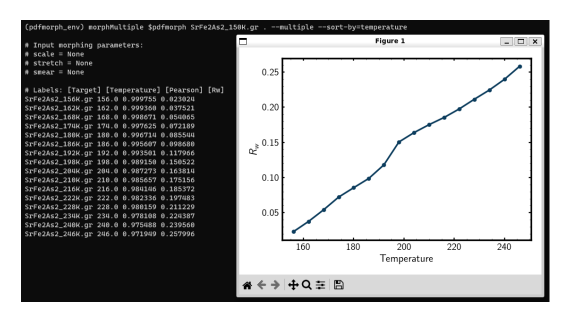


Figure 3.8: The R_W plotted against the temperature the target PDF was measured at.

5. Between 192K and 198K, the R_W has a sharp change, which may be indicative of a phase change. To be more certain, let us apply morphs to take into account isotropic expansion and differences in incident flux (stretching and scaling).

```
pdfmorph SrFe2As2_150K.gr . --scale=1 --stretch=0 --multiple-targets \ --sort-by=temperature
```

The change in R_W has become more pronounced.

- Note that we are not applying the smear morph in this example as it takes a long time to refine and does not significantly change the R_W values in this example.
- 6. We can also change what is being plotted in the ordinate using --plot-parameter. In our case, it is useful to look at the stretch factor

```
pdfmorph SrFe2As2_150K.gr . --scale=1 --stretch=0 --multiple-targets \ --sort-by=temperature --plot-parameter=stretch
```

We can see that the stretch factor generally increases, but from 192K to 198K, there is no increase. This means PDFmorph found that isotropic lattice expansion due to latent thermal effects (indicated by an increase in the stretch factor) does not account for the change in R_W in this region. This is highly suggestive of an alternative source of the dissimilarity such as a phase transition. In fact, there is a structural phase transition from orthorhombic below 192K to tetragonal above 198K! More sophisticated analysis can be done with PDFgui (https://www.diffpy.org/products/pdfgui).

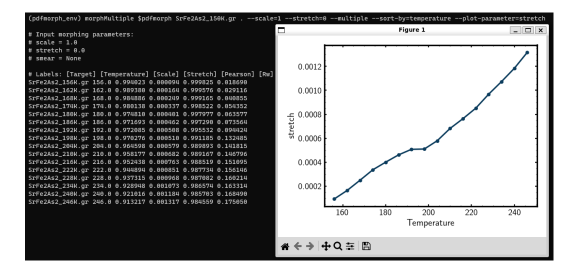


Figure 3.9: The refined stretch factor for each target PDF. There is very little change between 192K and 198K.

7. Finally, let us save all the morphed PDFs into a directory named savedMorphs.

```
pdfmorph SrFe2As2_150K.gr . --scale=1 --stretch=0 --multiple-targets \
--sort-by=temperature --plot-parameter=stretch \
--save=savedMorphs
```

Entering the directory with cd and viewing its contents with ls, we see a file named Morph_Reference_Table.txt with data about the input morph parameters and refined output parameters and a directory named Morphs containing all the morphed PDFs. See the --save-names-file option to see how you can set the names for these saved morphs!

3.2.2 Nanoparticle shape effect

A nanoparticle's finite size and shape can affect the shape of its PDF. We can use PDFmorph to morph a bulk material PDF to simulate these shape effects. Currently, the supported nanoparticle shapes include only spheres and spheroids.

• Within the additionalData directory, cd into the morphShape subdirectory. Inside, you will find a sample Nickel bulk material PDF Ni_bulk.gr. This PDF comes from Atomic Pair Distribution Function Analysis: A Primer (https://github.com/Billingegroup/pdfttp_data). Also in the directory are multiple .cgr, which are calculated Nickel nanoparticle PDFs.

```
(pdfmorph_env) PDFmorph_tutorial_directory $cd additionalData/
(pdfmorph_env) additionalData $ls
additionalData
(pdfmorph_env) additionalData $cd additionalData/
(pdfmorph_env) additionalData $ls
morphMultiple morphShape
(pdfmorph_env) additionalData $cd morphShape
(pdfmorph_env) morphShape $ls
Ni_bulk.gr Ni_nano_sphere.cgr Ni_nano_spheroid.cgr
(pdfmorph_env) morphShape $
```

Figure 3.10: The files within the morphShape directory.

• In the following sections, we apply shape effect morphs on the bulk material to reproduce the calculated PDFs.

Note that there are also support for morphing a nanoparticle PDF into bulk. For more information see Chapter 5 [PDFmorph options], page 19. When applying these inverse morphs it is recommended to set --rmax=psize where psize is the longest diameter of the nanoparticle as data beyond psize is noise.

3.2.2.1 Spherical shape

1. The Ni_nano_sphere.cgr file contains a generated spherical nanoparticle with unknown radius. First, let us plot Ni_bulk.gr against Ni_nano_sphere.cgr.

```
pdfmorph Ni_bulk.gr Ni_nano_sphere.cgr
```

- 2. Despite the two being the same material, the R_W is quite large. To reduce the Rw, we will apply spherical shape effects onto the PDF. However, in order to do so, we first need the radius of the spherical nanoparticle.
- 3. A nanoparticle cannot have bonds longer than the particle size. Thus, for r beyond the spherical diameter, we expect the PDF amplitude to be close to zero. On our plot, the nanoparticle PDF amplitude falls to zero around r = 22. Thus, the nanoparticle radius should be about half of that, or around 11.

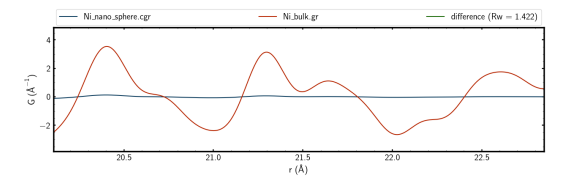


Figure 3.11: The PDF amplitude goes to zero between r = 21 and r = 22.

4. We are ready to perform a morph applying spherical shape effects. To do so, we use the --radius parameter

pdfmorph Ni_bulk.gr Ni_nano_sphere.cgr --radius=11 -a

We can see that the R_W value has decreased significantly from $R_W = 1.422$ to $R_W = 0.085232$.

5. Finally, let us refine by running the same command without the -a tag. The refined radius should be radius=12.183129.

3.2.2.2 Spheroidal shape

1. The Ni_nano_spheroid.cgr file contains a calculated spheroidal Niickel nanoparticle. Again, we can begin by plotting the bulk material against our nanoparticle.

pdfmorph Ni_bulk.gr Ni_nano_spheroid.cgr

2. The nanoparticle shape of the calculated PDF is an oblate spheroid with equitorial radius of about 12 and polar radius of about 6 (this information is contained within the Ni_nano_spheroid.cgr file). To apply the spheroidal shape effects onto the bulk, run

pdfmorph Ni_bulk.gr Ni_nano_spheroid.cgr --radius=12 --pradius=6 -a

- The --radius option corresponds to the equitorial radius.
- The --pradius option corresponds to the polar radius.
- 3. Run the same command without -a to refine. Refining should give radius=12.183129 and pradius=6.279252.

4 Available morphs

In this section, we detail the available morphs and the theory behind when they can (and should) be applied. For specifics on how to use these options in PDFmorph, check out Chapter 5 [PDFmorph options], page 19, and the Chapter 3 [Tutorials], page 4.

4.1 Temperature-related morphs

Comparing two PDFs of the same material measured at different temperatures can produce large R_W s and signals in the difference curves. Though this can be due to a structural phase transition across the measurements, structure-preserving changes such as isotropic expansion/compression and thermal peak broadening/thinning can produce just as large R_W values.

The following morph options are related to our discussion:

- --stretch=STRETCH Stretch the abscissa by a factor 1 + STRETCH.
- --scale=SCALE Scale the plotted function by a factor SCALE.
- --smear=SMEAR Broaden the PDF peaks with a Gaussian smear of width (standard deviation) SMEAR.

4.1.1 Isotropic expansion

The effects of isotropic expansion/compression can be accounted for by scaling and stretching the PDF. To prove this, we will make use of the (total) radial distribution function (RDF), denoted R(r). This function is related to the PDF through

$$G(r) = \frac{R(r)}{r} - 4\pi r \rho_0 \gamma_0(r),$$

where ρ_0 is the atomic number density the and $\gamma_0(r)$ is the nanoparticle form factor (see Section 4.2 [Shape-related morphs], page 17). A partial RDF $R_i(r)$ is defined such that $R_i(r)dr$ gives the number of atoms in the spherical shell bounded by r and r + dr centered at atom i. The total RDF for an atomic system is the average of the partial RDFs of each atom in the system¹.

$$R(r) = \frac{1}{\# \ atoms} \sum_{i}^{atoms} R_i(r).$$

Therefore, the integral of the RDF from a to b gives the number of atomic pairs per atom with a separation distance between a and b.

When a material expands isotropically by a factor α , all distances between pairs of atoms increase by a factor of α (expansion by a factor of $0 < \alpha < 1$ is considered compression). Therefore, the number of atomic pairs with separation distance between a and b before the expansion should equal the number of atomic pairs with separation distance between αa and αb after. Defining R(r) to be the RDF pre-expansion and R'(r) to be that post-expansion,

$$\int_{a}^{b} R(r)dr = \int_{\alpha a}^{\alpha b} R'(r)dr.$$

¹ Farrow, C.L. and Billinge, S.J.L. (2009), Relationship between the atomic pair distribution function and small-angle scattering: implications for modeling of nanoparticles. Acta Cryst. A, 65: 232-239. https://doi.org/10.1107/S0108767309009714

A change of variables tells us

$$\int_{a}^{b} R(r)dr = \int_{\alpha a}^{\alpha b} \frac{R(r/\alpha)}{\alpha} dr,$$

and since these relations hold for all choices of a < b,

$$R'(r) = \frac{R(r/\alpha)}{\alpha}.$$

The corresponding PDFs are

$$G(r) = \frac{R(r)}{r} - 4\pi r \rho_0 \gamma_0(r)$$

pre-expansion, and

$$G'(r) = \frac{R'(r)}{r} - 4\pi r \rho_0' \gamma_0'(r) = \frac{R(r/\alpha)}{\alpha r} - 4\pi r \rho_0' \gamma_0'(r)$$

post-expansion. Due to the expansion, the volume of the material has increased by α^3 , while the total number of atoms remains the same, meaning

$$\rho_0' = \frac{1}{\alpha^3} \rho_0,$$

and the nanoparticle form function is scaled

$$\gamma_0'(r) = \gamma_0(r/\alpha)$$

(see the bottom of Section 4.2 [Shape-related morphs], page 17).

Finally, we can conclude that the PDF after expansion follows

$$G'(r) = \frac{R(r/\alpha)}{\alpha r} - 4\pi r \frac{\rho_0}{\alpha^3} \gamma_0(r/\alpha) = \frac{G(r/\alpha)}{\alpha^2},$$

which is the original PDF scaled by a factor $1/\alpha^2$ and stretched by α .

4.1.2 Thermal broadening

Peaks in the radial distribution functions (see Section 4.1.1 [Isotropic expansion], page 15) obtained from measured PDFs have approximately Gaussian shapes due to Debye-Waller effects. The variance of each peak is the mean square atomic displacement factor (ADP), denoted $\bar{u^2}$ which can depend on dynamic (temperature-dependent) and static factors. Models, such as the Debye model, generally separate the two: $\bar{u^2} = A(T) + A_{static}$, where A(T) increases with temperature.

When a material consists of atoms with similar masses, the A(T) at each peak is approximately the same at a fixed temperature (motivated below). Therefore, an increase in temperature only serves to increase the ADP (and thus the variance of each Gaussian peak) by some fixed constant ζ^2 . PDFmorph simulates this effect by converting the morphed PDF into an RDF, convolving the RDF with a Gaussian of variance ζ^2 centered at r = 0, and

converting back to a PDF. The convolution step increases the variance of each peak by ζ^2 exactly².

Using the Debye model³, we can motivate the statement that A(T) is similar for a material composed of similar-mass atoms. The model shows

$$A(T) = \frac{3h^2T^2}{4\pi^2Mk_B\theta_D^3} \int_0^{\theta_D/T} \frac{x}{e^x - 1} dx + \frac{3h^2}{16\pi^2Mk_B\theta_D},$$

where M is the mass of the oscillating atom, θ_D is the Debye temperature of the (crystal) material, and h and k_B are Planck's constant and Boltzmann's constant respectively. Thus, when the M for each atom is similar, A(T) is also similar. Note also that A(T) is monotonically increasing as a function of temperature.

4.2 Shape-related morphs

The shape and size of a nanoparticle can affect its electronic and optical properties⁴. PDFmorph contains tools to help a researcher identify the shape and size of a nanoparticle PDF given a PDF of a bulk sample. The researcher should select a shape-related morph (listed below) associated with a particular shape and provide the bulk sample PDF as the morphed PDF and nanoparticle PDF as the target. PDFmorph will then multiply the nanoparticle form factor $\gamma(r)$ for that particular shape to the bulk PDF and refine the parameters (e.g. the radius for a spherical shape) to best match the target. Significant difference curve signals or R_W s indicate large deviations from the desired shape, while small signals allow the user to extract size parameters (e.g. the radius of the sphere) from the fit.

This approach has been used to estimate diameters of spherical CdSe nanoparticles consistent with those obtained from transmission electron microscopy, ultraviolet-visible spectroscopy, and photoluminescense measurements⁵.

The available shape morphs are listed below:

- --radius=RADIUS Multiply the PDF by the nanoparticle form factor for a sphere of radius RADIUS. If used with --pradius, multiply the PDF by the nanoparticle form factor for a spheroid of equitorial radius RADIUS and polar radius PRADIUS.
 - The sphere form factor was computed by Kodama et al.⁶.

² Bromiley, P. (2003). Products and Convolutions of Gaussian Distributions.

Dinnebier, R.E. and Billinge, S.J.L. (2018). Overview and principles of powder diffraction. In International Tables for Crystallography (eds C.P. Brock, T. Hahn, H. Wondratschek, U. Müller, U. Shmueli, E. Prince, A. Authier, V. Kopský, D.B. Litvin, E. Arnold, D.M. Himmel, M.G. Rossmann, S.R. Hall, B. McMahon, M.I. Aroyo, C.J. Gilmore, J.A. Kaduk, H. Schenk, C.J. Gilmore, J.A. Kaduk and H. Schenk). https://doi.org/10.1107/97809553602060000935

⁴ Singh, M., Goyal, M., & Devlal, K. (2018). Size and shape effects on the band gap of semiconductor compound nanomaterials. Journal of Taibah University for Science, 12(4), 470–475. https://doi.org/10.1080/16583655.2018.1473946

Masadeh, A. S., Božin, E. S., Farrow, C. L., Paglia, G., Juhas, P., Billinge, S. J. L., Karkamkar, A., & Kanatzidis, M. G. (2007). Quantitative size-dependent structure and strain determination of CdSe nanoparticles using atomic pair distribution function analysis. Phys. Rev. B, 76(11), 115413. https://doi.org/10.1103/PhysRevB.76.115413

⁶ Kodama, K., Iikubo, S., Taguchi, T., & Shamoto, S. (2006). Finite size effects of nanoparticles on the atomic pair distribution functions. Acta Crystallographica Section A, 62(6), 444-453. https://doi.org/10.1107/S0108767306034635

- --pradius=PRADIUS Multiply the PDF by the nanoparticle form factor for a spheroid of equitorial radius RADIUS and polar radius PRADIUS.
 - The spheroid form factor was computed by Lei et al. 7.
- --iradius=IRADIUS Divide the PDF by the nanoparticle form factor for a sphere of radius IRADIUS. If used with --ipradius, divide the PDF by the nanoparticle form factor for a spheroid of equitorial radius IRADIUS and polar radius IPRADIUS.
- --ipradius=IPRADIUS Divide the PDF by the nanoparticle form factor for a spheroid of equitorial radius IRADIUS and polar radius IPRADIUS.

In the rest of this section, we will detail how the nanoparticle form factor is computed. We first need to define the shape function $s\vec{r}$, which is 1 within the nanoparticle and 0 outside. The unaveraged particle form factor $\gamma_0(\vec{r})$ is the autocorrelation of this shape function averaged over the nanoparticle volume V.

$$\gamma_0(\vec{r}) = \frac{1}{V} \int \int \int s(\vec{r}') s(\vec{r}' + \vec{r}) d\vec{r}'.$$

In this manual, we refer to the angle-averaged version, denoted $\gamma_0(r)$, as the nanoparticle form factor.

$$\gamma_0(r) = \frac{\int d\phi \int d\theta \sin(\theta) r^2 \gamma_0(\vec{r})}{\int d\phi \int d\theta r^2 \sin(\theta)}.$$

This form factor is particularly useful as one can approximate the nanoparticle PDF $G_{nano}(r)$ by multiplying the proper nanoparticle form factor with the bulk PDF $G_{bulk}(r)^8$ as follows

$$G_{nano}(r) = \gamma(r)G_{bulk}(r).$$

Finally, an important property of the nanoparticle form factor $\gamma_0(r)$ that we use in Section 4.1.1 [Isotropic expansion], page 15, is that, after undergoing isotropic expansion by a factor α , the new form factor $\gamma'(r)$ is a stretched version of the original form factor: $\gamma'_0(r) = \gamma_0(r/\alpha)$. To show this, consider the new shape function $s'(\alpha \vec{r}) = s(\vec{r})$ as each point \vec{r} pre-expansion is mapped to a point $\alpha \vec{r}$ post-expansion. The volume of the nanoparticle also increases by a factor α^3 , so

$$\gamma_0'(\vec{r}) = \frac{1}{\alpha^3 V} \int \int \int s'(\vec{r}') s'(\vec{r}' + \vec{r}) d\vec{r}'.$$

Aplying a change of variables $\vec{r}' \to \vec{r}'/\alpha$ gives

$$\gamma_0'(\vec{r}) = \frac{1}{\alpha^3 V} \int \int \int s'(\alpha \vec{r}') s'(\alpha \vec{r}' + \vec{r}) \alpha^3 d\vec{r}',$$

and substituting $s'(\alpha \vec{r}) = s(\vec{r})$ gives

$$\gamma_0'(\vec{r}) = \frac{1}{V} \int \int \int s(\vec{r}') s(\vec{r}' + \vec{r}/\alpha) d\vec{r}' = \gamma_0(\vec{r}/\alpha).$$

Taking the angle-averaged integrals gives the desired relation $\gamma_0'(r) = \gamma_0'(r/\alpha)$.

⁷ Lei, M., de Graff, A. M. R., Thorpe, M. F., Wells, S. A., & Sartbaeva, A. (2009). Uncovering the intrinsic geometry from the atomic pair distribution function of nanomaterials. Phys. Rev. B, 80(2), 024118. https://doi.org/10.1103/PhysRevB.80.024118

⁸ Farrow, C.L. and Billinge, S.J.L. (2009), Relationship between the atomic pair distribution function and small-angle scattering: implications for modeling of nanoparticles. Acta Cryst. A, 65: 232-239. https://doi.org/10.1107/S0108767309009714

5 PDFmorph options

In this section we describe all the options available to use in the PDFmorph program. Without options, the PDFmorph program is run with the command

pdfmorph <morphed_PDF> <target_PDF>

where morphs will be applied to the morphed PDF in attempt to minimize the residual with the target PDF.

-h, --help

Show a brief summary of all the options listed in this section.

-V, --version

Show the program version (does not run the program).

-s NAME, --save=NAME

Save the morphed PDF into a file named NAME. You can use – to save to stdout instead. When you have the –-multiple-targets tag enabled, multiple morphed PDFs are generated. Using this command with the –-multiple-targets tag will save all these morphs into a directory named NAME as well as a .txt file summary of refined morph parameters (if applicable) and R_W for each morph done. To specify names for each saved PDF file, use the –-save-names-file option.

-v, --verbose

Increase the amount of information saved to the file(s) generated by --save. Without this option enabled, only the morphed PDF table will be saved (the table of r and corresponding G(r) values). When enabled, information about the input and refined morph parameters (when applicable) and the R_W and Pearson correlation coefficient will be saved as parameters above the PDF table.

--rmin=RMIN

The minimum r-value to use for PDF comparisons and manipulations.

--rmax=RMAX

The maximum r-value to use for PDF comparisons and manipulations.

--pearson

When enabled, refinement will seek only to maximize the Pearson correlation coefficient between the two PDFs.

--addpearson

When enabled, refinement will seek to both maximize the Pearson correlation coefficient and minimize the residual between the two PDFs.

Morphs and manipulations

These options select the morphs and manipulations that are to be applied to the morphed PDF. The passed values will be refined unlesss specifically excluded with the --apply or --exclude options. If no morphs are selected, the morphed PDF will be unchanged and plotted against the target PDF.

-a, --apply

Apply the morph but do not refine the morph parameter value. When this is not enabled, PDFmorph will automatically refine the morph parameters for all morph options below.

-x MORPH, --exclude=MORPH

Do not refine the morph named MORPH. Note that the input MORPH must be lower case

and match exactly the name of the morph option that you wish not to refine. For example, running

pdfmorph <MORPHED> <TARGET> --scale=2 --smear=0.1 --exclude=scale will scale the morphed PDF by exactly 2, but will refine the parameter 0.1 for smear. However, --exclude=Scale will not stop --scale=2 from being refined.

--scale=SCALE

Apply a scale factor SCALE to the function being plotted. For instance, scaling G(r) by SCALE will return SCALE * G(r).

--smear=SMEAR

Smear the PDF peaks with a Gaussian of width (standard deviation) abs(SMEAR). This input can be negative, but will have the same effect as its absolute value. This function is designed only for use on PDFs. This operation assumes the RDF (see Chapter 4 [Available morphs], page 15) peaks are approximately Gaussian and works by converting the PDF to an RDF, convoluting a Gaussian of width SMEAR with the RDF, and converting back to the PDF. This conversion requires a parameter BASELINESLOPE which will be refined if not provided by the user in the --slope option.

--stretch

Stretch the function being plotted along the abscissa by a factor of 1 + STRETCH. For example, if the original function is G(r), the stretch returns g(r/(1 + STRETCH)). The returned PDF will always only be defined between RMIN and RMAX (see --rmin and --rmax). The STRETCH coefficient must be larger than -1 and values in the range (-1,0) will compress the function and set the remaining portion of the function up to RMAX as zero.

--slope=BASELINESLOPE

The slope of the baseline used when applying the smear factor. Unless excluded using --apply or --exclude, it will be refined whenever --smear is used, even if --smear is not being refined. The equation BASELINESLOPE = $4\pi r \rho_0 \gamma_0$ can be used if the atomic number density ρ_0 and the nanoparticle form factor γ_0 is known for the morphed PDF.

--qdamp=QDAMP

Dampen the PDF by a factor QDAMP. A limited Q-resolution of the diffractometer requires us to apply a Gaussian dampening envelope centered at r=0 with width (standard deviation) QDAMP. This envelop is directly multiplied to the PDF.

--radius=RADIUS

Apply the nanoparticle form factor γ_0 for a sphere of radius RADIUS. If PRADIUS is also specified, instead apply the characteristic function of a spheroid with equitorial radius RADIUS and polar radius PRADIUS.

--pradius=PRADIUS

If RADIUS is also specified, see --radius. Otherwise, apply the characteristic function of a sphere with radius PRADIUS.

--iradius=IRADIUS

Apply the inverse characteristic function $1/\gamma_0$ of a sphere of radius IRADIUS/ If IPRADIUS is also specified, instead apply the characteristic function of a spheroid with equitorial radius IRADIUS and polar radius PRADIUS.

--ipradius=IPRADIUS

If IRADIUS is also specified, see --iradius. Otherwise, apply the characteristic function of a sphere with radius IPRADIUS.

Plot options

These options control plotting. The morphed and target PDFs will be plotted against each other with a difference curve shown below. The following changes occur when --multiple-targets is enabled. (1) The R_W for each morphed PDF compared to the target will be plotted unless another parameter is specified by --plot-parameter. (2) The plot will be a bar chart where the abscissa names are the file names of the target PDFs unless otherwise specified by --sort-by. If the parameter given to --sort-by has numerical value, the plot will be a line chart; otherwise, a bar chart the parameter values as the abscissa names will be plotted. By default, ASCII sort order is used for the bar chart abscissa names, but --reverse can be used to reverse the order.

-n, --noplot

Do not display a plot if enabled. Otherwise, show a plot.

--mlabel=MLABEL

Set the label for the morphed PDF to MLABEL on the plot. If not specified, the morphed file name will be used as the label.

--tlabel=TLABEL

Set the label for the target PDF to TLABEL on the plot. If not specified, the target file name will be used as the label.

--pmin=PMIN

The minimum r-value (abscissa) to plot. If not specified, RMIN will be used.

--pmax=PMAX

The maximum r-value (abscissa) to plot. If not specified, RMAX will be used.

--maglim=MAGLIM

The function G(r) will be magnified by MAG for r > MAGLIM. This may be useful as PDF amplitude can get very small for large r. No magnification will take place if --maglim is not specified.

--mag=MAG

See --maglim. No magnification will take place if --maglim is not specified.

--lwidth=LWIDTH

Set the line thickness of the plotted curves. If not specified, it will default to 1.5.

Multiple morphs

PDFmorph allows one to morph one PDF against multiple different targets when --multiple-targets is enabled. See -s and the description under "Plot options" for how saving and plotting change when --multiple-targets is enabled.

--multiple-targets

Changes usage of PDFmorph to

```
pdfmorph <MORPHED_FILE> <TARGET_DIRECTORY>
```

where the PDF in the file MORPHED_FILE will be morphed with each (PDF) file in the directory TARGET_DIRECTORY as the target. Files in TARGET_DIRECTORY will be sorted in ASCII sort order order unless a sorting parameter is specified by sort-by.

--sort-by=FIELD

Used with --multiple-targets. Sort the files in TARGET_DIRECTORY by some parameter named FIELD. Parameters can be specified within each target PDF file by lines of the form

<PARAM_NAME> = <PARAM_VALUE> in the header (anywhere above the r versus G(r) data table). PDFmorph will attempt to find a parameter named FIELD using a case-insensitive search. Numerical PARAM_VALUE will be sorted in ascending order and non-numerical ones will be sorted in ASCII sort order.

--reverse

Used with --multiple-targets. Sort the files in TARGET_DIRECTORY in reverse ASCII sort order. If a parameter is given by --sort-by, reverse the order given by --sort-by.

--serial-file=SERIALFILE

Used with --multiple-targets and --sort-by. Look for FIELD in a serial file named SERIALFILE instead. Only serial file types supported by diffpy.utils (https://www.diffpy.org/diffpy.utils) such as .json are allowed.

--save-names-file=NAMESFILE

Used with --multiple-targets and -s. Specify names for each manipulated PDF when saving using a serial file named NAMESFILE. The format of NAMESFILE should be as follows: (1) Each target PDF file name is an entry in NAMESFILE. (2) For each entry, there should be a key save_morph_as whose value specified the name to save the manipulated PDF as. An example .json NAMESFILE is below.

```
{
  "target1.gr": {
      "save_morph_as": "name.extension"
},
  "target2.gr": {
      "save_morph_as": "another_name.extension",
      "other_information": "[optional]"
},
  "target3.gr": {
      "SaVe_MoRph_As": "this_also_works.extension",
      "note": "Capitalization does NOT matter. This is case insensitive."
}
```

Only serial file types supported by diffpy.utils (https://www.diffpy.org/diffpy.utils) such as .json are allowed.

--plot-parameter=PLOTPARAM

Used with --multiple-targets and when plotting is enabled. Choose a parameter PLOTPARAM to plot for each morph. When not specified, the R_W values for each morphed PDF compared to the target PDFs will be plotted. This option is not case sensitive meaning --plot-parameter=Scale and --plot-parameter=scale will both cause the parameter scale to be plotted.

Index

| \mathbf{A} | P |
|---|----------------------------|
| acknowledgments2availability2available morphs15 | pdfmorph options |
| ${f E}$ | Q quick start tutorial |
| extra tutorials | \mathbf{R} |
| | requirements |
| ${f I}$ installation | S shape-related morphs |
| $ \begin{array}{cccccccccccccccccccccccccccccccccccc$ | spherical shape morph |
| | ${f T}$ |
| N | temperature-related morphs |
| nanoparticle shape effect | tutorials |