```
(* +----- evaluate_density.nb ------ *)
(* +
            evaluate_density.nb calculates for a data set with xyz coordinates
            the local density of "atoms" around each site.
(* +
            evaluate order.nb creates a three dimensional temperature map for the + *)
            best match of each "atom" in then xyz data file with a database item + *)
(* +
            The "lower" the temperature in the map the better the match.
            INPUT: xyz coodinates of the cluster
            OUTPUT: Graphics file with a color map that shows the density
                   for each "atom" in the xyz data file.
            USAGE: (1) Delete All Output (from the 'Cell' menu)
                   (2) Fill the section between 'BEGIN USER INPUT' and
(* +
                      'END USER INPUT'
                   (3) Evaluate Notebook (from the 'Evaluate' menu)
            DEPENDENCIES: None
(* +
(* +
            AUTHOR: L. Houben, Weizmann Institute of Science
                   lothar.houben(at)weizmann.ac.il
            COPYRIGHT: This software is licensed under the GNU GENERAL PUBLIC
(* +
(* +
                   LICENSE Version 3
(* +----- BEGIN USER INPUT -----+ *)
(* +------*)
(* baseDir is the base folder for program input and output.
(* dataDir is a subfolder that holds the data file.
(* outDir is a subfolder for the program output files.
(* Please make sure that the folders exist.
baseDir = StringJoin[$HomeDirectory, "/Desktop/diOPA"];
dataDir = "data";
outDir = "out";
(* datafileName is name of the data file with the cluster xyz coordinates *)
datafileName = "Ferritin-agglomerate.cel";
Nfcc = {1.6, 18.}; (* shell radius in units of the nearest neighbour distance, number of nearest neighbours in the shell in the bulk structure *)
(* Nfcc[[1]] is the cutoff in units of the nearest neighbour distance for the density calculation, i.e. the resolution of the density map *)
(* Nfcc[[2]] is required to normalize the result to the bulk density of the material *)
(* Note that Nfcc[[2]] depends on the symmetry of the coordination *)
(* +----- *)
(* +----+ *)
celfile = StringJoin[baseDir, "/", dataDir, "/", datafileName]
/Users/lothar/Desktop/diOPA/data/Ferritin-agglomerate.cel
(* (3) define output files
```

```
graphfileprefix = StringJoin[baseDir, "/", outDir, "/", FileBaseName[datafileName], "_Density"]
logfilename = StringJoin[baseDir, "/", outDir, "/", FileBaseName[datafileName], "_Density.log"]
/Users/lothar/Desktop/diOPA/out/Ferritin-agglomerate_Density
/Users/lothar/Desktop/diOPA/out/Ferritin-agglomerate_Density.log
(* define plot parameters
                                                                          *)
axespos = {0, 0, 0};
viewvc = {50, -300, 100};
imsz = {512, 512};
axstyle = {Thick, Thick, Thick};
labSz = 14; (* set to zero if no labels, otherwise 14 or alike *)
colscalelabSz = 14;
ImMagfact = 1.5;
plotopac = 0.5; (* bubble plot opacity *)
(* END PARAMETER DEFINITION
(* START FUNCTION DEFINITION
                                                                                                                                                     *)
(* Function for Reading a celfile *)
(* J. Barthel, RWTH-Aachen University, Germany *)
(* Email: ju.barthel-at-fz-juelich.de *)
(* 28.08.2015 *)
(* *)
(* Reads super cell structure data in CEL file format from *)
(* The specified file and stores the information in global arrays: *)
(* celdims = dimensions of the super-cell *)
(* celangs = angles between the super-cell axes *)
(* natoms = number of atoms in the super-cell *)
(* *)
ReadCelFile[filename ] := Block[{celstrs},
  (* - init *)
  Clear[celdims, celangs, celatms, natoms];
  celdims = {0., 0., 0.}; celangs = {90., 90., 90.}; celatms = {}; natoms = 0;
  (* - read data from file *)
  celstrs = ReadList[filename, "String"];
  (* - analyse / extract numeric data *)
  celdims = Take[ReadList[StringToStream[celstrs[[2]]], "Number"], {2, 4}];
  celangs = Take[ReadList[StringToStream[celstrs[[2]]], "Number"], {5, 7}];
  celatms = Table[
    ReadList[StringToStream[celstrs[[i]]], Join[{"Word"}, Table["Number", {8}]]][[1]]
    , {i, 3, Length[celstrs] - 1}];
  natoms = Length[celatms];
  (* - report *)
  Print["- INPUT CEL file: ", InputForm[filename]];
  Print["- OUTPUT (celdims) dimensions (a,b,c) [nm]: ", celdims];
  Print["- OUTPUT (celangs) axes angles (\alpha, \beta, \gamma) [deg]: ", celangs];
  Print["- OUTPUT (natoms) number of atoms
                                                      : ", natoms];
  If[natoms > 0,
   Print["- OUTPUT (celatms[[1]]) first atom
                                                       : ", celatms[[1]]];
  If[natoms > 1,
   Print["- OUTPUT (celatms[[", natoms, "]]) last atom
                                                                  : ", celatms[[natoms]]];
  ];
```

```
QuickDistance[model_, nlist_] := Block[{SortedEntry = {}, tmplist = model, selected = {}, x = {}},
  (* - init *)
  Clear[EuclDist];
  EuclDist = 0;
  (* - analyse / extract numeric data *)
  For[k = 1, k <= coordination, k++,</pre>
   x = Nearest[tmplist → Automatic, nlist[[k]], 1];
   selected = N[tmplist[[x[[1]]]]];
   AppendTo[SortedEntry, selected]; tmplist = Drop[tmplist, x]
  EuclDist = EuclideanDistance[nlist, SortedEntry];
GetNeighbours[atomindex_, nneighbours_] := Block[{k = {}},
  (* Extract atom atomindex and evaluate its n nearest neighbours *)
  (* the function nearest returns the test element itself, therefore we start with n+1 neighbours *)
  (* centre the list around the selected atom, the selected atom is the origin of the coordinate system *)
  (* Initialize global variable: neighbours , type: list *)
  Clear[neighbours];
  neighbours = {};
  (* *)
  k = Nearest[q, q[[atomindex]], nneighbours + 1];
  neighbours = k[[2;; nneighbours + 1]];
  neighbours[[All, 1]] = neighbours[[All, 1]] - k[[1, 1]];
  neighbours[[All, 2]] = neighbours[[All, 2]] - k[[1, 2]];
  neighbours[[All, 3]] = neighbours[[All, 3]] - k[[1, 3]];
GetDistMeas[atomindex_] := Block { DistMap = { } } ,
  (* Calculate a distance measure to the closest model item in the DB \star)
  (* *)
  (* Returns a global variable: Dist, type: list *)
  (* Dist = {atomnr, DBindex, {angle,angle,distance}} *)
  (* Example: *)
  (* In: GetDistMeas[41]; Dist *)
  (* Out: \left\{41,256,\left\{\frac{\pi}{5},\frac{\pi}{10},0.468556698631839\right\}\right\} *)
  Clear[Dist];
  Dist = {};
  (* Create a distance map *)
  (* Here we need permutations because the list of nearest neighbours is not sorted *)
  (* Finding the minimum over all possible permutations is the right way to go, yet very slow *)
  (* therefore we sort the list of data base model atoms *)
  (* for each atom in the neighbour list we search for the closest in the model \star)
  (* note that this gives the minimum euclidian distance is achieced only if model and neighbour list are close *)
  (* there might be a solution with a better compromise if the match is not close *)
  GetNeighbours[atomindex, coordination]
   For[ind = 1, ind \( \) Length[ModelDB], ind++, QuickDistance[ModelDB[[ind, 3]], neighbours]; AppendTo[
     DistMap, {ModelDB[[ind, 2, 1]], ModelDB[[ind, 2, 2]], EuclDist}]];
  WhereMin = Ordering[DistMap[[All, 3]], 1];
  Dist = {atomindex, WhereMin[[1]], DistMap[[WhereMin]][[1]]};
```

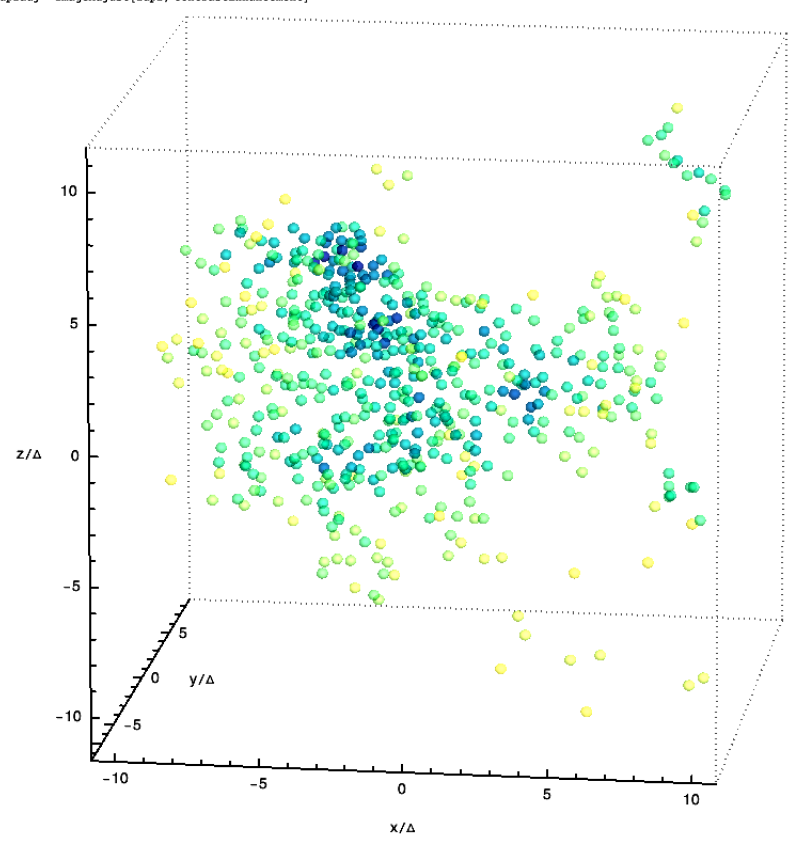
```
(* BoundingBox *)
(* returns box coordinates around the centre *)
(* in a list variable cutoff *)
(* INPUT: xyz_= \{\{x1,y1,z1\},\{x2, y2, z2\}, ...\} *)
          margin = margin size to add
(*
(* OUTPUT: bbox ={{xmin-margin,xmax+margin},{ymin-mrgin,ymax+margin}{zmin-margin,zmax+margin}} * *)
BoundingBox[xyz_, margin_] := Block[{tmp},
  (* - init *)
  Clear[bbox];
  bbox = \{\{0, 0\}, \{0, 0\}, \{0, 0\}\};
  bbox[[1, 1]] = Min[xyz[[All, 1]]] - margin;
  bbox[[2, 1]] = Min[xyz[[All, 2]]] - margin;
  bbox[[3, 1]] = Min[xyz[[All, 3]]] - margin;
  bbox[[1, 2]] = Max[xyz[[All, 1]]] + margin;
  bbox[[2, 2]] = Max[xyz[[All, 2]]] + margin;
  bbox[[3, 2]] = Max[xyz[[All, 3]]] + margin;
  axespos = {bbox[[1, 1]], bbox[[2, 1]], bbox[[3, 1]]};
  Print["- BoundingBox
                               : ", bbox];
  Print["- Axes Position
                               : ", axespos];
(* CentreBox *)
(* Centres xyz around 0 *)
(* in a list variable cutoff *)
(* INPUT: xyz_= \{\{x1,y1,z1\},\{x2, y2, z2\}, \ldots\} * \}
(*
(*
(* OUTPUT: centrelist = list with vectors centred around 0 *)
CentreBox[xyz_] := Block[{tmp},
  (* - init *)
  Clear[centrelist];
  centrelist = xyz;
  centrelist[[All, 1]] -= Min[xyz[[All, 1]]] + 0.5 * (Max[xyz[[All, 1]]] - Min[xyz[[All, 1]]]);
  centrelist[[All, 2]] -= Min[xyz[[All, 2]]] + 0.5 * (Max[xyz[[All, 2]]] - Min[xyz[[All, 2]]]);
  centrelist[[All, 3]] -= Min[xyz[[All, 3]]] + 0.5 * (Max[xyz[[All, 3]]] - Min[xyz[[All, 3]]]);
  Print["- Centred List "];
(*
(* END FUNCTION DEFINITION
           (* START PROCESSING
(* Open log file *)
logfile = OpenWrite[logfilename];
WriteString[logfile, "Filename:", celfile, "\n"];
ReadCelFile[celfile];
Clear[coordatms, f, q];
(* Extract relative atomcoordinates only *)
coordatms = celatms[[All, {2, 3, 4}]];
(* multiply atom relative coordinates with cell dimensions *)
(* multiply atom relative coordinates with cell dimensions *)
f[x_] := \{celdims[[1]] * x[[1]], celdims[[2]] * x[[2]], celdims[[3]] * x[[3]]\}
q = Map[f, coordatms];
 \text{ListPointPlot3D[q, BoxRatios} \rightarrow \{1, 1, 1\}, \, \text{PlotStyle} \rightarrow \text{PointSize[0.01], AxesLabel} \rightarrow \{\text{"x}/\Delta\text{", "y}/\Delta\text{", "z}/\Delta\text{"}\}, \, \text{Lighting} \rightarrow \text{"Neutral", LabelStyle} \rightarrow \text{Directive[Bold]]} 
WriteString[logfile, "Number of atoms:", ToString[Length[q]], "\n"];
```

```
- INPUT CEL file: "/Users/lothar/Desktop/diOPA/data/Ferritin-agglomerate.cel"
- OUTPUT (celdims) dimensions (a,b,c) [nm]: {19.6635, 16.056, 21.3574}
- OUTPUT (celangs) axes angles (\alpha, \beta, \gamma) [deg]: {90., 90., 90.}
- OUTPUT (natoms) number of atoms
                                         : 559
- OUTPUT (celatms[[1]]) first atom
                                         : {Fe, 0., 0.514721, 0.544393, 1., 0.005, 0.1, 0.1, 0.1}
- OUTPUT (celatms[[559]]) last atom
                                           : {Fe, 1., 0.21619, 0.073572, 1., 0.005, 0.1, 0.1, 0.1}
      20
       15
                    \mathbf{X}/\Delta
     (* Color Map Definitions *)
     colortbl = "Rainbow";
     gamma = 1.6; (* gama factor to stretch the contrast *)
     ContrastEnhancement = 1.;
     colnorm[x_] := ((x - CScaleMin) / (CScaleMax - CScaleMin)) ^gamma; (* color normailzation, map a ramge of values CMin..CMaxx to 0..1 *)
     revcolnorm[x] := (1 - (x - CScaleMin) / (CScaleMax - CScaleMin)) ^gamma; (* inverted color normailzation map a ramge of values CMin..CMaxx to 0..1 *)
     (* DENSITY MAP calculation
     (* +------ + *)
     (* INPUT: q=atomlist *)
     (* use the cutoff list and calculate for each point of it the euclidian distances to the points in the reference list *)
     (* this will give Length[cutofflist] times Length[xyz] values *)
     distances = With[{tr = Transpose[q]}, Function[point, Sqrt[Total[(point - tr)^2]]] /@ q]; // AbsoluteTiming
     {0.019066, Null}
```

**6** | evaluate\_density.nb

```
ddata = {};
                      For [1 = 1, 1 \le Length[q], 1++, neighbourcount = BinCounts[distances[[1]], Nfcc[[1]]]; AppendTo[
                                   ddata, {q[[1, 1]], q[[1, 2]], q[[1, 3]], (neighbourcount[[2]] - 1) / Nfcc[[2]]}]];
                      CentreBox[ddata]
                      ddata = centrelist;
                      BoundingBox[ddata, 1]
                      CScaleMin = 0;
                      CScaleMax = 1;
                      CMean = Mean[ddata[[All, 4]]];
                      CMedian = Median[ddata[[All, 4]]];
                      CMin = Min[ddata[[All, 4]]];
                      CMax = Max[ddata[[All, 4]]];
- Centred List
                                                                                        : {{-10.8318, 10.8318}, {-8.78118, 8.78118}, {-11.6715, 11.6715}}
- BoundingBox
- Axes Position
                                                                                        : {-10.8318, -8.78118, -11.6715}
                      colortbl = "BlueGreenYellow";
                      colddata = ddata;
                      colddata[[All, 4]] = Map[revcolnorm, ddata[[All, 4]]];
                      ldp1 = ListPointPlot3D[List /@ Most /@ colddata, PlotRange → bbox, BoxStyle → Directive[Thick, Dotted],
                                  \textbf{AxesOrigin} \rightarrow \textbf{axespos}, \ \textbf{AxesLabel} \rightarrow \{ \texttt{"x}/\Delta \texttt{"}, \ \texttt{"y}/\Delta \texttt{"}, \ \texttt{"z}/\Delta \texttt{"} \}, \ \textbf{Lighting} \rightarrow \texttt{"Neutral"}, \ \textbf{AxesStyle} \rightarrow \textbf{axstyle}, \ \textbf{BoxRatios} \rightarrow \{1, 1, 1\}, \ \textbf{ViewCenter} \rightarrow \{0.5, 0.5, 0.5\}, \ \textbf{AxesLabel} \rightarrow \texttt{(axeslabel)} \rightarrow \texttt{(axesl
                                   LabelStyle → Directive[Bold, FontFamily → "Helvetica", FontSize → labSz], PlotStyle → ({Opacity[plotopac], AbsolutePointSize[1], ColorData[colortbl][(#+0) / 1]} & /@ Last /@ colddata),
                                   ViewCenter \rightarrow \{0.5, 0.5, 0.5\}, ViewVector \rightarrow viewvc, ViewVertical \rightarrow \{0, 0, 1\}, ImageSize \rightarrow imsz * ImMagfact];
                      ldp2 = ldp1 /. Point[x_] \Rightarrow (Sequence@{EdgeForm[], Lighting} \rightarrow "Neutral", Sphere[#, .2]) &@({x} /. {{a_, b_, c_}}) \Rightarrow {{a, b, c}}));
```

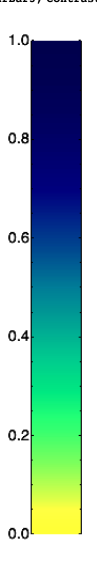
ldp2adj = ImageAdjust[ldp2, ContrastEnhancement]



colourBar3 = DensityPlot[(1 - v) ^gamma, {u, 0, 1}, {v, CScaleMin, CScaleMax}, ColorFunction → ColorData[colortb1], Frame → {False, True},

LabelStyle → Directive[Bold, FontFamily → "Helvetica", FontSize → colscalelabSz], AspectRatio → 10, PlotRangePadding → 0, ImagePadding → 25, ImageSize → imsz, FrameTicks → {None, Automatic}];

colourBar3adj = ImageAdjust[colourBar3, ContrastEnhancement]



```
Print["- Average density (% of bulk): ", CMean]
    Print["- Median density (% of bulk): ", CMedian]
    Print["- Minimum density (% of bulk): ", CMin]
    Print["- Maximum density (% of bulk): ", CMax]
    Print["- Contrast cut off (Min,Max): (", CScaleMin, ",", CScaleMax, ")"]

- Average density (% of bulk): 0.274299

- Median density (% of bulk): 0.277778

- Minimum density (% of bulk): 0

- Maximum density (% of bulk): 0.777778

- Contrast cut off (Min,Max): (0,1)

    WriteString[logfile, "- Average density (% of bulk): ", ToString[CMean], "\n"];
    WriteString[logfile, "- Median density (% of bulk): ", ToString[CMedian], "\n"];
    WriteString[logfile, "- Minimum density (% of bulk): ", ToString[CMin], "\n"];
    WriteString[logfile, "- Maximum density (% of bulk): ", ToString[CMax], "\n"];
    WriteString[logfile, "- Maximum density (% of bulk): ", ToString[CScaleMin], "\n"];
    WriteString[logfile, "- Contrast cut off (Min,Max): (", ToString[CScaleMin], ",", ToString[CScaleMax], ")", "\n"];
```