```
In[1]:= Print[" Figure3cd(Layers2L30nmRIn2d1nm.m"];
    Print[];
            The source of the data of the manuscript"];
    Print["
            'Structure and energetics of carbon, "];
    Print["
            hexagonal boron nitride, and"];
    Print["
    Print[" carbon/hexagonal boron nitride"];
            single-layer and bilayer nanoscrolls' "];
    Print[" / A.I. Siahlo, N.A. Poklonski, A.V. Lebedev,"];
    Print[" I.V. Lebedeva, A.M. Popov, S.A. Vyrko, "];
    Print[" A.A. Knizhnik, Yu.E. Lozovik "];
    Print[" // Phys. Rev. Materials.- 2018.- V. 2,"];
    Print[" № 3.- P. 036001 (9 pp.)."];
    Print[" [DOI: 10.1103/PhysRevMaterials.2.036001] "];
    Print[" -----"];
    NoL1 = 1; NoL2 = 2;
    NoLp = NoL2;
    Print[" I.O The Units (nm, meV, AA)"];
    "nm=10^(-9)m;";
    nm = 10^{(-9)} m;
    AA = 10^{(-10)} m;
    JJkgms = kg m^2/s^2;
    C1 = Ampers;
    "eV=JJ Electronp;";
    JJ = eV / Electron;
    JJms = (kg m^2) / s^2;
    meV = N[eV/1000];
    Print["----"];
    Print[" I.1. All Input Parameters and Constants-----"];
    Print[" I.1.1. The sampling parameters"]
    npRIn1 = 1000;
    Print["npRIn1=", npRIn1];
    Print[" I.1.2. The Input Geometry Parameters of the system"];
    L14d839nm = 14.839nm;
    L12d709nm = 12.709 nm;
    L129d678nm = 29.678 nm;
    L1p = L14d839nm;
    L1p = L129d678nm;
    Print[" The carbon nanoribbon length L1=", L1p/nm, "nm"];
    Lw11d8nm = 11.8 nm;
    Lwp = Lw11d8nm;
    Print[" The carbon nanoribbon width Lw=", Lwp/nm, "nm"];
    Print["----"];
    Print[" Number of the layers in carbon nanoscroll NoL=", NoLp];
    Print[" The length of a carbon nanoribbon L1=", L1p/nm, "nm"];
    Lw1nm = 1. nm; Lwp = Lw1nm;
    Print[" The carbon nanoribbon width Lw=", Lwp/nm, "nm"];
    RIn1d1nm = 1.1 nm;
    RIn1d2nm = 1.2 nm;
    RIn1d14nm = 1.14 nm:
    RIn2nm = 2.047 nm;
    RIn2d1nm = 2.1 nm;
    RIn2d2nm = 2.2 nm:
    RIn2d3nm = 2.3 nm;
```

```
RIn2d4nm = 2.4 nm;
RIn2d5nm = 2.5 nm:
RIn2d6nm = 2.6 nm;
RIn1p = RIn2d5nm;
RIn1p = RIn2d3nm;
RIn1p = RIn2d2nm;
RIn1p = RIn1d14nm;
RIn1p = RIn2d1nm;
Print[" The inner radius of the nanoscroll RIn1=", RIn1p/nm, "nm"];
Print[" I.1.2. The Input Energy Constants"];
Print[" epsVdW - the interlayer interaction energy per one atom of"];
Print[" the nanoscroll:"];
epsVdW35 = 35.0 meV / atom; epsVdWp = epsVdW35;
Print[" epsVdW=", epsVdWp/(eV/atom), "eV/atom"];
Print[" C - the bending elastic constant:"];
C201 = 2.01 eV AA^2/atom;
CBN1328 = 1.328 eV AA^2 / atom;
CCp = C201;
CBNp = CBN1328;
CBNp = CCp;
Print[" CCelast=", CCp / (eV AA^2 / atom), "eV AA^2 / atom"];
Print[" CCBNelast=", CBNp / (eV AA^2 / atom), "eV AA^2 / atom"];
Print[" I.1.3.The Input Geometry constants-----"];
Print[" The interatomic distance aCC and the interlayer distance h"];
aCC142AA = 1.42 AA; aCCp = aCC142AA;
h335nm = 0.3354 nm; hp = h335nm;
Print["h=", hp/nm, " nm (Interlayer distance)"];
Print[" aCC=", aCCp/nm, "nm, h=", hp/nm, "nm"];
NatomsInCell2 = 2; NatomsInCellp = NatomsInCell2;
Print["NatomsInCell=", NatomsInCellp];
Print[" dPhi12 - The difference of the inner angles of the spirales"];
Print["
         of the Layers"];
dPhi12eq0 = 0.0 Pi;
dPhi12eqPi = 1.0 Pi;
dPhi12p = 1.0 Pi;
dPhi12p = 0.5 Pi;
dPhi12p = 0.0 Pi;
Print[" I.5.The parameters for the visualisation"];
RIn1MinMonoScroll = hp / 5;
RIn1MinBiScroll = hp / 5;
RIn1MaxMonoScroll = 4 nm;
RIn1MaxBiScrol1 = 8 nm;
PlotRangeMonoScroll = {-4 eV / atom, 12 eV / atom};
PlotRangeBiScroll = {-10 eV / atom, 30 eV / atom};
ShowSpirales = True;
ShowThePlot = True;
Print[" I.6. The parameters of the output file"];
NanoscrollNamep = StringJoin["Nanoscroll", ToString[NoLp], "L", ToString[Llp/nm], "nm"];
Print[" NanoscrollName=", NanoscrollNamep];
CarbonNanoscrollEnergyVsRInFileName = StringJoin[NanoscrollNamep, ".txt"];
Print[CarbonNanoscrollEnergyVsRInFileName];
Print[" (The output of the data to a file Is Not Performed)"];
npRIn1 = 1000;
Print[" The number of the output points = ", npRIn1];
Print[" I.7. The Input Numerical Constants used in the programm"];
Print[" The Indexes used for the work with EVdW[...] function"];
```

```
iEVdW = 1; iEVdW1Un1 = 2; iEVdW1Ov1 = 3; iEVdW1Un2 = 4; iEVdW1Ov2 = 5;
iEVdW2Un1 = 6; iEVdW2Ov1 = 7;
Print[" ------"];
AA = 0.1 nm; PhiIn := \varphiIn; PhiOut := \varphiOut;
Print["-----"];
Print[" II. The derivated parameters and the functions required"];
Print[" II.1. The derivated parameters"];
fSa[aCC_] := aCC^23Sqrt[3]/4; fSa[aCCp]; Sap = fSa[aCCp];
Print[" The cell area Sa=", fSa[aCC], "=", Sap/nm^2, "nm^2"];
Print[" II.2. The required functions-----"];
Print[" II.2.1. The function fSpiraleLen[", NoLp, ",\varphiIn, \varphiOut, h] defines"];
Print[
 " the Length of a Spirale with the inner agle \varphiIn and the outer angle \varphiOut(>=\varphiIn):"];
fSpiraleLen[NoLv_, PhiInv_, PhiOutv_, hv_] :=
(1 / (4 Pi) hv NoLv (-PhiInv Sqrt[1 + PhiInv^2] +
   PhiOutv Sqrt[1 + PhiOutv^2] - ArcSinh[PhiInv] +
   ArcSinh[PhiOutv]));
Print[" fSpiraleLen[", NoLp,
  ", φIn, φOut, h]=", fSpiraleLen[NoLp, PhiIn, PhiOut, h], "."];
Print[" II.2.2. The function fElast[\varphiIn,\rhoOut] is
    required to calculate an nanoscrollelastic energy: "];
fElast[PhiInv_,
 PhiOutv_] := (Sqrt[PhiInv^2 + 1] / PhiInv -
 Sqrt[PhiOutv^2 + 1] / PhiOutv - ArcSinh[PhiInv] + ArcSinh[PhiOutv]);
Print[" fElast[φIn,φOut] = ", fElast[PhiIn, PhiOut], "."];
Print[" II.2.3. Define the function fPhiOutvsPhiInLh[", NoLp, ", ", PhiIn, ", L, h]."]
fPhiOutvsPhiInLh[NoLv_, PhiInv_, Lv_, hv_] :=
Sqrt[4 \pi Lv / (NoLv hv) + PhiInv^2];
Print[" The function fPhiOutvsPhiInLh[", NoLp,
  ",", PhiIn, ",L,h]=", fPhiOutvsPhiInLh[NoLp, PhiIn, L, h], " is a
 good approximation to obtain the value of \varphiOut for the defined \varphiIn,L,h."];
fPhiInvsPhiOutLh[NoLv_, PhiOutv_, Lv_, hv_] :=
Sqrt[PhiOutv^2 - 4 \pi Lv / (NoLv hv)];
Print[" The inverse function fPhiInvsPhiOutLh[", NoLp, ", φOut, L, h]]=",
fPhiInvsPhiOutLh[NoLp, PhiOut, L, h]];
Print[" could be used in the program applications
    if ROut (instead of RIn) is the input parameter of the system."];
Print[" II.2.4. The functions
    fSpirale1Under(Over)Spirale1Length[NoLv,PhiIn1v ,PhiOut1v,hv]"];
fSpirale1UnderSpirale1Length[NoLv_, PhiIn1v_, PhiOut1v_, hv_] :=
  fSpiraleLen[NoLv, PhiIn1v, PhiOut1v - 2 Pi, hv];
fSpirale1OverSpirale1Length[NoLv_, PhiIn1v_, PhiOut1v_, hv_] :=
  fSpiraleLen[NoLv, PhiIn1v + 2 Pi , PhiOut1v, hv];
fSpirale1UnderSpirale2Length[NoLv_, PhiIn1v_, PhiOut1v_, hv_, dPhi12v_] :=
  fSpiraleLen[NoLv, PhiIn1v, PhiOut1v - 2 Pi / NoLv, hv];
fSpirale10verSpirale2Length[NoLv_, PhiIn1v_, PhiOut1v_, hv_, dPhi12v_] :=
  fSpiraleLen[NoLv, PhiInlv + 2 Pi/NoLv + dPhi12v, PhiOut1v, hv];
fSpirale2UnderSpirale1Length[NoLv_, PhiIn1v_, PhiOut1v_, hv_, dPhi12v_] :=
  fSpiraleLen[NoLv, PhiIn1v, PhiOut1v - 2 Pi / NoLv, hv];
fSpirale2OverSpirale1Length[NoLv_, PhiIn1v_, PhiOut1v_, hv_, dPhi12v_] :=
  fSpiraleLen[NoLv, PhiIn1v - dPhi12v + 2 Pi/NoLv, PhiOut1v - dPhi12v, hv];
Print[" These functiona are not required, dut could be helpful),"];
```

```
If[NoLp == 1,
Print["fSpirale1UnderSpirale1Length[1,PhiIn1v ,PhiOut1v,hv]="];
Print[" =fSpiraleLen[NoLv,PhiIn1v ,PhiOut1v-2Pi,hv]=",
   fSpiraleLen[NoLv, PhiIn1v, PhiOut1v - 2 Pi, hv], ";"];
Print[" fSpirale1UnderSpirale1Length[NoLp,PhiIn1p
     ,PhiOutlp,hp]=fSpirale1UnderSpirale1Length[", NoLp, ",",
   PhiIn1p / (2 Pi), "(2Pi),", PhiOut1p / (2 Pi), "(2Pi),", hp /nm, "nm] ="];
Print[" =", fSpirale1UnderSpirale1Length[NoLp, PhiIn1p, PhiOut1p, hp] /nm, "nm."];
Print["fSpirale1OverSpirale1Length[1,PhiIn1v ,PhiOut1v,hv]="];
Print[" =fSpiraleLen[NoLv,PhiIn1v+2Pi ,PhiOut1v,hv]=",
   fSpiraleLen[NoLv, PhiIn1v + 2 Pi , PhiOut1v, hv], ";"];
Print[" fSpirale1OverSpirale1Length[NoLp,PhiIn1p
     ,PhiOutlp,hp]=fSpirale1OverSpirale1Length[", NoLp, ",",
   PhiIn1p/(2 Pi), "(2Pi),", PhiOut1p/(2 Pi), "(2Pi),", hp/nm, "nm] ="];
Print[" =", fSpirale1OverSpirale1Length[NoLp, PhiIn1p, PhiOut1p, hp] /nm, "nm."];
];
If[NoLp == 2,
Print[" fSpirale1UnderSpirale2Length[1,PhiIn1v ,PhiOut1v,hv,dPhi12v]="];
Print["
         fSpiraleLen[NoLv, PhiIn1v, PhiOut1v -2 Pi/NoLv, hv] = ",
   fSpirale1UnderSpirale2Length[1, PhiIn1v , PhiOut1v, hv, dPhi12v], ";"];
Print["fSpirale1UnderSpirale2Length[NoLp,PhiIn1p
     ,PhiOut1p,hp,dPhi12p]=fSpirale1UnderSpirale2Length[",
   NoLp, ",", PhiInlp/(2 Pi), "(2Pi),", PhiOutlp/(2 Pi), "(2Pi),", hp/nm,
   "nm,", dPhi12p/(2 Pi), "(2Pi)] ="];
Print["
        =", fSpirale1UnderSpirale2Length[NoLp, PhiIn1p, PhiOut1p, hp, dPhi12p]/nm,
   "nm."];
\label{print} {\tt Print["fSpirale1OverSpirale2Length[1,PhiIn1v],PhiOut1v,hv,dPhi12v]="];}
Print[" =fSpiraleLen[NoLv,PhiIn1v+Pi ,PhiOut1v,hv]=",
   fSpiraleLen[NoLv, PhiIn1v + Pi , PhiOut1v, hv], ";"];
Print["fSpirale10verSpirale2Length[NoLp,PhiIn1p
     ,PhiOut1p,hp]=fSpirale1OverSpirale1Length[", NoLp, ",",
   PhiIn1p / (2 Pi), "(2Pi),", PhiOut1p / (2 Pi), "(2Pi),", hp /nm, "nm] ="];
Print[" =", fSpirale1OverSpirale2Length[NoLp, PhiIn1p, PhiOut1p, hp, dPhi12p] / nm,
   "nm."];
Print[""];
Print[" fSpirale2UnderSpirale1Length[1,PhiIn1v ,PhiOut1v,hv]="];
Print[" fSpiraleLen[NoLv,PhiIn1v, PhiOut1v -2 Pi/NoLv,hv]=",
   fSpirale1UnderSpirale2Length[1, PhiIn1v, PhiOut1v, hv, dPhi12v], ";"];
Print["fSpirale2UnderSpirale1Length[NoLp,PhiIn1p
     ,PhiOutlp,hp,dPhi12p]=fSpirale2UnderSpirale1Length[",
   NoLp, ",", PhiInlp/(2 Pi), "(2Pi),", PhiOutlp/(2 Pi), "(2Pi),", hp/nm,
   "nm,", dPhi12p/(2 Pi), "(2Pi)] ="];
Print[" =", fSpirale1UnderSpirale2Length[NoLp, PhiIn1p, PhiOut1p, hp, dPhi12p]/nm,
   "nm."];
Print["
        fSpirale1OverSpirale2Length[1,PhiIn1v ,PhiOut1v,hv]="];
        =fSpiraleLen[NoLv,PhiIn1v+Pi ,PhiOut1v,hv]=",
Print["
   fSpiraleLen[NoLv, PhiIn1v + Pi , PhiOut1v, hv], ";"];
Print["fSpirale1OverSpirale2Length[NoLp,PhiIn1p
     ,PhiOutlp,hp]=fSpirale1OverSpirale2Length[", NoLp, ",",
   PhiIn1p/(2 Pi), "(2Pi),", PhiOut1p/(2 Pi), "(2Pi),", hp/nm, "nm] ="];
Print[" =", fSpirale10verSpirale2Length[NoLp, PhiIn1p, PhiOut1p, hp, dPhi12p] /nm,
   "nm."];
1:
Print[" II.2.4. The function fRIn1Sharp[NoLv,L1v,hv]"]
```

```
fRIn1Sharp[NoLv_{,} L1v_{,} hv_{]} := (L1v/(2Pi) - (NoLvhv/2));
Print["fRIn1Sharp[NoLv,L1v,hv]=", fRIn1Sharp[NoLv, L1v, hv]];
Print["is a good approximation to obtain
    the value of the sharp in the dependence ScrollEnergy[RIn]"];
Print["fRIn1Sharp[", NoLp, ", ", L1p/nm, "nm, ", hp/nm, "nm] = ",
  fRIn1Sharp[NoLp, L1p, hp] / nm, "nm"];
Print["-----"];
Print[" III. Begin of Calculation "];
If[NoLp == 1,
Print[" III.1. The inner and the outer angle of the spirale of the layer:"]];
If[NoLp == 2,
Print[" III.1. The inner and the outer angles of the spirales of the layers:"]];
 \texttt{Print}[" \varphi \texttt{In1}=", \texttt{RIn1} 2 \texttt{Pi} / (\texttt{NoLp} \ \texttt{h}) \ , ", \varphi \texttt{Out1}=\texttt{fPhiOutvsPhiInLh}[", \texttt{NoLp}, ", \varphi \texttt{In1}, \texttt{L1}, \texttt{h}] \ ; "]; 
fPhiIn1[NoLv_, RIn1v_, hv_] := RIn1v 2 Pi / (NoLv hv);
PhiIn1p = fPhiIn1[NoLp, RIn1p, hp];
fPhiOut1[NoLv_, L1v_, RIn1v_, hv_] :=
fPhiOutvsPhiInLh[NoLv, fPhiIn1[NoLv, RIn1v, hv], L1v, hv];
Print[" For RIn1=", RIn1p/nm, "nm,h=", hp/nm, "nm:"];
PhiOutlp = fPhiOutl[NoLp, Llp, RInlp, hp];
ROut1p = PhiOut1p NoLp hp / (2 Pi);
 {\tt Print[" $\phi$In1=", $PhiIn1p/(2 Pi), "(2Pi), $\Psi$Out1=", $PhiOut1p/(2 Pi), "(2Pi)."]; } 
fPhiIn2[NoLv_, RIn1v_, hv_, dPhi12v_] :=
fPhiIn1[NoLv, RIn1v, hv] + dPhi12v;
PhiIn2dPhi12p =
fPhiIn2[NoLp, RIn1p, hp, 0]; (*www orig 2022.10*)
PhiIn2dPhi12p =
fPhiIn2[NoLp, RIn1p, hp, dPhi12p]; (* for dPhi12p!=0, checked 2022.10*)
PhiIn2dPhi12Pip =
fPhiIn2[NoLp, RIn1p, hp, Pi];
fPhiOut2[NoLv_, L1v_, RIn1v_, hv_, dPhi12v_] :=
fPhiOutvsPhiInLh[NoLv, fPhiIn2[NoLv, RIn1v, hv, dPhi12v], L1v, hv];
PhiOut2dPhi12p =
fPhiOut2[NoLp, L1p, RIn1p, hp, dPhi12p];
If[NoLp == 2, Print[" \varphi In2=", PhiIn2dPhi12p / (2 Pi) ,
   "(2Pi),\varphiOut2=", PhiOut2dPhi12p/(2Pi), "(2Pi)"];
PhiOut2dPhi12Pip =
fPhiOut2[NoLp, L1p, RIn1p, hp, Pi];
Print[" for d\phi12=Pi: PhiIn2=", PhiIn2dPhi12Pip/(2 Pi),
   "(2Pi), φOut2=", PhiOut2dPhi12Pip/(2Pi), "(2Pi)"];];
Print["L1=", L1p/nm, "nm, RIn1=", RIn1p/nm, "nm"];
If[NoLp == 1, Print[" Plot the Spirale of the layer:"]];
If[NoLp == 2, Print[" Plot Spirales of the layers:"]]; "for d\phi 12=0";
Spirale1Plot =
PolarPlot[(Phiv) NoLp hp / (2 Pi) /nm, {Phiv, PhiIn1p,
 PhiOut1p},
 PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
  1.1 ROut1p / nm } } , PlotStyle -> {Red, Thin} , Axes -> None];
If[NoLp == 1, Print[Show[Spirale1Plot]];];
If[NoLp > 1,
Print[" Plot the Spirale of the layers:"];
Spirale2Plot = PolarPlot[(Phiv - Pi) NoLp hp / (2 Pi) / nm,
   {Phiv, PhiIn2dPhi12p + Pi, PhiOut2dPhi12p + Pi},
 PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
   1.1 ROut1p / nm}}, PlotStyle -> {Blue, Thin}, Axes -> None];
Spirale2dPhi12PiPlot = PolarPlot[(Phiv - Pi) NoLp hp / (2 Pi) / nm,
   {Phiv, PhiIn2dPhi12p + Pi, PhiOut2dPhi12Pip + Pi},
```

```
PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
      Print[Show[{Spirale1Plot, Spirale2Plot}]];
If[NoLp == 1,
 Spirale1OverSpirale1Plot =
   If[PhiIn1p + 2 Pi < PhiOut1p,</pre>
   PolarPlot[(Phiv) NoLp hp / (2 Pi) /nm, {Phiv, PhiIn1p + 2 Pi,
    PlotRange \rightarrow \{\{-1.1\,ROutlp/nm,\ 1.1\,ROutlp/nm\},\ \{-1.1\,ROutlp/nm,\ 1.1\,ROutlp/nm\},\ \{-1.1\,ROutlp/nm,\ 1.1\,ROutlp/nm\},\ \{-1.1\,ROutlp/nm\},\ \{-1.1\,R
       1.1 ROut1p / nm } } , PlotStyle -> {Red, Thick}, Axes -> None], {}];
  Spirale1UnderSpirale1Plot =
   If[PhiIn1p < PhiOut1p - 2 Pi,</pre>
   PolarPlot[(Phiv) NoLphp/(2Pi)/nm, {Phiv, PhiIn1p,
     PhiOut1p - 2 Pi},
    PlotRange \rightarrow {\{-1.1 \text{ ROutlp/nm}, 1.1 \text{ ROutlp/nm}\}, \{-1.1 \text{ ROutlp/nm}, 1.1 \text{ ROutlp/nm}\}
       1.1 ROutlp/nm}}, PlotStyle -> {Red, Thick}, Axes -> None], {}];
 Print[" {Spirale1UnderSpirale1}, {Spirale1.Spirale1OverSpirale1}:"];
 Print[Show[{Spirale1Plot, Spirale1UnderSpirale1Plot}],
  Show[{Spirale1Plot, Spirale1OverSpirale1Plot}]];
1;
If[NoLp == 2,
Spirale1UnderSpirale2dPhi120Plot =
  If[PhiIn1p < PhiOut2dPhi12p - Pi,</pre>
   PolarPlot[(Phiv) NoLphp/(2Pi)/nm, {Phiv, PhiIn1p,
     PhiOut2dPhi12p - Pi}, PlotStyle -> {Red, Thick},
    PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
       1.1 ROut1p / nm } } ] , { } ];
Spirale10verSpirale2dPhi120Plot =
  If[PhiIn1p + Pi < PhiOut1p,</pre>
   PolarPlot[(Phiv) NoLp hp / (2 Pi) / nm, {Phiv,
     PhiIn1p + Pi +
      dPhi12p, PhiOut1p},
    PlotStyle -> {Red, Thick},
    PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
       1.1 ROut1p/nm}}], {}];
Spirale2UnderSpirale1dPhi120Plot =
  If[PhiIn2dPhi12p + Pi < PhiOut1p,</pre>
   PolarPlot[(Phiv - Pi) NoLp hp / (2 Pi) / nm, {Phiv,
     PhiIn2dPhi12p + Pi, PhiOut1p}, PlotStyle -> {Blue, Thick},
    PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
       1.1 ROut1p/nm}}], {}];
Spirale2OverSpirale1dPhi12OPlot =
  If[2 Pi + PhiIn2dPhi12p -
     dPhi12p <
    PhiOut2dPhi12p + Pi,
   PolarPlot[(Phiv - Pi) NoLphp/(2 Pi)/nm, {Phiv,
     2 Pi + PhiIn2dPhi12p -
      dPhi12p,
     PhiOut2dPhi12p + Pi}, PlotStyle -> {Blue, Thick},
    PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
       1.1 ROut1p/nm}}], {}];
Print[
      "Plot Spirales for dPhi12=Pi (could be NotRequired, dPhi12=0 in this program)"];
Spirale1UnderSpirale2dPhi12PiPlot =
  If[PhiIn1p < PhiOut2dPhi12Pip - Pi,</pre>
   PolarPlot[(Phiv) NoLphp/(2Pi)/nm, {Phiv, PhiIn1p,
```

```
PhiOut2dPhi12Pip - Pi}, PlotStyle -> {Red, Thick},
  PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
   1.1 ROutlp/nm}}], {}];
Spirale10verSpirale2dPhi12PiPlot =
 If[PhiIn1p + Pi +
  dPhi12p < PhiOut1p,
 PolarPlot[(Phiv) NoLp hp / (2 Pi) /nm, {Phiv,
  PhiIn1p + Pi +
   dPhi12p, PhiOut1p},
  PlotStyle -> {Red, Thick},
  PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
   1.1 ROutlp/nm}}], {}];
Spirale2UnderSpirale1dPhi12PiPlot =
 If[PhiIn2dPhi12p + Pi < PhiOut1p,</pre>
 PolarPlot[(Phiv - Pi) NoLp hp / (2 Pi) / nm, {Phiv,
  PhiIn2dPhi12Pip + Pi, PhiOut1p}, PlotStyle -> {Blue, Thick},
  PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
   1.1 ROut1p/nm}}], {}];
Spirale2OverSpirale1dPhi12PiPlot =
 If[2 Pi + PhiIn2dPhi12Pip -
  dPhi12p <
  PhiOut2dPhi12Pip + Pi,
 PolarPlot[(Phiv - Pi) NoLphp/(2 Pi)/nm, {Phiv,
  2 Pi + PhiIn2dPhi12Pip -
  dPhi12p,
  PhiOut2dPhi12Pip + Pi}, PlotStyle -> {Blue, Thick},
  PlotRange -> {{-1.1 ROutlp/nm, 1.1 ROutlp/nm}, {-1.1 ROutlp/nm,
   1.1 ROut1p / nm } } ] , { } ];
 Print[" {Spirale1,Spirale2,Spirale1UnderSpirale2,Spirale2UnderSpirale1},"];
             {Spirale1,Spirale2,Spirale1OverSpirale2,Spirale2OverSpirale1}"];
 Print[" for dPhi12=0: ", Show[Spirale1Plot, Spirale2Plot],
 Show[Spirale1Plot, Spirale2Plot, Spirale1UnderSpirale2dPhi120Plot,
 Spirale2UnderSpirale1dPhi120Plot],
 Show[Spirale1Plot, Spirale2Plot, Spirale1OverSpirale2dPhi12OPlot,
 Spirale2OverSpirale1dPhi120Plot]];
 Print[" for dPhi12=Pi: ", Show[Spirale1Plot, Spirale2dPhi12PiPlot],
 Show[Spirale1Plot, Spirale2dPhi12PiPlot, Spirale1UnderSpirale2dPhi12PiPlot,
 Spirale2UnderSpirale1dPhi12PiPlot],
 Show[Spirale1Plot, Spirale2dPhi12PiPlot, Spirale1OverSpirale2dPhi12PiPlot,
 Spirale2OverSpirale1dPhi12PiPlot]];
1;
Print[" III.2. The nanoscroll energy calculation"];
Print[" III.2.1. The elastic energy calculation"];
fEelastCC[NoLv_, Lwv_, Llv_, RInlv_, hv_, aCCv_, CCv_] :=
Module[{},
 Return[2 Pi CCv Lwv / (hv fSa[aCCv]) fElast[
   fPhiIn1[NoLv, RIn1v, hv],
   fPhiOut1[NoLv, L1v, RIn1v, hv]]];];
fEelastCBN[NoLv_, Lwv_, L1v_, RIn1v_, hv_, aCCv_, CBNv_] :=
Module[{},
 Return[2 Pi CBNv Lwv / (hv fSa[aCCv]) fElast[
   fPhiIn1[NoLv, RIn1v, hv],
   fPhiOut1[NoLv, Llv, RInlv, hv]]];];
EelastCCp = fEelastCC[NoLp, Lwp, L1p, RIn1p, hp, aCCp, CCp];
EelastCBNp = fEelastCBN[NoLp, Lwp, L1p, RIn1p, hp, aCCp, CBNp];
```

```
Print[" EelastC=", EelastCCp / (eV / atom), "eV / atom"];
If[NoLp == 2, Print[" EelastBN=", EelastCBNp / (eV /atom), "eV/atom"];];
Print[" III.2.2. The Van-der-Waals energy calculation"];
"The definition of the function ";
"'fEVdWLayer10verlap[NoLv,Lwv,L1v, RIn1v, hv, aCCv, epsVdWv]'";
"(Note: This function is omitted at calculations";
     but could be helpful at
   calculation of VdW ebergy of monoscroll at debugging;";
     for example,";
     fEVdWLayer1Overlap[NoL1,Lwp,15nm, 2nm, hp, aCCp, epsVdWp] ";
" and fEVdWLayersOverlap[NoL2,Lwp,L1p=15nm, 2nm, hp, aCCp, epsVdWp, 0]";
     give the same values";
fEVdWLayer1Overlap[NoLv_, Lwv_, L1v_, RIn1v_, hv_, aCCv_, epsVdWv_] := Module[
EVdW1Un1v = 0 (eV / atom) , EVdW1Ov1v = 0 (eV / atom) ,
Spirale1UnderSpirale1Length = 0 nm, Spirale1OverSpirale1Length = 0 nm,
PhiIn1v = fPhiIn1[NoLv, RIn1v, hv],
PhiOutlv = fPhiOutl[NoLv, Llv, RInlv, hv],
Spirale1OverSpirale1Length = fSpiraleLen[NoLv, PhiIn1v + 2 Pi, PhiOut1v, hv];
Spirale1UnderSpirale1Length = fSpiraleLen[NoLv, PhiIn1v, PhiOut1v - 2 Pi, hv];
"Note: Spirale1OverSpirale1Length>Spirale1UnderSpirale1Length";
EVdWlUnlv = -epsVdWvLwv/(2 fSa[aCCv]) SpiralelUnderSpiralelLength;
EVdW1Ov1v = -epsVdWv Lwv / (2 fSa[aCCv]) Spirale1OverSpirale1Length;
EVdWv = (EVdW1Un1v + EVdW10v1v);
Return[{EVdWv, EVdW1Un1v, EVdW1Ov1v}];
"The definition of the function";
"fEVdWLayersOverlap[NoLv_,Lwv_,Llv_, RInlv_, hv_, aCCv_, epsVdWv_, dPhi12v_]";
fEVdWLayersOverlap[NoLv_, Lwv_, Llv_, RInlv_, hv_, aCCv_, epsVdWv_, dPhil2v_] := Module[
EVdW1Un1 = 0 (eV /atom) , EVdW1Ov1 = 0 (eV /atom) ,
EVdW1Un2 = 0 (eV/atom), EVdW1Ov2 = 0 (eV/atom),
EVdW2Un1 = 0 (eV/atom), EVdW2Ov1 = 0 (eV/atom),
Spirale1UnderSpirale1Length = 0 nm, Spirale1OverSpirale1Length = 0 nm,
Spirale1UnderSpirale2Length = 0 nm, Spirale1OverSpirale2Length = 0 nm,
Spirale2UnderSpirale1Length = 0 nm, Spirale2OverSpirale1Length = 0 nm,
PhiIn1 = fPhiIn1[NoLv, RIn1v, hv],
PhiIn2 = fPhiIn2[NoLv, RIn1v, hv, dPhi12v],
PhiOut1 = fPhiOut1[NoLv, Llv, RIn1v, hv],
PhiOut2 = fPhiOut2[NoLv, L1v, RIn1v, hv, dPhi12v],
ReturnEnergiesv = {1, 2, 3, 4, 5, 6, 7}
},
If[NoLv == 1,
If[PhiIn1 < PhiOut1 - 2 Pi,</pre>
     Spirale1UnderSpirale1Length = fSpiraleLen[NoLv, PhiIn1, PhiOut1 - 2 Pi, hv];];
If[PhiIn1 + 2 Pi < PhiOut1, Spirale1OverSpirale1Length =</pre>
       fSpiraleLen[NoLv, PhiIn1 + 2 Pi, PhiOut1, hv];];
EVdW1Un1 = -epsVdWv Lwv / (2 fSa[aCCv]) Spirale1UnderSpirale1Length;
EVdW10v1 = -epsVdWv Lwv / (2 fSa[aCCv]) Spirale10verSpirale1Length;
EVdW = (EVdW1Un1 + EVdW1Ov1);
ReturnEnergiesv = {EVdW, EVdW1Un1, EVdW1Ov1};
If[NoLv == 2,
 If[PhiIn1 < PhiOut2 - Pi,</pre>
     Spirale1UnderSpirale2Length = fSpiraleLen[NoLv, PhiIn1, PhiOut2 - Pi, hv];];
 If[PhiIn1 + Pi + dPhi12v < PhiOut1, Spirale1OverSpirale2Length =</pre>
```

```
fSpiraleLen[NoLv, PhiIn1 + Pi + dPhi12v, PhiOut1, hv];];
 If[PhiIn1 + dPhi12v < PhiOut1 - Pi, Spirale2UnderSpirale1Length =</pre>
       fSpiraleLen[NoLv, PhiIn1 + dPhi12v, PhiOut1 - Pi, hv];];
  If[PhiIn1 - dPhi12v + Pi < PhiOut2 - dPhi12v, Spirale2OverSpirale1Length =</pre>
       fSpiraleLen[NoLv, PhiIn1 - dPhi12v + Pi, PhiOut2 - dPhi12v, hv];];
EVdW1Un2 = -epsVdWv Lwv / (2 fSa[aCCv]) Spirale1UnderSpirale2Length;
  EVdW1Ov2 = -epsVdWv Lwv / (2 fSa[aCCv]) Spirale10verSpirale2Length;
  EVdW2Un1 = -epsVdWv Lwv / (2 fSa[aCCv]) Spirale2UnderSpirale1Length;
  EVdW2Ov1 = -epsVdWv Lwv / (2 fSa[aCCv]) Spirale2OverSpirale1Length;
  EVdW = (EVdW1Un2 + EVdW1Ov2 + EVdW2Un1 + EVdW2Ov1);
  ReturnEnergiesv[[iEVdW]] = EVdW;
  ReturnEnergiesv[[iEVdW1Un2]] = EVdW1Un2;
  ReturnEnergiesv[[iEVdW10v2]] = EVdW10v2;
  ReturnEnergiesv[[iEVdW2Un1]] = EVdW2Un1;
  ReturnEnergiesv[[iEVdW2Ov1]] = EVdW2Ov1;
Return[ReturnEnergiesv];
EVdWdPhi12eq0allp =
 fEVdWLayersOverlap[NoL2, Lwp, L1p, RIn1p, hp, aCCp, epsVdWp, dPhi12eq0];
EVdWvardPhi12allp =
fEVdWLayersOverlap[NoLp, Lwp, L1p, R1n1p, hp, aCCp, epsVdWp, dPhi12p];
If[NoLp == 1,
Print[" EVdWvardPhi12allp[[iEVdW]]=",
   EVdWvardPhi12allp[[iEVdW]] / (eV / atom) , "eV/atom"];
Print["( EVdWvardPhi12allp[[iEVdW1Un1]]=",
   EVdWvardPhi12allp[[iEVdW1Un1]] / (eV / atom) , "eV/atom"];
Print[" EVdWvardPhi12allp[[iEVdW1Ov1]]=",
   EVdWvardPhi12allp[[iEVdW1Ov1]] / (eV / atom) , "eV/atom )"];
If[NoLp == 2,
Print[" for dPhi12=", dPhi12p/Pi, "Pi EVdWvardPhi12allp[[iEVdW]]=",
 EVdWvardPhi12allp[[iEVdW]] / (eV / atom) , "eV/atom"];
Print[" For dPhi12=", dPhi12eq0/Pi, "Pi:"];
Print[" EVdWvardPhi12allp[[iEVdW]]=",
 EVdWdPhi12eq0allp[[iEVdW]] / (eV / atom) , "eV/atom"];
 Print[" EVdWvardPhi12allp[[iEVdW1Un2]]=",
 EVdWdPhi12eq0allp[[iEVdW1Un2]] / (eV / atom) , "eV/atom"];
 Print[" EVdWvardPhi12allp[[iEVdW10v2]]=",
 EVdWdPhi12eq0allp[[iEVdW1Ov2]] / (eV / atom) , "eV/atom"];
 Print[" EVdWvardPhi12allp[[iEVdW2Un1]]=",
 EVdWdPhi12eq0allp[[iEVdW2Un1]] / (eV / atom) , "eV/atom"];
 Print[" EVdWvardPhi12allp[[iEVdW2Ov2]]=",
 EVdWdPhi12eq0allp[[iEVdW2Ov1]] / (eV / atom) , "eV/atom"];
EVdWdPhi12eqPiallp =
 fEVdWLayersOverlap[NoLp, Lwp, L1p, RIn1p, hp, aCCp, epsVdWp, dPhi12eqPi];
(**) Print[" For dPhi12=", dPhi12eqPi/Pi, "Pi:"];
 Print[" EVdWvatdPhi12allp[[iEVdW]]=",
 EVdWdPhi12eqPiallp[[iEVdW]] / (eV /atom) , "eV/atom"];
 Print[" EVdWvatdPhi12allp[[iEVdW1Un2]]=",
 EVdWdPhi12eqPiallp[[iEVdW1Un2]] / (eV / atom) , "eV/atom"];
 Print[" EVdWvatdPhi12allp[[iEVdW10v2]]=",
 EVdWdPhi12eqPiallp[[iEVdW1Ov2]] / (eV / atom) , "eV/atom"];
 Print[" EVdWvatdPhi12allp[[iEVdW2Un1]]=",
 EVdWdPhi12eqPiallp[[iEVdW2Un1]] / (eV / atom) , "eV/atom"];
 Print[" EVdWvatdPhi12allp[[iEVdW2Ov2]]=",
```

```
EVdWdPhi12eqPiallp[[iEVdW2Ov1]] / (eV / atom) , "eV/atom"];
EVdWEVdWdPhi12eq0p = EVdWdPhi12eq0allp[[iEVdW]];
Print[" EVdWdPhi12eq0allp=", EVdWdPhi12eq0allp/(eV/atom),
 "eV/atom"];
EVdWEVdWdPhi12eqPip = EVdWvardPhi12allp[[iEVdW]];
Print[" EVdWEVdWdPhi12eqPip=", EVdWEVdWdPhi12eqPip/(eV/atom),
 "eV/atom"l;
(**)
1;
If[NoLp == 2, Print[" III.3. The energy of flat planes "];];
fEnergyFlatPlanes[NoLv_, Lwv_, L1v_, aCCv_, epsVdWv_] :=
  If[NoLv == 2, -epsVdWv Lwv / fSa[aCCv] L1v, 0 eV / atom];
EnergyFlatPlanesp = fEnergyFlatPlanes[NoLp, Lwp, L1p, aCCp, epsVdWp];
If[NoLp == 2, Print[" EnergyFlatPlanes=-eps width/Sa L1(NoL-1) =",
EnergyFlatPlanesp / (eV / atom) , "eV/atom"];];
Print[" III.4. The total energy of the nanoscroll"];
fScrollEnergydPhi[NoLv_, Lwv_, Llv_,
   RIn1v_, hv_, aCCv_, epsVdWv_, CCv_, CBNv_, dPhi12v_] :=
Module[{ScrollEnergyv, EVdWv, EVdWnoDimv},
EVdWv = fEVdWLayersOverlap[NoLv, Lwv, L1v, RIn1v, hv, aCCv, epsVdWv, dPhi12v][[1]];
EVdWnoDimv = EVdWv / . \{eV \rightarrow 1, atom \rightarrow 1, nm \rightarrow 1\};
If[NoLv == 1,
  If[EVdWnoDimv == 0, ScrollEnergyv = fEelastCC[NoLv, Lwv, Llv, RInlv, hv, aCCv, CCv],
    ScrollEnergyv = EVdWv + fEelastCC[NoLv, Lwv, Llv, RIn1v, hv, aCCv, CCv];];
 1;
If[NoLv == 2,
If[EVdWnoDimv == 0,
   ScrollEnergyv = fEelastCC[NoLv, Lwv, Llv, RIn1v, hv, aCCv, CCv] +
   fEelastCBN[NoLv, Lwv, Llv, RInlv, hv, aCCv, CCv],
   ScrollEnergyv = EVdWv + fEelastCC[NoLv, Lwv, Llv, RInlv, hv, aCCv, CCv] +
   fEelastCBN[NoLv, Lwv, L1v, RIn1v, hv, aCCv, CCv];];
Return[ScrollEnergyv];
];
fScrollEnergyVdWandElast[NoLv_, Lwv_,
   L1v_, RIn1v_, hv_, aCCv_, epsVdWv_, CCv_, CBNv_] :=
Module[{ ScrollEnergyVdWandElastv, EVdWv},
(*If[NoLv == 1.
EVdWv=fEVdWLayer1Overlap[NoLv,Lwv,L1v,RIn1v,hv,aCCv,epsVdWv][[1]];
1;
If[NoLv == 2,
EVdWv=fEVdWLayersOverlap[NoLv,Lwv,Llv,RInlv,hv,aCCv,epsVdWv][[1]];
EVdWv = fEVdWLayer10verlap[NoLv, Lwv, L1v, RIn1v, hv, aCCv, epsVdWv][[1]];
If[NoLv == 1,
ScrollEnergyVdWandElastv = EVdWv + fEelastCC[NoLv, Lwv, Llv, RInlv, hv, aCCv, CCv];];
If[NoLv == 2,
ScrollEnergyVdWandElastv = EVdWv + fEelastCC[NoLv, Lwv, Llv, RInlv, hv, aCCv, CCv] +
       fEelastCC[NoLv, Lwv, Llv, RIn1v, hv, aCCv, CBNv];];
Return[ScrollEnergyVdWandElastv];
fScrollEnergy[NoLv_, Lwv_, L1v_, RIn1v_, hv_, aCCv_, epsVdWv_, CCv_, CBNv_] := Module[
{ScrollEnergyv = -10^20 eV / atom},
If[RIn1v/m <= fRIn1Sharp[NoLv, L1v, hv]/m,</pre>
```

```
ScrollEnergyv =
     fScrollEnergyVdWandElast[NoLv, Lwv, Llv, RInlv, hv, aCCv, epsVdWv, CCv, CBNv];
"note: the function fScrollEnergyVdWandElast[1,..] is analytycal";
"whereas the function fScrollEnergy[....] uses the 'If[..]'- function";
If[RIn1v/m>= fRIn1Sharp[NoLv, L1v, hv]/m,
If[NoLv == 1, ScrollEnergyv = fEelastCC[NoLv, Lwv, Llv, RInlv, hv, aCCv, CCv];];
If[NoLv == 2, ScrollEnergyv = fEelastCC[NoLv, Lwv, Llv, RIn1v, hv, aCCv, CCv] +
        fEelastCC[NoLv, Lwv, Llv, RInlv, hv, aCCv, CBNv];];
1;
Return[ScrollEnergyv];
ScrollEnergyp = fScrollEnergy[NoLp, Lwp, Llp, RInlp, hp, aCCp, epsVdWp, CCp, CBNp];
ScrollEnergyp = fScrollEnergy[1, Lwp, L1p, RIn1p, hp, aCCp, epsVdWp, CCp, CBNp];
Print["fScrollEnergy[1,Lwp,Llp, RIn1p,hp, aCCp, epsVdWp,CCp,CBNp]="];
Print["=fScrollEnergy[1, Lw=", Lwp/nm, "nm, L1=",
 L1p/nm, "nm, RIn1=", RIn1p/nm, "nm, h=", hp/nm, "nm,"];
Print[" aCC=", aCCp/nm, "nm, epsVdW=", epsVdWp/(eV/atom),
  "eV/atom, CC=", CCp/(eV AA^2/atom), "(eV AA^2/atom)="];
Print["
                  =", ScrollEnergyp/(eV/atom), "eV/atom"];
Print[" III.5. Determine the inner angles mismatch for the bi-layer nanoscroll
      for the high nanoribbon Length"];
Print[" For L1=", L1p/nm, "nm,RIn=",
 RIn1p/nm, "nm,h=", hp/nm, "nm and dPhi12=0:"];
Print[" ScrollEnergy=", ScrollEnergyp/(eV/atom), "eV/atom"];
Print[" For L1=", L1p/nm, "nm,RIn=",
 RIn1p/nm, "nm,h=", hp/nm, "nm and dPhi12=Pi:"];
Print[" ScrollEnergy=", ScrollEnergyp / (eV / atom) , "eV / atom"];
Print["-----"];
Print[" IV.The potential energy of the nanoscroll"];
Print[" as a function of the inner radius RIn"];
PlotRangep =
Switch[NoLp, 1, PlotRangeMonoScroll, 2, PlotRangeBiScroll];
RIn1Minp =
Switch[NoLp, 1, RIn1MinMonoScroll, 2, RIn1MinBiScroll];
RIn1Maxp =
Switch[NoLp, 1, RIn1MaxMonoScroll, 2, RIn1MaxBiScroll];
PlotRangep =
Switch[NoLp, 1, PlotRangeMonoScroll, 2, PlotRangeBiScroll];
Switch[NoLp, 1, RIn1MaxMonoScroll, 2, RIn1MaxBiScroll];
tL1 = Switch[NoLp, 1, {7.nm, 10.nm, 12.5nm, 15.nm},
 2, {15. nm, 20. nm, 25. nm, 30. nm}];
Print[" NoL=", NoLp];
Print[" epsVdW=", epsVdWp/(eV/atom),
  "eV/atom, C=", CCp/(eV nm^2/atom), "(eV nm^2/atom)",
 "(eV nm^2/atom),aCC=", aCCp/nm, "nm,h=", hp/nm, "nm"];
Print[" Plot ScrollEnergy[RIn1/nm]/(eV/atom) for L1=", L1p/nm,
"nm (NoL=", NoLp, ",Lw=", Lwp/nm, "nm"];
PlotScrollEnergyVsRIn1 =
 Plot[(fScrollEnergy[NoLp, Lwp, L1p, RIn1nmv nm, hp, aCCp, epsVdWp,
   CCp, CBNp]) / (eV /
   atom), {RIn1nmv, RIn1Minp/nm, RIn1Maxp/nm},
```

```
PlotRange -> PlotRangep / (eV / atom) ];
Print[PlotScrollEnergyVsRIn1];
Print[" Plot ScrollEnergy[RIn1/nm]/(eV/atom) for L1=", tL1/nm,
 "nm (NoL=", NoLp, ",w=", Lwp/nm, "nm)"];
PlotScrollEnergyVsRIn1L1th =
Plot[(fScrollEnergy[NoLp, Lwp, tL1[[1]], RIn1nmv nm, hp, aCCp, epsVdWp,
   CCp, CBNp])/(eV/
   atom), {RIn1nmv, RIn1Minp/nm, RIn1Maxp/nm},
   PlotRange -> PlotRangep / (eV / atom) ];
PlotScrollEnergyVsRIn1L2th =
Plot[(fScrollEnergy[NoLp, Lwp, tL1[[2]], RIn1nmvnm, hp, aCCp, epsVdWp,
   \texttt{CCp}, \texttt{CBNp}])/(eV/
   atom), {RIn1nmv, RIn1Minp/nm, RIn1Maxp/nm},
   PlotRange -> PlotRangep / (eV / atom) ];
PlotScrollEnergyVsRIn1L3th =
Plot[(fScrollEnergy[NoLp, Lwp, tL1[[3]], RIn1nmvnm, hp, aCCp, epsVdWp,
   CCp, CBNpl)/(eV/
   atom), {RIn1nmv, RIn1Minp/nm, RIn1Maxp/nm},
   PlotRange -> PlotRangep / (eV / atom) ];
PlotScrollEnergyVsRIn1L4th =
Plot[(fScrollEnergy[NoLp, Lwp, tL1[[4]], RIn1nmv nm, hp, aCCp, epsVdWp,
   CCp, CBNp]) / (eV /
   atom), {RIn1nmv, RIn1Minp/nm, RIn1Maxp/nm},
   PlotRange -> PlotRangep / (eV / atom) ];
Print[Show[{PlotScrollEnergyVsRIn1L1th, PlotScrollEnergyVsRIn1L2th,
 PlotScrollEnergyVsRIn1L3th, PlotScrollEnergyVsRIn1L4th}]];
Print["The examples of using of 'fScrollEnergy[..]' function:"]
Print["fScrollEnergy[NoLp,Lwp,tL1[[1]],RIn1p,hp, aCCp, epsVdWp,CCp, CBNp]=",
  fScrollEnergy[NoLp, Lwp, tL1[[1]], RIn1p, hp, aCCp, epsVdWp, CCp, CBNp]/(eV/atom),
  " eV/atom"];
Print["fScrollEnergy[NoLp,Lwp,tL1[[1]],1nm,hp, aCCp, epsVdWp,CCp, CBNp]=",
  fScrollEnergy[NoLp, Lwp, tL1[[1]], 1 nm, hp, aCCp, epsVdWp, CCp, CBNp]/(eV/atom),
  " eV/atom"];
Print["fScrollEnergy[NoLp,Lwp,7nm,1nm,hp, aCCp, epsVdWp,CCp, CBNp]=",
  fScrollEnergy[NoLp, Lwp, 7. nm, 1. nm, hp, aCCp, epsVdWp, CCp, CBNp] / (eV /atom),
  " eV/atom"];
Print["fEVdWLayer10verlap[1,Lwp,7.nm,1.nm,hp,aCCp,epsVdWp][[1]]=",
  fEVdWLayer1Overlap[1, Lwp, 7. nm, 1. nm, hp, aCCp, epsVdWp][[1]]/(eV/atom),
  "eV/atom (right !=0 value, because the layer overlaps"];
Print["fEVdWLayer10verlap[1,Lwp,7.nm,1.5nm,hp,aCCp,epsVdWp][[1]]=",
  fEVdWLayer10verlap[1, Lwp, 7. nm, 2.5 nm, hp, aCCp, epsVdWp][[1]]/(eV/atom),
  "eV/atom !=0, wrong value of the fEVdWLayer1Overlap[..]
    function because the layer does not not overlap"];
Print["The analytical expressions of the fEVdWLayer10verlap[..] function:"];
Print["--- fEVdWLayer1Overlap[NoL1,Lwv,Lpv,RInv,hv,aCCv,epsVdWv][[1]]: ---"];
Print[fEVdWLayer1Overlap[NoL1, Lwv, Lpv, RInv, hv, aCCv, epsVdWv][[1]]];
Print[];
Print["--- fEVdWLayer10verlap[NoL2,Lwv,Lpv,RInv,hv,aCCv,epsVdWv][[1]]: ---"];
Print[fEVdWLayer1Overlap[NoL2, Lwv, Lpv, RInv, hv, aCCv, epsVdWv][[1]]];
Print[];
Print["The Analytical expression of "];
Print["fScrollEnergyVdWandElast[",
 NoLp, ",Lwv,L1v,RIn1v,hv,aCCv, epsVdWv, CCv,CBNv]:"];
Print[fScrollEnergyVdWandElast[NoLp, Lwv, Llv, RIn1v, hv, aCCv, epsVdWv, CCv, CBNv]];
```

```
Print["V. Export the data of the plots of the nanoscroll energy"];
     Print["
              as a function of the inner radius"];
     Print["The parameters of the output file"];
     Print["The number of the output points = ", npRIn1];
     Print["Export the plot data to the files:"];
     tRIn1nmRegular =
       Table[(RIn1Minp + (RIn1Maxp - RIn1Minp) iiRin / (npRIn1)) / nm, {iiRin, 1, npRIn1}];
     tScrollEnergy = tRIn1nmRegular;
     tScrollEnergyeVatom = tRIn1nmRegular;
     tPlotEvsRin = Table[{}, {ii, 1, Length[tL1]}];
     AllPlotsEVsRin = {};
     For[iiL1 = 1, iiL1 <= Length[tL1], iiL1++,</pre>
     L1pi = tL1[[iiL1]];
     NanoscrollNamep =
        StringJoin["Nanoscroll", ToString[NoLp], "L", ToString[L1pi/nm], "nm"];
     Print["NanoscrollName=", NanoscrollNamep];
     ScrollEnergyFileName = StringJoin["EvsRIn1", NanoscrollNamep, ".dat"];
     Print["ScrollEnergyFileName=", ScrollEnergyFileName];
     For[iiRIn1 = 1, iiRIn1 <= npRIn1, iiRIn1++,</pre>
     RIn1pi = tRIn1nmRegular[[iiRIn1]] nm;
     tScrollEnergy[[iiRIn1]] = fScrollEnergy[NoLp, Lwp, L1pi, RIn1pi, hp, aCCp, epsVdWp,
        CCp, CBNp];
     tScrollEnergyeVatom[[iiRIn1]] = (tScrollEnergy[[iiRIn1]]) / (eV / atom);];
     tPlotEvsRin[[iiL1]] = ListPlot[Transpose[{tRIn1nmRegular, tScrollEnergyeVatom}],
         PlotRange -> PlotRangep / (eV / atom) ];
     Print[tPlotEvsRin[[iiL1]]];
     AllPlotsEVsRin = Join[{AllPlotsEVsRin, tPlotEvsRin[[iiL1]]}];
     CarbonNanoscrollEnergyVsRinFileName = StringJoin[NanoscrollNamep, "dat"];
     Export[ToFileName[NotebookDirectory[], ScrollEnergyFileName],
        Transpose[{Insert[tRIn1nmRegular, "RIn1[nm]", 1],
          Insert[tScrollEnergyeVatom, "E[eV/atom]", 1]}]]
     ];
     Print["Plot ScrollEnergy[RIn1/nm]/(eV/atom) for L1=", tL1/nm,
      "nm (NoL=", NoLp, ",Lw=", Lwp/nm, "nm)"];
     Print[Show[AllPlotsEVsRin]];
 Figure3cd(Layers2L30nmRIn2d1nm.m
  The source of the data of the manuscript
  'Structure and energetics of carbon,
  hexagonal boron nitride, and
  carbon/hexagonal boron nitride
  single-layer and bilayer nanoscrolls'
/ A.I. Siahlo, N.A. Poklonski, A.V. Lebedev,
I.V. Lebedeva, A.M. Popov, S.A. Vyrko,
A.A. Knizhnik, Yu.E. Lozovik
// Phys. Rev. Materials. - 2018. - V. 2,
N^9 3.- P. 036001 (9 pp.).
```

```
[DOI: 10.1103/PhysRevMaterials.2.036001]
 I.0 The Units (nm, meV, AA) \,
I.1. All Input Parameters and Constants-----
I.1.1. The sampling parameters
npRIn1=1000
I.1.2. The Input Geometry Parameters of the system
The carbon nanoribbon length L1=29.678nm
The carbon nanoribbon width Lw=11.8nm
Number of the layers in carbon nanoscroll NoL=2
The length of a carbon nanoribbon L1=29.678nm
The carbon nanoribbon width Lw=1.nm
The inner radius of the nanoscroll RIn1=2.1nm
 I.1.2. The Input Energy Constants
epsVdW - the interlayer interaction energy per one atom of
 the nanoscroll:
epsVdW=0.035eV/atom
C - the bending elastic constant:
CCelast=2.01eV AA^2/atom
CCBNelast=2.01eV AA^2/atom
I.1.3. The Input Geometry constants-----
The interatomic distance aCC and the interlayer distance h
h=0.3354 nm (Interlayer distance)
aCC=0.142nm, h=0.3354nm
NatomsInCell=2
dPhi12 - The difference of the inner angles of the spirales
  of the Layers
 I.5. The parameters for the visualisation
I.6. The parameters of the output file
NanoscrollName=Nanoscroll2L29.678nm
Nanoscroll2L29.678nm.txt
 (The output of the data to a file Is Not Performed)
The number of the output points = 1000
 I.7. The Input Numerical Constants used in the programm
The Indexes used for the work with EVdW[...] function
```

-----End of the Input-----

- II. The derivated parameters and the functions required
- II.1. The derivated parameters

The cell area
$$Sa = \frac{3\sqrt{3} \ aCC^2}{4} = 0.0261938 nm^2$$

- II.2. The required functions-----
- II.2.1. The function $fSpiraleLen[2, \varphiIn, \varphiOut, h]$ defines

the Length of a Spirale with the inner agle φ In and the outer angle φ Out(>= φ In):

$$\text{fSpiraleLen[2, φIn, φOut, h] = } \frac{\text{h}\left(-\varphi\text{In}\,\sqrt{1+\varphi\text{In}^2}\right. + \varphi\text{Out}\,\sqrt{1+\varphi\text{Out}^2}\right. - \text{ArcSinh}[\varphi\text{In}] + \text{ArcSinh}[\varphi\text{Out}]\right)}{2\,\pi}$$

II.2.2. The function $fElast[\varphi In, \rho Out]$ is required to calculate an nanoscrollelastic energy:

$$\texttt{fElast}[\varphi \texttt{In}, \varphi \texttt{Out}] \ = \ \frac{\sqrt{1 + \varphi \texttt{In}^2}}{\varphi \texttt{In}} - \frac{\sqrt{1 + \varphi \texttt{Out}^2}}{\varphi \texttt{Out}} - \texttt{ArcSinh}[\varphi \texttt{In}] + \texttt{ArcSinh}[\varphi \texttt{Out}] \,.$$

II.2.3. Define the function $fPhiOutvsPhiInLh[2, \varphi In, L, h]$.

The function fPhiOutvsPhiInLh[2, φ In,L,h] = $\sqrt{\frac{2 \text{ L} \pi}{\text{L}} + \varphi$ In² is a

good approximation to obtain the value of φOut for the defined $\varphi \text{In,L,h.}$

The inverse function fPhiInvsPhiOutLh[2,
$$\varphi$$
Out, L, h]] = $\sqrt{-\frac{2 \text{ L} \pi}{\text{h}} + \varphi \text{Out}^2}$

could be used in the program applications

if ROut (instead of RIn) is the input parameter of the system.

II.2.4. The functions fSpirale1Under(Over)Spirale1Length[NoLv,PhiIn1v ,PhiOut1v,hv]

These functiona are not required, dut could be helpful),

fSpirale1UnderSpirale2Length[1,PhiIn1v,PhiOut1v,hv,dPhi12v] =

fSpiraleLen[NoLv,PhiIn1v, PhiOut1v -2 Pi/NoLv,hv] =
$$\frac{1}{4 \pi}$$

$$\text{hv} \left(-\text{PhiIn1v} \sqrt{1 + \text{PhiIn1v}^2} + \sqrt{1 + \left(\text{PhiOut1v} - 2 \, \pi \right)^2} \right. \\ \left. \left(\text{PhiOut1v} - 2 \, \pi \right)^2 \right. \\ \left. \left(\text{PhiOut1v}$$

$${\tt ArcSinh[PhiIn1v] + ArcSinh[PhiOut1v - 2\ \pi]}) \textit{;}$$

fSpirale1UnderSpirale2Length[NoLp,PhiIn1p ,PhiOut1p,hp,dPhi12p]=fSpirale1UnderSpirale2Length[

$$2, \frac{\text{PhiIn1p}}{2\pi} (2\text{Pi}), \frac{\text{PhiOut1p}}{2\pi} (2\text{Pi}), 0.3354\text{nm}, 0.(2\text{Pi})] =$$

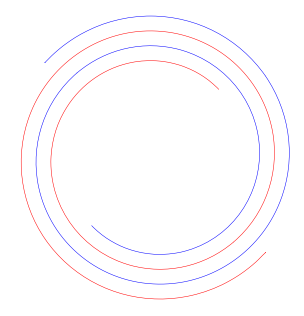
=0.0533806
$$\left(-PhiIn1p\sqrt{1+PhiIn1p^2}\right)$$
 +

$$\sqrt{\text{1 + (PhiOutlp} - \pi)^2} \text{ (PhiOutlp} - \pi) - \text{ArcSinh[PhiInlp]} + \text{ArcSinh[PhiOutlp} - \pi] \right) \text{nm.}$$

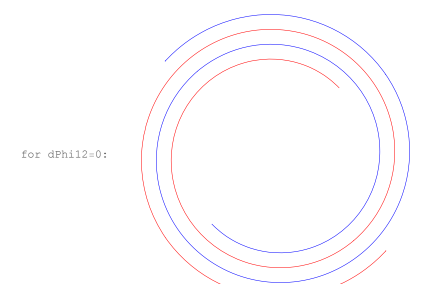
fSpirale1OverSpirale2Length[1,PhiIn1v ,PhiOut1v,hv,dPhi12v] =

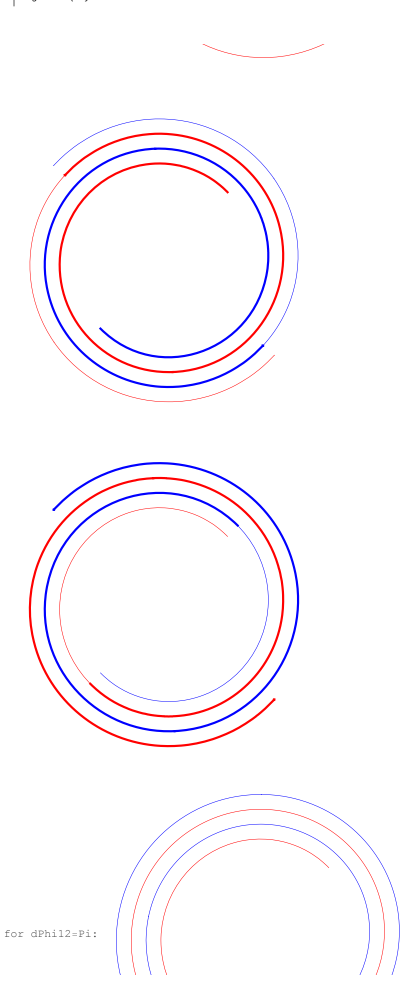
```
=fSpiraleLen[NoLv,PhiIn1v+Pi ,PhiOut1v,hv] = \frac{1}{4\pi}
   \texttt{hv}\,\,\texttt{NoLv}\,\left(\texttt{PhiOutlv}\,\sqrt{\texttt{1}+\texttt{PhiOutlv}^2}\,\,+\,\left(-\texttt{PhiInlv}-\pi\right)\,\sqrt{\texttt{1}+\left(\texttt{PhiInlv}+\pi\right)^2}\,\,+\,\right.
               ArcSinh[PhiOut1v] - ArcSinh[PhiIn1v + \pi];
fSpirale1OverSpirale2Length[NoLp,PhiIn1p ,PhiOut1p,hp]=fSpirale1OverSpirale1Length[
   2, \frac{\text{PhiIn1p}}{2\pi} (2\text{Pi}), \frac{\text{PhiOut1p}}{2\pi} (2\text{Pi}), 0.3354\text{nm}] =
       =0.0533806 \left( (-3.14159 - PhiIn1p) \sqrt{1 + (3.14159 + PhiIn1p)^2} + \right)
           \texttt{PhiOut1p} \sqrt{1 + \texttt{PhiOut1p}^2} - \texttt{ArcSinh} [\texttt{3.14159} + \texttt{PhiIn1p}] + \texttt{ArcSinh} [\texttt{PhiOut1p}] \\ \texttt{nm.}
        fSpirale2UnderSpirale1Length[1,PhiIn1v ,PhiOut1v,hv] =
        fSpiraleLen[NoLv, PhiIn1v, PhiOut1v -2 Pi/NoLv, hv] = \frac{1}{4\pi}
   \text{hv} \left( -\text{PhiIn1v} \sqrt{1 + \text{PhiIn1v}^2} + \sqrt{1 + \left( \text{PhiOut1v} - 2 \, \pi \right)^2} \right) \left( \text{PhiOut1v} - 2 \, \pi \right) - \frac{1}{2} \left( \frac{1}{2} \right) \left( \frac{1}{2
               ArcSinh[PhiIn1v] + ArcSinh[PhiOut1v - 2\pi];
 fSpirale2UnderSpirale1Length[NoLp,PhiIn1p ,PhiOut1p,hp,dPhi12p]=fSpirale2UnderSpirale1Length[
   2, \frac{\text{PhiInlp}}{2\pi} (2Pi), \frac{\text{PhiOutlp}}{2\pi} (2Pi), 0.3354nm, 0.(2Pi)] =
       =0.0533806 \left(-\text{PhiIn1p}\sqrt{1+\text{PhiIn1p}^2}\right)
               \sqrt{1 + (\texttt{PhiOut1p} - \pi)^2} \ (\texttt{PhiOut1p} - \pi) \ - \texttt{ArcSinh}[\texttt{PhiIn1p}] \ + \texttt{ArcSinh}[\texttt{PhiOut1p} - \pi] \ ) \\ \texttt{nm.}
        fSpirale1OverSpirale2Length[1,PhiIn1v ,PhiOut1v,hv]=
        =fSpiraleLen[NoLv,PhiIn1v+Pi ,PhiOut1v,hv] = \frac{1}{4}
   hv NoLv \left( \text{PhiOutlv} \sqrt{1 + \text{PhiOutlv}^2} + (-\text{PhiInlv} - \pi) \sqrt{1 + (\text{PhiInlv} + \pi)^2} \right)
               ArcSinh[PhiOutlv] - ArcSinh[PhiInlv + \pi];
 fSpirale1OverSpirale2Length[NoLp,PhiIn1p ,PhiOut1p,hp]=fSpirale1OverSpirale2Length[
   2, \frac{\text{PhiIn1p}}{2\pi} \text{(2Pi)}, \frac{\text{PhiOut1p}}{2\pi} \text{(2Pi)}, 0.3354\text{nm}] =
       =0.0533806 \left( (-3.14159 - PhiIn1p) \sqrt{1 + (3.14159 + PhiIn1p)^2} + \right)
           PhiOutlp \sqrt{1 + PhiOutlp^{2}} - ArcSinh[3.14159 + PhiInlp] + ArcSinh[PhiOutlp] \\ nm.
    II.2.4. The function fRIn1Sharp[NoLv,Llv,hv]
fRIn1Sharp[NoLv,L1v,hv] = -\frac{\text{hv NoLv}}{2} + \frac{\text{L1v}}{2\pi}
is a good approximation to obtain the value of the sharp in the dependence ScrollEnergy[RIn]
fRIn1Sharp[2, 29.678nm, 0.3354nm] = 4.388nm
```

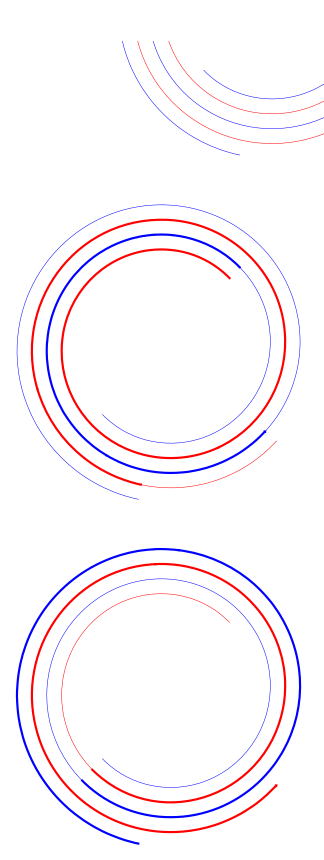
```
III.1. The inner and the outer angles of the spirales of the layers:
 \varphi \texttt{In1} = \frac{\pi \, \texttt{RIn1}}{\texttt{h}}, \ \ \varphi \texttt{Out1} = \texttt{fPhiOutvsPhiInLh[2, } \phi \texttt{In1, } \texttt{L1, } \texttt{h];}
 For RIn1=2.1nm, h=0.3354nm:
 \varphiIn1=3.13059(2Pi),\PhiOut1=4.88707(2Pi).
  \varphiIn2=3.13059(2Pi),\varphiOut2=4.88707(2Pi)
 for dφ12=Pi: PhiIn2=3.63059(2Pi),φOut2=5.2215(2Pi)
L1=29.678nm, RIn1=2.1nm
 Plot Spirales of the layers:
 Plot the Spirale of the layers:
```



Plot Spirales for dPhi12=Pi (could be NotRequired, dPhi12=0 in this program) {Spirale1, Spirale2, Spirale1UnderSpirale2, Spirale2UnderSpirale1}, {Spirale1, Spirale2, Spirale1OverSpirale2, Spirale2OverSpirale1}



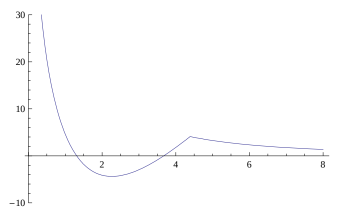




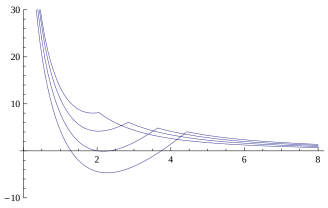
III.2. The nanoscroll energy calculation III.2.1. The elastic energy calculation EelastC=6.40779eV/atom

EelastBN=6.40779eV/atom

```
III.2.2. The Van-der-Waals energy calculation
for dPhi12=0.Pi EVdWvardPhi12allp[[iEVdW]]=-56.7797eV/atom
For dPhi12=0.Pi:
EVdWvardPhi12allp[[iEVdW]]=-56.7797eV/atom
EVdWvardPhi12allp[[iEVdW1Un2]]=-13.3111eV/atom
EVdWvardPhi12allp[[iEVdW10v2]]=-15.0787eV/atom
EVdWvardPhi12allp[[iEVdW2Un1]]=-13.3111eV/atom
EVdWvardPhi12allp[[iEVdW2Ov2]]=-15.0787eV/atom
For dPhi12=1.Pi:
EVdWvatdPhi12allp[[iEVdW]]=-53.3618eV/atom
EVdWvatdPhi12allp[[iEVdW1Un2]]=-17.6025eV/atom
EVdWvatdPhi12allp[[iEVdW1Ov2]]=-9.61051eV/atom
EVdWvatdPhi12allp[[iEVdW2Un1]]=-8.54618eV/atom
EVdWvatdPhi12allp[[iEVdW2Ov2]]=-17.6025eV/atom
EVdWEVdWdPhi12eqPip=-56.7797eV/atom
III.3. The energy of flat planes
EnergyFlatPlanes=-eps width/Sa L1(NoL-1) =-39.6556eV/atom
III.4. The total energy of the nanoscroll
fScrollEnergy[1,Lwp,L1p, RIn1p,hp, aCCp, epsVdWp,CCp,CBNp]=
=fScrollEnergy[1, Lw=1.nm, L1=29.678nm, RIn1=2.1nm, h=0.3354nm,
aCC=0.142nm, epsVdW=0.035eV/atom, CC=2.01(eV AA^2/atom)=
          =-15.3963eV/atom
III.5. Determine the inner angles mismatch for the bi-layer nanoscroll
      for the high nanoribbon Length
For L1=29.678nm, RIn=2.1nm, h=0.3354nm and dPhi12=0:
ScrollEnergy=-15.3963eV/atom
For L1=29.678nm, RIn=2.1nm, h=0.3354nm and dPhi12=Pi:
ScrollEnergy=-15.3963eV/atom
IV. The potential energy of the nanoscroll
as a function of the inner radius RIn
NoL=2
epsVdW=0.035eV/atom, C=0.0201(eV nm^2/atom)(eV nm^2/atom),aCC=0.142nm,h=0.3354nm
Plot ScrollEnergy[RIn1/nm]/(eV/atom) for L1=29.678nm (NoL=2,Lw=1.nm
```



 $\label{eq:plot_scrollenergy} Plot Scrollenergy[RIn1/nm]/(eV/atom) for L1=\{15., 20., 25., 30.\}nm \ (NoL=2, w=1.nm) \\$



The examples of using of 'fScrollEnergy[..]' function: fScrollEnergy[NoLp,Lwp,tL1[[1]],RIn1p,hp, aCCp, epsVdWp,CCp, CBNp]=7.85611 eV/atom fScrollEnergy[NoLp,Lwp,tL1[[1]],1nm,hp, aCCp, epsVdWp,CCp, CBNp]=13.4436 eV/atom fScrollEnergy[NoLp,Lwp,7nm,1nm,hp, aCCp, epsVdWp,CCp, CBNp]=13.1901 eV/atom fEVdWLayer10verlap[1,Lwp,7.nm,1.nm,hp,aCCp,epsVdWp][[1]]= 0.39374eV/atom (right !=0 value, because the layer overlaps fEVdWLayer1Overlap[1,Lwp,7.nm,1.5nm,hp,aCCp,epsVdWp][[1]]= 12.2479eV/atom !=0, wrong value of the fEVdWLayer10verlap[..] function because the layer does not not overlap

The analytical expressions of the fEVdWLayer1Overlap[..] function:

```
--- fEVdWLayer10verlap[NoL1,Lwv,Lpv,RInv,hv,aCCv,epsVdWv][[1]]: ---
```

$$-\frac{1}{6\sqrt{3}} \frac{\text{epsVdWv hv Lwv}}{\text{aCCv}^2 \pi}$$

$$\left[\sqrt{\frac{4 \text{ Lpv } \pi}{\text{hv}} + \frac{4 \pi^2 \text{ RInv}^2}{\text{hv}^2}} \sqrt{1 + \frac{4 \text{ Lpv } \pi}{\text{hv}} + \frac{4 \pi^2 \text{ RInv}^2}{\text{hv}^2}} + \left(-2 \pi - \frac{2 \pi \text{ RInv}}{\text{hv}}\right) \sqrt{1 + \left(2 \pi + \frac{2 \pi \text{ RInv}}{\text{hv}}\right)^2} - \frac{1}{6\sqrt{3} \text{ aCCv}^2 \pi}\right]$$

$$-\frac{1}{6\sqrt{3} \text{ aCCv}^2 \pi} \frac{\text{epsVdWv hv Lwv}}{\text{hv}}$$

$$\left[2 \pi \text{ RInv} \sqrt{1 + \frac{4 \pi^2 \text{ RInv}^2}{\text{hv}^2}} + \left(-2 \pi + \sqrt{\frac{4 \text{ Lpv } \pi}{\text{hv}} + \frac{4 \pi^2 \text{ RInv}^2}{\text{hv}^2}}\right) \sqrt{1 + \left(-2 \pi + \sqrt{\frac{4 \text{ Lpv } \pi}{\text{hv}} + \frac{4 \pi^2 \text{ RInv}^2}{\text{hv}^2}}\right)^2} - \frac{1}{\text{ArcSinh}} \left[2 \pi - \sqrt{\frac{4 \text{ Lpv } \pi}{\text{hv}} + \frac{4 \pi^2 \text{ RInv}^2}{\text{hv}^2}}}\right]$$

--- fEVdWLayer10verlap[NoL2, Lwv, Lpv, RInv, hv, aCCv, epsVdWv][[1]]: ---

$$\frac{1}{3\sqrt{3}\operatorname{accv}^{2}\pi}$$

$$\operatorname{epsVdWv}\operatorname{hv}\operatorname{Lwv}\left(\sqrt{\frac{2\operatorname{Lpv}\pi}{\operatorname{hv}}+\frac{\pi^{2}\operatorname{RInv}^{2}}{\operatorname{hv}^{2}}}\sqrt{1+\frac{2\operatorname{Lpv}\pi}{\operatorname{hv}}+\frac{\pi^{2}\operatorname{RInv}^{2}}{\operatorname{hv}^{2}}}+\left(-2\pi-\frac{\pi\operatorname{RInv}}{\operatorname{hv}}\right)\sqrt{1+\left(2\pi+\frac{\pi\operatorname{RInv}}{\operatorname{hv}}\right)^{2}}\right)$$

$$\operatorname{ArcSinh}\left[2\pi+\frac{\pi\operatorname{RInv}}{\operatorname{hv}}\right]+\operatorname{ArcSinh}\left[\sqrt{\frac{2\operatorname{Lpv}\pi}{\operatorname{hv}}+\frac{\pi^{2}\operatorname{RInv}^{2}}{\operatorname{hv}^{2}}}\right]\right]-\frac{1}{3\sqrt{3}\operatorname{accv}^{2}\pi}\operatorname{epsVdWv}\operatorname{hv}\operatorname{Lwv}$$

$$\left(-\frac{\pi\operatorname{RInv}\sqrt{1+\frac{\pi^{2}\operatorname{RInv}^{2}}{\operatorname{hv}^{2}}}}{\operatorname{hv}}+\left(-2\pi+\sqrt{\frac{2\operatorname{Lpv}\pi}{\operatorname{hv}}+\frac{\pi^{2}\operatorname{RInv}^{2}}{\operatorname{hv}^{2}}}}\right)\sqrt{1+\left(-2\pi+\sqrt{\frac{2\operatorname{Lpv}\pi}{\operatorname{hv}}+\frac{\pi^{2}\operatorname{RInv}^{2}}{\operatorname{hv}^{2}}}\right)^{2}}-\right.$$

$$\operatorname{ArcSinh}\left[\frac{\pi\operatorname{RInv}}{\operatorname{hv}}\right]-\operatorname{ArcSinh}\left[2\pi-\sqrt{\frac{2\operatorname{Lpv}\pi}{\operatorname{hv}}+\frac{\pi^{2}\operatorname{RInv}^{2}}{\operatorname{hv}^{2}}}\right]$$

The Analytical expression of

fScrollEnergyVdWandElast[2,Lwv,L1v,RIn1v,hv,aCCv, epsVdWv, CCv,CBNv]:

$$8 \operatorname{CBNv} \operatorname{Lwv} \pi \left(\frac{\operatorname{hv} \sqrt{1 + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}}}{\pi \operatorname{Rinlv}} - \frac{\sqrt{1 + \frac{e^{2} \operatorname{Liv} \pi}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}}}{\sqrt{\frac{2 \operatorname{Liv} \pi}{\operatorname{Rinlv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}}} - \operatorname{ArcSinh} \left[\frac{\pi \operatorname{Rinlv}}{\operatorname{hv}} \right] + \operatorname{ArcSinh} \left[\sqrt{\frac{2 \operatorname{Liv} \pi}{\operatorname{hv}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}}} \right] + \frac{3 \sqrt{3} \operatorname{aCCv}^{2} \operatorname{hv}}{3 \sqrt{3} \operatorname{aCCv}^{2} \operatorname{hv}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} - \frac{\sqrt{1 + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}}} - \operatorname{ArcSinh} \left[\frac{\pi \operatorname{Rinlv}}{\operatorname{hv}} \right] + \operatorname{ArcSinh} \left[\sqrt{\frac{2 \operatorname{Liv} \pi}{\operatorname{hv}} + \frac{\pi^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}} \right] - \frac{3 \sqrt{3} \operatorname{accv}^{2} \operatorname{hv}}{3 \sqrt{3} \operatorname{accv}^{2} \operatorname{hv}} + \frac{\pi^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}} - \operatorname{ArcSinh} \left[2 \pi + \frac{\pi \operatorname{Rinlv}}{\operatorname{hv}} \right] + \operatorname{ArcSinh} \left[\sqrt{\frac{2 \operatorname{Liv} \pi}{\operatorname{hv}} + \frac{\pi^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}} \right] - \frac{1}{3 \sqrt{3} \operatorname{accv}^{2} \pi} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}} - \operatorname{ArcSinh} \left[2 \pi + \frac{\pi \operatorname{Rinlv}}{\operatorname{hv}} \right] + \operatorname{ArcSinh} \left[\sqrt{\frac{2 \operatorname{Liv} \pi}{\operatorname{hv}} + \frac{\pi^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}}} \right] - \frac{1}{\operatorname{ArcSinh} \left[\frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} \right]} - \operatorname{ArcSinh} \left[\frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} \right] - \operatorname{ArcSinh} \left[\frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} \right] - \operatorname{ArcSinh} \left[\frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} \right] - \operatorname{ArcSinh} \left[\frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} \right] - \operatorname{ArcSinh} \left[\frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} \right] - \operatorname{ArcSinh} \left[\frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} \right] - \operatorname{ArcSinh} \left[\frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} + \frac{e^{2} \operatorname{Rinlv}^{2}}{\operatorname{hv}^{2}} \right] - \operatorname{ArcSinh} \left[\frac{e^{2$$

V. Export the data of the plots of the nanoscroll energy

as a function of the inner radius

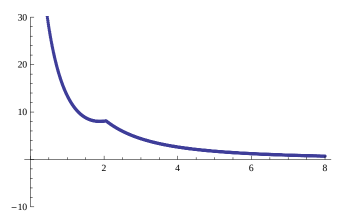
The parameters of the output file

The number of the output points = 1000

Export the plot data to the files:

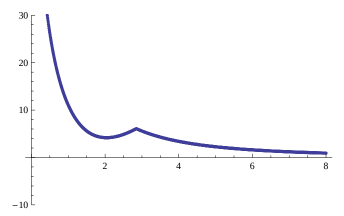
NanoscrollName=Nanoscroll2L15.nm

ScrollEnergyFileName=EvsRIn1Nanoscroll2L15.nm.dat



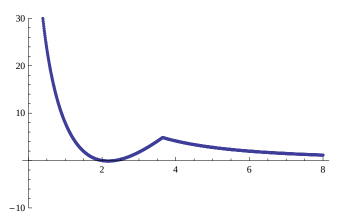
NanoscrollName=Nanoscroll2L20.nm

ScrollEnergyFileName=EvsRIn1Nanoscroll2L20.nm.dat



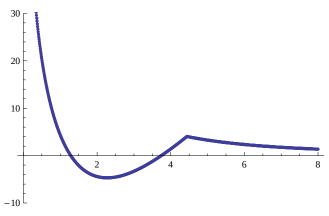
NanoscrollName=Nanoscroll2L25.nm

ScrollEnergyFileName=EvsRIn1Nanoscroll2L25.nm.dat



NanoscrollName=Nanoscroll2L30.nm

ScrollEnergyFileName=EvsRIn1Nanoscroll2L30.nm.dat



Plot ScrollEnergy[RIn1/nm]/(eV/atom) for L1= $\{15., 20., 25., 30.\}$ nm (NoL=2,Lw=1.nm)

