```
% 初始化工作区
clear
% 基础变量
Type = 'FCC';
element = 'Ni';
a0 = 3.524;
mass = 58.693:
% 定义晶胞原子
atom1 = [0.0 * a0, 0.0 * a0, 0.0 * a0];
atom2 = [0.5 * a0, 0.5 * a0, 0.0 * a0];
atom3 = [0.5 * a0, 0.0 * a0, 0.5 * a0];
atom4 = [0.0 * a0, 0.5 * a0, 0.5 * a0];
% 定义基矢量
ux = [1.0 * a0, 0.0, 0.0];
uy = [0.0, 1.0 * a0, 0.0];
uz = [0.0, 0.0, 1.0 * a0];
% 循环生成
id = 0:
for i = 0:10
   for j = 0:10
       for k = 0:10
          % 得到位移矢量
          vector = ux * i + uy * j + uz * k;
          % 由晶胞原子出发沿位移矢量添加原子
          id = id + 1;
          crystal(id, 1:3) = atom1 + vector;
          id = id + 1;
          crystal(id, 1:3) = atom2 + vector;
          id = id + 1;
          crystal(id, 1:3) = atom3 + vector;
          id = id + 1;
          crystal(id, 1:3) = atom4 + vector;
   end
end
% 裁切生成的晶格以符合仅10个胞
crystal = lattice_slicer(crystal, [0.0, 0.0, 0.0], [a0*10, a0*10, a0*10]);
% 更新晶格矩阵长度
id = length(crystal);
% 生成序号的随机全排列数列
rand_list = randperm(id);
% 按比例分配随机数号段
Ni_atoms_indices = rand_list(1:ceil(id / 2));
Al atoms indices = rand list((ceil(id / 2) + 1):id);
fprintf('Total\ amount\ of\ Ni\ atoms:\ %d\n',\ length(Ni\_atoms\_indices));
fprintf('Total\ amount\ of\ Al\ atoms:\ %d\n',\ length(Al\_atoms\_indices));
% 遍历查找并拾取对应原子
Ni_i = 1;
A1_id = 1;
for i = 1:id
   if find(Ni atoms indices == i)
       % 序号属于Ni号段
       Ni_crystal(Ni_id, 1:3) = crystal(i, 1:3);
       Ni_i = Ni_i + 1;
       % 序号属于A1号段
       Al crystal(Al id, 1:3) = crystal(i, 1:3);
      A1_id = A1_id + 1;
   end
end
% MATLAB作图
% 作图并保存句柄
Ni_plot = plot3(Ni_crystal(:, 1), Ni_crystal(:, 2), Ni_crystal(:, 3), 'o', 'MarkerFaceColor','g', 'MarkerSize', 10);
% 方正化坐标
axis square:
% 锁定作图
```

```
hold on:
Al plot = plot3(Al crystal(:, 1), Al crystal(:, 2), Al crystal(:, 3), 'o', 'MarkerFaceColor', 'b', 'MarkerSize', 10);
axis square:
% 取消锁定
hold off:
%添加图例,并定位在图右侧(East side)
legend([Ni_plot, Al_plot], 'Ni Atom', 'Al Atom', 'Location', 'eastoutside');
% 输出pdb和txt
% 打开文件并保存句柄
pdb_file_ID = fopen('FCC-NiAl lattice.pdb', 'w');
txt_file_ID = fopen('FCC-NiAl lattice.txt', 'w');
% 定义文件格式
rasmol_format_spec = '%-6s%-5d %-4s%-1s%-3s %-4d%-1s %-8.3f%-8.3f%-8.3f\n';
table_format_spec = '%-5s\t%-8.3f\t%-8.3f\t%-8.3f\n';
% 循环输出各原子坐标信息
num = 0:
for id = 1:length(Ni_crystal)
    num = num + 1:
    fprintf(pdb_file_ID, rasmol_format_spec, 'ATOM', num, element, '', '', 1, '', Ni_crystal(id, 1), Ni_crystal(id, 2), Ni_crystal(id, 3)); fprintf(txt_file_ID, table_format_spec, '1', Ni_crystal(id, 1), Ni_crystal(id, 2), Ni_crystal(id, 3));
num = 0:
for id = 1:length(Al_crystal)
    num = num + 1;
    fprintf(pdb_file_ID, rasmol_format_spec, 'ATOM', num, 'Al', ' ', ', 1, ' ', Al_crystal(id, 1), Al_crystal(id, 2), Al_crystal(id, 3));
    fprintf(txt\_file\_ID,\ table\_format\_spec,\ '2',\ Al\_crystal(id,\ 1),\ Al\_crystal(id,\ 2),\ Al\_crystal(id,\ 3));
% 关闭文件
fclose(pdb_file_ID);
fclose(txt_file_ID);
% 晶格裁切函数
function output_lattice = lattice_slicer(input_lattice, min_limit, max_limit)
    id = 1:
    for i = 1:length(input_lattice)
         if \ sum(input\_lattice(i, 1:3) >= min\_limit) == 3 \ \&\& \ sum(input\_lattice(i, 1:3) <= max\_limit) == 3 \\ \\
            output_lattice(id, 1:3) = input_lattice(i, 1:3);
            id = id + 1;
    end
end
```

Total amount of Ni atoms: 2316 Total amount of Al atoms: 2315

