MOTT TRANSITION IN STRONGLY CORRELATED MATERIALS: A REALISTIC MODELING USING LDA+DMFT

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ABSTRACT OF THE DISSERTATION

Mott transition in strongly correlated materials: a realistic modeling using LDA+DMFT

by Sahana Murthy

Dissertation Director: Professor Gabriel Kotliar

We study aspects of the Mott metal-insulator transition in simple models and in real materials. We first investigate the density-driven Mott transition in the degenerate Hubbard model within the framework of dynamical mean-field theory (DMFT). We demonstrate the divergence of compressibility near the finite temperature transition endpoint using quantum Monte Carlo simulations. We show that our results are relevant to the $\alpha$-$\gamma$ transition in Cerium.

In the latter part of the thesis, we use a combination of density functional methods with local density approximation (LDA) and many body techniques such as DMFT to realistically model two materials with strong correlations. We compute the band structure and spectra of YbRh$_2$Si$_2$ which has an antiferromagnetic ground state. YbRh$_2$Si$_2$ is known to have a strong anisotropy in its magnetic response with respect to its crystal structure. We determine magnetic anisotropy energy of YbRh$_2$Si$_2$ from its total energy. Using LDA+DMFT methods we calculate the spectra and equilibrium volume of Americium. We show that on applying pressure, a Mott metal-insulator transition takes place in Americium which is in accordance with experimental studies.
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# Table of Contents

Abstract ......................................................... ii  
Acknowledgements ........................................... iii  
List of Tables .................................................. viii  
List of Figures ................................................. ix  
1. Introduction .................................................. 1  
2. Finite temperature Mott transition in the degenerate Hubbard model 5  
   2.1. The Mott transition ...................................... 5  
   2.2. The Hubbard model ...................................... 9  
   2.3. Dynamical mean-field equations ......................... 9  
      2.3.1. Hirsch-Fye Quantum Monte Carlo algorithm .......... 11  
      2.3.2. Implementation .................................... 13  
   2.4. Results .................................................. 15  
      2.4.1. Effect of doping ................................... 15  
      2.4.2. Coexistence between metallic and insulating phases . 16  
      2.4.3. Convergence of solutions ............................ 21  
      2.4.4. Compressibility divergence ......................... 22  
   2.5. Phase diagram ......................................... 23  
   2.6. Discussion ............................................. 25  
   2.7. Summary ................................................. 30  
3. Density Functional Theory, LDA, LDA+U and LDA+DMFT ........ 31  
   3.1. Density functional theory ............................... 31
3.1.1. LDA .................................................. 31
3.1.2. LDA+U ........................................... 33
3.1.3. LDA+DMFT method ............................ 36
3.2. Structure of the LMTART program .............. 39
3.3. DMFT Loop ........................................... 42
  3.3.1. Impurity solver: Hubbard-I method ......... 46
3.4. Preparing control files for LMTART ............... 48
  3.4.1. *INI* file ..................................... 49
  3.4.2. *STR* file ..................................... 50
  3.4.3. Modifications for anti-ferromagnetic solution . 53
  3.4.4. *HUB* file ................................... 54
4. Electronic structure and properties of \( f \)-band materials: YbRh\(_2\)Si\(_2\) ........................................ 60
  4.1. Introduction ....................................... 60
  4.2. Properties of YbRh\(_2\)Si\(_2\) ....................... 61
  4.3. Band structure calculation of YbRh\(_2\)Si\(_2\) .......... 62
    4.3.1. LDA calculation ................................ 63
    4.3.2. LDA+U calculation ............................. 66
    4.3.3. Effect of the double counting term: .......... 70
    4.3.4. Comparison with other methods involving the double-counting term ......................... 71
    4.3.5. Effects of pressure .......................... 75
    4.3.6. Anti-ferromagnetic solution ................. 76
  4.4. Magnetic Anisotropy Energy ....................... 78
  4.5. Summary and Conclusions ......................... 82
5. Electronic structure and properties of \( f \)-band materials: Americium 85
  5.1. Introduction ....................................... 85
  5.2. LDA and LDA+U band structure calculation of Americium .................. 90
    5.2.1. Effect of the double counting term .......... 99
## List of Tables

<table>
<thead>
<tr>
<th>Table</th>
<th>Description</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.1</td>
<td>Input file nio.ini</td>
<td>51</td>
</tr>
<tr>
<td>3.2</td>
<td>Input file nio.str</td>
<td>52</td>
</tr>
<tr>
<td>3.3</td>
<td>Input file nio.ini for anti-ferromagnetic NiO</td>
<td>54</td>
</tr>
<tr>
<td>3.4</td>
<td>Input structure file nio.str for the anti-ferromagnetic case</td>
<td>55</td>
</tr>
<tr>
<td>3.5</td>
<td>Input hub file nio.hub</td>
<td>58</td>
</tr>
<tr>
<td>3.6</td>
<td>Partial occupancies for Ni in nio.hub</td>
<td>59</td>
</tr>
<tr>
<td>4.1</td>
<td>Input file ybrh2si2.str</td>
<td>63</td>
</tr>
<tr>
<td>4.2</td>
<td>Input file ybrh2si2.ini</td>
<td>64</td>
</tr>
<tr>
<td>4.3</td>
<td>Input file ybrh2si2.hub</td>
<td>65</td>
</tr>
<tr>
<td>4.4</td>
<td>Input file ybrh2si2.str</td>
<td>78</td>
</tr>
<tr>
<td>5.1</td>
<td>Input file am.str</td>
<td>90</td>
</tr>
<tr>
<td>5.2</td>
<td>Input file am.ini</td>
<td>91</td>
</tr>
<tr>
<td>5.3</td>
<td>Input file am.hub</td>
<td>92</td>
</tr>
<tr>
<td>5.4</td>
<td>Equilibrium volumes for Am in $\text{Å}^3$</td>
<td>100</td>
</tr>
</tbody>
</table>
List of Figures

2.1. Left: Temperature versus pressure phase diagram of $\kappa$-(BEDT – TTF)$_2$Cu[N(CN)$_2$Cl for different crystals. The grey circle near 200 kBar represents the critical point. Right: relative sound velocity (w.r.t. its value at 90K) as a function of temperature at various pressures. From D. Fournier et al. [28]. ......................................................... 6


2.3. Phase diagram of Hubbard model at the particle-hole symmetric point $\mu = U/2$. The solid lines denote $U_{c1}(T)$ (left) and $U_{c1}(T)$ (right). The dashed line denotes the first order transition line. The black circle at the top is the position of the second-order transition $(U_{MIT},T_{MIT})$. ....... 8

2.4. Hirsch-Fye Quantum Monte Carlo algorithm ................................................. 14

2.5. $n_f$ vs. $\mu$ curves in the half-filled case for a single band with $\beta = 1/T = 32, \Delta \tau = 0.5, D = 1$. The interaction has a value $U = 1.5$ (left) and $U = 3.2$ (right). ................................................................. 16

2.6. $n_f$ vs. $\mu$ curves at $\beta = 8$ ................................................................. 17

2.7. Coexisting solutions for $U=2.44, \beta = 64$ in the single band case. ...... 18

2.8. Imaginary part of Green’s functions for $U=2.44, \beta = 64$ ................. 18

2.9. Left: Occupation number for various values of $\beta$. Right: Occupation number for $\beta = 64$ metallic and insulating solution ...................... 19

2.10. Particle occupation $n$ function of $\mu$ for different temperatures in the 2-band model at $U=3.0$. The top panel shows curves with $T > T_{MIT}$ and in the bottom panel we have $T < T_{MIT}$. ................................. 20
2.11. Left: Metallic seed obtained from $U=2.8$. Right: Insulating seed obtained from $U=3.5$. 

2.12. Metallic and insulating Green’s function for $U = 3.0$, $\beta = 40$ (left) and $\beta = 64$ (right).

2.13. Convergence of metallic (left) and insulating (right) Green’s function.

2.14. 1-band model, $U=4.0$. Top: Compressibility as a function of chemical potential. Bottom: Inverse compressibility $\kappa^{-1} = (dn/d\mu)^{-1}$ at $U=4.0$ in the 1-band model as a function of $T$. The solid line is the function $\kappa^{-1} \sim (T - T_{C})^{b}$ with $b \sim 0.47$.

2.15. Left: Inverse compressibility $\kappa^{-1} = (dn/d\mu)^{-1}$ at $U=2.46$ in the 1-band model. The solid line is the function $\kappa^{-1} \sim (T - T_{C})^{b}$ with $b \sim 0.33$. Right: $\kappa^{-1}$ at a constant doping $n = 1.003$ as a function of $T$ in the 2-band model. The solid line is a fit to a power law with $\kappa \approx (T - T_{c})^{b}$ with the power $b \sim 1/3$. The intercept with the $T-$axis gives our estimate for $T_{c} = 0.0288$.

2.16. Schematic phase diagram for the 1-band Hubbard model. The cross sections are on the $T - \mu$ plane for different values of $U$ with $U$ increasing towards the top of the figure. The peaks are symmetric about $\mu = U/2$. $\mu_{c1}$ and $U_{c1}$ are the chemical potential and interaction respectively below which the insulating solution does not survive. $\mu_{c2}$ and $U_{c2}$ are those above which the metallic solution does not exist. The black circle denotes the second order transition point $(U_{MIT}, T_{MIT})$ at $n = 1$. The shaded regions denote the coexistence region between metallic and insulating phases.

2.17. Phase diagram for $U=4.0$. The black circles denote are obtained from QMC simulations. The dashed and the dotted lines denote $\mu_{c1}$ and $\mu_{c2}$ lines respectively.
2.18. Critical slowing down observed in the vicinity of $\mu_{c1}$ and $\mu_{c2}$ for the 1-band model at $U = 4.0, \beta = 64$. The inverse number of iterations required for the metallic solution to converge goes to zero as we near $\mu_{c2}$ and those of the insulating solution to converge goes to zero as we approach $\mu_{c1}$. The arrows indicate the directions in which the boundaries were approached.

2.19. Schematic phase diagram for the 2-band case. There is an asymmetry in the triangular peaks as compared to the 1-band case. The cross sections are on the $T - \mu$ plane for different values of $U$ as before. $\mu_{c1}$ and $U_{c1}$ are the chemical potential and interaction respectively below which the insulating solution does not survive. $\mu_{c2}$ and $U_{c2}$ are the chemical potential and interaction above which the metallic solution does not exist. $U_{MIT}$ is the value of the interaction at which the metal-insulator transition takes place. The shaded portions are the regions of coexistence between the metallic and the insulating solutions.

3.1. Flow diagram for LDA+DMFT scheme
3.2. Unit cell for NiO and translation vectors
3.3. Unit cell for anti-ferromagnetic case
4.1. Body centered tetragonal crystal structure for YbRh$_2$Si$_2$
4.2. Energy bands of YbRh$_2$Si$_2$ – Full Potential(left) and Atomic Sphere Approximation(right). The colored regions in are the fat bands for Yb:4f.
4.3. Energy bands of YbRh$_2$Si$_2$. The colored regions in are the fat bands for Yb:5d (left) and Rh(4d) and Si:3p(right).
4.4. Energy bands of Yb:4f $z(x^2 - y^2)$
4.5. Energy bands of Yb:LDA+U, $U=2,4,6,8$eV
4.6. Position of center of 4f-bands of Yb as a function of the interaction $U$
4.7. Position of center of 4f-band as a function of the double-counting term $\bar{U}$ for $U=4$eV (top) and $U=6$eV (bottom)
4.8. Energy bands(left) and DOS(right) of Yb:4f, LDA+U, $U=4$eV.
4.9. Energy bands(left) and DOS(right) of Yb:4f, LDA+U, U=6eV.  
4.10. Location of the 4f-bands in various schemes. The original double-counting where $U = \bar{U}$ and the one used by Mazin et.al [54] as described above do not predict a magnetic ground state. The empirical method described in section 4.3.3 does predict the correct ground state.  
4.11. Dependence of magnetic moment on relative volume  
4.12. Application of pressure on the magnetic solution  
4.14. $\chi$ vs T (from Trovarelli et.al. [59])  
4.15. from Trovarelli et.al. [58]  
4.16. Magnetization as a function of external magnetic field (along z-axis)  
5.1. Volumes of actinide elements  
5.2. Left: Photoemission spectra of Am by Naegele [60]. Right: Relative volume of americium as a function of pressure obtained by Lindbaum et. al. [62]  
5.3. Four phases of Americium (Lindbaum et. al [62])  
5.4. Energy bands of Am in LDA+U, no spin-orbit coupling. The colored regions are the fat bands of Am:5f  
5.5. Density of states of Am in LDA+U, no spin-orbit coupling. The black curve is the total DOS and the red curve is the partial 5f-DOS  
5.6. Energy bands of Am in LDA+U, with spin-orbit coupling. The colored regions are the fat bands of Am:5f. Note the initial separation (for $U = 0$) between the filled and empty states, which is due to spin-orbit coupling  
5.7. Density of states in LDA+U, no spin-orbit coupling. Note the initial separation (for $U = 0$) between the filled and empty states, which is due to spin-orbit coupling. The black curve is the total DOS and the red curve is the partial 5f-DOS  

xii
5.8. Position of center of the Am $5f$-band as a function of the interaction strength .................................................. 97
5.9. Gap between spin-up and spin-dn electrons as a function of the interaction ......................................................... 98
5.10. Occupancy of Am $5f$-band as a function of the interaction strength $U$ ............................................................. 98
5.11. Effect of $U_{dc}$: Position of the center of the $f$-band as a function of the $U_{dc}$, with $U$ kept at 4eV ................................................. 99
5.12. Total energy (-61011 Ry) as a function of relative volume $V/V_0$ for various values of interaction $U$ (in eV) ........................................... 101
5.13. Input file $lmt.am$ for treating $f$-electrons as core .......................................................... 102
5.14. Energy bands of Am treating $5f$ electrons as core .......................................................... 103
5.15. DOS of Am treating $5f$ electrons as core .......................................................... 103
5.16. Comparison of DOS : LDA and Tight binding methods .......................... 107
5.17. Comparison of DOS : LDA and Tight binding methods .......................... 108
5.18. Comparison of DOS : LDA and Tight binding methods .......................... 109
5.19. Comparison of DOS : LDA and Tight binding methods .......................... 110
5.20. Impurity level (w.r.t $E_F$) as a function of relative volume .......................... 111
5.21. Tight binding hoppings as a function of relative volume .......................... 111
5.22. Am: Density of States as a function of $U$ for the tight-binding model .......................... 112
5.23. Energy vs. relative volume ................................................. 113
5.24. Pressure vs. relative volume ................................................. 114
5.25. Am DOS in LDA ................................................. 115
5.26. Am DOS in LDA ................................................. 116
5.27. Am DOS in LDA ................................................. 117
5.28. Am DOS in LDA+DMFT: $U = U_{DC} = 4.5eV$ ................................................. 119
5.29. Am DOS in LDA+DMFT: $U = U_{DC} = 4.5eV$ ................................................. 120
5.30. Am DOS in LDA+DMFT: $U = U_{DC} = 4.5eV$ ................................................. 121
5.31. Am DOS in LDA+DMFT: $U = 4.5eV, U_{DC} = 4.1eV$ ................................................. 122
5.32. Am DOS in LDA+DMFT: $U = 4.5eV, U_{DC} = 4.1eV$ ................................................. 123
5.33. Am DOS in LDA+DMFT: $U = 4.5eV, U_{DC} = 4.1eV$ ................................................. 124
5.34. Density of states at Fermi level in LDA and LDA+DMFT . . . . . . . . 125
5.35. Ratio of 5f-DOS over total DOS . . . . . . . . . . . . . . . . . . . . . 126
5.36. Position of center of 5f-band . . . . . . . . . . . . . . . . . . . . . . 127
Chapter 1

Introduction

The understanding of the structure and properties of materials, the relationship between their phases and the mechanism that gives rise to transitions between the phases have been the focus of a number of theoretical and computational investigations in solid state physics. There have been two main approaches that have contributed to progress in determining the structure and properties of materials: studies of model Hamiltonians and first-principles or ab-initio calculations. We can learn aspects of universal behavior of systems by studying simple models. For example, critical exponents of measurable quantities such as heat capacity and susceptibility derived from the Ising model are applicable to a wide variety of systems. On the other hand, non-universal features require detailed modeling that involve particulars such as atomic charge and crystal structure. This thesis deals with the realistic modeling of strongly correlated materials using tools from both the approaches.

Model Hamiltonians have proved to be very useful in understanding materials with strong correlation effects. These are materials in which there is a competition between the kinetic energy and the interaction energy of electrons. Strong correlation effects arise due to partially filled $d$ and $f$ shells in the atoms constituting the materials. Strongly correlated materials exhibit interesting physical properties such as high temperature superconductivity, anomalous behavior in resistivity, specific heat and transport properties. Examples of strong correlation effects are seen in high temperature superconductors such as copper oxides, Uranium and Cerium based heavy fermion systems, colossal magnetoresistance materials such as manganites and carbon based compounds such as bucky balls and carbon nanotubes.

The Mott-Hubbard metal-insulator transition [1] is another important effect induced
by strong correlations. It has been observed in $\text{V}_2\text{O}_3$ [2], pyrites such as $\text{Ni(Se, S)}_2$ [3], and layered organic conductors [4]. It is a difficult problem to treat theoretically since the electron kinetic energy and interaction energy are of the same order. Since there is no natural small parameter, standard perturbative techniques fail. The dynamical mean-field theory (DMFT) [5] is a framework in which both localized bands in the insulating phase and quasi-particle bands in the metallic phase are treated on an equal footing. It is exact in the limit of infinite dimensions. In particular, DMFT studies of the Hubbard model have been successful in understanding some universal aspects of the physics of the Mott transition such as high temperature behavior near Mott endpoint [], phase diagram at integer occupation [] and transfer of spectral weight near the transition [6, 7]. However, the main drawback of studying models is that physical quantities such as interaction strength and doping need to be treated as parameters.

A different approach to studying electronic structure of solids is based on first-principles (or ab-initio) techniques. There are no empirically adjustable parameters here. The input to these calculations consists only of atomic charges and crystal structures, hence a detailed modeling of the solid is possible. The most widely used tool in ab-initio calculations is Density Functional Theory [8,9], especially in the Local Density Approximation (LDA) or Generalized Gradient Approximation (GGA) [10]. Density functional methods have been tremendously successful in treating weakly correlated materials such as simple metals and semi-conductors. They provide a simple conceptual picture of the ground state and correctly predict excitation spectra and transport properties in such materials. However, these methods often fail to capture the correct physics in strongly correlated materials.

In recent years, a combination of Local Density Approximation and dynamical mean-field theory (LDA+DMFT) has been successfully used to study strongly correlated materials [11, 12]. In this approach, the light $s, p$ (sometimes $d$) electrons which have extended wave functions are treated by LDA. The localized $d$ (or $f$) electrons are treated in the DMFT framework. This method can be used to calculate excitation spectra and total energy as well as correlation functions. Theoretically, the LDA+DMFT method can be constructed from an effective action point of view [13] where the free energy is
a functional of the local Green’s function and the density. Other extensions to LDA have also been tried: LDA+U [14–16], which has been shown to be a static limit of LDA+DMFT [17] and LDA+GW [18,19] which is a perturbative calculation starting from LDA spectra.

LDA+DMFT has proved to be a powerful tool for the realistic modeling of strongly correlated electron systems. There have been various successes: Phase diagram of plutonium [20], α-γ transition in the 4f electron system Cerium [21, 22], optical and transport properties of $\text{La}_{1-x}\text{Sr}_x\text{TiO}_3$ [23,24], Mott Hubbard metal-insulator transition in $\text{V}_2\text{O}_3$ [25] to name a few. This thesis is an attempt to understand the structure and properties of two strongly correlated materials using the above methods, with a focus on the metal-insulator Mott transition.

This thesis is organized as follows:

In chapter 2 we begin with the physical ideas behind the Mott transition. We introduce the the Hubbard model. We go through the mechanism behind dynamical mean-field theory and its mapping to the impurity model. We then describe one of various ways to solve for the properties of the impurity model. This is the quantum Monte Carlo technique, which is exact, but computationally difficult for accessing low temperatures. Studying the behavior of the system as a function of various parameters, we look into the details of the finite temperature doping driven Mott transition. We concentrate on the behavior of the charge compressibility near the transition.

Chapter 3 serves two purposes. In the first part, we briefly review density functional theory from an effective action point of view. We discuss existing density functional methods that go beyond the standard LDA technique, and can be used when strong correlations are present. These are the LDA+U and LDA+DMFT methods. We outline the numerical implementation of another impurity solver for the DMFT loop, the so-called Hubbard-1 approximation, which we will employ in a later calculation. The latter part of 3 is devoted to the details of our computation scheme. We describe the various input files that are required in the computer program.

We then turn to applications of the above methods to real materials in the lanthanide and actinide series. In chapter 4 we study the ground state and magnetic
properties of a heavy fermion compound YbRh$_2$Si$_2$. We also propose a procedure in the LDA+U implementation, that improves results for the ground state properties. The theoretical tools we discussed, the physical concepts behind the Mott transition and the insights we gained into the details of the band structure calculation all come together in chapter 5. We use the LDA+DMFT technique, with the Hubbard-1 impurity solver, to investigate the structure, equilibrium volume, and details of the Mott transition in elemental Americium.

We conclude in chapter 6 with a summary of the thesis, and directions for future work.
Chapter 2

Finite temperature Mott transition in the degenerate Hubbard model

The Mott transition is a metal-insulator transition that is driven by electron-electron interactions [1]. It occurs when the ratio of the electron interaction strength to the bandwidth is increased. Experimentally, this phenomenon is realized in many compounds such as $V_2O_3$ [2], $Ni(Se,S)_2$ [3, 26] and the family of organic conductors $\kappa$-BEDT [4, 27]. On the theoretical side, the Hubbard model is among the first, and perhaps the simplest model that can capture the essential physics of the transition. In recent years, tremendous theoretical progress has been made in understanding the Mott transition by applying the dynamical mean-field theory (DMFT) [5] to the Hubbard model. In this chapter, we study finite temperature aspects of the doping driven Mott transition in the Hubbard model, and the behavior of the compressibility near the Mott endpoint within the DMFT approach.

2.1 The Mott transition

Experimental evidence

The classical system in which a Mott transition is observed is $V_2O_3$ doped with Cr or Ti [2]. The transition can be driven by varying pressure, chemical potential (or doping) and temperature. Recently, a first order metal-insulator transition was observed in the organic superconductor $\kappa$-(BEDT $\sim$ TTF)$_2$Cu[N(CN)$_2$Cl, as a function of hydrostatic pressure [28]. The study shows a large softening of sound velocity of the material near the critical point. The sound velocity is proportional to the square root of compressibility, hence a sharp decrease in the velocity points to a divergence in the compressibility.
Fig. 2.1 shows the phase diagram and the dip in sound velocity in this system.

It has been suggested by B. Johansson [29] that the Mott transition concept is also relevant to lanthanide and actinide materials. The $\alpha\rightarrow\gamma$ transition in Cerium shows certain universal features that are common to other systems in which a metal-insulator transition is observed. For example, Beecroft and Swenson [30] measured a large decrease in the compressibility in the transition from the insulating low-pressure phase ($\gamma$-Ce) to the metallic high-pressure phase ($\alpha$-Ce). Fig. 2.2 shows the pressure-volume isotherms for the $\alpha\rightarrow\gamma$ transition in Ce.

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Figure 2.1: Left: Temperature versus pressure phase diagram of $\kappa$-(BEDT-TTF)$_2$Cu[N(CN)$_2$]Cl for different crystals. The grey circle near 200 kBar represents the critical point. Right: relative sound velocity (w.r.t. its value at 90K) as a function of temperature at various pressures. From D. Fournier et al. [28].

Figure 2.2: Relative volume as a function of pressure in cerium. From Beecroft and Swenson [30]
Theoretical understanding

From a theoretical point of view, the electronic structure of solids has been well understood in two limits. When the overlap between electron orbitals is large, a wave-like description is relevant, and band theory accounts for most of the properties of the solid. At the other end, when the electron orbitals are far apart, a real-space description in terms of localized atomic wave-functions is valid.

Strongly correlated materials fall away from either of these limits. Strong correlation effects arise due to the dual character of the electron: wave-like and particle-like. In other words, there is a competition between the kinetic energy and the electron-electron interaction energy. One of the earliest models that was proposed to capture the effects of strong correlations in $d$ and $f$-band materials was the Hubbard model [31]. However, it has been solved exactly only in one dimension, where there cannot be a finite temperature phase transition.

Standard methods such as perturbation theory fail in the strong correlation problem since a method that can simultaneously treat itinerant and localized nature of electrons is required. One such framework that can describe both metallic (itinerant) and insulating (localized) phases is dynamical mean-field theory [5]. DMFT can be regarded as the quantum analog of the classical mean-field theory for magnetism. The essence of the method is to replace the interactions of an electron at a lattice site with its neighbors by an effective interaction with an average field. This effective field represents the dynamic state of the neighbors.

In the DMFT framework, the metal-insulator transition (MIT) can be interpreted as arising from bifurcation points of a functional of the local Green’s function. The case of the correlation driven MIT at half-filling, is now well understood. At temperature $T = 0$ there are two bifurcation points, one denoted by $U_{c1}(T = 0)$ where the insulating solution disappears, and the other denoted by $U_{c2}(T = 0)$ where the metallic solution disappears [32]. It was found that in the $U$-$T$ phase diagram of the frustrated Hubbard model, there is a region where two mean-field solutions, one metallic-like and one insulating-like can be obtained. This region is delimited by the $U_{c1}(T)$ and $U_{c2}(T)$ lines.
as shown in Fig. 2.3. Within the region there is a first-order MIT line \([33,34]\) that ends in a finite temperature second-order critical point \((U_{MIT}, T_{MIT})\) with a rapid variation of the susceptibility connected to the double occupancy \([6,35]\). At higher temperatures the \(U_{c2}(T)\) and \(U_{c1}(T)\) lines become crossover lines.

![Figure 2.3: Phase diagram of Hubbard model at the particle-hole symmetric point \(\mu = U/2\). The solid lines denote \(U_{c1}(T)\) (left) and \(U_{c2}(T)\) (right). The dashed line denotes the first order transition line. The black circle at the top is the position of the second-order transition \((U_{MIT}, T_{MIT})\).](image)

The doping driven MIT at zero temperature was studied in \([36]\). It was shown that there are two solutions in an area bound by the curves \(\mu_{c1}(U)\), where the insulating solution disappears, and \(\mu_{c2}(U)\), where the metallic state disappears. The finite temperature aspects of the doping driven Mott transition will be the subject of this chapter. We will concentrate on the paramagnetic phase, and will not consider effects of long range order.

We will also discuss the behavior of the charge compressibility near the Mott endpoint at finite temperatures. The compressibility is known to diverge at the density driven MIT in two dimensions at \(T = 0\) \([37]\). This behavior has also been observed on other models of correlated electron systems such as the t-J model indicating that this phenomenon is quite general \([38]\). Simple models of the Mott transition, such as the Gutzwiller approximation or the slave-boson approach predict a finite compressibility \([39]\). It is important to understand the physical origin of this result, and to see if it
is realized in the DMFT solution of the Hubbard model. The previously investigated bifurcation points within the DMFT, have either a finite charge compressibility, such as in the $T = 0$ density driven Mott transition, or a vanishing charge compressibility, as in the $T = 0$ correlation driven transition.

### 2.2 The Hubbard model

We begin with a general Hubbard Hamiltonian that has $m$-band degeneracy. Since we will be working in the grand canonical ensemble, we introduce a chemical potential in the Hamiltonian, given by:

$$H = - \sum_{\{i,m,\sigma\}} t_{ij}^m c_{i,m,\sigma}^{\dagger} c_{i,m,\sigma} + \frac{1}{2} \sum_{i,m,m',\sigma} U n_{i,m,\sigma} n_{i,m',-\sigma} + \frac{1}{2} \sum_{i,m \neq m',\sigma} U n_{i,m,\sigma} n_{i,m',\sigma} - \mu \sum_{i,m,\sigma} n_{i,m,\sigma}$$

(2.1)

Here $c_{i,m,\sigma}^{\dagger}$ creates an electron in a state localized at site $i$ with orbital index $m$ and spin $\sigma = \uparrow, \downarrow$. The first term describes the hopping between nearest neighbor sites $\langle ij \rangle$ where the hopping integral $t$ has been assumed to be independent of the site indices. The parameter $U$ is the on-site Coulomb repulsion, namely, the energy cost associated with having a double occupancy on each site. $n_{i,m,\sigma} = c_{i,m,\sigma}^{\dagger} c_{i,m,\sigma}$ is the particle occupation number. $t$ and $U$ have been assumed to be independent of the band indices. This model is particle-hole symmetric when $\mu = \frac{(2m-1)U}{2}$.

### 2.3 Dynamical mean-field equations

The central idea of this method is to map a lattice model with many degrees of freedom, such as the Hubbard model in Eqn. (2.1), onto a one-site impurity model with fewer degrees of freedom [5]. This is supplemented by a self-consistency condition that determines the effective bath (effect of the other degrees of freedom) with which the impurity interacts. This approach can be shown to be exact in the limit of infinite dimensions $d \rightarrow \infty$, or large lattice coordination number $z$, with an appropriate scaling of parameters [40].

We consider the Hubbard model in (2.1) on a Bethe lattice with large connectivity $z$. A semi-circular density of states $\rho_\omega(\epsilon) = \frac{1}{2\pi\epsilon^2} \sqrt{4t^2 - \epsilon^2}$ is obtained where $D=2t$ is
the half-bandwidth. In this case, we can write the effective action for this model as:

$$S_{\text{eff}} = \int_0^\beta d\tau \int_0^\beta d\tau' \sum_{\langle ij \rangle, m, \sigma} c_{im\sigma}^\dagger(\tau) G_{0m\sigma}^{-1}(\tau - \tau') c_{jm\sigma}(\tau')$$

$$+ \frac{U}{2} \int_0^\beta d\tau \left( \sum_{i,m,m',\sigma} n_{im\sigma} n_{im'\sigma} - \sum_{i,m \neq m',\sigma} n_{im\sigma} n_{im'\sigma} \right)$$ \hspace{1cm} (2.2)

In the $d \to \infty$ limit the above lattice model is mapped on to the degenerate Anderson impurity model.

$$H_{AM} = H_0 + H_I$$ \hspace{1cm} (2.3)

$$H_0 = \sum_{k \geq 2, m, \sigma} \epsilon_{km} c_{kma}^\dagger c_{kma} + \sum_{k \geq 2, m, \sigma} (V_{km} c_{kma}^\dagger f_m + h.c) + \sum_m (E_f + (2m - 1)/2U) n_{m\sigma}$$ \hspace{1cm} (2.4)

$$H_I = U \sum_{m < m', \sigma} \left\{ n_{m\sigma} n_{m'\sigma} - \frac{1}{2} (n_{m\sigma} + n_{m'\sigma}) \right\}$$ \hspace{1cm} (2.5)

The Weiss function $G_{0m}$ in Eqn. (2.2) is given by a self-consistency condition

$$G_{0m}^{-1}(i\omega_n) = i\omega_n + \mu - t^2 G_m(i\omega_n)$$ \hspace{1cm} (2.6)

Here $G(i\omega_n)$ is the local Green’s function for the impurity. The conduction orbitals have index $k = 2, N$ and the impurity $f$–orbital corresponds to $k = 1$. $c_{km}$ and $c_{km}^\dagger$ are the band electron annihilation and creation operators, $V_{km}$ is the hybridization between the band and the impurity electrons. $n_m = f_{m}^\dagger f_m$ is the occupation number for the impurity electrons and $E_f$ is the f-level energy.

In terms of the hybridization function $\Delta_m(i\omega)$

$$\Delta_m(i\omega) = \sum_k \frac{V_{km}^2}{i\omega - \epsilon_{km}}$$

the DMFT equation for the Bethe lattice can be written as

$$t^2 G_m(i\omega)[\Delta] = \Delta_m(i\omega)$$ \hspace{1cm} (2.7)

The coupled problem of finding the impurity Green’s function $G$ and the Weiss field $G_0$ is solved iteratively until required convergence is reached. Impurity models have been intensely studied for the past few decades, and a number of analytical and numerical tools exist to tackle them. In the next section, we focus on one numerical method: the Quantum Monte Carlo (QMC) algorithm by Hirsch and Fye [41], to obtain the Green’s function $G$. 
2.3.1 Hirsch-Fye Quantum Monte Carlo algorithm

We discretize the time interval $[0, \beta]$ into $L$ slices of length $\Delta \tau$ such that
$\Delta \tau_l = l \Delta \tau, \quad l = 1, 2, \ldots, L$ and $\beta = L \Delta \tau$.

The partition function is

$$Z = \text{Tr} \exp\{-\beta H\} = \text{Tr} \Pi \exp\{-\Delta \tau (H_0 + H_I)\}$$  \hspace{1cm} (2.8)

Using the Trotter formula, the discretized partition function is

$$Z \simeq Z^{\Delta \tau} = \text{Tr} \Pi \exp\{-\Delta \tau H_0\} \exp\{-\Delta \tau H_I\}$$  \hspace{1cm} (2.9)

The discretized Green’s function is

$$G_{m,k_1,k_2}^{\Delta \tau}(\tau_1, \tau_2) = \left< T(c_{mk_1}^\dagger(\tau_1)c_{mk_2}(\tau_2)) \right>$$

$$\frac{1}{Z^{\Delta \tau}} \text{Tr} e^{-\beta H} c_{mk_1}(\tau_1)c_{mk_2}^\dagger(\tau_2)$$  \hspace{1cm} (2.10)

for $l_1 > l_2$. We then use a discrete Hubbard-Stratonovich transformation to decouple the quartic interaction in $H_I$ [42].

$$\exp \left[ -\Delta \tau U \left( n_m n_m' - \frac{1}{2} (n_m + n_m') \right) \right] = \frac{1}{2} \sum_{s_{mm'}^{\pm \pm}} \exp \left[ \lambda S_{mm'}(n_m - n_m') \right]$$  \hspace{1cm} (2.11)

where $\cosh \lambda = \exp(\Delta \tau U/2)$ and $S_{mm'}(\tau_l)$ are auxiliary Ising-like fields at each time slice. This transformation renders $H_{AM}$ quadratic in $S_{mm'}(\tau_l)$.

Using (2.11) we write the partition function as

$$Z^{\Delta \tau}[\{S_{mm'}\}] = \text{Tr}[S_{mm'}] \prod_m \det O_m[\{S_{mm'}\}]$$  \hspace{1cm} (2.12)

Here $O[\{S_{mm'}\}]$ is a $NL \times NL$ ($N$ being the number of conduction orbitals, and $L$ being the number of time slices) matrix defined as

$$(O_m)_{l,l} = I$$

$$(O_m)_{l,l-1} = -e^{(-\Delta \tau H_{0m})} e^{(V_m^l)(1-2\delta_{l,1})}$$  \hspace{1cm} (2.13)

$$(O_m)_{l,l'} = 0 \quad \text{otherwise}$$

with $V_m^l = \lambda \sum_{m \neq m'} S_{mm'}(\tau_l) |m\rangle \langle m| \text{ for } m < m'$. 
For the auxiliary spin dependent Green’s function, we have

\[ G^\Delta_r[{S_{mm'}}] = O_m^{-1}[{S_{mm'}}] \equiv G_m \tag{2.14} \]

We then get a relation between Green’s functions for two different configurations.

\[ G_m^{-1} = A_m G_m \tag{2.15} \]

where \( A_m = 1 + (1 - G_m)(e^{V'_m - V_m} - 1). \tag{2.16} \)

This is a \( L \times L \) matrix equation.

If we now make only one local change in the field at one time slice such that two configurations differ by a single spin

\[ S_{mm'} \rightarrow S'_{mm'} \text{ for } m > m' \]
\[ \rightarrow -S'_{mm'} \text{ for } m < m' \]

the above equations are simplified. \( \exp\{V'_m - V_m\} \) has only one non-zero diagonal element at the \( l \)-th part of the matrix \( O \). \( A^{-1}_{lk} = 0 \) for \( l \neq k \) and \( \det A_m = A_m^{ll} = 1 + (1 - G_m^{ll})(\exp\{V'^{ll}_m - V_m^{ll}\} - 1) \).

We would also need to calculate the Boltzmann ratio of two different configurations which for a single spin flip is given by:

\[ R = \prod_m R_m \]

\[ R_m = \frac{\det(O'_m)}{\det(O_m)} = \det \left[ I - (G_m^{ll} - I) \left( \exp\{V'_m - V_m\} - I \right) \right] \tag{2.17} \]

Again, for a single spin flip, this simplifies to

\[ R_m = R_m R_{m'} \]
\[ R_m = 1 - [G^{ll}_m(l,l) - 1][\exp(-2\lambda S_{mm'} - 1)] \]
\[ R_{m'} = 1 - [G^{ll}_m(l,l) - 1][\exp(2\lambda S_{mm'} - 1)] \]

The \( f \)-Green’s function simplifies to:

\[ G'^{ll}_{m'}(l_1,l_2) = G^{ll}_{m'}(l_1,l_2) + [G^{ll}_{m'}(l_1,l) - \delta_{11}][\exp(-2\lambda S_{mm'} - 1)] \frac{1}{R_{m'}} G^{ll}_{m'}(l_1,l_2) \tag{2.18} \]
\[ G'^{ll'}_{m'}(l_1,l_2) = G^{ll'}_{m'}(l_1,l_2) + [G^{ll'}_{m'}(l_1,l) - \delta_{11}][\exp(2\lambda S_{mm'} - 1)] \frac{1}{R_{m'}} G^{ll'}_{m'}(l_1,l_2) \tag{2.19} \]
The physical Green’s function is then calculated as an average of the spin-dependent Green’s functions.

\[ G_{m}^{\text{physical}} = \frac{1}{Z} \sum_{S_{mm'}} G_m[S_{mm'}](\tau_l) \det O_m[S_{mm'}] \]  

(2.20)

2.3.2 Implementation

The practical implementation of the above procedure involves the following steps:

- We start with a guess for \( G_{0m} \) and an arbitrary configuration of spins.
- In the first iteration we invert the matrix \( O_m \) to obtain \( G_m[S_{mm'}(\tau_l)] \).
- For the subsequent iterations we change the configuration using a single spin-flip. We calculate the ratio \( R \). If \( R \) is greater than a random number (between 0 and 1) chosen we accept the flip and calculate \( G_m[S_{mm'}(\tau_l)] \). Otherwise we generate the next configuration.
- Every 100 iterations we start the spin configuration chain again by equating all auxiliary spins to zero and calculating the inverse of \( O_m \). This is done to check that the precision has not degraded.
- We then use the self-consistency condition

\[ G_{0m}^{-1}(i\omega_n) = i\omega_n + \mu - t^2 G_m(i\omega_n) \]


to calculate the new Weiss field \( G_{0m}(\text{new}) \).

- The \( G_{0m}(\text{new}) \) thus calculated is fed back into the QMC algorithm in the next step of the iteration.
- This process is iterated till \( G_m \) and \( G_{0m} \) converge.
**HIRSCH-FYE QMC ALGORITHM**

\[ L = \beta \Delta t \]  
_HUBBARD-STRATANOVICH TRANSFORMATION_

\[ \text{FULL UPDATE OF } G(S_{nm'}) = O^{-1} \]

\[ \text{PICK N_RAN, FLIP A SPIN} \]
\[ \text{NEW CONFIGURATION} \]

\[ \text{BOLTZMANN RATIO } R \]

\[ R > N_{\text{RAN}}? \]

\[ \text{YES} \]
\[ \text{NO} \]

\[ \text{YES} \]
\[ \text{NO} \]

\[ \text{ITERATIONS } = \text{(MOD)100} ? \]

\[ \text{YES} \]
\[ \text{NO} \]

\[ \text{SWEEPS } > 100000? \]

\[ \text{YES} \]
\[ \text{NO} \]

\[ \text{PHYSICAL } G \text{ AS AN AVERAGE OF } G(S_{mm'}) \]

\[ G' = G + (G-I) \left[ \exp(-2 \lambda S_{mm'} - 1) \right] (1/R)G \]

Figure 2.4: Hirsch-Fye Quantum Monte Carlo algorithm
2.4 Results

We use the Hirsch-Fye QMC algorithm to obtain the impurity Green’s function and iteratively solve the DMFT equations (2.3)-(2.6). Different phases can be obtained depending on the values of the interaction parameters in the Hamiltonian (Eqn. 2.1). In this section, we investigate the finite temperature aspects of the doping driven metal-insulator transition.

2.4.1 Effect of doping

The total occupation number for the impurity electrons is computed from the Green’s function. After the QMC runs stabilize, the full Green’s function $G(i\omega)$ is checked to see whether we have metallic or insulating behavior. Curves of the impurity occupation number $n_f$ are plotted against the chemical potential.

\begin{align}
G_m(\tau_i, 0) &= + \left\langle T \left( c_m(\tau_i)c_m^\dagger(0) \right) \right\rangle \\
G_m(0^+) &\rightarrow c_m(0)c_m^\dagger(0) \\
&= 1 - c_m^\dagger(0)c_m(0) \\
&= 1 - n_m \tag{2.22}
\end{align}

\begin{align}
n_f^m &= 1 - G_m(0^+) \tag{2.23}
\end{align}

We concentrate on the paramagnetic case where $n_\uparrow = n_\downarrow$ for a given band index $m$. We plot the average occupation number $n_f$ as a function of the chemical potential $\mu$ for various values of the interaction $U$. For the 1-band model, $\Delta\tau = 0.5$ for the 2-band model, $\Delta\tau = 0.25$. all energies are in units of half-bandwidth $D = 1$.

The graph of occupation number $n_f$ versus the chemical potential $\mu$ at $T = 1/32$ and $U = 1.5$ for the 1-band model is shown in Fig. 2.5 (left) and that for $U = 3.2$ is shown in Fig. 2.5 (right). For smaller values of the interaction $U$ the occupation number is a smooth function of $\mu$, that is, the system has metallic-like behavior. As $U$ is increased, a gap opens up in the $n_f-\mu$ curve near half-filling, indicating the insulating-like behavior of the system. Due to particle-hole symmetry in the 1-band model, we use a shifted chemical potential $\tilde{\mu} = \mu - \frac{U}{2}$. 
The 2-band results are shown in Fig. 2.6. We concentrate on the region near $n_f = 1$ at $T = 1/8$. At this high temperature only one solution exists. However, as $T$ is decreased, we see a flat portion in the $n_f$-$\mu$ curve at which the occupation number remains close to 1 over a range of $\mu$.

### 2.4.2 Coexistence between metallic and insulating phases

**Coexistence in the 1-band model**

In the region of intermediate $U$, at sufficiently low temperatures, we expect the metallic and insulating solutions to exist simultaneously. As we start doping the system a little, the two phases should still coexist. However at high doping we expect the insulating phase to become unstable and disappear. The existence of two phases should result in two simultaneous values for $n_f$ at the same chemical potential. The $n_f$ versus $\mu$ curve in Fig. 2.7 shows this coexistence at $U = 2.44$ and $\beta = 1/T = 64$ in the 1-band model. At $\tilde{\mu} = 0$, both the metallic and the insulating solutions are seen. Away from half-filling, the two solutions give different values of $n_f$. When the doping becomes large, only the metallic solution survives.

In Fig. 2.7, to obtain the two different values of the occupation number, we start...
from two different initial guesses or seeds in the QMC algorithm. The occupation number represented by circles (metallic behavior) are calculated by using a seed from low $U$. We first start at high doping in a metallic state and continuously evolve towards integer filling. The solution always remains metallic. To obtain the other solution, represented by triangles, we begin with the insulating solution at $\tilde{\mu} = 0, n = 1$ using a seed from high $U$ and gradually increase the doping. This state is essentially incompressible as $n$ almost remains constant while increasing $\mu$. This can be continuously followed as $\mu$ is increased, until the eventual jump of $n$ towards the unique solution present at the higher values of $\mu$. This procedure is later used to determine the location of the coexisting region [43] in the phase diagram. The Green’s functions corresponding to these values of $n_f$ are distinct and are shown in Fig. 2.8.

As we increase the interaction to $U = 4.0$ we find that at $\tilde{\mu} = 0$ and $T = 1/\beta = 1/64$ only the insulating solution survives (Fig. 2.9. If we start from a large (hole) doping where only the metallic solution survives and decrease the magnitude of the chemical potential, we find that the metallic solution disappears at a value of the chemical potential $\mu \approx 1.07$. At the same time, if we start at $\tilde{\mu} = 0$ where $n_f = 1$ and increase the magnitude of doping, the insulating solution disappears at a chemical potential
Figure 2.7: Coexisting solutions for $U=2.44$, $\beta = 64$ in the single band case.

Figure 2.8: Imaginary part of Green’s functions for $U=2.44$, $\beta = 64$
\( \mu \approx 1.03 \). However at a higher temperature \( T = 1/\beta = 1/40 \) no coexistence between metallic and insulating solutions is seen.

**Figure 2.9:** Left: Occupation number for various values of \( \beta \). Right: Occupation number for \( \beta = 64 \) metallic and insulating solution

**Coexistence in the 2-band model**

We find coexisting metallic and the insulating solutions near \( n_f = 1 \) at \( T = 1/40 \). In Fig. 2.10 we see the \( n_f \) versus. \( \mu \) curves in the 2-band case for \( U = 3.0 \) at various temperatures. As \( T \) is decreased, we see that the curves begin to acquire a sigmoidal shape, which is a hallmark of the approach to a second-order critical point in Landau theory of phase transitions.

As discussed in the 1-band case, we followed a careful procedure to obtain coexisting solutions. At \( T = 1/40 \) two different solutions were found at the same chemical potential \( \mu = 1.2 \). The metallic solution (the circles in Fig. 2.10) was obtained using as starting point, the solution for \( U = 2.8 \) and the insulating solution (the triangles in the n versus \( \mu \) curve in Fig. 2.10) was obtained by using the Green’s function at \( U=3.5 \) as the seed (Fig. 2.11). Using these solutions the \( n_f \) versus. \( \mu \) curve slightly away from \( n_f = 1 \) was obtained. The insulating solution disappeared for increased doping and away from \( n_f = 1 \).
We then study the nature of the solutions as the temperature is raised. Each of the metallic and the insulating solutions at $T = 1/40$ was used as a seed to determine the solution at higher temperatures. It was found that the metallic solution disappeared as the temperature was raised (to $T = 1/32$) near $n_f < 1$ but the insulating solution survived. However at $\mu = 1.25$ for which $n_f > 1$ it was found that both solutions survived till $T = 1/32$. As the temperature was further raised, no coexistence was seen. The Green’s function for the two phases at $\mu = 1.2$ for $T = 1/40$ and $T = 1/64$ are shown in Fig. 2.12.

Figure 2.10: Particle occupation $n$ function of $\mu$ for different temperatures in the 2-band model at $U=3.0$. The top panel shows curves with $T > T_{MIT}$ and in the bottom panel we have $T < T_{MIT}$. 
Figure 2.11: Left: Metallic seed obtained from $U=2.8$. Right: Insulating seed obtained from $U=3.5$

Figure 2.12: Metallic and insulating Green’s function for $U = 3.0$, $\beta = 40$(left) and $\beta = 64$(right)

2.4.3 Convergence of solutions

We present a few technical details regarding the stability of the solutions, especially near a phase boundary. To determine the convergence of the self-consistent solutions, we examined the difference in the imaginary part of the self energy at the first Matsubara frequency $\text{Im}\Sigma(\omega)$ as $\omega \to 0$ between two successive iterations $i$. The iterations were halted when this difference $\|\text{Im}\Sigma^{i+1}(\omega \to 0) - \text{Im}\Sigma^i(\omega \to 0)\|$ became smaller than a threshold.

If we are away from the coexistence region, the solution converges in 12–15 iterations.
when the initial guess is close to the solution. At a lower value of \( U \) where there is only one solution, the Green’s function converges in \( \sim 20 \) iterations even with an initial guess far from the true solution. Each DMFT iteration consists of 100000 Monte Carlo sweeps. However, close to the coexistence region, the number of iterations required for convergence becomes large. At the boundaries of the region, the number of iterations is expected to become very large due to critical slowing down.

The convergence of the metallic Green’s function in Fig. 2.12 at \( U = 3.0, \mu = 1.225 \) starting from the metallic seed is shown in Fig. 2.13(left). The convergence of the insulating Green’s function in Fig.2.12 is shown in Fig. 2.13(right).

\[ \begin{align*}
\text{Figure 2.13: Convergence of metallic (left) and insulating (right) Green’s function}
\end{align*} \]

### 2.4.4 Compressibility divergence

From the \( n_f-\mu \) curves, we compute the numerical derivative of the particle number with respect to chemical potential, which is proportional to the compressibility. In the case of the finite temperature doping driven MIT, we observe a divergence in compressibility as we approach the second order transition point [44].

Fig. 2.14 shows the behavior of \( \kappa = \Delta n/\Delta \mu \) as a function of \( \mu \) and \( T \) for \( U=4.0 \) in the 1-band model. We can see that near \( \mu = 1.07 \), which we identify as \( \mu_c \), \( \kappa \) increases rapidly indicating a divergence near the phase boundary where the insulating solution disappears. (Figs. 2.15 shows that as temperature is decreased \( \kappa^{-1} \) decreases rapidly indicating a divergent compressibility as we pass through the MIT. The data for \( \kappa^{-1} \)
as a function of $T$ was fitted to a power law. It was found that a power law type of behavior $\kappa^{-1} \sim (T - T_c)^b$ with $b \approx 0.33$ for $U = 2.44$ matches the data.

![Graph showing compressibility as a function of chemical potential and inverse compressibility as a function of temperature.](image)

**Figure 2.14:** 1-band model, $U=4.0$. Top: Compressibility as a function of chemical potential. Bottom: Inverse compressibility $\kappa^{-1} = (dn/d\mu)^{-1}$ at $U=4.0$ in the 1-band model as a function of $T$. The solid line is the function $\kappa^{-1} \sim (T - T_c)^b$ with $b \approx 0.47$.

### 2.5 Phase diagram

From the above numerical results we propose a schematic phase diagram for the Hubbard model, which for the 1-band case is shown in Fig. 2.16. The regions of
Figure 2.15: Left: Inverse compressibility $\kappa^{-1} = (dn/d\mu)^{-1}$ at $U=2.46$ in the 1-band model. The solid line is the function $\kappa^{-1} \sim (T - T_C)^b$ with $b \sim 0.33$. Right: $\kappa^{-1}$ at a constant doping $n = 1.003$ as a function of $T$ in the 2-band model. The solid line is a fit to a power law using the functional form $\kappa \approx (T - T_c)^b$ with the power $b \sim 1/3$. The intercept with the $T-$axis gives our estimate for $T_c = 0.0288$.

Coexisting solutions are shown as cross sections of constant $U$ in the $(U, T, \mu)$ parameter space. The $\mu$-axis starts at zero doping, thus $n = 1$ on the $(U, T)$-plane where $\bar{\mu} = 0$, which for the 1-band model is $\bar{\mu} = \mu - U/2$. At larger values of $U$, the regions of coexisting solutions are two triangles, one for $n < 1$ and the other for $n > 1$. In the 1-band model, these triangular peaks are symmetric about $\mu = U/2$. These triangles are delimited by the $\mu_{c1}(T)$ and $\mu_{c2}(T)$ lines which correspond to the disappearance of the insulating and the metallic solutions respectively. As $U$ decreases, the triangular regions approach each other and fully merge at $U = U_{MIT}$. Further lowering $U$ makes the single triangular region diminish until $U = U_{c1}(T = 0)$ where it vanishes.

The coexistence region in Fig. 2.7 curve at $U = 2.44$ and $\beta = 1/T = 64$ near $\bar{\mu} = 0$ correspond to the region in Fig. 2.16 where $U_{MIT} < U < U_{c2}$. For the parameters in Fig. 2.7, we see that the metallic state exists all the way down to zero doping at $\bar{\mu} = 0$. This implies that for $U_{MIT} < U < U_{c2}$, the $\mu_{c2}(T)$ line does not go all the way to $T = 0$ in contrast to the case when $U > U_{c2}$ (Fig. 2.9).

The interaction $U = 4.0$ is greater than $U_{c2}$ in the 1-band model. Using our results for $U = 4.0$ and those obtained in [45], we can map out the part of the phase diagram
in Fig. 2.16 that corresponds to the parameter space where $U > U_{c2}$ and $n < 1$. In this case, at zero doping, only the insulating solution survives at low temperatures. The results are shown in Fig. 2.17.

The disappearance of the metallic or the insulating solutions is captured by the critical slowing down of the solutions as shown in Fig. 2.18. As we approach the phase boundaries of the coexistence region, it is seen that the number of QMC iterations required for the solutions to converge increase. In Fig. 2.18 it is seen that starting with the metallic solution at low doping, if we increase $\mu$, the number of iterations needed for the metallic solution to converge diverges, indicating that we are near the $\mu_{c2}$ boundary. Similarly starting with the insulating solution near half-filling, if we decrease $\mu$, i.e go far away from half-filling the number of iterations required for the convergence of the insulating solution diverges. This indicates that we are close to the $\mu_{c1}$ boundary.

The phase diagram for the 2-band model shown in Fig. 2.19. is qualitatively similar to the 1-band model except that the pairs of triangles at a given $U$ do not have the same height due to absence of particle-hole symmetry. The $n_f-\mu$ curves in Fig. 2.10 correspond to the region in the phase diagram where $U_{MIT} < U < U_{c2}$. As $T$ is decreased, the curves show a crossover from small to large compressibility.

2.6 Discussion

A common feature that emerges from the model is that, in the regions where two mean-field solutions exist, the system has two different values of $n$ for given $T$ and $\mu$. Furthermore, these two solutions have different free energies and the actual thermodynamic state of the system is that of minimum energy. Hence a jump in particle number is predicted at a first order line. The determination of this line implies a precise calculation of the free energy, which is technically difficult, and outside the scope of this work.

In the $T-\mu-U$ phase diagram, the two mean-field solutions exist within the triangular regions. A first order transition exists where the free energies cross. At finite $T$, this
Figure 2.16: Schematic phase diagram for the 1-band Hubbard model. The cross sections are on the $T - \mu$ plane for different values of $U$ with $U$ increasing towards the top of the figure. The peaks are symmetric about $\mu = U/2$. $\mu_{c1}$ and $U_{c1}$ are the chemical potential and interaction respectively below which the insulating solution does not survive. $\mu_{c2}$ and $U_{c2}$ are those above which the metallic solution does not exist. The black circle denotes the second order transition point $(U_{MIT}, T_{MIT})$ at $n = 1$. The shaded regions denote the coexistence region between metallic and insulating phases.
leads to a first order transition surface between an insulating and a metallic-like state. The intersections of this surface with the constant $U$ cross sections are first order lines that we denote by dashed lines in Fig. 2.16. At $T = 0$ within the coexistence regions, i.e. the base of the triangles, the metallic state is always stable, thus one can cross the first-order surface towards the insulator by either increasing $T$ or changing the chemical potential.

At finite $T$, the two solutions merge where the triangular regions end. Hence there is a line of second order transitions where the first order surface ends (thick double-line in Fig. 2.16. The doping is small but non-zero along this line except at $U = U_{MIT}$. Our numerical results show a divergence of compressibility on this line of second order transitions. The idea of divergent compressibility at the Mott endpoint has been shown to arise from general Landau theory arguments [46].

From the experimental viewpoint, we believe that our results highlight important
Figure 2.18: Critical slowing down observed in the vicinity of $\mu_{c1}$ and $\mu_{c2}$ for the 1-band model at $U = 4.0, \beta = 64$. The inverse number of iterations required for the metallic solution to converge goes to zero as we near $\mu_{c2}$ and those of the insulating solution to converge goes to zero as we approach $\mu_{c1}$. The arrows indicate the directions in which the boundaries were approached.
Figure 2.19: Schematic phase diagram for the 2-band case. There is an asymmetry in the triangular peaks as compared to the 1-band case. The cross sections are on the $T-\mu$ plane for different values of $U$ as before. $\mu_{c1}$ and $U_{c1}$ are the chemical potential and interaction respectively below which the insulating solution does not survive. $\mu_{c2}$ and $U_{c2}$ are the chemical potential and interaction above which the metallic solution does not exist. $U_{MIT}$ is the value of the interaction at which the metal-insulator transition takes place. The shaded portions are the regions of coexistence between the metallic and the insulating solutions.
aspects in the $\alpha$-$\gamma$ transition in Ce. The divergence of compressibility in the Ce $\alpha$-$\gamma$ transition has an electronic origin and can be understood from model calculations. The decrease in compressibility that we found during the transition from the insulating to the metallic phase is similar to what has been measured by Beecroft and Swenson [30].

2.7 Summary

We have shown that within the dynamical mean-field theory of the Hubbard model, there is a region where the paramagnetic metallic and paramagnetic insulating phases coexist. We have explored the coexistence region for the 1-band Hubbard model both at the particle-hole symmetric point and away from it. For the 2-band Hubbard model we have studied the region in phase space near the occupation number $n_f = 1$. We calculated the compressibility from the $n_f$ versus $\mu$ curve. We find numerical evidence for a divergent compressibility near the finite temperature Mott endpoint. Based on our numerical studies we have proposed a phase diagram for the doped 1- and 2-band Hubbard model. Our results are relevant to the Ce $\alpha$-$\gamma$ transition.
Chapter 3

Density Functional Theory, LDA, LDA+U and LDA+DMFT

In this chapter, we first briefly review the density functional theory (DFT) local density approximation (LDA). We then discuss methods that go beyond the LDA to tackle strongly correlated systems: LDA+U and LDA+DMFT. We also discuss the structure of the LMTART program that is based on DFT methods.

3.1 Density functional theory

3.1.1 LDA

Density functional theory is a powerful tool to study weakly interacting systems. It is the basis of the LDA and the LDA+U methods which are used to calculate the band structures of materials. A brief description of DFT from the effective action point of view is presented here [8], [9], [47].

Consider a fermionic system that is coupled to an external source $J(x)$. The Hamiltonian is

$$H = H_0 + \int dx^3 \psi^\dagger(x)J(x)\psi(x)$$

(3.1)

where $H_0$ is the Hamiltonian without the source. The partition function is

$$Z = \exp[-W[J]] = \int D[\psi\psi^\dagger] \exp - \int dt L$$

(3.2)

where $L$ is the Lagrangian.

For the system consists of electrons moving in a crystal potential $V_c(x)$ and interacting via Coulomb interactions $V$, in the presence of an external source $J$ coupled to
the electron density, the partition function is

\[ Z = \int D[\psi \psi^\dagger] \exp \left[ -\int dx \psi^\dagger(x)[\partial_x - \frac{\nabla^2}{2m}] \psi(x) \right] \]

\[ \frac{1}{2} \int dx dx' \psi^\dagger(x)\psi^\dagger(x')V(x-x')\psi(x')\psi(x) - J(x)\psi^\dagger(x)\psi(x) \] (3.3)

Here \( x = (x, \tau) \) denotes the space–imaginary time coordinates. By performing a Legendre transform of \( \Gamma \) we can eliminate the potential in favor of the density \( \rho \).

\[ \Gamma[\rho] = W[J] - \int J(x)\rho(x) \] (3.4)

The minimum of this functional gives the true density and the total energy.

To construct approximations to the functional \( \Gamma \) it is very useful to introduce the Kohn–Sham potential, \( V_{KS} \), which is defined as the potential such that when added to the non–interacting kinetic energy, it produces the given density in a reference system of non–interacting particles . i.e.

\[ \rho(r) = T \sum \sum_{\omega_n} \langle \sigma r \mid (i\omega_n + \nabla^2/2 - V_{KS})^{-1} \mid \sigma r \rangle e^{i\omega r^+} \] (3.5)

The exact functional can now be viewed as a functional of two variables

\[ \Gamma(\rho, V_{KS}) = -T \sum_{\omega_n} \text{tr} \log[i\omega_n + \nabla^2/2 - V_{KS}] - \int V_{KS}(r)\rho(r) dr + \frac{1}{2} \int \frac{\rho(r)\rho(r')}{|r - r'|} dr dr' + \int V_{ext}(r)\rho(r) dr + E_{xc}[\rho] \] (3.6)

\( \Gamma(\rho) \) is obtained by substituting \( V_{KS}(\rho) \) obtained by solving eqn.(3.5) (which makes eqn.(3.6) stationary) into \( \Gamma(\rho, V_{KS}) \). \( E_{xc}[\rho] \) is the exchange–correlation energy which is a functional of the density and not of the external potential.

Extremizing eqn.(3.6) with respect to \( \rho \) gives

\[ V_{KS}(r)[\rho] = \int \frac{\rho(r') dr'}{|r - r'|} + V_{xc}(r)[\rho] + V_{ext}(r)[\rho] \] (3.7)

where \( V_{xc}(r) \) is the exchange-correlation potential obtained as

\[ \frac{\delta E_{xc}}{\delta \rho(r)} = V_{xc}(r) \] (3.8)

If we restrict ourselves to zero–temperature and interpret the Fermi functions as step functions, Eqn. (3.5) can be rewritten as

\[ [-\nabla^2/2 + V_{KS}(r)]\psi_{kj}(r) = \epsilon_{kj}\psi_{kj}(r) \] (3.9)
\[ \rho(\mathbf{r}) = \sum_{k_j} f(\epsilon_{k_j}) \psi_{k_j}^*(\mathbf{r}) \psi_{k_j}(\mathbf{r}) \] (3.10)

and \( V_{KS} \) is given as an explicit function of the density.

The total energy of the crystal is given as

\[ E_{tot} = \sum_{k_j} f(\epsilon_{k_j}) \epsilon_{k_j} + \frac{1}{2} \int \frac{\rho(\mathbf{r}) \rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r} d\mathbf{r}' + \int V_{ext}(\mathbf{r}) \rho(\mathbf{r}) d\mathbf{r} + \int \epsilon_{xc}[\rho(\mathbf{r})] \rho(\mathbf{r}) d\mathbf{r} + E_{dc} \] (3.11)

where

\[ E_{dc} = - \int V_{KS}(\mathbf{r}) \rho(\mathbf{r}) d\mathbf{r} \] (3.12)

simply subtracts the interaction energy from the Kohn–Sham eigenvalues which are explicitly included in the Hartree and exchange–correlation term to avoid double counting.

Since \( E_{xc}[\rho] \) is not known explicitly, this method is useful only because of successful approximations to the exchange energy. In the Local Density Approximation (LDA) the exchange energy functional is assumed to be local:

\[ E_{xc}[\rho] = \int \epsilon_{xc}(\rho(\mathbf{r})) \rho(\mathbf{r}) d\mathbf{r} \] (3.13)

with \( \epsilon_{xc}[\rho(\mathbf{r})] \) being the energy density of the uniform electron gas.

Here \( \rho(\mathbf{r}) \) is uniquely expressed in terms of the orbitals \( \psi_{k_j}(\mathbf{r}) \). In order to truncate the DFT, we introduce a finite basis set \( \chi_{\alpha}^k(\mathbf{r}) \) and expand

\[ \psi_{k_j}(\mathbf{r}) = \sum_{\alpha} \chi_{\alpha}^k(\mathbf{r}) A_{\alpha}^{k_j} \] (3.14)

keeping a finite set of \( \alpha \). This truncation restricts the active part of the multiplicative operator associated with the potential \( V_{KS} \) to have a form

\[ \hat{V} = \sum_k |\chi_{\alpha}^k\rangle V_{\alpha\beta} \langle \chi_{\alpha}^k| \] (3.15)

For a known potential \( V_{KS} \) this construction can be done once and for all. However, since \( V_{KS} \) depends on the density, the basis \( |\chi_{\alpha}^k\rangle \) is adapted iteratively to the self–consistent solution.

### 3.1.2 LDA+U

The LDA method is not very successful when strong electron correlations are present as the electrons are localized due to the interactions. The LDA+U method [14], [15]
is a technique that is proposed when strong correlations are present in the system. For simplicity we will deal with spin unrestricted formalism so that the total energy is invariant with respect to orientation. This allows us to choose a specific quantization axis.

We introduce a set of localized orbitals \( \phi_a(\mathbf{r} - \mathbf{R}) \) which are used to build an occupancy matrix. If \( \phi_a \) are identified with the correlated electrons, we can represent the correlated part of the electron density by

\[
n_{ab}^\sigma = \sum_{k} f(\epsilon_{kj}) \int \psi_{kj}^*(\mathbf{r})\phi_a(\mathbf{r})d\mathbf{r} \int \psi_{kj}(\mathbf{r}')\phi_b^*(\mathbf{r}')d\mathbf{r}' \tag{3.16}
\]

The total energy is now a functional of \( \rho(\mathbf{r}) \) and of \( n_{ab}^\sigma \). The LDA+U functional is

\[
\Gamma_{\text{LDA}+U}[n_{ab}^\sigma, \lambda_{ab}^\sigma, V_{KS}^\sigma, \rho^\sigma] = -T \sum_{\omega_n} \text{tr} \log[i\omega_n + \nabla^2 / 2 - V_{KS}^\sigma - \sum_{ab} \lambda_{ab}^\sigma \phi_a(\mathbf{r})\phi_b^*(\mathbf{r})] \\
- \sum_{\sigma} \int V_{KS}^\sigma(\mathbf{r})\rho^\sigma(\mathbf{r})d\mathbf{r} - \sum_{\sigma} \sum_{ab} \lambda_{ab}^\sigma n_{ab}^\sigma + \frac{1}{2} \int \frac{\rho(\mathbf{r})\rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}d\mathbf{r}d\mathbf{r}' \\
+ E_{\text{dc}}^{\text{LDA}}[\rho^\sigma] + E_{\text{Model}}[n^\sigma] - E_{\text{dc}}[n^\sigma] 
\tag{3.17}
\]

where \( \lambda_{ab}^\sigma \) are Lagrange multipliers.

\[
E_{\text{Model}}[n^\sigma] = \frac{1}{2} \sum_{\sigma} \sum_{abcd} U_{abcd} n_{ab}^\sigma n_{cd}^\sigma + \frac{1}{2} \sum_{\sigma} \sum_{abcd} (U_{abcd} - J_{abcd}) n_{ab}^\sigma n_{cd}^\sigma 
\tag{3.18}
\]

is the contribution from the Coulomb energy in the shell of correlated electrons. \( E_{\text{dc}}[n^\sigma] \) is the double-counting term which is subtracted as part of the energy in \( E_{\text{Model}} \) has been taken into account in the LDA part.

The form of the double counting term [14] used here is

\[
E_{\text{dc}} = \frac{1}{2} \tilde{U} \tilde{n}(\tilde{n} - 1) - \frac{1}{2} \tilde{J}[\tilde{n}^\dagger(\tilde{n}^\dagger - 1) + \tilde{n}^\dagger(\tilde{n}^\dagger - 1)], 
\tag{3.19}
\]

where

\[
\tilde{U} = \frac{1}{(2l + 1)^2} \sum_{ab} \langle ab \rangle \frac{1}{r}|ab \rangle \tag{3.20}
\]

\[
\tilde{J} = \tilde{U} - \frac{1}{2l(2l + 1)} \sum_{ab} \langle \langle ab \rangle \rangle \frac{1}{r}|ab \rangle \tag{3.21}
\]

Here \( \tilde{n}^\sigma = \sum_a n_{aa}^\sigma \), and \( \tilde{n} = \tilde{n}^\dagger + \tilde{n}^\dagger \).
We get the self-consistency equations by extremizing the functional with respect to $V_{KS}$.

$$\rho^\sigma(r) = T \sum_{\omega_n} \left| \left[ i\omega_n + \nabla^2/2 - V_{KS}^\sigma - \sum_{ab} \lambda_{ab}^\sigma \phi_a(r)\phi_b^*(r') \right]^{-1} r \right| e^{i\omega_n 0^+} = \sum_{kj} f(c_{kj\sigma}) |\psi_{kj\sigma}(r)|^2 \tag{3.22}$$

with

$$[-\nabla^2/2 + V_{KS}^\sigma + \sum_{ab} \lambda_{ab}^\sigma \phi_a(r)\phi_b^*(r')]|\psi_{kj\sigma} = c_{kj\sigma} |\psi_{kj\sigma}$$ \tag{3.23}

$V_{KS}^\sigma$ is obtained by extremizing the functional w.r.t. $\rho(x)$:

$$V_{KS}(r)[\rho] = \int \frac{\rho(r') dr'}{|r-r'|} + V_{xc}(r)[\rho] + V_{ext}(r)[\rho] \tag{3.24}$$

We get the correction to the potential $\lambda_{ab}^\sigma$ by extremizing the functional w.r.t. $n_{cd}^\sigma$

$$\lambda_{ab}^\sigma = \sum_{cd} U_{abcd} n_{cd}^\sigma + \sum_{cd} (U_{abcd} - J_{abcd}) n_{cd}^\sigma - \frac{dE_{dc}[n^\sigma]}{dn_{ab}^\sigma} \tag{3.25}$$

The double counting term generates the correction to the potential in the form:

$$\lambda_{ab}^\sigma = \sum_{cd} U_{abcd} n_{cd}^\sigma + \sum_{cd} (U_{abcd} - J_{abcd}) n_{cd}^\sigma - \bar{U}(\bar{n} - \frac{1}{2}) + J(\bar{n}^\sigma - \frac{1}{2}) \tag{3.26}$$

The interaction matrices are defined as:

$$U_{abcd} = \langle ac \ | \ bd \rangle = \int \phi_a^*(r) \phi_c^*(r') v_C(r-r') \phi_b(r) \phi_d(r') dr dr' \tag{3.27}$$

$$J_{abcd} = \langle ac \ | \ db \rangle = \int \phi_a^*(r) \phi^*_c(r') v_C(r-r') \phi_d(r) \phi_b(r') dr dr' \tag{3.28}$$

where $\phi(r)$ are the set of projectors and $v_C(r-r')$ is the Coulomb interaction that has to take into account the effects of screening by conduction electrons.

These matrices are expressed in terms of Slater integrals. Representing $\phi_{tm}(r) = \phi_t(r) Y_{lm}(\hat{r})$ and for $a \equiv lm, b \equiv lk, c \equiv l'm', d \equiv l'k'$ we can express $U$ and $J$ as

$$\langle ml'm' \ | \ l'k' \rangle = \sum_{l''=0,2,...}^{\min(2l', 2l'')} \frac{4\pi}{2l''+1} l''^{l''} C^{l''l''}_l m'' m' = -k C^{l''l''}_l m'' m' = k' - m' \delta_{k'-m', m-k} \tag{3.29}$$

$$\langle ml'm' \ | \ l'k' \rangle = \sum_{l''=0,2,...}^{\min(2l', 2l'')} \frac{4\pi}{2l''+1} l''^{l''} C^{l''l''}_l k' \delta_{m'' m' = k - m'} \delta_{m-k', k'-m} \tag{3.30}$$
where the quantities $F^{(u)}$ and $F^{(j)}$ are given by the following radial integrals

$$F_{l''}^{(u)} = \int \frac{r^{l''}}{r^{l''+1}} \phi_1^2(r) \phi_{l''}^2(r') dr dr' \quad (3.31)$$

$$F_{l''}^{(j)} = \int \frac{r^{l''}}{r^{l''+1}} \phi_1(r) \phi_{l''}^2(r') \phi_l(r') dr dr' \quad (3.32)$$

If $l = l'$, $F^{(u)}$ and $F^{(j)}$ are equal and are reduced to constants. For $d$-electrons, there are three constants $F^{(0)}$, $F^{(2)}$, $F^{(4)}$ and for $f$-electrons, there are four constants $F^{(0)}$, $F^{(2)}$, $F^{(4)}$, $F^{(6)}$. The Slater integrals can be linked to $U$ and $J$ via $U = F^{(0)}$ and $J = (F^{(2)} + F^{(4)})/14$ for $d$-electrons with $F^{(2)}/F^{(4)} = 0.625$. For $f$-electrons, $U = F^{(0)}$ and $J = (286F^{(2)} + 194F^{(4)} + 250F^{(6)})/6435$.

### 3.1.3 LDA+DMFT method

In the LDA+U method, the Coulomb interaction in eqn.(3.18) are treated in a Hartree-Fock approximation, thus it fails to describe the many body nature of the electron-electron interaction in strongly correlated materials. Also, in the LDA+U method, we do a Legendre transformation with respect to a part of the density, which lacks a clear physical significance. A method that has been developed to treat correlated materials, is LDA combined with the Dynamical Mean-Field Theory (DMFT). The DMFT has been an extremely successful method to treat model Hamiltonians that describe materials with strong correlations. However, the model Hamiltonians used have parameters. By combining LDA+DMFT, we would like to introduce microscopic details to the approach and make the calculation first-principles [47].

The following section contains a brief description of the LDA+DMFT method from an effective action point of view [13,48].

In DMFT, we deal with functionals of the local Green’s function, which is a well defined object. We start with the LDA+DMFT functional $\Gamma_{LDA+DMFT}(\rho, V_{KS,\chi}, A)$
which has the form
\[
\Gamma_{LDA+DMFT}(\rho, V_{KS}, \chi, A) = -T \sum_{i\omega_n} tr \log[i\omega_n + \nabla^2/2 - V_{KS} - \Sigma_{ab}(i\omega_n)\phi_a(r)\phi_b^*(r')] -
\int V_{KS}(r) \rho(r) dr - \sum_{i\omega_n} \sum_{ab} \Sigma_{ab}(i\omega_n) A_{ba}(i\omega_n) +
\int V_{ext}(r) \rho(r) dr + \frac{1}{2} \int \frac{\rho(r) \rho(r')}{|r - r'|} dr dr' + E_{xc}^{LDA}[\rho] +
\sum_j \Phi[A_{ab}(j, i\omega)] + \Phi_{DC}
\] (3.33)

Φ is the sum of all two–particle irreducible graphs constructed with the local part of the interaction and Φ_{DC} is taken to have the same form as in LDA+U method as described later.

In a fixed tight–binding basis, \(-\nabla^2 + V_{KS}\) reduces to \(H^{TB}(k)\). The functional \(\Gamma_{LDA+DMFT}\) for a fixed density and truncated to a finite basis set takes the form of the DMFT functional
\[
\Gamma[A[\Delta], \chi[\Delta]] =
W_0[\Delta] - Tr \log A - \sum_k \sum_{i\omega_n} Tr \log[i\omega_n - \epsilon(k) - \chi] - \sum_{i\omega_n} [\chi - i\omega_n + \bar{\epsilon} + \Delta] A
\] (3.34)

Its minimization leads to the set of equations with Kohn–Sham potential as in the LDA method, and
\[
\Sigma_{ab}(i\omega_n) = \frac{\delta \Phi}{\delta A_{ab}(i\omega_n)} + \epsilon_{DC}^{ab}
\] (3.35)

which identifies Σ as a self–energy of a generalized Anderson impurity model in a bath characterized by a matrix of levels
\[
\epsilon_0^{ab} = \epsilon_{DC}^{ab} + \sum_k H_{ab}^{TB}(k)
\] (3.36)

and a hybridization function \(\Delta_{ab}(i\omega_n)\) obeying a self–consistency condition
\[
i\omega O_{ab} - \epsilon_0^{ab} - \Delta_{ab}(i\omega_n) = \Sigma_{ab}(i\omega_n) + \left[\sum_k (i\omega_n O - \epsilon^0 - t(k) - \Sigma(i\omega_n))^{-1}\right]^{-1}
\] (3.37)

Finally, minimizing eqn.(3.33) with respect to \(V_{KS}\) indicates that \(\rho(r)\) should be computed as
\[
\rho(r) = T \sum_{i\omega_n} \left< r \left| [i\omega_n + \nabla^2/2 - V_{KS} - \sum_{ab} \Sigma_{ab}(i\omega_n)\phi_a(r)\phi_b^*(r')]^{-1} r \right| e^{i\omega_n 0^+} \right>
\] (3.38)
which as indicated before, when truncating in a fixed set of orbitals becomes

$$\rho(r) = T \sum_{\omega_n} \chi_\alpha(r) \left[ (i\omega_n - H^{TB}(k) - \Sigma(i\omega_n))^{-1} \chi_\beta^*(r) \right]$$  \hspace{1cm} (3.39)$$

We express the functional in terms of the Weiss field, i.e. the hybridization function that added to the atom produces the exact spectral function. This allows us to eliminate the functional $\Phi$. We can express the LDA+DMFT functional as:

$$\Gamma_{\text{LDA+DMFT}}(\rho, V_{KS}, A, \Delta) = -Tr \log[\omega_n + \nabla^2/2 - V_{KS} - \Sigma_{ab}(i\omega_n, r, r')(\chi)] -$$

$$\int V_{KS}(r)\rho(r)dr - \sum_{i\omega_n} \sum_{ab} Tr[\Sigma_{ab}(i\omega_n)A_{ba}(i\omega_n)] +$$

$$\int V_{ext}(r)\rho(r)dr + \frac{1}{2} \int \frac{\rho(r)\rho(r')}{|r-r'|} d^3r d^3r' + E_{LDA}[\rho] +$$

$$+ \Phi_{DC} + W_{at}[\Delta[A]] - Tr \Delta[A]A - Tr \log A + Tr A_{at}^{-1} A$$

\hspace{1cm} (3.40)

Here

$$\Sigma_{ab}(i\omega_n, r, r')(\chi) = \frac{1}{N_s} \sum_{\alpha\beta R_p R_l R_q} \chi_\alpha(r-R_p)O^{-1}(R_p-R_l)a_{\alpha\beta}\chi_\beta^*(r-R_q)$$

where $N_s$ is the number of lattice sites and $W_{at}[\Delta]$ is simply the free energy of the atom (described by the atomic Hamiltonian $h_{at}$) in the presence of a hybridization field.

$$S_{at}[\Delta] = \int_{\tau, \tau'} \sum_{i,\sigma} c_{i\sigma}^\dagger(\tau) \left[ \delta(\tau - \tau') \frac{\partial}{\partial \tau'} + \Delta^{ln}(\tau - \tau') \right] c_{i\sigma}(\tau') + \int d\tau h_{at}$$

\hspace{1cm} (3.41)

$$W_{at}[\Delta] = - \log \int dc^+ dc e^{-S_{at}[\Delta]-\sum_{ab} \int_{\tau, \tau'} \Delta(\tau, \tau')c_{a\tau}^\dagger(\tau)c_{a\tau}(\tau')d\tau d\tau'} \hspace{1cm} (3.42)$$

$$\Phi_{dc_{Model}} = \frac{1}{2} \tilde{U} \bar{n}(\bar{n} - 1) - \frac{1}{2} \tilde{J}[\bar{n}^\dagger(\bar{n}^\dagger - 1) + \bar{n}^\dagger(\bar{n}^\dagger - 1)]$$

\hspace{1cm} (3.43)

with $U$ and $J$ as defined in eqns.(3.20) and (3.21).

Here $\bar{n}^\sigma = \sum_{a} n_{a\sigma}^a$, and $\bar{n} = \bar{n}^\dagger + \bar{n}^\dagger$. $n$ in this case should be viewed as a functional of the local Greens function $n = tr A = T\sum_{\omega_n, a} e^{i\omega_n} A(\omega_n)_{aa}$

This generates the correction to the potential in the form:

$$\lambda_{ab}^\sigma = \sum_{cd} U_{abcd}n_{cd}^\sigma + \sum_{cd} (U_{abcd} - J_{abcd})n_{cd}^\sigma - U(\bar{n} - \frac{1}{2}) + J(\bar{n}^\sigma - \frac{1}{2}) \hspace{1cm} (3.44)$$

To solve the impurity model, various techniques such as Quantum Monte Carlo, Gutzwiller method or approximations such as the Hubbard-I method can be used.
3.2 Structure of the LMTART program

In this section we shall describe the overall structure and details of certain parts of the LMTART program, which is based on the equations in section 3.1.

The LMTART package contains Fortran 90 programs which can be grouped according to the function they perform. Furthermore, the programs are named such that the first three letters of the program name in each group are the same. The names of the programs in the LMTART package can be thought of as having two parts such as: `abc_*`.f where the first part `abc_` denotes the group it can be classified into, and the second part (denoted by the wildcard character `*`) is different for each program in the group. We first provide a list (alphabetical) that contains a description of the various programs grouped according to the task they perform. Following that, Fig. 3.1 shows an overview of the schematic flow diagram of the LDA+DMFT implementation.

Programs with the name:

- `bnd_*`.f
  Form the core of the solution to the generalized eigenvalue problem in the LMTO method. This includes preparing Hamiltonian matrix and overlap matrix (after factoring in contributions due to the spin-orbit coupling), calculating eigenvalues and eigenvectors, and preparing wave functions. Also calculating the fat-bands when required.

- `chi_*`.f
  Calculate the susceptibility.

- `dmf_*`.f
  Implement DMFT loop. Solve impurity model to find Green’s function. Execute DMFT self-consistency condition.

- `frc_*`.f
  Calculate total energy and band energy in LDA, and various contributions to force by computing the derivative of total energy w.r.t. nuclear displacements.

- `fth_*`.f
Calculate Hamiltonian, overlap matrices and all physical quantities such as total energy, in the tight-binding scheme.

In particular, see \texttt{ftb\_ftbands.f} which is the control module to solve the eigenvalue problem using the tight-binding method.

- \texttt{hop\_*.f}
  
  Read hopping integrals from HOPFILE, prepare hopping matrix, tight-binding hopping integrals are calculated by integrating over Brillouin zone, also calculate screened structure constants.

- \texttt{hub\_*.f}
  
  Read HUBFILE which contains Slater integrals necessary for constructing U-matrix for the LDA+U method, compute potential in the LDA+U scheme from Hubbard-U parameters, calculate occupation matrices needed for LDA+U method.

- \texttt{ini\_*.f}
  
  Control the initial procedures to be performed before the main solution of the eigenvalue problem begins.

  - \texttt{ini\_get\_*.f}: read data for various physical quantities, \textit{e.g.} \texttt{ini\_getchi.f} reads susceptibility data, \texttt{ini\_getbnd.f} reads fatband data.
  
  - \texttt{ini\_make\_*.f}: controls initial calculation of crystal group, k-mesh.
  
  - \texttt{ini\_read\_*.f}: reads data, parameters from initial files defined by the user.

  - In particular, see \texttt{ini\_channels.f}: Allocation of different channels inside the program, \textit{i.e.} describes the channel numbers for all the input and output files.

- \texttt{lib\_*.f}
  
  Contain libraries to perform standard mathematical operations such as computing spline approximations, evaluating spherical and cubic harmonics, calculating derivatives, computing Pade coefficients etc.

- \texttt{man\_*.f}
  
  This set of files form the main control module and contain information on which
subroutines will be implemented depending on which options are given by the user.

- *main.f*: Main module of the entire program. Reads inputs, directs to either ASA self-consistent method or PLW full potential self-consistent method or Tight-binding method.

- *lmtscf.f*: Main control module for self-consistency stream calls various routines depending on options given by the user.

- *ftbscf.f*: Main control module for tight-binding stream, calls various programs depending on options given by user.

- *artout.f*: writes outputs.

- *opt_*.*f*
  Calculate optical matrix elements and optical properties.

- *pot_*.*f*
  Calculate Coulomb potential, exchange-correlation energy and potential, multipoles.

- *qmc_*.*f*
  Contain routines for Quantum Monte Carlo solution of the impurity model.

- *rho_*.*f*
  Implement operations with charge density files: *e.g.* renormalizing core density, admixture in scf loop, Broyden mixing.

- *sgm_*.*f*
  Perform calculations and implement procedures involving self-energy.

- *str_*.
  Prepare structure constants and derivatives, generate sites within a sphere, generate vectors in direct and reciprocal space.
Calculate weight functions using tetrahedron method, search for Fermi energy, calculate velocities inside tetrahedron.

Fig. 3.1 shows the schematic flow diagram of the LDA+DMFT implementation.

### 3.3 DMFT Loop

In this section, we describe in detail the sub-routines that implement the DMFT loop. The DMFT equations have a self-energy $\Sigma$, (eqn.3.35), which is the self-energy of a generalized Anderson impurity model in a bath characterized by a matrix of levels

$$
\epsilon_{ab}^0 = \epsilon_{ab}^{DC} + \sum_k H_{ab}^{TB}(k)
$$

(3.45)

The hybridization function $\Delta_{ab}(i\omega_n)$ obeys a self-consistency condition

$$
i\omega O_{ab} - \epsilon_{ab}^0 - \Delta_{ab}(i\omega_n) = \Sigma_{ab}(i\omega_n) + \left[\sum_k (i\omega_n O - \epsilon^0 - t(k) - \Sigma(i\omega_n))^{-1}\right]^{-1}
$$

(3.46)

There are various possibilities for the double-counting term $\epsilon_{ab}^{DC}$. One choice is to fix $\epsilon_{ab}^{DC}$ to coincide with $\Sigma(\infty)$, which is equivalent to $U_n$, the Hatree-Fock limit. A different possibility is to fix the double counting term so that the Luttinger theorem is fulfilled, i.e. to choose $\epsilon_{ab}^{DC} = -\frac{1}{N_{lqg}} \delta_{ab} \sum_{ab} \Sigma_{ab}(0)$. 


Synthesize LMTO Hamiltonian and Initial charge density

Overlap matrices $G; G_0 \equiv G_1$

Flow Diagram: LDA+DMFT

Figure 3.1: Flow diagram for LDA+DMFT scheme
The exact sub-routines involved are described below.

**Start DMFT loop**

Input to DMFT loop: $H_{k}^{LDA}, O_{k}$, i.e. LDA Hamiltonian and overlap matrices from temporary file $IUHAM$.

- **Subroutine grfsig**
  
  *Input:* $\Sigma(i\omega)$ from temporary file $IUSGM$.
  
  *Task:* Read $\Sigma(i\omega)$ from IUSGM ($\Sigma = 0$ in 1st iteration)

- **Subroutine grfpot**
  
  *Input:* $\Sigma(i\omega)$.
  
  *Task:* Choose and set double counting $\epsilon_{DC} = \Sigma(0)$ or $\Sigma(\infty)$

- **Subroutine grfbnd**
  
  *Input:* $H_{k}^{LDA}$ and $\Sigma(i\omega)$.
  
  *Task:* Construct $H_{LDA} + \Sigma(i\omega) - \Sigma_{DC}$, compute eigenvectors $A^{L}, A^{R}$ and eigenvalues $\epsilon_{kj}$.

- **Subroutine grffrm**
  
  *Input:* Eigenvalues $\epsilon_{kj}$ of $[H_{LDA} + \Sigma(i\omega) - \Sigma_{DC}]$, filling of electrons from control file.
  
  *Task:* Find Fermi level $\epsilon_{F}$.

\[
\sum_{kj} \sum_{n} \frac{1}{i\omega_{n} + \epsilon_{F} - \epsilon_{kj}} = N_{electrons}
\]

- **Subroutine grfwgtx**
  
  *Input:* Eigenvectors $A^{L}, A^{R}$ and eigenvalues $\epsilon_{kj}$ of $[H_{LDA} + \Sigma(i\omega) - \Sigma_{DC}]$, Fermi level $\epsilon_{F}$.
  
  *Task:* Find weights for k-summation

\[
\sum_{k} \frac{A^{L}A^{R}}{i\omega_{n} + \epsilon_{F} - \epsilon_{k}} = \sum_{k} W_{k}A^{L}A^{R}
\]
• Subroutine **grflev**
  
  **Input:** Eigenvectors $A^L, A^R$ and eigenvalues $\epsilon_{kj}$ of $[H_{LDA} + \Sigma(i\omega) - \Sigma_{DC}]$, Overlap matrix $O_k$, weights for k-summation.
  
  **Task:** Set matrix of impurity levels
  
  $$E_{imp} = \left( \sum_k O_k^{-1} \right)^{-1} \left[ \sum_k O_k^{-1} (H_k^{LDA} + \Sigma(\infty) - \Sigma_{DC}) \sum_k O_k^{-1} \right] \left( \sum_k O_k^{-1} \right)^{-1}$$

• Subroutine **implev**
  
  **Input:** Impurity levels $E_{imp}$.
  
  **Task:** Diagonalizes matrix of impurity levels: used later in impurity solver if diagonal representation is selected.

• Subroutine **grfloc**
  
  **Input:** Eigenvectors $A^L, A^R$ and eigenvalues $\epsilon_{kj}$ of $[H_{LDA} + \Sigma(i\omega) - \Sigma_{DC}]$, Overlap matrix $O_k$, weights for k-summation, Fermi level.
  
  **Task:** Calculate local Green’s function
  
  $$G(i\omega_n) = \left( \sum_k [(i\omega_n + \epsilon_F)O(k) - (H_k^{LDA} + \Sigma(i\omega_n) - \Sigma_{DC})] \right)^{-1}$$

• Subroutine **grfhyb**
  
  **Input:** Fermi level, overlap matrix $O_k$, impurity levels $E_{imp}$, $\Sigma(i\omega)$, $\Sigma_{DC}$, local Green’s function $G(i\omega)$.
  
  **Task:** Calculate hybridization function
  
  $$\Delta(i\omega_n) = (i\omega_n + \epsilon_F)O - (E_{imp} - \Sigma(\infty)) - G^{-1}(i\omega_n) - \Sigma(i\omega_n)$$

• Subroutine **impmod**
  
  **Input:** Hybridization $\Delta(i\omega)$, impurity levels $E_{imp}$, overlap $O_k$.
  
  **Task:** Impurity solver

• Subroutine **grfpot**
  
  **Task:** Set double counting with new $\Sigma(i\omega)$.
  
  $$\epsilon_{DC} = \Sigma(0) \quad \text{or} \quad \Sigma(\infty)$$
Check if number of DMFT iteration is equal to NDIT (In control file: *hubfile*, controls maximum number of DMFT iterations). If so, proceed to next file *grfden* to calculate charge density. If not, use $\Sigma(i\omega)$ from Impurity solver as an input to the next iteration in the self-consistency condition.

- **Subroutine grfden**

  *Input:* New Green’s function constructed from $\Sigma(i\omega)$ calculated by the impurity solver, eigenvectors of $[H_{LDA} + \Sigma(i\omega) - \Sigma_{DC}]$.

  *Task:* Calculate charge density

  $$
  \rho(r) = \sum_{\omega_n} \sum_{ab} \chi_a(r) G_{ab}(i\omega_n) \chi_b(r)
  $$

  from which the outer LDA loop begins again.

### 3.3.1 Impurity solver: Hubbard-I method

The impurity model can be solved using different solvers – exact methods such as Quantum Monte Carlo (computationally expensive for low temperatures) or approximate methods such as Hubbard-I or Gutzwiller approximation. In this section we describe the details of one such method that is based on the Hubbard-I approximation [49, 50].

The equations can be tailored to fit the symmetry of the system in hand. A simple case is when different states for a fixed number of electrons in an atom are considered equivalent. The equations we describe here are suited for a system in which spin-orbit effects are significant. In particular, we deal with an actinide system with 14 $f$-levels that are split into two sub-bands due to spin-orbit coupling. In chapter 5, we discuss the results of this calculation for Americium.

The starting point of the Hubbard-I approximation is the atomic limit. The self-energy describes the two atomic levels and the Green’s function consists of two poles at $i\omega - \mu$ and $\omega - \mu + U$. We first find the Green’s function for the atomic Hamiltonian using the equation of motion method for the Hubbard operators. We then express the impurity Green’s functions in terms of the atomic Green’s function and the hybridization $\Delta(i\omega)$. 
We start with a multi-band impurity model that has the form $H = H_{\text{atom}} + H_{\text{band}} + H_{\text{hyb}}$.

\[
H_{\text{atom}} = \sum_{l,l'=5,7/2}^{m=1,...(2l'+1)} \epsilon_{ll'mm'} f_{lm}^{\dagger} f_{l'm'} + U \sum_{l,l'=5,7/2}^{m,m'} f_{lm}^{\dagger} f_{l'm'} f_{l'm'} f_{lm'}
\]

\[
H_{\text{band}} = \sum_{lk} \epsilon_{lk} c_{lk}^{\dagger} c_{lk}
\]

\[
H_{\text{hyb}} = \sum_{ll'lk} V_{ll'k}(f_{lm}^{\dagger} c_{lk} + \text{h.c.})
\] (3.47)

Here $l, l'$ label the two bands that are split due to spin-orbit coupling. These two bands have the $j$-quantum number as $5/2$ or $7/2$. Notation: The band with $l = 5/2$ and is 6-fold degenerate and that with $l = 7/2$ and is 8-fold degenerate. Thus $m, m'$ label the degeneracy and run from 1 to $(2j+1)$. Since the impurity level matrix $\epsilon_{ll'mm'}$ has off-diagonal terms as zero, the first term simplifies to $\sum_{lm} \epsilon_{l} f_{lm}^{\dagger} f_{lm}$. $H_{\text{hyb}}$ is due to the hybridization between the conduction electrons and the localized $f$ electrons. The hybridization function can be written as $\Delta_l(i\omega) = \sum_{lk} V_{lk}^2/(i\omega - \epsilon_{lk})$.

Due to the above degeneracy, we have a $SU(6) \times SU(8)$ symmetry. With this, the impurity Green’s function in the Hubbard-1 approximation becomes

\[
[G_{l}^{\text{imp}}(i\omega)]^{-1} = [G_{l}^{\text{at}}(i\omega)]^{-1} - \Delta_l(i\omega)
\] (3.48)

The atomic Green’s function $G_{l}^{\text{at}}(i\omega)$ is given by

\[
G_{l}^{\text{at}}(i\omega) = \sum_{n=0}^{14} \sum_{n_1=0}^{n} \left( \frac{C_{n_1}^{6} C_{n-n_1}^{8} \left[ e^{-E_{ln}/T} - e^{-E_{n}/T} \right]}{i\omega + \mu - E_{ln} + E_{n}} \right) / Z
\] (3.49)

where $E_{ln}$ is the energy of an atom with $n$ total number of electrons, $n_1$ in level $l = 1$ and $n-n_1$ in level $l = 2$ is given by:

\[
E_{ln} = E_n + \epsilon_1^f + Un
\] (3.50)

\[
E_n = \frac{U(n-1)n}{2} + n_1 \epsilon_1^f + (n-n_1) \epsilon_2^f
\] (3.51)

The numerator in (3.49) denotes the probability of finding an atom with $n$ total number of electrons, $n_1$ in level $l = 1$ and $n-n_1$ in level $l = 2$. $C_{n_1}^{6}$ and $C_{n-n_1}^{8}$ are the
combinatorial coefficients. \( Z \) is a normalization factor for the probability given by

\[
Z = \sum_{n=0}^{14} \sum_{n_1=0}^{n} \sum_{n_1 \leq 6}^{n_1 \leq 8} C_{n_1}^{n} C_{n-n_1}^{8} e^{-E_n/T} \tag{3.52}
\]

Using the Hubbard-I approximation to solve for the impurity Green’s function, we get a splitting of the upper and lower Hubbard bands for arbitrarily small but finite \( U \).

In the LMTART program the above procedure is implemented in the routine `impatomsun_so` in the file `dmf_imp_atomso.f`. This in turn calls upon `cal_sigma_atomso.f` and `cal_green_atomso.f` to calculate the self-energy and Green’s function respectively. The output is the Green’s function and the probability distribution of finding an atom with \( n \) electrons. At large \( U \) we expect this distribution to be peaked at a single value \( n \) which should also describe the number of electrons calculated from the Green’s function.

### 3.4 Preparing control files for LMTART

In this section we briefly describe the control files that need to be prepared for the implementation of the LMTART program. For LDA calculations only, there are two essential input files – the `ini` and the `str` files. The `ini` file contains details about the various approximation schemes such as ‘full-potential plane wave’ or ‘atomic sphere approximation’ as well as those about the exchange-correlation functional. It also contains information about the different atoms in the unit cell. The structure file or the `str` file has information about the crystal structure. In addition, for LDA+U or LDA+DMFT calculations, we need the `hub` file that contains the interaction parameters and the DMFT options.

Each control file is divided into different sections, separated by `(SECTION =)`, each containing a certain set of data. There are certain common features in each control file:

- The first line `(FILE = * * *FILE, INPUT = MODERN)` where ‘***’ stands for the type of control file – i.e., INI or STR or HUB, must be present in each control file. This line is required by the main program in the LMTART program to call the corresponding subroutine to read the control file.
• The first section in the control files contains the title of the material – this section is optional.

• A comment that usually contains the description of the parameter and some options for it can be placed on each line after a ‘!’ mark.

• Many parameters have default values. If a default value exists, this parameter can be omitted from the file.

3.4.1 INI file

• (SECTION = CTRL)

The parameters in this section control the choice of the scheme we would like to use in the implementation of LMTART.

  – FullPot This parameters allows us to choose the approximation for the potential seen by an atom in the solid. The choices are a Full Potential plane wave approximation (PLW) or the atomic sphere approximation (ASA) where the potential in the interstitial region is approximated by a constant. We can also describe the Hamiltonian in a tight-binding representation (FTB) and solve the eigenvalue problem.

• (SECTION = ITER)

Details about the iterational loop are contained in this section.

  – (SECTION = MAIN)

This section contains information about the number of atoms and spins in the unit cell.

  – Nsort: The number of non-equivalent atoms in the unit cell. This parameter becomes important in anti-ferromagnetic calculations as explained in section 3.4.3.

  – Is: This labels the different kinds of basis atoms. The order must correspond to that in (SECTION = BASS) in the str file. An example is provided in Table 3.3.
- **Par0:** The lattice parameter in atomic units needs to be provided, which is used, along with the crystal structure to calculate the energy in Rydberg units.

- **(SECTION = SORT)**
  
  This section contains information about the atoms. Once the name and atomic number of each element is entered, the LMTART routine searches for relevant information for the atom in the directory /atomdat/.

  **Note:** The set of parameters above, in (SECTION = SORT) must be entered for each non-equivalent atom (the number of non-equivalent atoms being indicated in the parameter Nsort).

### 3.4.2 STR file

The structure file contains all the information about the crystal structure. To prepare the file, we need to know about the unit cell of the lattice – the parameters describing the geometry of the cell, the primitive translation vectors and the basis atoms. This information can be obtained from experimental values.

Table 3.2 describes a sample str file for NiO which has a FCC structure with two basis atoms, i.e. Ni and O. The unit cell for this is schematically shown in Fig. 3.2 In the notation we use, the primitive vectors of the unit cell are expressed as $A\hat{x} + B\hat{y} + C\hat{z}$ as shown in Fig. 3.2.

- **(SECTION = CTRS)**

  - **Natom:** The total number of basis atoms in the unit cell. For the above example of NiO it is 2. *(no default)*

  - **B to A:** The ratio of the length along the $y$-direction to that along the $x$-direction. In the above example, we have a cubic cell, hence this ratio is 1. *(default is 1.0)*
FILE=INFILE, INPUT=MODERN

SECTION=HEAD
  title = NiO

SECTION=CTRL
  Lmto = Bare
  FulPot = ASA
  LDA = Vosko

SECTION=EXCH
  LDA = Vosko

SECTION=ITER
  Niter1 = 40
  Lbroy = 1
  Admix1 = 0.05000

SECTION=MAIN
  Natom = 2
  Nsort = 2
  Nspin = 1
  Norbs = 1
  Par0 = 7.92600
  Is(:) = 1 2

SECTION=SORT
  Name = Ni
  Znuc = 28.0000
  Smts = 2.17900
  Split = 0.50000

SECTION=SORT
  Name = O
  Znuc = 8.0000
  Smts = 1.78300
  Split = 0.50000
  Ndiv(:) = 6 6 6

Table 3.1: Input file nio.ini
(FILE=STRFILE, INPUT=MODERN)

**************************************************************
(SECTION=HEDS)
Structure Title: Slab1=NiO
(SECTION=CTRS)
! Control Structure:
Natom = 2  ! # of atoms
BtoA = 1.00000  ! b over a ratio
CtoA = 1.00000  ! c over a ratio
(SECTION=TRAN)
! Primitive Translations:
1/2, 1/2, 0.0  ! Ax,Ay,Az
1/2, 0.0, 1/2  ! Bx,By,Bz
0.0, 1/2, 1/2  ! Cx,Cy,Cz
(SECTION=BASS)
! Basis Atoms:
0.0, 0.0, 0.0  ! Ni@1
1/2, 1/2, 1/2  ! O@2

Table 3.2: Input file nio.str

- **C to A**: The ratio of the length along the z-direction to that along the x-direction. In the above example, we have a cubic cell, hence this ratio is

  1. *(default is 1.0)*

- (SECTION = TRAN) *(no default)*

  This section contains the primitive translation vectors of the lattice. For example, for the FCC lattice the set of vectors can be written as \(\left(\frac{1}{2}, \frac{1}{2}, 0\right), \left(0, \frac{1}{2}, \frac{1}{2}\right), \left(\frac{1}{2}, 0, \frac{1}{2}\right)\) and for the BCC lattice the set is \(\left(\frac{1}{2}, \frac{1}{2}, -\frac{1}{2}\right), \left(-\frac{1}{2}, \frac{1}{2}, \frac{1}{2}\right), \left(\frac{1}{2}, -\frac{1}{2}, \frac{1}{2}\right)\).

- (SECTION = BASS) *(no default)*

  This section contains the positions of the basis atoms in the unit cell, that, along with the primitive translations from the above section will generate the full lattice.

  **Note**: The order in which the basis atoms are written must correspond to the order in which the atoms are written in (SECTION = SORT) in the ini file.
3.4.3 Modifications for anti-ferromagnetic solution

If we need to compute the anti-ferromagnetic solution for a given material, we need to make appropriate changes in the ini as well as str files. To have an anti-ferromagnetic solution, we pick alternate magnetic atoms in the lattice to have up and down spins respectively. The unit cell thus has to be doubled, and contains two non-identical magnetic atoms due to different spins, even though the atomic numbers of the two atoms remain the same.

The main modifications in the ini file are in (SECTION = MAIN) as shown in a part of the ini file for anti-ferromagnetic NiO in Table 3.3. Note, especially, the parameter Is($) which sorts the different atoms. Even though Atom 1 and Atom 2 are both nickel, we treat them differently due to their different spins. (SECTION = SORT) contains information about the different spins in the parameter Split which has different signs for potential for up and down spins.

To determine the crystal structure we treat the lattice as a new structure with different primitive translations and basis atoms and re-construct the str file. For the same example of NiO as above, we begin with Fig. 3.3 to generate the primitive

Translation vectors:
- $\overrightarrow{T_1} = \frac{1}{2} x + \frac{1}{2} y + 0z$
- $\overrightarrow{T_2} = \frac{1}{2} x + 0y + \frac{1}{2} z$
- $\overrightarrow{T_3} = 0x + \frac{1}{2} y + \frac{1}{2} z$
translations and basis atoms. The \textit{str} file for this case is shown in Table 3.4.

```plaintext
(SECTION=MAIN) ! MAIN ATOMIC DATA:
Natom = 4 ! # of atoms in the unit cell
Nsort = 3 ! # of sorts in the unit cell
Nspin = 2 ! # of spins
Norbs = 1 ! 1-without/2 -with spin orbit coupling
Par0 = 7.02600 ! lattice parameter in a.u.
Is(:, :) = 1 2 3 3 ! atom-to-sort pointer array
(SECTION=SORT) ! SORT DATA:
Name = Ni1 ! atom label
Znuc = 28.0000 ! nuclear charge
Smts = 2.17900 ! non-overlapping MT-sphere
Split = 0.50000 ! initial splitting
Name = Ni2 ! atom label
Znuc = 28.0000 ! nuclear charge
Smts = 2.17900 ! non-overlapping MT-sphere
Split = -0.50000 ! initial splitting
Name = O ! atom label
Znuc = 8.0000 ! nuclear charge
Smts = 1.78300 ! non-overlapping MT-sphere
Split = 0.50000 ! initial splitting
Ndiv(:, :) = 6 6 6 ! Tetrahedron mesh
```

Table 3.3: Input file nio.ini for anti-ferromagnetic NiO

More working examples of the \textit{str} file are given in chapter 4.3.1 YbRh$_2$Si$_2$ and in section 5.2 for Am. Also, 3.4.3 contains the \textit{str} file for the anti-ferromagnetic case for YbRh$_2$Si$_2$, where the unit cell has to be doubled to contain two magnetic Yb atoms.

### 3.4.4 \textit{HUB} file

The main data in this file that need to be entered by the user are the choice of scheme to treat the correlated orbitals (LDA+U or LDA+DMFT) and the values of the interaction parameters. After the first iteration, additional data regarding occupancy matrices, self-energies and potential terms are written into this file.
Figure 3.3: Unit cell for anti-ferromagnetic case

Table 3.4: Input structure file nio.str for the anti-ferromagnetic case
A sample hub file for the anti-ferromagnetic case in NiO is shown in Table 3.5. All the energy values are in Rybergs. The various sections are explained below:

- **(SECTION = CTRL)**
  - **Scheme:** The details about the scheme to be used to treat the correlated orbitals, i.e. LDA+U or LDA+DMFT. A list of all the possible schemes that can be used is present in the file hub_readhub.f and hub_hubpot.f.
  - **Ncrl:** The number of states which appear in the correlated term. Descriptions and options of the other parameters in this section appear as comments in the hub file in Table 3.5.

- **(SECTION = DMFT)**
  This section needs to be included only if the option for Scheme in the above section is chosen as LDA+DMFT. The details of the number of Matsubara frequencies to be used and the bandwidth for the grid are contained here.
  - **solver:** The impurity solver used in the DMFT loop. A list of the solvers (e.g. QMC, Hubbard-I, Gutzwiller) appears in the file hub_readhub.f.
  - **Kstates:** This is a pointer to the states in the correlated orbital that are treated similarly. For example, in NiO, there are 10 correlated states from the 3d orbital – hence we need 10 labels. The $t_2g$ spin-up states in are treated similarly, thus they all carry the label 1. Similarly the $e_g$ spin-up states all carry the label 2, and so on.

- **(SECTION = CORR)**
  This section has the core information about the interaction terms.
  - **Cstate:** This points to the atom that has correlations, its position in (SECTION = BASS) in the str file and the correlated orbital.
  - **F0, F2, F4, F6:** These are parameters that appear in the Slater integrals as explained in equations 29-32. For d-electrons, three constants F0, F2 and F4 are needed and for f-electrons, there are four constants F0, F2, F4 and
These can be linked to $U$ and $J$ via $U = F_0$ and $J = (F_2 + F_4)/14$ for d-electrons with $F_2/F_4 = 0.625$. For f-electrons, $U = F_0$ and $J = (286F_2 + 194F_4 + 250F_6)/6435$.

**Note:** The set of parameters above, in (SECTION = CORR) must be entered for each correlated state (the number of correlated states being defined by the parameter Ncrl).

- (SECTION = DHUB)

From this section onwards, data about occupancy matrices, self-energies etc are written into the hub file after the first iteration. These quantities are self-consistently calculated at each iteration and are rewritten. A sample occupancy matrix for the 3d state of Ni in NiO is shown in Table 3.6.

We have now established the theoretical groundwork for the LDA+DMFT method to realistically treat strongly correlated materials. We shall turn to two applications of this method: in Chapter 4, we study the heavy fermion compound YbRh$_2$Si$_2$ and in Chapter 5, we discuss the Mott transition issue in Americium.
**FILE=HUBFILE, INPUT=MODERN**

**SECTION=CTRL**

- **Scheme=LDA+U1.1**
  - LDA+U1.# / LDA+C / LDA+DMFT
- **Yharm=cubic**
  - Cubic/Spherical harmonics (output)
- **Iharm=cubic**
  - Cubic/Spherical harmonics (input)
- **Rorbs=Both**
  - One/Both orbits to read
- **Rspin=Both**
  - One/Both spins to read
- **Ncr1 = 2**
  - # of correlated states

**SECTION=DMFT**

- **Nmsb = 50**
  - # of Matsubara frequencies
- **Obnd = 0.5000000**
  - Effective bandwidth for the grid
- **Nmsl = 2048**
  - # of linear Matsubara frequencies
- **Nmsb = 1**
  - # of Matsubara frequencies
- **Ndit = 20**
  - # of DMFT iterations
- **Efermi= 0.7278130**
  - Fermi energy (Ry)
- **Solver =Hub1**
  - Impurity solver choice
- **kStates(:)=1 1 1 2 2 3 3 3 4 4**

**SECTION=CORR**

- **Cstate =Ni1@1::3d**
  - Correlated state pointer
  - F0 = 0.5880000
  - Slater integrals
  - F2 = 0.6012300
  - Slater integrals
  - F4 = 0.3787700
  - Slater integrals
  - F6 = 0.000000
  - Slater integrals
- **Cstate =Ni2@2::3d**
  - Correlated state pointer
  - F0 = 0.5880000
  - Slater integrals
  - F2 = 0.6012300
  - Slater integrals
  - F4 = 0.3787700
  - Slater integrals
  - F6 = 0.000000
  - Slater integrals

**SECTION=DHUB**

- **cState=Ni1@1:3d**
  - spin up/dn-up/dn data are:

---

Table 3.5: Input hub file nio.hub
<table>
<thead>
<tr>
<th>cState=Ni1@1:3d</th>
<th>! PARTIAL OCCUPANCIES:</th>
</tr>
</thead>
<tbody>
<tr>
<td>yz</td>
<td>zx</td>
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Table 3.6: Partial occupancies for Ni in nio.hub
Chapter 4

Electronic structure and properties of $f$-band materials: $\text{YbRh}_2\text{Si}_2$

4.1 Introduction

In this chapter, we present our calculations of the band structure and electronic properties of $\text{YbRh}_2\text{Si}_2$, a heavy-fermion compound. This material is experimentally well studied and has an anti-ferromagnetic phase transition at low temperature ($T < 10 \text{ K}$) and ambient pressure. Susceptibility measurements on the material show a strong magnetic anisotropy [53]. Using DFT methods, we would like to study the various magnetic phases of this compound and compute its magnetic properties such as magnetic anisotropy energy.

We start our calculations with the standard LDA method. However, since $\text{YbRh}_2\text{Si}_2$ is a material with a strongly localized $f$-band, the LDA method is inadequate to explain the magnetic ground state of the material, as we expect. We then employ the LDA+$U$ method which includes electron-electron interaction terms in the energy functional. However, we find that in this case, LDA+$U$ does not predict the correct magnetic ground state. The problem is traced to the double-counting term in the LDA+$U$ functional, which includes the part of the total energy that have already been taken into account in the LDA Hamiltonian. This term is to be subtracted from the total energy, so that only the correction ot the LDA mean-field solution is left. However, this procedure is not unique and there are various ideas previously implemented such as those by Mazin [54], Anisimov [15], Czyzyk and Sawatzky [55].

Our approach to the double-counting issue was to use an empirical method to determine which choice of parameters correctly predict the ground state. Once we found
the suitable ground state solution, we proceeded to calculate properties such as magnetization and magnetic anisotropy energy. With the application of pressure, it should be possible to drive the non-magnetic phase of YbRh₂Si₂ into a magnetically ordered ground state. In our calculation, we study the evolution of magnetism in this material as a function of various parameters such as pressure and Coulomb interaction.

In section 4.2 we describe the motivation and the experimentally known properties of this material. Section 4.3 contains the results of our calculation to reproduce the magnetic ground state of the material using various techniques. In section 4.4, we use our calculated band structure to compute the magnetic anisotropy energy of YbRh₂Si₂. Finally, in section 4.5, we summarize the various techniques we used in the study of the compound, and draw our conclusions.

4.2 Properties of YbRh₂Si₂

Experimentally YbRh₂Si₂ is known to have an anti-ferromagnetic quantum critical point at ambient pressure [53]. It is an undoped and atomically well-ordered material. X-Ray diffraction studies showed that YbRh₂Si₂ has a body centered tetragonal phase as shown in Fig. 4.1 with the lattice parameters \( a = 4.007 \text{ Å} \) and \( c = 9.858 \text{ Å} \) [56]. The valence shell of Yb is in the \( 4f^{13} \) configuration.

At high temperatures, the magnetic susceptibility shows a Curie-Weiss like behavior with magnetic moment \( \mu_{\text{eff}} = 4.5\mu_B \) which is almost equal to that of the free Yb\(^{3+}\) ion. The magnetic susceptibility along the plane of the tetragonal structure is measured to be much larger than that along the \( c\)–axis. This anisotropy in the magnetic response indicates that the Yb\(^{3+}\) moments form a 2-dimensional lattice perpendicular to the \( c\)–axis. At ambient pressure and low temperature \( T_N = 65mK \), YbRh₂Si₂ develops a long range anti-ferromagnetic order. One of the goals of our band structure computation is to obtain the different magnetic phases of YbRh₂Si₂, and to calculate it magnetic properties. Further, we would like to compute properties of this material that might arise as a result of the strong magnetic anisotropy.

YbRh₂Si₂ shows many deviations from Fermi liquid behavior at low temperatures
such as linear temperature dependence of electrical resistivity and a logarithmic dependence of specific heat \( C/T \propto -\ln T \). These unusual properties arise due to the interactions between the localized \( f \)-electrons in \( \text{Yb}^{3+} \) and the conduction band formed by the \( s,p \) and \( d \) electrons. We would like to use the band structure calculation to understand how localized bands and strong correlations in the \( 4f \)-electrons affect the properties of the material.

### 4.3 Band structure calculation of \( \text{YbRh}_2\text{Si}_2 \)

The computation of the band structure is done using the \( \text{LmtART} \) code [57] based on the linear muffin-tin orbital density functional method as described in the previous chapter. We start with the LDA method. Since \( \text{YbRh}_2\text{Si}_2 \) exhibits strong electron-electron correlations we expect an LDA calculation to fail to reproduce the magnetic ground state of the material. We recalculate the structure using the LDA+U technique. We will show that LDA+U also does not work well, due to problems with the double counting term in the functional.
4.3.1 LDA calculation

We first describe some technical details of the numerical simulation:

The program and the database associated with it provides the Hamiltonian and the overlap matrix of the valence orbitals as well as the necessary space group information and Brillouin zone data. The unit cell of YbRh$_2$Si$_2$ has five basis atoms at the following positions: Yb(0,0,0), Rh1(0,1/2,1/4), Rh2(1/2,0,1/4), Si1(0,0,3/8), Si2(0,0,5/8). This information is contained in the input structure file *ybrh2si2.str* shown in Table 4.1. Details about the atomic numbers of elements in the solid, the lattice constant, the schemes used for self-consistency (ASA) and exchange-correlation, and information about the iterations in the self-consistent loop, in are contained the input file *ybrh2si2.ini* in Table 4.2. Finally, for the LDA+U calculation, the details of the interactions U and J, and the correlations are contained in the file *ybrh2si2.hub* in Table 4.3.

---

Table 4.1: Input file ybrh2si2.str
FILE=INIFILE, INPUT=MODERN

SECTION=HEAD
   title = YbRh2Si2

SECTION=CTRL
   Lmto = Bare
   FulPot = ASA

SECTION=EXCH
   LDA = Vosko

SECTION=ITER
   Niter1 = 200
   Lbroy = -1
   Admix1 = 0.05000

SECTION=MAIN
   Natom = 5
   Nsort = 3
   Nspin = 2
   Norbs = 2
   Par0 = 7.57500
   Is(:) = 1 2 3 3

SECTION=SORT
   Name = Yb
   Znuc = 70.0000
   Smts = 3.66200
   Split = 0.0000

SECTION=SORT
   Name = Rh
   Znuc = 45.0000
   Smts = 2.26700

SECTION=SORT
   Name = Si
   Znuc = 14.0000
   Smts = 2.17800

SECTION=FFTS
   Ndiv(:) = 6 6 6

Table 4.2: Input file ybrh2si2.ini
\[ \{ \text{FILE=HUBFILE, INPUT=MODERN} \} \]

***************************************************************************

\{ \text{SECTION=CTRL} \}

\text{Scheme=LDA+U1.1} \quad \text{! LDA+U1.1 / LDA+C / LDA+CU1.1 / LDA+G1}
\text{Yharm =spherical} \quad \text{! Cubic/Spherical harmonics (output)}
\text{Iharm =spherical} \quad \text{! Cubic/Spherical harmonics (input)}
\text{Rorbs =Both} \quad \text{! One/Both orbits to read}
\text{Rspin =Both} \quad \text{! One/Both spins to read}
\text{Format=complex} \quad \text{! Real/Complex input/output}

\{ \text{SECTION=CORR} \}

\text{Ncrl = 1} \quad \text{! # of correlated states}
\text{Cstate =Yb@1::4f} \quad \text{! Correlated state pointer}
\text{OutSys = local} \quad \text{! global/local coordinate system}
\text{OutAxis = -1,1,0} \quad \text{! rotational axis}
\text{OutAngle= 0*pi} \quad \text{! rotational angle}
\text{OutInv =no} \quad \text{! apply inversion after rotation}
\text{InpSys = local} \quad \text{! global/local coordinate system}
\text{InpAxis = -1,1,0} \quad \text{! rotational axis}
\text{InpAngle= 0*pi} \quad \text{! rotational angle}
\text{InpInv =no} \quad \text{! apply inversion after rotation}
\text{F0 = 0.4411765} \quad \text{! Slater integrals}
\text{F2 = 0.000000} \quad \text{! Slater integrals}
\text{F4 = 0.000000} \quad \text{! Slater integrals}
\text{F6 = 0.000000} \quad \text{! Slater integrals}

\{ \text{SECTION=DHUB} \}

\text{cState=Yb@01::4f} \quad \text{! spin up/dn-up/dn up data are :}

\begin{table}
\centering
\caption{Input file ybrh2si2.hub}
\end{table}
Both the full potential plane-wave (PLW) and the atomic sphere approximation (ASA) schemes have been used. The results for the energy bands (using the PLW scheme) show a narrow localized set of bands below the Fermi level $E_F$ which is near 13.7eV (Fig. 4.2, left). On investigating the character of these localized bands, which are near energies of $\sim 13eV$, it is seen that these correspond to the localized 4$f$ bands of Yb, as expected. Self-consistency for the energy was obtained in $\sim 40$ iterations. We find that the there are 14 fully occupied 4$f$-bands below $E_F$ at the $\Gamma$ point. The LDA calculation predicts that all the 4$f$-bands are filled, that is, the occupancy of the 4$f$-bands of Yb in YbRh$_2$Si$_2$ is 14.

On performing the same calculation using the ASA method to compute the potential, we find that the main differences are that the overall energies and the Fermi level have shifted downwards with the Fermi energy $E_F \sim 13eV$, and that the position of the localized 4$f$ bands of Yb are pushed further below to 12eV (Fig. 4.2, right). Unless otherwise mentioned, all further calculations are in the ASA method. A large contribution to the bands crossing the Fermi level comes from the 5$d$-bands of Yb and the 4$d$-bands of Rh (Fig. 4.3).

According to the experimental evidence, Yb$^{3+}$ in YbRh$_2$Si$_2$ is in a 4$f^{13}$ configuration. Thus we need to improve upon the simple LDA method in order to check if one of the Yb$^{3+}$ 4$f$-bands lies above the Fermi energy. Since the electrons on the Yb$^{3+}$ ion are strongly correlated we will use the LDA+U method next to obtain the electronic structure.

### 4.3.2 LDA+U calculation

We used the LDA calculation to estimate the interaction energy required to push one of the 4$f$-bands from below to above the Fermi level. From the LDA calculation in the ASA case, we located the 4$f$ sub-band (amongst all the 4$f$-bands) that has the highest energy at the $\Gamma$-point. It is most likely that it is this band that would be unfilled upon the inclusion of $U$. From Fig. 4.4, we can see that this band is the 4$f : \{z(x^2 - y^2)\}$ sub-band. Thus to obtain only 13 filled 4$f$-bands we use the correct initial partial occupancies in the DHUB matrix. We input a partial occupancy of 0
Figure 4.2: Energy bands of YbRh$_2$Si$_2$ – Full Potential(left) and Atomic Sphere Approximation(right). The colored regions in are the fat bands for Yb:4f.

Figure 4.3: Energy bands of YbRh$_2$Si$_2$. The colored regions in are the fat bands for Yb:5d (left) and Rh(4d) and Si:3p(right).
for the $4f : \{z(x^2 - y^2)\}$–spin-up (an arbitrary choice, as spin-up and spin-dn bands displayed the same behavior) and a partial occupancy of 1 for the other sub-bands.

We find that even on increasing the value of $U$, we get all the 14 $4f$-bands of Yb to be filled. The results for $U = 2, 4, 6$ and 8 eV and are shown in Fig. 4.5 Self-consistency for the energy was obtained in 100 iterations.

Under the simple LDA calculation, all the 14 $4f$-bands of Yb are filled. We want to locate the magnetic solution with one of the $4f$-bands to be pushed above the Fermi level. Fig. 4.6 shows the behavior of the position of the center of the $f$-band w.r.t. $U$. Thus in order to push one of the bands towards and above the Fermi level, we propose that the value of $U$ used in the double counting term is different from that in
Fat bands of Yb 4f-up and 4f-dn in YbRh$_2$Si$_2$
LDA+U (=2eV)

Fat bands of Yb 4f-up and 4f-dn in YbRh$_2$Si$_2$
LDA+U (=4eV)

Fat bands of Yb 4f-up and 4f-dn in YbRh$_2$Si$_2$
LDA+U (=6eV)

Fat bands of Yb 4f-up and 4f-dn in YbRh$_2$Si$_2$
LDA+U (=8eV)

Figure 4.5: Energy bands of Yb:LDA+U, U=2,4,6,8eV
the LDA+U energy functional for the model.

4.3.3 Effect of the double counting term:

Our prescription for obtaining the correct band structure for this material is that we treat the interaction $U$ that is used in the model and the interaction $\tilde{U}$ used in the double counting term as two separate parameters. The former controls the position of the band below the Fermi level while the latter controls the splitting in energy between the occupied and the unoccupied bands, on which the magnetic moment depends. To test this hypothesis, we run the LDA+U calculation using different values of $U$ and $\tilde{U}$.

In Fig. 4.7 we plot the position of the 4$f$-level as a function of $\tilde{U}$ for the interaction $U = 4.0eV$ and $U = 6.0eV$. Initially we keep $\tilde{U} = U$. As we decrease $\tilde{U}$ (to $\tilde{U} = 5.5eV$ for $U = 6eV$ and to $\tilde{U} = 3.4eV$ for $U = 4eV$) we find that the system has a magnetic
solution. These results are shown in Figs. 4.8 and 4.9 where we see that a part of the 4f-band of Yb is unoccupied and the Yb-ion shows a non-zero magnetic moment.

However, when we calculate the total magnetic moment, we find that it is less than the experimental value of $\sim 4\mu_B$. For this solution, our initial condition had, for the 4f : $\{z(x^2 - y^2)\}$ sub-band, an occupation number of 0 (for the spin-up electron), and for the other 4f-bands, an occupation number of 1. This led to a stable converged solution with a total occupation of 13.3 for the f-bands. However, magnetic moment (in units of Bohr magnetons $\mu_B$) for this state was lower than the experimental one. For $U = 6eV, \bar{U} = 5.5eV$:

- Orbital moment $M_z = 0.4$
- Spin moment = 1.16
- Total moment = 1.56.

The difference is due to the contribution from the orbital angular moment. In the above state, the contribution came from the orbitals $l_z = +2, -2$. In order to obtain the correct angular moment, the initial state has to have the $l_z = +3$ orbital empty and the other orbitals filled. With this initial condition, we get a band structure as shown in Fig. 4.9. The magnetization for $U = 6eV, \bar{U} = 5.5eV$ is:

- Orbital moment $M_z = 2.97$
- Spin moment = 1.14
- Total moment = 4.11

Thus, using the correct value of the interaction $\bar{U}$ in the double-counting term, and the correct initial conditions in the occupancy matrix, we obtain a band structure for YbRh$_2$Si$_2$ that has the correct magnetic solution.

### 4.3.4 Comparison with other methods involving the double-counting term

In order to describe the physics of correlated metals, which fall in between materials which have fully localized shells, and those which have uniform electron occupancies, Mazin et.al [54] proposed an alternate form of the double counting term in the LDA+U
Figure 4.7: Position of center of 4f-band as a function of the double-counting term $\bar{U}$ for $U=4\text{eV}$ (top) and $U=6\text{eV}$ (bottom)
Figure 4.8: Energy bands(left) and DOS(right) of Yb:4f, LDA+U, U=4eV.

Figure 4.9: Energy bands(left) and DOS(right) of Yb:4f, LDA+U, U=6eV.
functional (c.f. Eqns. 3.18 and 3.19):

\[
E_{LD\Delta+U}^{DFT} = -\frac{(U - J)}{2} \sum_{\sigma} \left[ Tr(\delta\rho^\sigma \cdot \rho^\sigma) - (2l + 1) \left[ \alpha_\sigma n_\sigma^2 + (1 - \alpha_\sigma) n_\sigma \right] \right]
\]

(4.1)

\[
0 \leq \alpha_\sigma = \frac{Tr(\delta\rho^\sigma \cdot \delta\rho^\sigma)}{(2l + 1) n_\sigma (1 - n_\sigma)} \leq 1
\]

(4.2)

The correction to the potential is

\[
\lambda_{LD\Delta+U}^{DFT}(ab, \sigma) = -(U - J) \left[ \rho^\sigma_{ab} - \alpha_\sigma n_\sigma - \frac{1 - \alpha_\sigma}{2} \right]
\]

(4.3)

Here, the orbital occupation matrix, \(\rho^\sigma_{ab} = -1/\pi \int E dE G_{ab}(E)\) where \(G_{ab}\) is the one-electron Green’s function, \(n_\sigma = Tr\rho^\sigma/(2l + 1)\) and \(\delta\rho^\sigma_{ab} = \rho^\sigma_{ab} - n_\sigma.\) \(\alpha_\sigma\) is a system dependent constant between 0 and 1 that is calculated using the self-consistent occupation matrix at each iteration as in Eqn. 4.2 According to this scheme, our LDA+U calculation has a double counting term in eqn.3.19 as \((\rho^\sigma_{ab} - \frac{1}{2} \delta_{ab})\) that is equivalent to keeping the value of \(\alpha = 0.\)

We tested the above functional for our calculation on YbRh\(_2\)Si\(_2\). We replace the original double-counting term (eqn.3.19) in the file hubpot.f by the one above. We find that this double-counting term does not produce a magnetic ground state for YbRh\(_2\)Si\(_2\). Fig. 4.10 shows how the 4\(f\)-bands of Yb in YbRh\(_2\)Si\(_2\) behave as a function of \(U\) using the scheme described above. From the self-consistent procedure we get a value of \(\alpha = 0.22.\) Also included in the figure are the location of the bands using the original double-counting with \(U = \bar{U}\), as well as the result from our empirical method of treating \(U\) and \(\bar{U}\) as different parameters. We see that using the functional described by Mazin, the center of the bands move in the correct direction, however we still do not obtain the correct magnetic ground state. For that we must follow the method we described in section 4.3.3 where we treat the interaction \(\bar{U}\) as a different parameter from the interaction \(U\) in the model.

In yet another functional for the total energy proposed by Czyzyk and Sawatzky [55], the double counting term to be subtracted from the potential is given by \((\rho^\sigma_{ab} - n^\sigma \delta_{ab})\) which corresponds to the limit of uniform occupancy in LDA, i.e. \(\rho^\sigma_{ab}(LDA) = \delta_{ab} n^\sigma.\) For correlated materials, the limit of uniform occupancy is not correct, hence this method of treating the double counting term produces the incorrect ground state for YbRh\(_2\)Si\(_2\).
4.3.5 Effects of pressure

If we apply pressure, we expect that the f-level moves closer to the Fermi level. If sufficient pressure is applied, we expect some of the f-bands to cross the Fermi level so the material has a magnetic solution. To incorporate the effects of pressure, we decrease the volume compression factor $V/V_0$ from its original value of 1. We find that there is an overall increase in all the energies observed, including the Fermi level. When pressure is increased by $\sim 10\%$, the center of the 4f-band moves up by $\sim 12\%$ but the Fermi level increases by about as much. Hence we do not observe a magnetic solution.
with the application of pressure alone.

Once we have obtained the correct magnetic solution by using the appropriate value of the double-counting term and the interaction term, we find that on applying pressure, the gap between the occupied and the unoccupied states decreases, leading to a decrease in the magnetization. As pressure is applied, the volume of the muffin-tin spheres decreases which causes a decrease in the effective interaction. The Hubbard bands come closer and the magnetic moment decreases. Fig. 4.11 shows this trend. The band structures for the magnetic solution on applying pressure are shown in Fig. 4.12.

![Magnetic moment vs relative volume](image)

Figure 4.11: Dependence of magnetic moment on relative volume

### 4.3.6 Anti-ferromagnetic solution

Experimentally YbRh$_2$Si$_2$ is in an anti-ferromagnetic state. To obtain the anti-ferromagnetic solution and the band structure, we have to first incorporate the correct crystal structure in the input files. In the ferromagnetic case, each unit cell has one Yb atom, which is located at the origin. To have an anti-ferromagnetic solution, we have to double the unit cell so that it contains two Yb atoms. Adjacent Yb atoms have opposite spins. The following table shows the input structure file that shows the basis atoms and the
Figure 4.12: Application of pressure on the magnetic solution
positions of the atoms. (Compare with structure file in Table 4.1)

(FILE=STRFILE)
**************************************
(SECTION=HEDS)
Slabl =YbRh2Si2 ! STRUCTURE TITLE:
(SECTION=CTRS)
Natom =10 ! # of atoms
BtoA = 1.00000 ! b over a ratio
CtoA = 2.46019 ! c over a ratio
(SECTION=TRAN)
1 , 1 , 0 ! Ax,Ay,Az
1 , -1 , 0 ! Bx,By,Bz
1/2 , 1/2 ,1/2 ! Cx,Cy,Cz
(SECTION=BASS)
0.0 , 0.0 , 0.0 ! Yb1
1.0 , 0.0 , 1.0 ! Yb2
0.0 , 1/2 , 1/4 ! Rh1
1/2 , 0.0 , 1/4 ! Rh2
1.0 , 1/2 , 1/4 ! Rh3
3/2 , 0.0 , 1/4 ! Rh4
0.0 , 0.0 , 3/8 ! Si1
0.0 , 0.0 , 5/8 ! Si2
1.0 , 0.0 , 3/8 ! Si3
1.0 , 0.0 , 5/8 ! Si4

Table 4.4: Input file ybrh2si2.str

The interaction parameters we used were $U = 6\text{eV}$, $\bar{U} = 5.5\text{eV}$. The magnetic moment (in units of Bohr magnetons) in this case is as follows:

- Orbital magnetic moment for 1st Yb atom (spin-up): 2.959937
- Orbital magnetic moment for 2nd Yb atom (spin-dn): 2.667827
- Spin magnetic moment for 1st Yb atom (spin-up): 1.112074
- Spin magnetic moment for 2nd Yb atom (spin-dn): 0.923587

Fig. 4.13 shows the energy bands of YbRh$_2$Si$_2$ in the anti-ferromagnetic calculation.

4.4 Magnetic Anisotropy Energy

YbRh$_2$Si$_2$ shows a highly anisotropic magnetic response along different crystallographic directions. The experimentally measured magnetic susceptibility (Fig. 4.14 from [59])
Figure 4.13: Energy bands of Yb in the anti-ferromagnetic case. Top: Bands of Yb-1 atom. Bottom: Bands of Yb-2 atom
along the plane of the tetragonal structure is $\approx 20$ times larger than that measured along the perpendicular $c$ axis at 2 $K$ and up to 100 times larger at 0.1$K$ according to Trovarelli et. al. [53], [59]. The magnetic anisotropy energy (MAE) is defined as the difference of the total energies with the orientation of the magnetization pointing in different crystalline directions. Since it depends on the total energy, the MAE is a ground state property and hence can be calculated using density functional theory.

To calculate the MAE, we calculate the total energy with the magnetic axis pointing in two different directions – [100] corresponding to the $z$-axis in the crystal and [110], corresponding to the X-Y plane. In practice, we implement this as explained below.

We first start with the converged paramagnetic solution that we obtain with LDA+U. To obtain the solution with the magnetization along the $c$-axis of the crystal, which is also our $z$-direction, we set the parameter $AxMag = 0, 0, 1$ in the input file $ybrh2si2.ini$. We also use the correct initial values in the occupancy matrix with the magnetic axis in the [100] crystallographic direction. We calculate the total energy with these conditions as $E_c$. The magnetization obtained is in the $z$-direction.

To calculate the total energy in the [110] crystallographic direction, that is, along the basal $a$-$b$ plane of the crystal, we set $AxMag = 1, 1, 0$ in the input file $ybrh2si2.ini$. We also have to rotate the initial occupancy matrix, so that the orbital and the spin magnetic moments both point along the x-y plane. We calculate the total energy $E_a$ in this configuration. The difference of the energies in the above two configurations is the MAE.

We first compute the magnetic anisotropy energy using the ferromagnetic solution. We use 12 k-points per dimension in the Brillouin zone.

For the magnetization along the $c$-axis, total energy $E_c = -48409.504501$ Ry.

Here, magnetic moment $M_z = 1.08\mu_B$ and $M_x, M_y << M_z$.

For the magnetization along the $a$-$b$ plane, total energy $E_a = -48409.504513$ Ry.

Magnetic moment $M_x = M_y = 0.48202 \mu_B$ and $M_z << M_x, M_y$.

The magnetic anisotropy energy per Yb atom is

$$MAE = E_a - E_c = -1.32 \times 10^{-5} Ry \simeq -179.5 \mu eV$$
Thus $E_a < E_c$, that is, the configuration where the spins point along the x-y plane is more stable, as confirmed by experiment.

Experimentally we know that the magnetic susceptibility $\chi_a \simeq 20\chi_c$ at a field of 5T and temperature of 2K (Fig. 4.14 from [59]). Here $\chi_a$ is the susceptibility along the plane of the tetragonal structure and $\chi_c$ is that perpendicular to the plane, i.e. along the c-axis. To compare the magnetic anisotropy energies in our LDA+U calculation, as compared to the experimental value, we do a calculate the experimental value of the magnetic anisotropy energy given the susceptibility.

We have

$$E = \frac{1}{2} \frac{M^2}{\chi},$$

where $E$ is the energy, $M$ is the magnetization and $\chi$ is the susceptibility. The magnetic anisotropy energy $MAE = E_a - E_c$.

From the plots in Figs. 4.14 and 4.15 obtained from [58] and [59] at a field of 5 T and temperature 2 K we have:

$$M_a = 0.73\mu_B/\text{atom}, \quad M_c = 0.1\mu_B/\text{atom}$$

$$\chi_a = 1.0 \times 10^{-6} \text{m}^3/\text{mol}, \quad \chi_c = 0.1 \times 10^{-6} \text{m}^3/\text{mol}$$

Converting all quantities so that the value are per atom, we have

$$\chi_a = \frac{1.0 \times 10^{-6} \text{m}^3/\text{mol}}{6.023 \times 10^{23} \text{atoms/mol}} = 1.66 \times 10^{-30} \text{m}^3/\text{atom}$$

$$\chi_c = \frac{0.1 \times 10^{-6} \text{m}^3/\text{mol}}{6.023 \times 10^{23} \text{atoms/mol}} = 1.66 \times 10^{-31} \text{m}^3/\text{atom}$$

MAE per atom $= E_a - E_c = \frac{1}{2} \left( \frac{M_a^2}{\chi_a} - \frac{M_c^2}{\chi_c} \right)$

$$= \frac{1}{2} \left( \frac{(0.73\mu_B/\text{atom})^2}{1.66 \times 10^{-30} \text{m}^3/\text{atom}} - \frac{(0.1\mu_B/\text{atom})^2}{1.66 \times 10^{-31} \text{m}^3/\text{atom}} \right)$$

$$= \frac{1}{2} \times \left( 3.166 \times 10^{29}(\mu_B)^2/\text{m}^3 - 6.024 \times 10^{28}(\mu_B)^2/\text{m}^3 \right)$$

$$= -1.282 \times 10^{29}(\mu_B)^2/\text{m}^3$$

(4.7)
Using the value of the Bohr magneton, \(1 \mu_B = 9.274 \times 10^{-24} \text{J/T}\), and the conversion \(1(\mu_B)^2/m^3 = (9.274 \times 10^{24} \text{J/T})^2 \times \mu_0 = 1.079 \times 10^{-52} \text{J}\) we get

\[
\text{MAE per atom} = E_a - E_c = -1.384 \times 10^{-23} \text{J/atom} = -8.65 \times 10^{-5} \text{eV/atom}
\]

(4.8)

When we compute the total energy for the \textbf{anti-ferromagnetic} solution, by doubling the unit cell, we get total energy \(E_c = -96819.024913\) when the magnetization is along the \(c\)-axis of the crystal. With the magnetization pointing along the \(a\)-\(b\) plane, we get the total energy \(E_a = -96819.024942\). Since each unit cell has two Yb atoms, the magnitude of the MAE per Yb atom is \(|E_a - E_c|/2 = (2.9 \times 10^{-5})/2 = 1.45 \times 10^{-5} \text{Ry} \simeq 200 \mu \text{eV}\).

The typical order of magnitude of magnetic anisotropy energies are a few \(\mu \text{eV}\) per atom. For example, MAE per atom at low temperature for Fe is 1.4 \(\mu \text{eV}\), for Ni is 2.7 \(\mu \text{eV}\) and for Co is 60 \(\mu \text{eV}\). In comparison, from our LDA+U calculation, we have, MAE per Yb atom in YbRh\(_2\)Si\(_2\) to be \(\simeq 200 \mu \text{eV}\) and the experimental value for YbRh\(_2\)Si\(_2\) per Yb atom is \(\simeq 86.5 \mu \text{eV}\) with \(E_a < E_c\), that is, the configuration where the magnetization is along the \(a\)-\(b\) plane being more stable.

We also calculate the exchange energy using the total energies per atom for the ferromagnetic and the anti-ferromagnetic case in the two directions.

Parallel to \(a\)-\(b\) plane:

\[
E_{a, \text{Ferro}} - E_{a, \text{Antiferro}} = -48409.504513 - (-48409.512471) = 0.07958 \text{Ry}
\]

Perpendicular to \(a\)-\(b\) plane, i.e. along \(c\)-axis:

\[
E_{c, \text{Ferro}} - E_{c, \text{Antiferro}} = -48409.504501 - (-48409.512456) = 0.07955 \text{Ry}
\]

4.5 Summary and Conclusions

We used LDA and LDA+U methods to compute the band structure of YbRh\(_2\)Si\(_2\). Both methods predict an incorrect non-magnetic structure with all the above 4f-shells occupied. Experimentally YbRh\(_2\)Si\(_2\) is in an anti-ferromagnetic state with a total magnetic
Figure 4.14: $\chi$ vs T (from Trovarelli et.al. [59])

Figure 4.15: from Trovarelli et.al. [58]
moment of $\sim 4.5\mu_B$. In order to obtain the correct magnetic solution, we proposed that the value of $U$ used in the double counting term, $U_{dc}$, is different from that in the energy functional. When $U_{dc}$ is decreased, keeping $U$ fixed, we do observe that the $4f : \{z(x^2 - y^2)\}$ sub-band of Yb gets unoccupied as per the initial condition.

The magnetic anisotropy energy of YbRh$_2$Si$_2$ is calculated using the difference of the total energies with the magnetization pointing in two different directions. Our calculations show that YbRh$_2$Si$_2$ is a highly anisotropic material, as was observed in experiments.
Chapter 5

Electronic structure and properties of $f$-band materials: Americium

5.1 Introduction

In this chapter, we use LDA+U and LDA+DMFT methods described in chapter 3, to study the electronic structure and properties of Americium, another material which has occupation in the $f$-shell.

**Experiment and motivation:**

Americium is a metal that is nearly in the center of the actinide series, which have occupation in the $5f$ shell. The actinides show interesting behavior in terms of crystal structure and atomic volumes as one moves across the series. The lighter actinides until Pu are known to have delocalized $5f$-electrons that participate in bonding. They have open low-symmetry structures. On the other hand, the $5f$-electrons in the heavier actinides are localized and the materials have closed structures.

Fig. 5.1 shows the volume change as one moves across the actinide series. One can see that Americium is at a pivotal position. There is a sudden change in the atomic volume as one moves from Pu to Am across the actinide series. This jump in volume is argued to be a Mott transition with the $5f$-electrons localized in Am, and itinerant in the preceding element, Plutonium. A natural question we can ask at this point is whether an external parameter such as pressure can change the character of the $5f$-electrons in Am.

Am has strong spin-orbit coupling due to which it is in a non-magnetic $J = 0$ ground state. Experimentally, the photoemission spectra of Am has been studied by Nagele [60], which shows a localized $5f$-peak (Fig. 5.2,left).
Figure 5.1: Volumes of actinide elements

Figure 5.2: Left: Photoemission spectra of Am by Naegele [60]. Right: Relative volume of americium as a function of pressure obtained by Lindbaum et al. [62]
Experimental studies have indicated [61], [62] that pressure induces major changes in the nature of the $5f$ electrons of Americium. At normal pressure, the $5f$ electrons of Americium are localized. With the application of pressure, the $f$ electrons acquire an itinerant character. There has been considerable debate in literature about the volume of Americium in its various phases. From the high pressure studies using synchrotron radiation as described by Heathman et al. in Ref. [61] Americium has 4 phases. At ordinary pressures Americium has a double hexagonal close packed (dhcp) structure which under application of pressure gets converted to the Americium II fcc structure at 6 GPa. This transition was reported to occur at higher pressures of 10 GPa by Benedict et al. [63]. With additional pressure Americium II transforms to the Americium III phase accompanied by a volume collapse of 7% [61] which indicates that the $f$ electrons begin to get delocalized (Fig. 5.2, right). The goal of this study is to test the predictions of the LDA+DMFT technique regarding the electronic structure, equilibrium atomic volume and the Mott transition in Am.

**Theoretical studies:**

Previous density functional electronic structure calculations in Americium [64], [65], [66] have shown that Americium has localized $5f$-electrons at normal pressure with a FCC crystal structure. Application of high pressure leads to a phase with open, low symmetry structures which are accompanied by a delocalization of the $5f$-electrons. However there has been some debate about the exact structure that Americium is calculated to have at high pressures. Two contenders have been the orthorhombic ($\alpha$-Uranium) structure [64] or the monoclinic ($\alpha$-Plutonium) structure [65], [66]. A discrepancy in all the previous theoretical studies is that the calculated volume collapse for the transition is much higher than the experimental value.

Eriksson and Wills [64], calculate the total energy versus volume of Americium in FCC and the $\alpha$-U structures using relativistic full-potential LMTO methods. The basis set used in this computation contains $6s, 6p, 7s, 7p$ and $5f$ orbitals. On applying pressure, they calculate a transition from Am-II (FCC) to Am-III ($\alpha$-U) accompanied by a volume collapse of 34%.
According to Soderlind et al [65], however, the stable phase at high pressure is determined to be the monoclinic, i.e. the $\alpha$-Pu structure. In this calculation, the total energies of six different crystal structures of Americium is determined as a function of volume. A full-potential LMTO method using the generalized gradient approximation for the exchange-correlation term is used. Spin degeneracy is assumed for all the structures except the FCC structure. The results show that on applying pressure, there is a transition from the FCC-Americium to monoclinic-Am accompanied by a volume collapse of 25%.

The theoretical calculations of Eriksson [64] and Soderlind et al [65] underestimate the equilibrium volume of Americium at normal pressure. The volume of FCC-Americium is calculated to be 15% less than the experimentally determined volume by
Benedict [63]. Soderlind et. al also compute the total volume of Americium by treating the 5f electrons as core: in this calculation, the equilibrium volume at normal pressure is closer to the experimental volume, but still differs by 10%.

In the study by Pénicaud [66], the total energy versus atomic volume of Am is calculated in various structures. The normal phase is the FCC structure as in the previous calculations. At high pressures, the stable phase is calculated to be the α-Pu phase. On further applying pressure, a transition to the Am-IV structure is observed, which remains stable up to very high pressures. The equilibrium volume of Am at normal pressure in this study is close to the experimental value. The volume collapse for the FCC-α-Pu transition is calculated to be 31%.

In our calculation, we use the LMTO method in the atomic sphere approximation. We calculate the band structure for FCC-Americium in the LDA+U scheme, and we find that for a certain interaction $U$ and double counting term, we can correctly predict the position photoemission peak. Further, our LDA+U computation of the equilibrium volume is in close agreement with experiment. However, using the LDA+U method, we do not get the correct magnetic moment of zero. We then use the LDA+DMFT technique using the Hubbard-I impurity solver to compute the band structure of Am, which should also predict the correct magnetic moment.

In section 5.2, we describe the details of the LDA+U computation of the band structure and density of states. Section 5.2.2 deals with the equilibrium volume calculation. In section 5.3 we discuss the LDA+DMFT method, and apply it to calculate the bands in Am. Finally, our conclusions are stated in section 5.4.
5.2 LDA and LDA+U band structure calculation of Americium

The computation of the band structure is done using the LmtART code. The details of the calculation are as follows:

There are two input files required for the LDA calculation and an additional input file for the LDA+U calculation. The files are shown in the following tables. Table 5.1 has details of the crystal structure including the positions of the basis atoms and the lattice vectors. We used the FCC structure of Americium throughout our calculation. Table 5.2 contains the scheme used to calculate the self-consistency, the number of iterations in the self-consistent loop, atomic numbers and the lattice constant in atomic units. We used the atomic sphere approximation (ASA) scheme to achieve self-consistency. In order to obtain a well converged charge density and magnetization, we used the Broyden mixing scheme. For the LDA+U calculation, the details of the correlated orbitals and the interaction U are contained in Table 5.3. The band structure of Americium was calculated at various values of the interaction strength $U$.

---

Table 5.1: Input file am.str

```plaintext
( FILE=STRFILE,INPUT=MODERN )
************************************************
( SECTION=HEDS )
Slabl =Am
( SECTION=CTRS )
Natom =1
BtoA = 1.00000
CtoA = 1.00000
( SECTION=TRAN )
0.0, 1/2, 1/2
1/2, 0.0, 1/2
1/2, 1/2, 0.0
( SECTION=BASS )
0.0, 0.0, 0.0
```

---
We start by calculating the Americium band structure using the non spin-polarized LDA method. As we expect, this method predicts the $5f$ bands to lie at the Fermi level. Once we include the effects of $U$, we see the formation of Hubbard bands separated by an energy $U$. The $5f$-bands and the density of states are plotted in Figs. 5.4 and 5.5 respectively for various values of $U$. In Americium, spin-orbit effects are significant, which lead to a non-magnetic ground state with total $J = 0$. When we include the effects of spin-orbit coupling in our LDA calculation, the bands for $U = 0\text{eV}$ that lie near the Fermi level are split by an energy equal to the spin-orbit coupling. When we increase the interaction $U$, the localized $5f$-band moves further away from the Fermi level, and the energy separation between the filled and the empty bands increases by

Table 5.2: Input file am.ini

We start by calculating the Americium band structure using the non spin-polarized LDA method. As we expect, this method predicts the $5f$ bands to lie at the Fermi level. Once we include the effects of $U$, we see the formation of Hubbard bands separated by an energy $U$. The $5f$-bands and the density of states are plotted in Figs. 5.4 and 5.5 respectively for various values of $U$. In Americium, spin-orbit effects are significant, which lead to a non-magnetic ground state with total $J = 0$. When we include the effects of spin-orbit coupling in our LDA calculation, the bands for $U = 0\text{eV}$ that lie near the Fermi level are split by an energy equal to the spin-orbit coupling. When we increase the interaction $U$, the localized $5f$-band moves further away from the Fermi level, and the energy separation between the filled and the empty bands increases by

```plaintext
(FILE=INFILE, INPUT=MODERN )

(SECTION=HEAD ) ! PROJECT HEAD:
Title =Am

(SECTION=CTRL ) ! CONTROL PARAMETERS:
Lmto =Bare
FulPot=ASA

(SECTION=EXCH ) ! EXCHANGE-CORRELATION:
LDA =Vosko
GGA =none

(SECTION=ITER ) ! ITERATIVE PROCEDURES:
Niter1=200
Lbroy =-1

(SECTION=MAIN ) ! MAIN ATOMIC DATA:
Natom =1
Nsort =1
Nspin =2
Par0 = 8.72023
Is(:) =1

(SECTION=SORT ) ! SORT DATA:
Name = Am
Znuc = 95.0000

(SECTION=FFTS ) ! FFT GRIDS:
Ndiv(:)=8 8 8

Table 5.2: Input file am.ini
```

We start by calculating the Americium band structure using the non spin-polarized LDA method. As we expect, this method predicts the $5f$ bands to lie at the Fermi level. Once we include the effects of $U$, we see the formation of Hubbard bands separated by an energy $U$. The $5f$-bands and the density of states are plotted in Figs. 5.4 and 5.5 respectively for various values of $U$. In Americium, spin-orbit effects are significant, which lead to a non-magnetic ground state with total $J = 0$. When we include the effects of spin-orbit coupling in our LDA calculation, the bands for $U = 0\text{eV}$ that lie near the Fermi level are split by an energy equal to the spin-orbit coupling. When we increase the interaction $U$, the localized $5f$-band moves further away from the Fermi level, and the energy separation between the filled and the empty bands increases by
The bands are seen in Fig. 5.6 and the density of states corresponding to the above bands are seen in Figs. 5.7. In all further calculations, we include the effects of spin-orbit coupling.

---

Table 5.3: Input file am.hub

The occupancy matrices and the LDA+U potential are written in (SECTION = DHUB) in the HUB file.

In Fig. 5.8 we plot the position of the center of the 5f-level (w.r.t the Fermi level)
Figure 5.4: Energy bands of Am in LDA+U, no spin-orbit coupling. The colored regions are the fat bands of Am:5f
Figure 5.5: Density of states of Am in LDA+U, no spin-orbit coupling. The black curve is the total DOS and the red curve is the partial 5f-DOS.
Figure 5.6: Energy bands of Am in LDA+U, with spin-orbit coupling. The colored regions are the fat bands of Am:5f. Note the initial separation (for $U = 0$) between the filled and empty states, which is due to spin-orbit coupling.
Figure 5.7: Density of states in LDA+U, no spin-orbit coupling. Note the initial separation (for $U = 0$) between the filled and empty states, which is due to spin-orbit coupling. The black curve is the total DOS and the red curve is the partial 5f-DOS.
Figure 5.8: Position of center of the Am $5f$-band as a function of the interaction strength as a function of the interaction $U$, as obtained from our LDA+U calculations. The $5f$-band moves away from the Fermi level as $U$ is increased, and the center of the occupied $5f$-band linearly decreases. Experimentally, according to photoemission data (Fig. 5.2, left) obtained by Naegele et. al [60], the conduction band spectrum of Americium shows a peak at 2.8eV below the Fermi energy which corresponds to the localized $5f$-band. From our calculations of the spectra as a function of $U$, (Fig. 5.8) we see that at $U = 3eV$ the $5f$ level of Americium is centered around 2.8eV below $E_{Fermi}$.

In the LDA calculation, the filled and the empty states are separated by an energy gap of the order of spin-orbit coupling. As we increase the interaction $U$, the gap increases linearly with $U$. When we add correlations, the gap between the occupied and unoccupied states is the LDA-energy gap plus an energy equal to $U$. In Fig. 5.9 we see the gap between the spin-up and the spin-down electrons of Americium plotted as a function of $U$.

The LDA+U calculation however fails to predict the correct occupancy and correct magnetic moment of Americium. The LDA calculation predicts the occupancy of the Am $5f$-band to be 6.3. As we increase $U$, the cost of putting two electrons on the same
Figure 5.9: Gap between spin-up and spin-down electrons as a function of the interaction

Figure 5.10: Occupancy of Am $5f$-band as a function of the interaction strength $U$
site goes up, and the occupancy of the 5f-band increases. In the LDA+U calculation with $U = 4\text{eV}$, it is close to 6.8. Fig. 5.10 shows the occupancy of the Americium 5f-band as a function of $U$. However, the occupancy of the 5f states of Americium is known to be 6 (Ref. [67]). In section 5.3 we will show that the LDA+DMFT calculations predict the correct occupancy of 6, and a zero magnetic moment for Americium.

5.2.1 Effect of the double counting term

In the fat bands and density of states (Figs. 5.6 and 5.7), we kept the interaction $\bar{U}$ in the double counting term in the same as that used in the model, $U$. Decreasing the value of $\bar{U}$ however shifts the 5f peak away from the Fermi level. Fig. 5.11 shows the position of the localized 5f-level as a function of the double counting interaction $U_{dc}$, or $\bar{U}$ at $U = 3\text{eV}$. In the LDA+DMFT calculation in section 5.3 we will use $U_{dc}$ as a parameter to fix the 5f peak at its experimental value.

![Graph showing the center of the 5f-band as a function of $U_{dc}$ with $U = 3\text{eV}$](image)

Figure 5.11: Effect of $U_{dc}$: Position of the center of the 5f-band as a function of the $U_{dc}$, with $U$ kept at $4\text{eV}$.

In the next section, we turn to the details of the calculation of equilibrium volume in LDA+U, which gives excellent agreement with experiment.
5.2.2 Volume behavior

We calculate the equilibrium atomic volume of Americium by plotting the total energy from our LDA+U calculations as a function of relative volume $V/V_0$, for various values of $U$ (Fig. 5.12). From the minimum in these plots, the equilibrium volume for Americium is calculated. In each of the methods, the fcc structure is used as the crystal structure for Am. In Fig. 5.12, we use the experimental equilibrium atomic volume of 29.3 Å\(^3\) [70] for $V_0$. Our LDA+U calculation predicts the correct equilibrium volume at $U = 4.5$ eV.

We also performed a calculation by considering the 5\(f\) electrons as core electrons. In order to do this we change one of the input files lmt.am that contains all the atomic data. We remove the 5\(f\)-electrons from the basis and set them as the core electrons. We set $l_{\text{max}}=2$, and consider 8 valence electrons only from the 6\(p\) and 7\(s\) subshells. The 6\(s\) electrons are considered as semi-core electrons. The input file lmt.am that we enter the above data in is shown in Fig. 5.13. However, in this case, the self-consistent equations fail to give a bound state solution with 5\(f\) electrons in the core. The band structure and the density of states for Americium in this scheme are shown in Figs. 5.14 and 5.15 respectively. According to similar calculations by Soderlind, et.al [65] the volume obtained by treating the 5\(f\)-electrons as core electrons is only 26.6 Å\(^3\), which is about 10% less than the experimental volume.

Table-5.4 summarizes the results we obtain:

<table>
<thead>
<tr>
<th>Method</th>
<th>Equilibrium volume (in Å(^3))</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDA</td>
<td>20.3682</td>
</tr>
<tr>
<td>LDA+U, $U=1$eV</td>
<td>23.6460</td>
</tr>
<tr>
<td>LDA+U, $U=2$eV</td>
<td>26.1092</td>
</tr>
<tr>
<td>LDA+U, $U=3$eV</td>
<td>27.5871</td>
</tr>
<tr>
<td>LDA+U, $U=4$eV</td>
<td>28.7118</td>
</tr>
<tr>
<td>LDA+U, $U=4.5$eV</td>
<td>29.2</td>
</tr>
<tr>
<td>LDA w/ 5(f) in core</td>
<td>27.97</td>
</tr>
<tr>
<td>Experiment</td>
<td>29.3</td>
</tr>
</tbody>
</table>

Table 5.4: Equilibrium volumes for Am in Å\(^3\)
Figure 5.12: Total energy (-61011 Ry) as a function of relative volume $V/V_0$ for various values of interaction $U$(in eV).
* ELEMENT: Am, Configuration 6s2/6p6/7s2/6d0/5f7

name =Am ! element name
znuc =95 ! nuclear charge
zval =8 ! # of valence electrons
zsem =2 ! # of semicore electrons
amas =243 ! atomic mass
lmaxb =2 ! Lmax for LMTO basis

(subsection=lmto)
! s p d f g ! States
Mqn(:) 7 6 6 5 5 ! Main quantum numbers
Bas(:) 1 1 1 0 0 ! LMTO basis set
Mnu(:) 3 3 3 0 0 ! Choice of Enu
Enu(:) +0.5,+0.5,+0.5,+0.5,+0.5 ! Initial Enu
Dnu(:) -1.0,-2.0,-3.0,-4.0,-5.0 ! Initial Dnu

(subsection=semi)
 nsem =1 ! # semicore states
 lsem =6s ! n,l

(subsection=conf)
Orbital  n  l  j  #el. Levels(Ry)
1s1/2  1 0 1/2 2 -7850.4727
2s1/2  2 0 1/2 2 -1447.8048
2p1/2  2 1 1/2 2 -1394.6574
2p3/2  2 1 3/2 4 -1155.9321
3s1/2  3 0 1/2 2 -360.94632
3p1/2  3 1 1/2 2 -336.82037
3p3/2  3 1 3/2 4 -282.31668
3d3/2  3 2 3/2 4 -243.61142
3d5/2  3 2 5/2 6 -232.36490
4s1/2  4 0 1/2 2 -89.627118
4p1/2  4 1 1/2 2 -78.886649
4p3/2  4 1 3/2 4 -64.904609
4d3/2  4 2 3/2 4 -47.651517
4d5/2  4 2 5/2 6 -45.061490
4f5/2  4 3 5/2 6 -22.427601
4f7/2  4 3 7/2 8 -21.774721
5s1/2  5 0 1/2 2 -19.277471
5p1/2  5 1 1/2 2 -15.228066
5p3/2  5 1 3/2 4 -12.070072
5d3/2  5 2 3/2 4 -6.1802559
5d5/2  5 2 5/2 6 -5.7243980
5f5/2  5 3 5/2 6 -0.1227601
5f7/2  5 3 7/2 1 -0.1227601
6s1/2  6 0 1/2 2 -3.0330305
6p1/2  6 1 1/2 2 -1.9080418
6p3/2  6 1 3/2 4 -1.4078676
7s1/2  7 0 1/2 2 -0.3556830

Figure 5.13: Input file lmt.am for treating f-electrons as core.
Figure 5.14: Energy bands of Am treating 5f electrons as core

Figure 5.15: DOS of Am treating 5f electrons as core
5.3 LDA+DMFT method

The equilibrium volume of Americium predicted by LDA+U is in good agreement with experiment. However it fails to predict the correct zero magnetic moment. We now apply the LDA+DMFT method described in chapter 3 to calculate the band structure of americium.

To solve the impurity model, we use a method that is based on the Hubbard-I approximation. We do a full self-consistent calculation, where the charge density is recomputed after the DMFT loop and is fed back to the routine that calculates the potential and then the new LDA Hamiltonian.

5.3.1 Hubbard-1 method

The starting point of the Hubbard-I approximation is the atomic limit. We first find the Green’s function for the atomic Hamiltonian using the equation of motion method for the Hubbard operators. We then express the impurity Green’s functions in terms of the atomic Green’s function and the hybridization $\Delta(i\omega)$.

We start with an impurity model that has the form $H = H_{\text{atom}} + H_{\text{band}} + H_{\text{hyb}}$.

\[
H_{\text{atom}} = \sum_{l,l'=5/2,7/2} \epsilon_{ll'\text{mm'}} f_{l'm'}^\dagger f_{lm} + U \sum_{l,l'=5/2,7/2} f_{l'm'}^\dagger f_{lm} f_{l'm'} f_{lm'}
\]

\[
H_{\text{band}} = \sum_{lk} \epsilon_{lk} c_{lk}^\dagger c_{lk}
\]

\[
H_{\text{hyb}} = \sum_{l'l' \text{mk}} V_{l'l'k} (f_{lm} c_{l'k} + \text{h.c.})
\]

(5.1)

Here $l, l'$ label the two bands that are split due to spin-orbit coupling. These two bands have the $j$-quantum number as $5/2$ or $7/2$. Notation: The band with $l = 5/2$ and is 6-fold degerate and that with $l = 7/2$ and is 8-fold degenerate. Thus $m, m'$ label the degeneracy and run from 1 to $(2j + 1)$. Since the impurity level matrix $\epsilon_{ll'\text{mm'}}$ has off-diagonal terms as zero, the first term simplifies to $\sum_{lm} \epsilon_{l} f_{lm} f_{lm}^\dagger$. $H_{\text{hyb}}$ is due to the hybridization between the conduction electrons and the localized $f$ electrons. The hybridization function can be written as $\Delta_{l}(i\omega) = \sum_{k} V_{l'k}^2 / (i\omega - \epsilon_{l'k})$. 
In the case of Americium, due to the above degeneracy, we have a $SU(6) \times SU(8)$ symmetry. With this, the impurity Green’s function in the Hubbard-1 approximation becomes

$$[G^{imp}_i(i\omega)]^{-1} = [G^{at}_i(i\omega)]^{-1} - \Delta_i(i\omega) \quad (5.2)$$

The atomic Green’s function $G^{at}$ is given by

$$G^{at}_i(i\omega) = \sum_{n=0}^{14} \sum_{n_1=0}^{n} \frac{(C^6_{n_1} C^8_{n-n_1} e^{-E_{ln}/T} - e^{-E_n/T})}{i\omega + \mu - E_{ln} + E_n} / Z \quad (5.3)$$

where $E_{ln}$ is the energy of an atom with $n$ total number of electrons, $n_1$ in level $l = 1$ and $n-n_1$ in level $l = 2$ is given by :

$$E_{ln} = E_n + \epsilon_1^f + Un \quad (5.4)$$

and

$$E_n = \frac{U(n-1)n}{2} + n_1\epsilon_1^f + (n-n_1)\epsilon_2^f \quad (5.5)$$

The numerator in (5.3) denotes the probability of finding an atom with $n$ total number of electrons, $n_1$ in level $l = 1$ and $n-n_1$ in level $l = 2$. $C^6_{n_1}$ and $C^8_{n-n_1}$ are the combinatorial coefficients. $Z$ is a normalization factor for the probability given by

$$Z = \sum_{n=0}^{14} \sum_{n_1=0}^{n} \frac{C^6_{n_1} C^8_{n-n_1} e^{-E_n/T}}{(n-n_1) \leq 8} \quad (5.6)$$
5.3.2 Tight-binding calculation

We first do a tight-binding calculation for the band structure of americium. We generate the hop file that gives the hopping integrals and positions of the impurity levels. We then run the LMTART program in the tight-binding mode and calculate the bands and density of states.

Experimentally, Americium has zero magnetic moment with 6 electrons in the $j = 5/2$ state. Thus from the tight-binding calculation, we choose a model with two bands – band 1 corresponding to $j = 5/2$ and band 2 that has $j = 7/2$:

$$H_{TB} = t_1 c_1^\dagger c_1 + t_2 c_2^\dagger c_2 + \epsilon_1 f_1^\dagger f_1 + \epsilon_2 f_2^\dagger f_2$$

where $c_1$ and $c_2$ are the operators for the conduction electrons in bands 1 and 2 respectively, and $f_1$ and $f_2$ are the operators for the $f$-electron. The parameters $t_1, t_2, \epsilon_1$ and $\epsilon_2$ are chosen so that the bands from the tight-binding calculation match the bands from the fully self-consistent LDA calculation. We find that the other hoppings are negligible compared to $t_1$ and $t_2$. Figs. 5.16-5.19 show a comparison between the density of states obtained from the tight-binding calculation and those obtained from the LDA self-consistent calculation.

Figs. 5.20 and 5.21 shows how the tight-binding parameters change as a function of pressure. In the calculation, we reduce the relative volume $V/V_0$, thus applying pressure on the system. As the pressure is increased we find that the parameters increase.

We then perform a DMFT calculation on the above model, using the Hubbard-I impurity solver described in Chapter 3. A plot containing the density of states for different values of $U$ obtained from LDA+DMFT on the above model is shown in Fig. 5.22. As $U$ is increased, we see the increase in the gap between the occupied and the unoccupied states.
Figure 5.16: Comparison of DOS: LDA and Tight binding methods
Figure 5.17: Comparison of DOS : LDA and Tight binding methods
Figure 5.18: Comparison of DOS: LDA and Tight binding methods
Figure 5.19: Comparison of DOS: LDA and Tight binding methods
Figure 5.20: Impurity level (w.r.t $E_F$) as a function of relative volume

Figure 5.21: Tight binding hoppings as a function of relative volume
Figure 5.22: Am: Density of States as a function of $U$ for the tight-binding model
5.3.3 LDA+DMFT Results

The above tight-binding model for Americium correctly describes its natural state. We now perform a full self-consistent LDA+DMFT calculation on Am, with the effects of spin-orbit coupling included. We use the Hubbard-I solver explained in section 5.3.1 as the impurity solver.

In Figs. 5.23 we plot the total energy (-61102Ry) as a function of relative volume at $U = 4.5eV$.

The volume $V_0$ is chosen to be the experimental volume.

![Figure 5.23: Energy vs. relative volume](image)

The equilibrium volume is predicted correctly by the LDA+DMFT method, similar to the equilibrium calculation in the LDA+U method. The curve with circles is obtained from the high volume side, gradually decreasing the volume and using the self-consistent results from the previous run for the next. The curve with squares is obtained starting from the low volume side, and gradually increasing the volume. We find a region near $V/V_0 = 0.7$ where there are two different energies for the same volume. The pressure,
i.e., the derivative of energy w.r.t. volume $\partial E/\partial V$ is plotted as a function of relative volume in Fig. 5.24.

The region of the where there is a discontinuity in the solutions can be thought of as a coexistence region for the metallic-like and the insulating-like phases of the material. This is a strong indication of the a Mott transition where the $5f$ electrons undergo a change of character from localized to delocalized as we increase the pressure.

![Figure 5.24: Pressure vs. relative volume](image)

We now turn to how the Americium $5f$-density of states behave as we apply pressure on the system. The DOS calculated from LDA is shown in Fig. 5.27. As we saw in sections 5.2 and 5.2.2, LDA neither correctly predicts the position of the $5f$ peak nor the equilibrium volume. However, as we apply pressure we see that the DOS start broadening.

We need to include the effects of $U$ to correctly predict the $5f$-peak and the equilibrium volume. At $U = 4.5eV$, the value of the interaction which gives the correct equilibrium volume, we find that the position of the $5f$-peak is at $3.8eV$ below the Fermi
Figure 5.25: Am DOS in LDA
Figure 5.26: Am DOS in LDA
Figure 5.27: Am DOS in LDA
level. In order to fix the 5f-peak at its experimental value, we use the interaction in the double counting term $\bar{U}$, or $U_{DC}$ as a parameter. Fig. 5.28 shows the evolution of the DOS with pressure at $U = 4.5eV$ when $U_{DC}$ is kept at the same value as $U$. It turns out that to have the 5f-peak at its experimental value of 2.8eV below the Fermi level, we need $U_{DC}$ to have a value slightly less than $U$, namely, $U_{DC} = 4.1eV$. In Fig. 5.31 we see how the DOS evolves with pressure for the parameters $U = 4.5eV$ and $U_{DC} = 4.1eV$.

At the equilibrium volume $V/V_0 = 1.00$, we see two Hubbard bands separated by an energy difference that accounts for $U$ as well as spin-orbit coupling. The filled 5f-band below the Fermi level has 6 electrons and the unoccupied band above the Fermi level contains 8 electrons.

We have plotted the total density of states as well as the 5f density of states. We can see that under pressure the highly localized 5f-bands start widening and merging into each other. When $V/V_0 = 1.00$ we see that the 5f states do not contribute much at the Fermi level. Most of the weight at low pressures is due to the s,p and d electrons, shown in grey in the figure. As the pressure is increased, the ratio of 5f states to the total DOS near the Fermi level increases. Near $V/V_0 = 0.75$ we observe a spreading out of the DOS that signals a transition to a metallic state. Since the Hubbard-1 approximation essentially starts with an atomic solution, it always predicts the existence of two Hubbard bands for an arbitrarily small $U$. It fails to predict a sharp quasiparticle peak for a metallic state. However we do see a finite and increasing but wide distribution of states at the Fermi level as we increase pressure. In Fig. 5.34 we see the total and partial 5f-density of states at the Fermi energy, and in Fig. 5.35 we see the ratio between the 5f-DOS and the total DOS plotted as a function of the relative volume.
Figure 5.28: Am DOS in LDA+DMFT: $U = U_{DC} = 4.5eV$
Figure 5.29: Am DOS in LDA+DMFT: $U = U_{DC} = 4.5eV$
Figure 5.30: Am DOS in LDA+DMFT: $U = U_{DC} = 4.5\, \text{eV}$
Figure 5.31: Am DOS in LDA+DMFT: $U = 4.5eV, U_{DC} = 4.1eV$
Figure 5.32: Am DOS in LDA+DMFT: $U = 4.5\, eV, U_{DC} = 4.1\, eV$
Figure 5.33: Am DOS in LDA+DMFT: $U = 4.5\text{eV}, U_{DC} = 4.1\text{eV}$
Figure 5.34: Density of states at Fermi level in LDA and LDA+DMFT
Figure 5.35: Ratio of 5f-DOS over total DOS
Figure 5.36: Position of center of 5f-band
5.4 Summary and Conclusions

We have calculated the electronic band structure of Americium using the LDA+U method and the LDA+DMFT methods. The LDA+U method gives consistent results with photoemission spectra and equilibrium volume of Am. However we fail to get the correct magnetic moment in Americium using LDA+U. Americium is in a $J = 0$ non-magnetic ground state due to spin-orbit coupling. However, the LDA+U computation predicts a half-filled shell and a magnetic moment of $7\,\mu_B$.

We then use the LDA+DMFT method to compute the structure and volume of Am. We can correctly predict the position of the 5f-peak as well as the correct equilibrium volume of Am. On applying pressure, we observe a hysteresis loop in the energy-volume curve which strongly indicates the proximity to a Mott transition.
Chapter 6
Summary and conclusions

In this thesis we have numerically studied aspects of the Mott transition in model Hamiltonians as well as in real materials.

In chapter 2, we introduced the idea of the Mott transition, that is, a first order metal-insulator transition driven by electron correlations. We applied the dynamical mean-field theory (DMFT) to study a simple model Hamiltonian: the degenerate Hubbard model. Based on numerical calculations of the behavior of the particle occupation number as a function of doping, we mapped out the phase diagram of the Hubbard model in the paramagnetic regime. We found a region of coexistence between metallic and insulating phases at finite temperatures. Furthermore, we showed that at small but finite doping, the compressibility diverges at the Mott endpoint. Our model calculations were found to be relevant to the $\alpha$-$\gamma$ transition in Ce.

We briefly shifted gears in chapter 3 and gave an overview of various density functional methods, which are first-principles methods in the sense that no empirical parameters (other than atomic charges and lattice structure) are needed to predict physical properties of materials. Density functional theory in the local density approximation (LDA) has had considerable success in understanding electronic structure of weakly interacting solids. We then moved on to describe methods that go beyond LDA, in order to deal with strongly correlated materials. We focused on two such methods: LDA+U, where U is the Coulomb interaction, and the LDA+DMFT which uses LDA to describe the light $s, p$ (or $s, p, d$) electrons and DMFT to treat correlated $d$ and $f$ electrons.

Chapters 4 and 5 contain applications of the two theoretical approaches we discussed so far: a many body Hamiltonian approach combined with first-principles calculations.
We used these techniques to study the electronic structure and properties of two materials that have strong correlations due to $f$-band electrons.

In chapter 4, we investigated YbRh$_2$Si$_2$, a heavy fermion compound that exhibits many deviations from Fermi liquid behavior due to interactions between the localized $4f$ electrons in the Yb$^{3+}$ ion and the conduction band formed by the $s$, $p$ and $d$ electrons. We used the LDA+U technique to determine the electronic structure and magnetic properties of YbRh$_2$Si$_2$. In the process, we did a careful study of the so-called double-counting term, that is, the energy that needs to be subtracted from the LDA+U functional, since part of the correlation energy is already present in the LDA functional. As the double-counting term was too large in the standard LDA+U functional, we failed to obtain the correct magnetic ground state of YbRh$_2$Si$_2$. We presented a prescription to rectify the situation: we used the interaction $U_{dc}$ in the double counting term as a different parameter compared to the interaction $U$ in the LDA+U functional. Keeping $U_{dc}$ slightly smaller than $U$ predicted the correct band structure and magnetic moment of YbRh$_2$Si$_2$. We also successfully calculated the magnetic anisotropy energy in this material, which is very large.

In chapter 5, we returned to the idea of the Mott transition. We used LDA+U and LDA+DMFT methods to examine the delocalization-localization transition of $5f$ electrons in elemental Americium. This material, which borders Plutonium in the periodic table, has been the subject of a number of theoretical and experimental investigations. We put into practice the insights we gained from chapter 4 regarding the double counting term to correctly predict the photo-emission spectra. The LDA+U calculation predicted the correct equilibrium volume but not the correct magnetic moment. We then applied the LDA+DMFT technique, using the Hubbard-1 method as the impurity solver for the DMFT loop. We obtained strong evidence for the existence of a Mott transition without a large volume collapse.

**Outlook**

The Mott transition in Americium can be better understood if we use different structures for the different phases. In our calculations we used an impurity solver that
is computationally fast, but the price we paid is perhaps an oversimplification of the physical picture. The Quantum Monte Carlo method, that we studied in chapter 2, is exact, but expensive in terms of computation time. A better impurity solver that is also fast is required in order to investigate complex materials. In addition, a first principles approach to determine the interaction parameters and the double-counting corrections will complete the picture of ab-inito many body calculations.
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