### **Abstract**

In this thesis we discuss tools, results and techniques pertaining to discussions around unconventional superconductors of chiral p-wave symmetry. This symmetry exists in the superfluid A-phase of He<sup>3</sup> and has been suggested to describe the superconductivity of  $Sr_2RuO_4$ .

In Paper I we use field-integral techniques to investigate the effects of spin-orbit coupling on the coefficients of the phenomenological Ginzburg-Landau theory of chiral *p*-wave superconductors. We find that these coefficients have a non-linear anisotropic dependence on the spin-orbit coupling strength and direction in spin-space. This dependence necessitates two independent phenomenological parameters for the mixed gradient terms and the mixed component terms respectively, even in the weak-field limit when written using dimensionless variables.

In Paper II we use large-scale Monte-Carlo simulations to investigate the vortex-matter of a superconducting system that can be modelled by a Ginzburg-Landau theory with chiral p-wave symmetry such as the one investigated in Paper I, but now in the limit of vanishing spin-orbit coupling. We find that a square vortex lattice consisting of single-quanta vortices is stable at high temperatures close to  $T_c(B)$ . The single-quanta vortices merge into double-quantum vortices at lower temperature which together then stabilizes a triangular vortex lattice.

In Paper III we investigate a  $\mathbb{Z}_2$  Ising transition resulting from spontaneously broken time-reversal symmetry in the neutral sector of chiral p-wave symmetric superconductors subjected to zero external field. We find that this transition is irrevocably tied to the superconducting

transition for all realistic values of the phenomenological parameters in our model.

### **Preface**

This thesis is submitted in partial satisfaction of the requirements of the degree Philosophiae Doctor (PhD) at the Norwegian University of Science and Technology, in Trondheim Norway.

The work that this thesis presents started in September 2015 and ended in early spring 2021 at the Center for Quantum Spintronics (QuSpin), NTNU. During this time, one year of accumulated time was dedicated to teaching duties at the Department of Physics, and half a year was devoted to completion of courses (30 ECTS) as pr. the requirements of the degree. The research was supervised by Prof. Asle Sudbø as main supervisor, and Prof. Jacob Linder as co-supervisor.

Computation-time was granted at the Vilje and Fram supercomputers through the UNINETT  $\int$  igma2 e-infrastructure. The code was written in the Julia programming language. The figures and plots were produced by the use of Julia and Inkscape. The thesis was written in LaTeX based on a template by J. A. Ouassou available under CC BY 4.0 license, that was heavily modified.

Fredrik Nicolai Krohg Oslo, March 2021

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In the department of physics at NTNU there are many others who deserve recognition and praise for their role in creating a friendly and interesting social environment. To mention everyone by name in such a long PhD as I have had would prove excessive and frankly a boring read. Therefore I will not attempt to make an exhaustive list but mention only a select few.

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tantly: a willingness to join me for Beatz! Another co-conspirator in that endeavour has been Akash, a man I learned a great deal from not only about physics and its thinly veiled politics but also in areas of life, philosophy and sociology, one who has been integral in the culture of QuSpin and who it's been an absolute pleasure knowing.

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## List of papers

### Paper I:

Fredrik Nicolai Krohg and Asle Sudbø

Derivation of Ginzburg-Landau free energy density containing mixed gradient terms of a p+ip superconductor with spin-orbit coupling

Physical Review B **98**, 014510 (2018) [1]

#### Paper II:

Fredrik Nicolai Krohg, Egor Babaev, Håvard Homleid Haugen and Asle Sudbø

 $Thermal\ fluctuations\ and\ vortex\ lattice\ structures\ in\ chiral\ p\mbox{-wave\ superconductors:}$   $robustness\ of\ double\mbox{-quanta\ vortices}$ 

arXiv:2007.09161 (2020) [2]

#### Paper III:

Håvard Homleid Haugen, Egor Babaev, Fredrik Nicolai Krohg and Asle Sudbø

Zero field phase transitions in a chiral two-component superconductor pre-print

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### Introduction

Research into superconductors holds a vital key in the development of technologies that can reduce global emissions of greenhouse gasses and thus prevent large economic as well as human losses due to the effects of the climate crisis. In IPCC's special report, they state that in order to have no or limited overshoot in global temperature from the goal of 1.5 °C, the global net anthropogenic emissions of CO<sub>2</sub> needs to decline by 45% compared to such emission levels in 2010, and this has to happen by 2030. The emission levels must then continue to decline, reaching net zero around 2050 [3]. In order for the member nations of the Paris Agreement to meet this goal, the NDC Synthesis report [4] highlights the need for further increase in the nations contributions compared to those that are currently declared. Measures mentioned by member nations for mitigating the release of greenhouse gasses include renewable energy generation, electrification of the transport sector and more efficient electrical grids. Because of the non-traditional properties of superconductors, such materials could potentially be of great benefit in further strengthening such mitigation strategies.

In aircraft travel, designs for hybrid electric aircraft such as NASA's N3-X are underway. Analysis shows that fully utilizing high temperature superconductors in the propulsion system could provide as much as 3.5 times higher power-to-weight ratio than previous designs due to superconductors high current-densities [5].

In Norway there has recently been a debate about the development of wind turbine parks close to population centers and in vulnerable natural habitats. Moving the wind power production from land to sea solves some of the debated issues but needs effective turbines. Including high temperature superconductors into the design of such offshore wind-turbines is beneficial for much the same reasons as for aircraft design: the high power density makes for a compact, lightweight and efficient construction [6, 7].

Other examples of future applications of superconductors include their use in more efficient power grids [8, 9], sustaining the high magnetic fields needed for nuclear fusion [10, 11] and for the operation of a particle collider more powerful than the LHC [12], faster, more efficient electronics for digital logic and memory devices and more robust quantum computers [13].

Superconductors also currently have numerous important applications. In the Chūō Shinkansen magnetic levitation line, which is currently under construction, the interaction between superconducting coils in the train and copper coils on both sides of the track provides levitation and guidance of the train at high speeds [14]. Superconductors are essential for generating the high strength magnetic fields needed in MRI imaging. They are also used in other medical settings such as measurements of the electrical currents in the heart (magnetocardiography), in measuring the concentration of iron stored in the liver (biomagnetic liver susceptometry) and cancer treatments through their role in particle accelerators [15].

All of this is the product of fundamental research into the electronic properties of metals and other materials that has shown that for some of them, at a critical temperature  $T_c$ , the electrical resistivity of the material suddenly vanishes and any external magnetic field is expelled. These are the two main properties that we associate with the phase of superconductivity. Zero resistivity means that electricity can travel through the material without losing any energy, in contrast to a normal conductor where energy is usually lost through heat. The expulsion of magnetic fields is called the Meissner effect and is in a sense the more fundamental of the two properties. On a microscopic level it is due to electrons forming paired states that share certain features in such a way that different pairs can behave as one. Because a macroscopic

number of states share these features, the quantum mechanical nature of such states which is usually only significant for tiny particles, becomes apparent through these non-classical macroscopically measurable effects.

### 1.1 A brief history of superconductivity

Superconductivity was first discovered in mercury at  $T_c \approx -268.99\,^{\circ}\mathrm{C}$  by Heike Kamerlingh Onnes in the Netherlands in 1911 [16]. The Meissner effect was then discovered in 1933 by W. Meissner and R. Ochsenfeld [17]. These discoveries happened without any previous theoretical prediction or explanation. Theoretical description was then gradually developed, first by a simple thermodynamic two-fluid model of electron densities by Garter and Casimir and then in 1935 by the phenomenological theory of the electromagnetic properties by H. and F. London [18]. WWII came and went and then a significant improvement on the London-model was published by V. L. Ginzburg and L. D. Landau in 1950 [19], which built on Landau's previous description [20] of a second order phase-transition by an order-parameter quantity. Based on this theory, Abrikosov introduced the concept of a type-II superconductor in 1952, which has negative surface energy and a mixed phase at non-zero magnetic field [21].

An attempt at a microscopic theory was given by Frölich in 1950 based on electron-phonon interaction [22]. Even though the perturbation theory he derived failed to predict important superconducting properties such as the Meissner effect, his Hamiltonian later became well known as a fruitful starting point for the application of field theoretic methods. In 1953 Pippard introduced a second length scale, the coherence length  $\xi$ , through a non-local modification to the Londonmodel [23]. This length scale was a measure of the width of the interface between normal and superconducting regions. Although not a theory of superconductivity itself, Landau's fermi liquid theory which came in 1956 would prove crucial in the development of a microscopic theory and describes the electronic properties of many metals that at lower temperature become superconducting [24]. A complete microscopic theory of superconductivity was published by J. Bardeen, L. N. Cooper and J. R. Schrieffer in 1957 [25, 26]. The BCS-theory was based on the idea that fermi-liquid quasiparticles with opposite momentum

could form an attractive interaction through an intermediate interaction with a phonon. This would then lead to the formation of pairs that could form a condensate, and which implied an energy gap  $\Delta$  between the energies of paired electrons and energies of normal quasiparticle states in the Fermi-sea. This year Abrikosov also published his prediction of the existence of a lattice of vortices in the mixed state of type-II superconductors  $\cite{Dallo}$ 

A separate form of a microscopic theory appeared in 1958 by N. N. Bogoliubov in a series of papers [28–30]. This methodology of solving the Frölich Hamiltonian was presented in a book [31] by P. G. de Gennes and has since become known as the Bogoliubov-de Gennes or BdG equations.

The diagrammatic methods developed for high-energy physics was first applied by Gor'kov to the problem of superconductivity in 1958 when he calculated Green's functions based on the ideas of BCS-theory that reproduced its results [32]. He then in 1959 used these methods to prove that the Ginzburg-Landau theory follows from the BCS theory in the limit  $T \to T_c$  [33]. The application of field theoretic methods was extensively developed by the work of Nambu, published in 1960, where he introduced the Nambu-spinor for calculating the Gor'kov Green's functions. A perturbation theory for these Green's functions were calculated by Éliashberg in the same year following a similar approach as Nambu, which later become known as Éliashberg theory [34].

The understanding of the effects of impurities got an important contribution in 1959 by what is known as Anderson's theorem [35]. It says that any instability of the Fermi-surface that does not lift the Kramer degeneracy of time-reversed paired quasi-particles do not affect the mean-field transition temperature [36]. The idea of an energy gap in the excitation spectrum, which was integral to the BCS theory, was given strong experimental backing by the tunneling experiments of I. Giæver in 1960 [37]. Such experiments were given a theoretical understanding by B. D. Josephson in 1962, through what is now known as the Josephson effect [38].

From the framework of the Gor'kov Green's functions, a set of transport equations were derived for type-II superconductors in 1968 by

Eilenberger [39]. These equations were further simplified for the case of a dirty superconductor by Usadel in 1970 [40].

From the perspective of our work, the discovery of a new phase in  ${\rm He^3}$  by Osheroff et al. in 1972 was of particular importance. Although a superfluid and not a superconducting phase, the A-phase of this system has an unconventional anisotropic pairing symmetry which was famously described by Leggett in his 1975 review article [41]. This is the same symmetry that we have considered in our work.

Superconductivity was found in the first heavy-fermion system  $CeCu_2Si_2$  in 1979 by Steglich et al. [42]. A class of systems where the superconducting state consists of quasiparticles that are fermions with a large effective mass and where the superconducting order is of an unconventional character. For a review see Ref. [43].

The first high- $T_c$  superconductor was discovered in the form of  ${\rm La_2CuO_4}$  by Bednorz and Müller in 1986 [44]. This was followed up one year later by the discovery of  ${\rm YBa_2Cu_3O_{7-x}}$  by M. K. Wu et al. [45]. These discoveries ushered in an era of superconductivity research dominated by cuperates — ceramic compounds consisting of metal oxides between planes of  ${\rm CuO_2}$ . These are truly unconventional superconductors in that their pairing symmetry is demonstrably non-isotropic. Through a group-theoretical approach, a vast array of unconventional symmetries and their Ginzburg-Landau theories and physical properties were enumerated by Sigrist and Ueda in 1991 [46].

In 1994, superconductivity was discovered in the perovskite structure of  $\rm Sr_2RuO_4$  by Y. Maeno et al. [47]. This proved that copper was not a necessary ingredient for superconductivity in layered perovskite crystal structures, and would be the starting point of a still-standing debate about its pairing symmetry which served as the immediate backdrop to our own research.

A strong case was made by D. J. Scalapino in 1995 that for cuperates, the pairing had  $d_{x^2-y^2}$  symmetry caused by antiferromagnetic spin-fluctuations [48]. Above the superconducting transition temperature but below a characteristic temperature  $T^*$  there is a hitherto undiscovered phase in such compounds where the electronic density of states near the Fermi-surface continue to be suppressed by an energy gap  $\Delta_{\rm PG}$ . This phase was named the pseudogap phase by Ding et al. in 1996 [49], and its origin continues to be a hotly debated topic.

2008 marked the beginning of the "iron age" of superconductivity research by the discovery of the first iron-based superconductor  ${\rm La}[{\rm O}_{1-x}{\rm F}_x]{\rm FeAs}$  by Y. Kamihara et al. [50]. The materials in this family of superconductors, called iron-pnictides, feature high  $T_c$  and several other exotic properties including nematic order [51]. For a review see Ref. [52].

Lastly one could argue that we now have entered a "hydrogen age", as an increasing number of hydrogen-rich compounds are approaching room-temperature superconductivity as long as they are put under insane pressures [53, 54]. It could also be argued that we are currently in a "topological age" as topological edge states in superconductors are field under intense study [55] that potentially has far reaching consequences through their immediate application to quantum computing. Others again, would surely argue that we are in a "graphene age" as novel forms of superconductivity have been observed in twisted layers of graphene [56]. Which age we are in, I suspect, depends on what field the researcher you are asking works on, and a clear answer will have to be postponed until seen through the coarse-grained eyes of history.

#### 1.2 About this work

In the last section we saw how the research into the phenomena of superconductivity has blossomed into a myriad of different directions and sub-fields. Through all this research, an implicit motivation has been the search of one day finding a theory or a specific system of a material that is superconducting at room temperature. This has become a goal like the philosopher's stone that we still have not quite reached, but whose continued pursuit itself has borne numerous fruits.

As for this thesis, we have focused on a branch of unconventional superconductivity that pertains to the description of phases with  $k_x \pm i k_y$  chiral p-wave pairing symmetry. The p in p-wave implies that the pairing states internal angular momentum has l=1, i.e. it has a 1st order (linear) dependence on its internal angular momentum. As we mentioned, this is the same kind of state that describes the real world sys-

The letter-convention stands for "principal" and comes historically from the study of atomic emission spectra that result when electrons jump between different orbitals.

tem of the  ${\rm He^3}$  superfluid A-phase. In that context it is often referred to as the ABM-state after Anderson Brinkman and Morel who first described this pairing state in the context of the BCS-theory [57] in 1961, and then demonstrated how this state was stable in the A-phase of  ${\rm He^3}$  in 1973 [58]. It was for a long time thought that the unconventional superconducting phase of the perovskite compound  ${\rm Sr_2RuO_4}$  had such a pairing symmetry, with one of the chief reasons being that it clearly features spontaneously broken time-reversal symmetry. This formed in part the motivation for much of our work.

The determination of pairing symmetries is an important step in understanding what type of superconductivity is in a system because it leads to distinct experimental consequences. Roughly speaking one can think of the pairing symmetry as determining the  $\boldsymbol{k}$  dependence of the gap function

$$\Delta(\mathbf{k}) = \sum_{m} \eta_{m} b_{m}(\mathbf{k}), \tag{1.1}$$

where  $b_m({m k})$  are basis functions that depend on the point group symmetry of the system and  ${m k}$  determines a point on the Brillouin zone. Depending on these basis functions then,  $\Delta({m k})$  could have points or lines in the Brillouin zone where it vanishes, so-called point- or linenodes. The existence of such nodes imply distinct signatures such as in the temperature dependence of the specific heat. Other examples of experimental signatures of unconventional symmetry include the Knight-shift, any Kerr-angle rotation and the possible symmetry of lattices of magnetic vortices. For an introduction to superconductors with unconventional pairing symmetries we highly recommend the lecture notes by Sigrist in Ref. [59] and [60].

Importantly, one may derive the form of the Ginzburg-Landau theory of the superconductor based solely on the pairing-symmetry as was done by Sigrist and Ueda for a large number of different symmetries in 1991 [46]. The Ginzburg-Landau theory is a phenomenological theory, meaning that it explains the effective phenomenon observable in superconductors without necessarily knowing all the microscopic details. As such, any correct microscopic theory should then reduce to the phenomenological Ginzburg-Landau theory in the limit of  $T \to T_c$ . As we mentioned, Gor'kov first did this for a conventional s-wave superconductor in 1959 through a Green's function approach

[33]. One fruitful starting point for such a microscopic theory is the Hubbard-Hamiltonian

$$\hat{H} = \sum_{ijss'} H_{ij;ss'} \hat{c}_{is}^{\dagger} \hat{c}_{js'} + \mathcal{O}(\hat{c}^4), \tag{1.2}$$

which itself can be viewed as an effective theory of the underlying many-body quantum mechanics. In a Hubbard-theory, electrons can occupy sites in an atomic crystal lattice and hop from one site to another. Any long-range interaction such as the Coulomb interaction is then written in terms of how electrons at different sites interact through nearest neighbor terms, next-nearest neighbor terms, etc. In Chapter 3 we will describe some of the tools useful in deriving an effective Ginzburg-Landau theory from such a microscopic starting point. We then in Chapter 4 present the group-theory needed for deriving the form of such a microscopic theory that results in a sought-after pairing-symmetry.

In the last chapter of Chapter 7, we present some tools useful when investigating magnetic vortices and vortex-lattices in chiral p-wave superconductors.

As was said by Leggett in his nobel lecture, there are very few things that can be proved rigorously in condensed matter physics, by which he referred to analytic arguments [61]. To get around this difficulty we have used Monte-Carlo techniques that relies on the power of computers to simulate physical consequences of theoretical models, specifically for our case: a Ginzburg-Landau theory. Such techniques have a long history in our group of being successfully able to simulate superconductive systems. For a few examples see [62–66]. Although similar techniques have been used for centuries, their modern form was first developed in the context of nuclear research in the Manhattan project. After the researchers had become familiar with the Monte Carlo casino in Monaco, they named it after the casino because of the technique's reliance on random or pseudo-random numbers to calculate multidimensional integrals [67].

In our use of Monte-Carlo techniques, which we present in Chapter 6, they are used to calculate thermal averages of statistical-mechanical observables. The rough procedure is that first, the theory under investigation is discretized to a corresponding lattice model as described in Chapter 5, such that the probability of any configuration of fields

on this numerical lattice can be computed. The theory then implies a probability distribution for how likely certain field-configurations are to materialize in a real system. Then, from a predetermined starting configuration, small changes are made incrementally to the numerical configuration in such as way as to yield numerous statistical samples of field configurations that follow the theoretical probability distribution after sufficient number of incremental changes have been done. These samples of field configurations finally are used to calculate statistical averages that correspond to taking thermal averages in statistical mechanics of the observables we are interested in.

We begin this thesis in Chapter 2 with a brief review of some relevant aspects of statistical mechanics that should refresh what is meant by thermal averages of observables, how these are tied to probability distributions of system configurations and a brief introduction to Landau and Ginzburg-Landau theory, as it pertains to phase transitions.



### **Statistical Mechanics**

In statistical mechanics we attempt to describe an ensemble of particles that may be interacting to extract not precise information about what each and every particle is doing, but statistical information about what most of the particles are doing. One can imagine this process as zooming out from a detailed view of individual entities and viewing the resulting collection through squinted eyes. This amounts to treating the collection in a course-grained manner. Through such eyes, only the most significant behavior is perceptable such that it can be understood and described in a simplified way.

### 2.1 Canonical ensemble and the partition function

Most of the business of statistical mechanics is about calculating what is known as the partition- function. Once this function is known, all the heavy lifting is done since most important statistical quantities can be extracted from it following already established systematic steps. To calculate the partition function is theoretically very simple: we sum the quantity  $e^{-\beta E_i}$  over all the possible states of the system. Every state is a particular configuration of things in the system and since all things in the system have a certain energy, the energy of a system state is given by the sum of all these things' individual energy, pluss any energy given by interactions between then. If we label each state of the system with the index i, then we can denote the energy of each

state  $E_i$ . The definition of the partition function Z in the canonical ensemble can then be written

$$Z = \sum_{i} e^{-\beta E_i},\tag{2.1}$$

where  $\beta = 1/(k_{\rm B}T)$  and  $k_{\rm B}$  is the Boltzmann constant,<sup>1</sup> given that the number of different possible states is countable.

As we can see, the essential ingredients needed to calculate the partition function is one: to be able to enumerate all possible states i of the system, and two: to be able to calculate their corresponding energies  $E_i$ . Since we have used the summation sign  $\sum_i$  in Eq. (2.1), we have assumed that there exists a countable number of different states. However if there is one thing in the system that can change in a continuous fashion, which we would measure using the set of real numbers  $\mathbb R$  and some unit, then the number of states is infinite and uncountable. In this case we sum over the different numbers of states by simply integrating over the things that are continuous and the unit of the partition function becomes the product of the units of the continuous variables (things). In most cases, it is the position r and momentum p of particles in the system that are continuous, hence the definition of the partition function becomes

$$Z \sim \sum_{i} \int d^3r \int d^3p \ e^{-\beta E_i(\boldsymbol{r}, \boldsymbol{p})}.$$
 (2.2)

The canonical partition function is directly related to the Helmholtz free energy F (often only referred to as the free energy) of the system through a simple exponential

$$Z = e^{-\beta F}. (2.3)$$

Because of the exponentials inherent in the definition of Z, calculating realistic values often results in excessively high numbers. This one of the reasons why it is more useful to work with F than Z itself since inverting Eq. (2.3)  $F = -\ln(Z)/\beta$ , reducing the value of Z through a logarithm.

<sup>1.</sup> In SI units, the value of the Boltzmann constant is given by  $k_{\rm B} \approx 1.380\,649\times 10^{-23}\,{\rm JK^{-1}}$ 

<sup>2.</sup> The reason why there is a  $\sim$  sign in Eq. (2.2) is that technically there is a factor of Planck's constant h in the denominator for each dr dp in the integral measure since this makes the partition function dimensionless and thus consistent with the definition in terms of countable number of states in Eq. (2.1).

#### 2.2 Calculating observables

An observable in statistical mechanics is a quantity that we can both calculate from the statistical theory, and (at least in principle) go out and measure in the real world. In quantum mechanics, observables are restricted to operators that have real expectation-values, as opposed to the complex values it usually deals with. This is to enforce the connection between observables and measurements, which we intuitively understand to always be reducible to a series of real values, i.e. points on a line.

Since we are interested again in ensembles of many particles we are restricting our attention to statistical information about this ensemble. To get this information we need some kind of probability distribution of the particular states of the system. We are imagining that we for each such state (indexed by j) can calculate a real number for the thing (observable) we are interested in measuring. Let's call this observable O. Then O is a statistical variable which takes a particular value  $o_j$  in the state j of the system. If we now let  $P_j$  be the probability distribution, i.e. the probability that the system exists in state j, then we know from probability theory that the expectation value of O is

$$\langle O \rangle = \sum_{j} o_{j} P_{j}. \tag{2.4}$$

The probability distribution  $P_j$  is in the canonical ensamble given by a Maxwell-distribution

$$P_j = e^{-\beta E_j} / Z, \tag{2.5}$$

normalized by the partition function Z. Inserting this we get

$$\langle O \rangle = \sum_{j} \frac{o_{j} e^{-\beta E_{j}}}{Z}.$$
 (2.6)

The observable of specific heat at constant volume  $C_v$  is particularly important in the study of phase transitions since its thermal behavior can be used to classify these transitions into particular categories. The specific heat is the measure of how much energy must be transferred to the system for its temperature to change by an infinitesimal amount

given constant volume of the system. Thus it is defined by the equation  $C_v=(\frac{\partial E}{\partial T})_V$ . In the canonical ensamble, this quantity is calculated by

$$C_v = k_{\rm B} \beta^2 (\langle E^2 \rangle - \langle E \rangle^2) = -k_{\rm B} \beta^2 \Big( 2 \frac{\partial F}{\partial \beta} + \beta \frac{\partial^2 F}{\partial \beta^2} \Big). \tag{2.7}$$

From this form of the specific heat we see that it can be interpreted as a measure of the variance or width of the distribution of energies, and also that it is related to the second derivative of the free energy. Any discontinuity in the second derivative of the free energy, thus implies a discontinuity in the specific heat.

### 2.3 Ginzburg-Landau model

The experimental discovery of superconductivity was a surprise to the scientists at the time. No theoretical model had so far predicted the properties that the experimentalists were measuring. The theoretical models in use at that time predicted a decrease in resistivity as the temperature was lowered, but its sudden disappearance was completely unprecedented and impossible to explain classically in a convincing manner. Thus superconductivity seemed to demand a radically different understanding of how electrons moved inside atomic structures.

#### 2.3.1 Landau Model

Before such an understanding had been developed, Ginzburg and Landau took a shortcut and came up with a theory that could describe the phenomenon of superconductivity without knowing its microscopic origin. In other words they treated superconductivity as a black box and instead of asking what was inside to give the box's output, he used the output to determine a small set of *material parameters* which could then be used to predict how the box would react to a large range of stimuli or conditions. The merit of the theory was that he used symmetry arguments to reduce this set as much as possible.

The Ginzburg-Landau (GL) theory of superconductivity is based on Landau's previous work on a theory of general second order phase-transitions.<sup>3</sup> The approach is given by two ideas. The first is simply

<sup>3. 2</sup>nd order phase-transitions are phase transitions of systems whose free energy has a discontinuous second order derivative at the transition point, but is con-

that the phase transition should be able to be characterized by the appearance of some kind of measurable order that can be described by a function  $\psi$  which we call the *order parameter*. In the liquid water to ice transition, it is the position of the molecules that become ordered in a lattice, in the magnetization of a metal it is the individual spins that become ordered along a particular direction.

The second idea is that at the phase transition, it is the appearance of this order that should dominate the behaviour of the system, to the exclusion of all other effects. Thus the system should be described in terms of the order-parameter and since this is infinitesimally small close to the transition, the free energy<sup>5</sup> can be expanded in a Maclaurinseries with respect to this parameter as

$$F = F_0 + c_1 \psi + c_2 \psi^2 + c_3 \psi^3 + c_4 \psi^4 + c_5 \psi^5 + \dots$$
 (2.8)

The real constants  $F_0$  and  $c_i$  constitute the set of material parameters of the theory and this set can then be reduced by any symmetries that we suspect should be inherent in the underlying theory. For example, if  $\psi$  should represent the order parameter of magnetization of a system of ising spins, which can point either up or down, then the free energy should be invariant of this global choice, i.e. we need to enforce that the free energy be invariant with respect to the transformation  $\psi \mapsto -\psi$ . Then all the constants  $c_j$  for odd j vanish.

In the case of superconductivity the order parameter  $\Psi$  represents the probability amplitude of the collective state of the superfluid of Cooper paired electrons such that  $|\Psi|$  can be interpreted as the density of such electron pairs. Since  $\Psi$  is complex it has to be combined with its complex conjugate  $\Psi^*$  in ways that yield real numbers to produce terms that are valid in the free energy since F itself should be a real number. Furthermore, the phenomenon of superconductivity is

tinuous for lower orders. Since the specific heat is given by the second order derivative, then the specific heat is discontinuous in this case.

<sup>4.</sup> The astute reader might have noticed that this example is a first order phase transition because of the existence of latent heat. Actually first order phase transitions is also able to be described by a modified Landau theory, however we will here focus on the second order kind.

<sup>5.</sup> In this case we are talking about the Helmholtz free energy which is related to the partition function through a logarithm while at the same time the Legendre transformation of the internal energy of the thermodynamic system.

produced as a result of the breaking of U(1) symmetry, so F needs also to be U(1) symmetric, i.e. it has to be invariant under the transformation  $\Psi\mapsto e^{i\phi}\Psi$  for  $\phi\in\mathbb{R}$ . These restrictions result in the free energy

$$F = F_0 - a|\Psi|^2 + b|\Psi|^4, \tag{2.9}$$

when keeping the lowest order terms that produce a phase transition. Thermodynamic equilibrium is reached at the minimum of free energy. This restricts  $b \geq 0$  since negative b yields a free energy with no definite minimum. The minimum is then found by the condition

$$\frac{\partial F}{\partial \Psi^*} = (-a + 2b|\Psi|^2)\Psi = 0, \tag{2.10}$$

which yields the possibilities  $|\Psi|=0$  or  $|\Psi|=\sqrt{a/2b}$ . The first case gives the energy  $F=F_0$ , while the second gives  $F=F_0-a^2/(4b)$ . We see that the second case is energetically favorable, but only exists and is different from the first case when a>0. Furthermore, the second case represents the ordered state since in this case the order-parameter  $|\Psi|\neq 0$ , in the conventional Landau theory.<sup>6</sup>

It is the thermodynamic parameter of temperature that traditionally determines whether a system is in one phase or another. Looking at the free energy in Eq. (2.9), the order parameter  $\Psi$  is the dynamical variable of the theory while the explicit temperature dependence lies in the material parameters a and b. Denoting the the critical temperature where the phase transition happens  $T_c$ , the dimensionless parameter  $t=(T-T_c)/T_c$  is small close to the critical point which means it can be used to expand the temperature dependence of the material parameters such that

$$\begin{split} a(T) &= a_0 + a_1 t + \dots \\ b(T) &= b_0 + b_1 t + \dots \end{split} \tag{2.11}$$

Now we argue for what terms to keep in these expansions. Since a(T) should change sign at t=0 based on the discussion of Eq. (2.10),

<sup>6.</sup> Actually this only represents when Cooper-pairs are forming and the real onset of superconductivity is determined by the point in parameter-space where the gauge-mass becomes non-zero, which is closely related but not exactly the same as where the density of Cooper-pairs becomes non-zero. The real onset of superconductivity is thus more related to when the phase of the wave-function settles on a value.

then we only keep odd terms. Since we need b(T)>0 for the theory to be thermodynamically stable it seems that  $b_0$  is the important term that needs to be larger than any negative contributions from the other terms. Keeping only lowest order terms then the expansions reduce to  $a(T)=a_1(T-T_c)/T_c$  and  $b(T)=b_0$ . Since the ordered state is the solution of the theory when a>0 and this ordered state exists at temperatures  $T< T_c$  then  $a_1<0$  and the final temperature dependence of a becomes  $a(T)=-|a_1|(T-T_c)/T_c$ . From this temperature dependence, it is straightforward to derive critical exponents, the specific heat etc.

#### 2.3.2 Gradient Terms

The simple Landau theory described above is a type of mean field theory in that there is no spatial dependence in the solution, and thus gives a simplified picture that can only be valid far away from any defects or boundaries. This simple approach can be extended to include spatial variation by allowing terms with gradients of the order parameter in the free energy through a gradient expansion

$$F = \int d^3r f(\Psi, \nabla \Psi, \nabla^2 \Psi, \nabla^3 \Psi, \dots). \tag{2.12}$$

Keeping only the lowest order in this expansion that is invariant under U(1) symmetry we get the term  $|\nabla\Psi|^2$ .

Perhaps the single most important phenomenon of superconductivity from a theoretical stand-point is the fact that it expels magnetic fields, hence it is clear that any theory that attempts to explain superconductivity needs to have some way the superconducting order interacts with magnetic fields. The standard way to achieve this is through the recipe of  $\it minimal\ coupling$ , where the vector potential  $\bf A$  times a constant is subtracted with from any momentum in the previously neutral theory. Specifically  $\bf p\mapsto p-q/cA$  where q is the charge of the particle and c is the speed of light. Using this trick, then the free energy density becomes

$$f = f_0 - a|\Psi|^2 + b|\Psi|^4 + K|(\nabla + ig\mathbf{A})\Psi|^2, \tag{2.13}$$

by letting  $g=q/\hbar c$ , which is the form of the free energy in the Ginzburg-Landau theory of conventional s-wave superconductivity. This form

of the gradient resulting from minimal coupling is called the covariant derivative and is defined as

$$D_{\mu} = \partial_{\mu} + igA_{\mu}. \tag{2.14}$$

For unconventional symmetries, the form of the gradient terms can vary substantially from that in Eq. (2.13). Provided that the transition can be described by a single component, i.e. a single function  $\Psi$ , then an unconventional symmetry could still lead to anisotropies in the gradient terms such that K becomes directionally dependent. An example is single-component pairing in a tetragonal crystal: the rotational symmetry in the xy-plane makes this gradient isotropic, but because of the lack of symmetry in the z-direction the gradient terms take the general form

$$K_{1} \sum_{\mu=x,y} |(\partial_{\mu} + igA_{\mu})\Psi)|^{2} + K_{2}|\partial_{z} + igA_{z})\Psi|^{2}. \tag{2.15}$$

Even more complex gradient terms are possible when the order-parameter consists of multiple components, i.e. there are degenerate states that all give significant contributions to the physics at the phase-transition. A particular case of this is when the pairing state is a irreducible representation of the crystal symmetry-group that is multi-dimensional (more about group theory and irreducible representations in Chapter 4). In this case, instead of a single complex function  $\Psi$  describing the order, we need several complex functions  $\eta_i$ .

A chiral p-wave superconductor has a pairing state that is such a two-dimensional irreducible representation which comes from the  $\Gamma_5$  irreducible representation of the tetragonal symmetry group  $D_{4h}$ . This representation consist of two components  $\eta_x$  and  $\eta_y$ , which combine to form the general gradient terms

$$\begin{split} &K_{1}\big[|D_{x}\eta_{x}|^{2}+|D_{y}\eta_{y}|^{2}\big]+K_{2}\big[|D_{x}\eta_{y}|^{2}+|D_{y}\eta_{x}|^{2}\big]\\ &+K_{3}\big[(D_{x}\eta_{x})^{*}(D_{y}\eta_{y})+\text{h.c.}\big]+K_{4}\big[(D_{x}\eta_{y})^{*}D_{y}\eta_{x}+\text{h.c.}\big]\\ &+K_{5}\big[|D_{z}\eta_{x}|^{2}+|D_{z}\eta_{y}|^{2}\big]. \end{split} \tag{2.16}$$

This is the general expression of the gradient terms in the model we have investigated in our work.

CHAPTER 3

# **Field Theory Methods**

In this chapter we will give a short introduction to the use of grassmann variables and complex numbers in the calculation of the fieldintegrals on the statistical-mechanics partition function and how they can be used to transform the expression for the action through the Hubbard-Stratonovich transformation.

A field theoretic expression for the quantum mechanical partition function  $\mathcal Z$  is obtained by using a coherent state basis. A coherent state is the eigen-state of an annihilation operator, thus it produces an eigenvalue when operated on by the annihilation operator. Letting  $\hat H$  be the quantum mechanical Hamiltonian of the system for which we are interested in calculating the partition function,  $\mu$  the the chemical potential and  $\hat N$  is the number operator then the partition function

$$\mathcal{Z} = \text{Tr}(e^{-\beta(\hat{H} - \mu\hat{N})}). \tag{3.1}$$

Inserting a basis of coherent states  $\{|\xi\rangle\}$  when calculating the trace, we obtain a functional integral over the coherent state eigenvalues  $\xi_{\alpha}$  and  $\xi_{\alpha}^{*}$  which substitute the annihilation and creation operators  $c_{\alpha}$  and  $c_{\alpha}^{\dagger}$  respectively. Here  $\alpha$  symbolize the collection of quantum-numbers needed to specify a state. The functional integral then takes the form

$$\mathcal{Z} = \int \mathcal{D}[\xi^* \, \xi] e^{-\int_0^\beta d\tau \, \sum_\alpha [\xi_\alpha^*(\partial_\tau - \mu)\xi_\alpha + H(\xi_\alpha, \xi_\alpha^*)]}. \tag{3.2}$$

The integration variable au is the imaginary time and a au dependence is implicit in the notation such that  $\xi_{lpha}=\xi_{lpha}( au)$ . This path-integral no-

tation is a shorthand for a more involved expression where the imaginary time-dependence of  $\tau$  is split into a collection of time-indexed coherent state eigenfunctions  $\xi_{\alpha,\tau}$  and the integration measure is a product over these indices and the quantum-state indices  $\alpha$ . For further detail we refer to Ref. [68] which we will follow for a large part of this chapter.

#### 3.1 Quadratic Fermionic Field Integrals

Because of the anti-commuting property of the fermion annihilation operators, any coherent state<sup>1</sup> has to have eigenvalues that anti-commute as well. This leads to the partition function written in the convenient basis of the coherent states being constructed with Graßmann numbers as the central variables.

#### 3.1.1 Graßmann algebras

A Graßmann algebra is constructed on a set of generators  $\{\xi_{\alpha}\}$  such that a specific product of the generators  $\xi_{\alpha_1}\xi_{\alpha_2}\cdots\xi_{\alpha_n}$  together with a complex coefficient  $\phi$  constitute a number in the algebra and the generators anti-commute such that  $\xi_{\alpha}\xi_{\beta}=-\xi_{\beta}\xi_{\alpha}$ . On such an algebra, differentiation can be defined such that

$$\frac{\mathrm{d}}{\mathrm{d}\xi_{\alpha_m}}\,\phi\,\xi_{\alpha_1}\cdots\xi_{\alpha_n}=(-1)^m\phi\,\xi_{\alpha_1}\cdots\xi_{\alpha_{m-1}}\xi_{\alpha_{m+1}}\cdots\xi_{\alpha_n},\qquad(3.3)$$

provided the generator  $\xi_{\alpha_m}$  is in the number and 0 otherwise. The factors of (-1) comes from anti-commuting the generator  $\xi_{\alpha_m}$  such that it is next to the differentiation operator. In Graßmann algebra, integration can be (perhaps a little non-intuitively) be defined such that it acts in the same way as differentiation, i.e. generators have to be anti-commuted until they are next to the symbolic differential symbol  $\mathrm{d}\xi_{\alpha}$ , and then

$$\int d\xi \; \xi = 1,\tag{3.4}$$

while

$$\int d\xi \ 1 = 0. \tag{3.5}$$

<sup>1.</sup> An eigenstate of the annihilation operator

On an algebra consisting of 2n generators we define conjugation as a map from the first half of the generators  $\{\xi_{\alpha_i}\}_{i=1}^n$  to the other half  $\{\xi_{\alpha_i}^*\}_{i=1}^n$  and in such a way that when applied to a particular number

$$(\phi \xi_{\alpha} \xi_{\beta})^* = \phi^* \xi_{\beta}^* \xi_{\alpha}^*. \tag{3.6}$$

#### 3.1.2 Nambu Spinor

In the Nambu notation we group spin-dependent Graßmann numbers  $\xi_{\uparrow}$  and  $\xi_{\downarrow}^*$ , which correspond to the annihilation- and creation-operators  $\hat{c}_{\uparrow}^{\dagger}$  and  $\hat{c}_{\downarrow}$ , in a vector called a Nambu spinor

$$\boldsymbol{\xi} = \begin{pmatrix} \xi_{\uparrow} \\ \xi_{\downarrow}^* \end{pmatrix}. \tag{3.7}$$

A sesquilinear form can then be created with this vector and its adjoint such that

$$\boldsymbol{\xi}^{\dagger} S \boldsymbol{\xi} = S_{11} \xi_{\uparrow}^* \xi_{\uparrow} + S_{22} \xi_{\downarrow}^* \xi_{\downarrow} + S_{12} \xi_{\uparrow}^* \xi_{\downarrow}^* + S_{21} \xi_{\uparrow} \xi_{\downarrow}. \tag{3.8}$$

This allows any action that contains spin-dependent terms of the form of the right hand side of Eq. (3.8) to be put on sesquilinear form. Assuming this is the case, then the partition function in the field-integral representation takes the form

$$\mathcal{Z} = \int \mathcal{D}[\xi^* \, \xi] \, e^{-\int_0^\beta d\tau \, \boldsymbol{\xi}_{\gamma}^{\dagger} S_{\gamma \delta} \boldsymbol{\xi}_{\delta}}. \tag{3.9}$$

In this equation, the indices  $\gamma$  and  $\delta$  is an arbitrary collection of quantum numbers needed to specify a state other than spin, for example they could be momentum indices  $\gamma=k, \delta=k'$ , and summation over these repeated indices is implicitly understood.

Splitting the integral over  $\tau$  into M imaginary time-slices and expanding the path integral measure into a product of individual integrals over specific quantum numbered and time-sliced Graßmann variables such that

$$\int \mathcal{D}[\xi^* \, \xi] \, \propto \, \lim_{M \to \infty} \int \prod_{\tau=1}^{M} \prod_{\alpha} \mathrm{d} \xi_{\alpha,\tau}^* \mathrm{d} \xi_{\alpha,\tau}, \tag{3.10}$$

the path-integral in Eq. (3.9) can be evaluated by the Gaussian Graßmann integral identity

$$\int \prod_{i} (\mathrm{d}\xi_{i}^{*} \mathrm{d}\xi_{i}) \ e^{-\xi_{i}^{*} S_{ij} \xi_{j}} = \det S, \tag{3.11}$$

for which a derivation is found in Ref. [68]. This identity holds for any Hermitian matrix S, even if it is not positive definite. The result is then that the partition function in Eq. (3.9) becomes  $\mathcal{Z}=\det S$ . To calculate this determinant one has to consider the matrix S as also a matrix with time-slice indices. This is perhaps most easily accomplished using the Matsubara formalism in which the  $\tau$  dependence is substituted with a dependence on Matsubara frequencies through a Fourier-like transform.

#### 3.1.3 Extended Nambu Spinor

From Eq. (3.8) we see that the Nambu spinor sesquilinear product fails to accommodate terms in a Hamiltonian that mix creation and annihilation operators of differing spins, e.g. a term  $\propto \hat{c}_{\uparrow}^{\dagger}\hat{c}_{\downarrow}$ . In general, a quadratic Hamiltonian can contain any combination of spin-indices of the form  $\hat{c}_{s_1}\hat{c}_{s_2}$ ,  $\hat{c}_{s_1}^{\dagger}\hat{c}_{s_2}$ ,  $\hat{c}_{s_1}^{\dagger}\hat{c}_{s_2}^{\dagger}$  and  $\hat{c}_{s_1}^{\dagger}\hat{c}_{s_2}^{\dagger}$ . This gives in total 16 different combinations and to accommodate them all we thus need a  $4\times 4$  matrix. Exchanging to Graßmann numbers we define the vector

$$\boldsymbol{\xi}_{\gamma} = \begin{pmatrix} \boldsymbol{\xi}_{\gamma,\uparrow}^* \\ \boldsymbol{\xi}_{\gamma,\uparrow} \\ \boldsymbol{\xi}_{\gamma,\downarrow}^* \\ \boldsymbol{\xi}_{\gamma,\downarrow} \end{pmatrix}, \tag{3.12}$$

where all quantum numbers except spin is included in the index  $\gamma$ . Writing the elements of this vector  $(\boldsymbol{\xi}_{\gamma})_i = \tilde{\boldsymbol{\xi}}_{\gamma,i}$  regardless of whether it is a conjugate or not, we can write all quadratic terms of a Hamiltonian on the bilinear form

$$\boldsymbol{\xi}_{\gamma}^{\mathsf{T}} S_{\gamma \delta} \boldsymbol{\xi}_{\delta} = \tilde{\xi}_{\gamma, i} S_{\gamma i; \delta j} \tilde{\xi}_{\delta, j}, \tag{3.13}$$

where  $S_{\gamma\delta}$  is a  $4\times 4$  antisymmetric<sup>2</sup> matrix, and  $S_{\gamma i;\delta j}$  denotes its elements. Let there be n number of different quantum numbers, now

<sup>2.</sup> To see why this matrix can always be said to be antisymmetric lets first simplify the notation and write the bilinear product as  $\tilde{\xi}_i S_{ij} \tilde{\xi}_j$ . Then the matrix S=

including spin. Then there must be 2n different Graßmann generators  $\tilde{\xi}_{\gamma,i}$ . All of these are integrated over in the discrete version of the partition function field integral

$$\mathcal{Z} = \int \mathcal{D}[\xi^* \, \xi] \, e^{-\int_0^\beta d\tau \, \tilde{\xi}_{\gamma,i} S_{\gamma i;\delta j} \tilde{\xi}_{\delta,j}}. \tag{3.14}$$

Even though this superficially looks like the field integral in Eq. (3.9), we now have a bilinear and not a sesquilinear form, and S is now a  $2n \times 2n$  matrix and not an  $n \times n$  matrix. This means that we can not use the integral in Eq. (3.11) to evaluate the integral but instead have to rely on the more general Gaussian Graßmann integral

$$\int \prod_{i} (d\tilde{\xi}_{i}) e^{-\frac{1}{2}\tilde{\xi}_{i}S_{ij}\tilde{\xi}_{j}} = Pf(S), \qquad (3.15)$$

which applies for any antisymmetric matrix S. The right hand side is called the Pfaffian Pf(S) of the matrix S and is defined for any antisymmetric matrix to be given by

$$Pf[S] = \frac{1}{2^n n!} \sum_{P \in S_n} (-1)^P S_{P_1 P_2} \cdots S_{P_{n-1} P_n},$$
 (3.16)

where P is a permutation in the finite group  $S_n$  of all possible permutations of n numbers. This matrix function is related to the determinant by the relation  $Pf(S)^2 = \det(S)$ .

Applying the integral identity in Eq. (3.15) to the partition function<sup>3</sup> in Eq. (3.14) after applying the proper discretization of the imaginary

 $(S+S^{\mathsf{T}})/2+(S-S^{\mathsf{T}})/2$ , such that we can write it as a symmetric matrix  $\mathcal{S}=(S+S^{\mathsf{T}})/2$  and an antisymmetric matrix  $\mathcal{A}=(S-S^{\mathsf{T}})/2$ . Considering only the symmetric part of the bilinear form we get

$$\tilde{\xi}_i \mathcal{S}_{ij} \tilde{x} i_j = -\tilde{\xi}_j \mathcal{S}_{ij} \tilde{\xi}_i = -\tilde{\xi}_i \mathcal{S}_{ji} \tilde{\xi}_j = -\tilde{\xi}_i \mathcal{S}_{ij} \tilde{\xi}_j.$$

Hence  $\tilde{\xi}_i\mathcal{S}_{ij}\tilde{\xi}_j=0$  and all that remains is the antisymmetric bilinear form.

3. In relating the discrete version of Eq. (3.15) to (3.14) we have to make sure that the spinor elements  $\tilde{\xi}_i$  are defined in terms of  $\xi_i$  and  $\xi_i^*$  in such a way as to get a correspondence to the sequence of Graßmann generators  $\mathrm{d}\xi_i^*\mathrm{d}\xi_i$  in the measure to avoid any sign errors. One solution is to set  $\xi_i^* = \tilde{\xi}_{2i-1}$  and  $\xi_i = \tilde{\xi}_{2i}$  as we have done in Eq. (3.12). With this definition, then the measure  $\int \prod_i \mathrm{d}\xi_i^*\mathrm{d}\xi_i$ , which results from the discretized version of the field-integral measure, becomes equal to  $\int \prod_{i=1}^{2n} \mathrm{d}\tilde{\xi}_i$  such that Eq. (3.15) can be directly applied.

time then yields the result

$$\mathcal{Z} = \int \mathcal{D}[\xi^* \, \xi] \, e^{-\int_0^\beta d\tau \, \tilde{\xi}_{\gamma,i} S_{\gamma i;\delta j} \tilde{\xi}_{\delta,j}} = \sqrt{\det(S)}. \tag{3.17}$$

We have chosen the positive result in  $\mathrm{Pf}(S)=\pm\sqrt{\det(S)}$  since the partition function  $\mathcal Z$  needs to be positive on physical grounds. The matrix S to be taken the determinant of on the right hand side of Eq. (3.17) is the full matrix one gets after discretizing the imaginary-time in slices which we usually have done through the Matsubara-frequency formalism.

Now that we know the partition function is given in terms of the determinant of the action-matrix S we can use this information to manipulate the definition of  $\tilde{\xi}_{\gamma,i}$  so that we can still write the action as a sesquilinear form. In particular, switching the position of  $\xi_{\gamma,s}^*$  and  $\xi_{\gamma,s}$  for both spins in the transposed vector on the left of the bilinear form  $\xi_{\gamma}^{\mathsf{T}}S_{\gamma\delta}\xi$ , the transposed vector becomes the adjoint vector. This affects the matrix S by switching two pairs of rows. Denoting the matrix where the rows are switched S' then we have rewritten the bilinear form such that

$$\boldsymbol{\xi}_{\gamma}^{\mathsf{T}} S_{\gamma\delta} \boldsymbol{\xi}_{\delta} = \boldsymbol{\xi}_{\gamma}^{\dagger} S_{\gamma\delta}' \boldsymbol{\xi}_{\delta}. \tag{3.18}$$

Now the integral over the exponent has not changed since all we have done is simply re-ordering its terms. However, since exchange of rows in a determinant at most produces a minus sign and we do this twice we get that  $\det S = \det S'$  and we can write

$$\mathcal{Z} = \int \mathcal{D}[\xi^* \, \xi] \, e^{-\int_0^\beta d\tau \, \boldsymbol{\xi}_{\gamma}^{\dagger} S_{\gamma\delta}' \boldsymbol{\xi}_{\delta}} = \sqrt{\det(S')}. \tag{3.19}$$

where we need to remember that S' is the row switched matrix of an antisymmetric matrix S.

#### 3.2 Matsubara formalism

The Matsubara formalism<sup>4</sup> is a way of handling the imaginary time  $\tau$  dependence of the coherent state eigenvalue fields  $\xi_{\alpha}(\tau)$ , where  $\alpha$  denotes a collection of quantum numbers sufficient to specify a state, without having to go back to the time-sliced path-integral. It also lets

<sup>4.</sup> Named after the Japanese physicist Matsubara, Takeo.

us automatically satisfy the imaginary-time boundary conditions  $\xi_{\alpha}(0)=\zeta\xi_{\alpha}(\beta)$ , where  $\zeta=+1$  for bosons and  $\zeta=-1$  for fermions. Imagining that  $\tau$  is a continuous variable as suggested in the path-integral notation, we define two countable infinite sets of new field-variables through the Fourier-transforms

$$\xi_{\alpha,n} = \frac{1}{\sqrt{\beta}} \int_0^\beta d\tau \, e^{i\omega_n \tau} \xi_\alpha(\tau), \tag{3.20a}$$

$$\xi_{\alpha,n}^* = \frac{1}{\sqrt{\beta}} \int_0^\beta d\tau \, e^{-i\omega_n \tau} \xi_\alpha^*(\tau). \tag{3.20b}$$

The frequencies  $\omega_n$  are called Matsubara frequencies and are defined by  $\omega_n=(2n+1)\pi/\beta$  with  $n\in\mathbb{Z}$  for fermions. For bosons we use instead the notation  $\nu_n$  where  $\nu_n=2n\pi/\beta$ . The inverse relations are given by

$$\xi_{\alpha}(\tau) = \sum_{n \in \mathbb{Z}} e^{-i\omega_n \tau} \xi_{\alpha,n}, \tag{3.21a}$$

$$\xi_{\alpha}^{*}(\tau) = \sum_{n \in \mathbb{Z}} e^{i\omega_{n}\tau} \xi_{\alpha,n}^{*}.$$
 (3.21b)

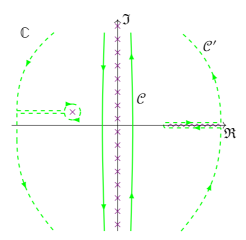
#### 3.2.1 Matsubara sums

When the Matsubara formalism  $\xi_{\alpha,n}$  is used for the field variables in the action of a partition-function field integral we will often need to evaluate infinite sums of Matsubara frequencies of the form

$$\sum_{n} h(if_n), \tag{3.22}$$

where  $f_n$  is either a fermionic or bosonic Matsubara frequency, to evaluate the field integral. A useful strategy in such evaluations is to transform the sum to a complex integral by using reverse residue integration. We consider the complex contour integral along a path  $\mathcal C$  running counter-clockwise around the complex plane infinitesimally close to the imaginary axis as shown in Fig. 3.1. The integrand we consider is given by the product of the summand and the complex continuation of the Fermi-Dirac- or Bose-Einstein-distribution function

$$n_{\zeta}(z) = (e^{\beta z} - \zeta)^{-1}, \quad z \in \mathbb{C}, \tag{3.23}$$



**Figure 3.1:** Integration contour for the Matsubara sum  $\sum_n h(if_n)$ . The contour is marked by a solid line and is imagined to continue to  $\pm i\infty$ . The crosses along the imaginary axis symbolizes the simple poles of the Fermi-Dirac distribution-function. A deformed integration contour is shown with dashed lines that is imagined to cross the real axis at  $\pm \infty$ . This contour then encloses a simple pole on the left and a branch cut on the right belonging to the summand.

depending on whether the Matsubara frequency in the sum is of fermionic  $(\zeta=+1)$  or bosonic  $(\zeta=-1)$  nature. This function has simple poles  $^5$  at  $z=if_n$  and thus integration around the contour results in a sum of residues of the integrand at these poles such that we get

$$\sum_{n} h(if_n) = \frac{\zeta \beta}{2\pi i} \oint_{\mathcal{C}} dz \ h(z) n_{\zeta}(z), \tag{3.24}$$

given that h(z) does not contain any poles at these points. The contour can now be continuously deformed at will as long as it does not cross any singularities which can greatly facilitate the calculation of the integral. The default approach is to see if the integrand vanishes as  $|z| \to \infty$ , in which case it is usually useful to expand the contour as much as possible as illustrated by the deformed contour  $\mathcal{C}'$  in Figure 3.1.

<sup>5.</sup> That the poles are simple, i.e. 1st order, is easily seen by expanding the exponential around  $if_n$  to leading order.

Using the method outlined above, we may calculate the sums

$$\sum_{n\in\mathbb{Z}}\frac{1}{if_n-x}=-\zeta\beta n_\zeta(x), \tag{3.25a}$$

$$\sum_{n\in\mathbb{Z}}\frac{1}{(i\omega_n-x)(i\omega_n-y)}=\frac{\beta}{x-y}\bigg(\frac{1}{1+e^{\beta x}}-\frac{1}{1+e^{\beta y}}\bigg),\quad (3.25b)$$

$$\sum_{n\in\mathbb{Z}}\ln[\beta(i\omega_n+x)]=\ln(1+e^{-\beta x}). \tag{3.25c}$$

#### 3.3 Hubbard-Stratonovich transformation

The Hubbard-Stratonovich transformation is a transformation in the fields of a theory where a new complex field is introduced in order to convert a term that is square and thus non-linear in an existing field variable, into a linear term in this variable that is coupled to the new field. This is particularly useful when the existing field is Fermionic and thus a Graßmann variable since it makes it possible to consider low energy exitations of the theory using e.g. a saddle-point approximation. It is however important to point out that the transformation itself is not in any way approximative, but is an exact transformation that maintains all information in the original theory.

In technical terms, the Hubbard-Stratonovich transformation can be viewed simply as the solution of a complex multivariate integral. Let A have a strictly positive Hermitian part and  $\boldsymbol{J}$  be a vector of coefficients that could contain Graßmann variables or complex variables. Then

$$e^{\mathbf{J}^{\dagger}A\mathbf{J}} = \det A^{-1} \int_{\mathbb{C}} \prod_{i} \left[ \frac{\mathrm{d}z_{i}^{*} \mathrm{d}z_{i}}{2\pi i} \right] e^{-(\mathbf{z}^{\dagger}A^{-1}\mathbf{z} + \mathbf{z}^{\dagger}\mathbf{J} + \mathbf{J}^{\dagger}\mathbf{z})}, \qquad (3.26)$$

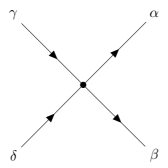
exchanges a quadratic term in  ${\bf J}$  with an integration over the complex  ${\bf z}$  variable. Since  ${\bf J}$  usually represents some field in a field theory, the new  ${\bf z}$  is called the auxiliary field or the conjugate field because of its linear coupling to  ${\bf J}$ . In the less general case that A is an Hermitian matrix, this formula is proved simply by completing the square, then diagonalizing A by a unitary transformation and calculating the resulting integrals by the formula  $\int_{\Gamma} {\rm d}z^* {\rm d}z \, e^{-azz^*} = 2\pi i/a$ .

From Eq. (3.26) we see that what we have to do to perform the Hubbard-Stratonovich transformation is first to make a choice for what to interpret as part of the matrix A and what to interpret as part of J.

We then have to check that this definition of A leads to its Hermitian part having only positive eigenvalues. Finally we need to know an analytical expression for its inverse. It is usually the first step that is the most difficult since this dictates the low energy excitation a subsequent saddle point approximation or a stationary phase approximation will produce. Typically we are interested in transforming a Fermionic interaction potential of the form

$$V = \frac{1}{2} \sum_{\alpha\beta\gamma\delta} V_{\alpha\beta\gamma\delta} \xi_{\alpha}^* \xi_{\beta}^* \xi_{\delta} \xi_{\gamma}, \tag{3.27}$$

where  $\xi_{\alpha}$  are Graßmann variables, which can be sketched in the way of single-vertex diagram in Figure 3.2. The HS-transformation is clas-



**Figure 3.2:** Generic two-body interaction.

sified into being done in a specific *channel* depending on which pair of Graßmann variables are considered to be part of  ${\pmb J}$  and consequently  ${\pmb J}^\dagger.$  The direct channel<sup>6</sup> is given by the identification  $J_i \sim \xi_\alpha^* \xi_\gamma$ , the Cooper channel<sup>7</sup> is defined by the identification  $J_i \sim \xi_\delta \xi_\gamma$  while the exchange channel is given by the identification  $J_i \sim \xi_\alpha^* \xi_\delta$ . Depending on exactly how  ${\pmb J}$  is chosen, the Gaussian integral in Eq. (3.26) might have to be modified. For example in the case of the direct- and exchange-channel, the exponential argument on the left side will have the form  ${\pmb J}^{\rm T} A {\pmb J}$  which necessitates the Gaussian integral identity

$$e^{-\frac{1}{2}\boldsymbol{J}^{\mathsf{T}}A\boldsymbol{J}} = \sqrt{\det A^{-1}} \int_{\mathbb{R}} \prod_{i} \left[ \frac{\mathrm{d}x_{i}}{\sqrt{2\pi}} \right] e^{-\frac{1}{2}\boldsymbol{x}^{\mathsf{T}}A^{-1}\boldsymbol{x} - i\boldsymbol{J}^{\mathsf{T}}\boldsymbol{x}}, \quad (3.28)$$

<sup>6.</sup> Also known as the density-density channel.

<sup>7.</sup> Also known as the particle-particle channel.

where the auxiliary field x now is a real conjugate field.

## 3.3.1 Transformation in symmetry channels

In the Cooper-channel of the Hubbard-Stratonovich transformation, the complex field z is conjugate to some combination of pairs of annihilation operators  $\hat{c}_{\delta}\hat{c}_{\gamma}$  (or their corresponding Graßmann variables). The symmetry of the specific combination in turn then determines the symmetry of any low energy field theory obtained through a subsequent stationary phase approximation. By diagonalizing the interaction potential  $\hat{V}$  into its particular irreducible representations as we did in Section 4.8 then a Hubbard-Stratonovich transformation in a specific symmetry channel is done by identifying J with irrep. basis functions.

Let's take the case of a BCS theory of superconductivity where the interaction can be written in terms of basis functions  $d_{s_1s_2}^{(b),m}({\pmb k})$  such that the diagonalized interaction takes the form

$$\hat{V} = \sum d_{s_1 s_2}^{(b),m}(\mathbf{k})^* v_{(b)} d_{s_1' s_2'}^{(b),m}(\mathbf{k}') c_{\frac{\mathbf{q}}{2} + \mathbf{k} s_1}^{\dagger} c_{\frac{\mathbf{q}}{2} - \mathbf{k} s_2}^{\dagger} c_{\frac{\mathbf{q}}{2} - \mathbf{k}' s_2'}^{\phantom{\dagger}} c_{\frac{\mathbf{q}}{2} + \mathbf{k}' s_1'}^{\phantom{\dagger}}, \tag{3.29}$$

where  $\sum$  indicates the sum over the indices,  ${\pmb k}, {\pmb k}', {\pmb q}, s_1, s_2, s_1', s_2', b$  and m. Here b specifies the irreducible representation while m enumerates the representation basis. Identifying

$$\hat{J}_{\mathbf{q}}^{(b_m)} = \sum_{\mathbf{k}s_1 s_2} d_{s_1 s_2}^{(b), m}(\mathbf{k}) \hat{c}_{\frac{\mathbf{q}}{2} - \mathbf{k}, s_1} \hat{c}_{\frac{\mathbf{q}}{2} + \mathbf{k}, s_2}, \tag{3.30}$$

the interaction potential is simply written

$$\hat{V} = \sum_{q,b,m} \hat{J}_{q}^{(b_{m})\dagger} v^{(b)} \hat{J}_{q}^{(b_{m})}.$$
(3.31)

In the path-integral representation of the partition function, the annihilation operators become Graßmann variables which we denote by writing J instead of  $\hat{J}$  such that the contribution from the interaction potential results in the exponential

$$\mathcal{Z}_{I} = e^{-\int_{0}^{\beta} d\tau \sum_{q,b,m} J_{q}^{(b_{m})\dagger} v^{(b)} J_{q}^{(b_{m})}}.$$
 (3.32)

Now it is straightforward to use the Hubbard-Stratonovich formula

$$e^{\int_0^\beta \mathrm{d}\tau \sum_{ij} J_i^* A_{ij} J_j} = \int \mathcal{D}[\eta_i^* \eta_i] e^{-\int_0^\beta \mathrm{d}\tau \left(\eta_i^* A_{ij}^{-1} \eta_j + J_i^* \eta_i + J_i \eta_i^*\right)}, \quad (3.33)$$

which is a path integral version of Eq. (3.26), to transform each pair of irreducible representation basis vectors to individual conjugate fields. In the notation of Eq. (3.33) implicit summation over repeated indices is used and each index i is a collection  $i=(b,m,\boldsymbol{q})$  of indices. Comparing Eq. (3.33) and (3.32) we gather that

$$A_{ij} = A_{b,m,q;b',m',q'} = -\delta_{qq'}\delta_{mm'}\delta_{bb'}v^{(b)},$$
(3.34)

which is trivially Hermetic and positive definite provided  $v^{(b)} < 0$ . In this case we say that the irreducible representation b is an attractive channel. A is in this case also trivially invertible with  $A_{ij}^{-1} = -\delta_{ij}/v^{(b)}$ . Writing out all the indices we finally arrive at the Hubbard-Stratonovich transformation of the interaction potential in individually attractive symmetry channels

$$\mathcal{Z}_{I} = \int \mathcal{D}[\eta_{\mathbf{q}}^{(b_{m})*} \eta_{\mathbf{q}}^{(b_{m})}] e^{\int_{0}^{\beta} d\tau \sum_{\mathbf{q}bm} \left[ \frac{|\eta_{\mathbf{q}}^{(b_{m})}|}{v^{(b)}} - \left( J_{\mathbf{q}}^{(b_{m})*} \eta_{\mathbf{q}}^{(b_{m})} + J_{\mathbf{q}}^{(b_{m})} \eta_{\mathbf{q}}^{(b_{m})} \right) \right]}, \tag{3.35}$$

where

$$J_{\mathbf{q}}^{(b_m)} = \sum_{\mathbf{k} s_1 s_2} d_{s_1 s_2}^{(b), m}(\mathbf{k}) \xi_{\frac{\mathbf{q}}{2} - \mathbf{k}, s_1} \xi_{\frac{\mathbf{q}}{2} + \mathbf{k}, s_2}$$
(3.36)

in terms of Graßmann variables  $\xi$ . We note that this derivation does not assume either odd or even basis functions for the irreducible representations and works just as well for either.

# 3.4 Field theory approximations

# 3.4.1 Stationary phase and the one loop expansion

Let Z be the partition function

$$Z(l) = \int \mathcal{D}[\eta_{\alpha}^* \, \eta_{\alpha}] \, e^{-lS(\eta_{\alpha}^*, \eta_{\alpha})}, \tag{3.37}$$

given in terms of bosonic fields  $\eta_{\alpha}^*$ ,  $\eta_{\alpha}$  and where l is some large parameter  $l \gg 1$ . In what is called the stationary phase approximation<sup>8</sup>

<sup>8.</sup> The name is misleading in the case of the many-particle partition function since we do not have a strict phase in the exponent but in general a complex function because  $\eta_{\alpha} \in \mathbb{C}$ . Technically it is the method of steepest descent in the case of an exponent with stationary points that is used in this case.

of a bosonic field integral we create an expansion of the free energy around the field configuration  $\{\eta_{\alpha}^c\}$  where the action S is stationary and a minimum. This configuration is the main contribution to the integral in Eq. (3.37) since it provides the maximum of the exponent, and determines the leading order asymptotic behavior as  $l\to\infty$ . It also corresponds in a sense to a classical solution and gives in the case of the feynman path integral for the evolution operator of a single particle in an external potential, the classical Euler-Lagrange equations. The configuration is found in the path-integral notation by varying the fields in the action such that

$$\frac{\delta S}{\delta \eta_{\alpha}} = 0 \quad \wedge \quad \frac{\delta S}{\delta \eta_{\alpha}^*} = 0. \tag{3.38}$$

As an example, the Hubbard-model can be expressed by a bosonic field integral through a Hubbard-Stratonovich transformation. The stationary field configuration  $\{\eta^c_q(\tau)\}$  is in this case given by a imaginary-time- and spatially-independent field-configuration  $\eta^c$ . Assuming such a solution, the action reduces to

$$S(\eta^{c*}, \eta^c) = \frac{\beta N}{U} |\eta^c|^2 - \sum_{q} \ln \left[ \left( 1 + e^{\beta E_q} \right) \left( 1 + e^{-\beta E_q} \right) \right], \quad (3.39)$$

where  $E_{\boldsymbol{q}} = \sqrt{(\epsilon(\boldsymbol{q}) - \mu)^2 + |\eta^c|^2}$ , N is the number of hopping sites,  $\epsilon(\boldsymbol{q})$  is the Fourier transformed kinetic hopping energy,  $\mu$  is the chemical potential, and U is the repulsive Hubbard interaction strength. The stationary action condition in Eq. (3.38) then yields the equation

$$|\eta^c| \left(\frac{1}{U} - \frac{1}{N} \sum_{\mathbf{q}} \frac{1}{2E_{\mathbf{q}}} \tanh \frac{\beta E_{\mathbf{q}}}{2}\right) = 0, \tag{3.40}$$

which has two solutions: one given by  $\eta^c=0$  and one given by setting the terms in the parenthesis equal to 0. The last non-trivial solution is a form of the BCS solution and represents the order-parameter in an s-wave superconductor. Because of the parameter  $\beta$ , this solution will be temperature dependent and disappears at some critical temperature at which the two solutions for  $\eta_c$  converge.

The stationary phase expansion is then the expansion resulting from expanding the action around the stationary solution. Setting  $\tilde{\eta}_{\alpha}=$ 

 $\sqrt{l}(\eta_{\alpha}-\eta_{\alpha}^{c})$  we get in general the expansion

$$\begin{split} S(\tilde{\eta}_{\alpha},\tilde{\eta}_{\alpha}^{*}) &= S_{c} + \frac{1}{l} \bigg[ \frac{1}{2!} \frac{\delta^{2}S}{\delta\eta_{\alpha}\delta\eta_{\beta}} \Big|_{\eta_{c}} \tilde{\eta}_{\alpha}\tilde{\eta}_{\beta} \\ &+ \frac{1}{2!} \frac{\delta^{2}S}{\delta\eta_{\alpha}^{*}\delta\eta_{\beta}^{*}} \Big|_{\eta_{c}} \tilde{\eta}_{\alpha}^{*}\tilde{\eta}_{\beta}^{*} + \frac{\delta^{2}S}{\delta\eta_{\alpha}\delta\eta_{\beta}^{*}} \Big|_{\eta_{c}} \tilde{\eta}_{\alpha}\tilde{\eta}_{\beta}^{*} \bigg] + \mathcal{O}\bigg( \frac{1}{l\sqrt{l}} \bigg) \\ &= \sum_{n_{1}+n_{2} \geq 2} \frac{1}{n_{1}!n_{2}!} \frac{\delta^{n_{1}+n_{2}}S}{\delta\tilde{\eta}_{\alpha}^{n_{1}}\delta\tilde{\eta}_{\beta}^{*}} \tilde{\eta}_{\alpha}^{n_{1}}\tilde{\eta}_{\beta}^{*} \frac{n_{2}}{\eta_{c}} \Big|_{\eta_{c}} \frac{1}{l^{(n_{1}+n_{2})/2}}. \end{split} \tag{3.41}$$

We have used implicit summation over repeated indices and on the last line we have assumed that new indices should be introduced and summed for each  $n_1$  and  $n_2$ . This is simply the multivariate Taylor expansion around  $\eta_c$  where we have treated  $\eta$  and  $\eta^*$  as independent variables.

A simpler expansion can be found when the bosonic fields correspond to order parameters in a system close to a phase transition. In this case we can assume the fields  $\eta_\alpha$  to be small in general such that S simply can be expanded about 0. If the system is fermionic, then the bosonic field integral of the order-parameters will have resulted from a Hubbard-Stratonovich transformation, in which case we will have a contribution to the integral of the form

$$\sqrt{\det G^{-1}(\eta_{\alpha},\eta_{\alpha}^{*})} = e^{\frac{1}{2}\operatorname{Tr}\ln G^{-1}}, \tag{3.42}$$

where  $G^{-1}$  is the result of the integration of a quadratic fermionic action, and will in general depend on the auxiliary bosonic fields  $\eta_{\alpha}$  and  $\eta_{\alpha}^{*}$  in a linear way. Then the matrix  $G^{-1}$  can be decomposed in a matrix  $G_{0}^{-1}$  that results purely from fermionic integration, and a matrix  $\phi$  that is dependent on the auxiliary fields  $\eta_{\alpha}$  and importantly vanishes as  $\eta_{\alpha} \to 0$ , such that  $G^{-1} = G_{0}^{-1} + \phi$ . When the system is close the the phase transition such that the auxiliary fields are small, then this allows for the expansion of the logarithm in Eq. (3.42) such that

$$\begin{split} \frac{1}{2} \operatorname{Tr} \ln(G_0^{-1} + \phi) &= \frac{1}{2} \Big( \operatorname{Tr} \ln G_0^{-1} + \operatorname{Tr} \ln(1 + G_0 \phi) \Big) \\ &= \frac{1}{2} \Big( \operatorname{Tr} \ln G_0^{-1} - \sum_{n=1}^{\infty} \frac{\operatorname{Tr} (-G_0 \phi)^n}{n} \Big). \end{split} \tag{3.43}$$

This is known as the one-loop expansion since in terms of perturbation theory,  $G_0$  is the fermionic propagator such that the sum in the last line of Eq. (3.43) corresponds to a series of propagators and interactions that are connected in a closed loop by the trace.

## 3.4.2 Gradient expansion

The gradient expansion rests on the assumption that the fields are sufficiently smooth such that progressively higher order derivatives with respect to the field parameters, are progressively smaller, i.e. we assume

$$l_{\alpha}^{n}|\partial_{\alpha}^{n}\eta_{\alpha}|\gg l_{\alpha}^{n+1}|\partial_{\alpha}^{n+1}\eta_{\alpha}|, \tag{3.44}$$

where  $l_{\alpha}$  is some appropriate length scale such as to make  $l_{\alpha}\partial_{\alpha}$  dimensionless. In practice this usually means that given a momentum-dependent action density,  $S(\eta_{{\bf q},\alpha}^*,\eta_{{\bf q},\alpha};{\bf q})$ , we assume  $q_i$  small compared to the lattice spacing  $^9$   $a_i$ . Then we expand the explicit  ${\bf q}$  dependence in the action density in the small parameters  $a_iq_i$  as a Maclaurin series. In the following notation, we will assume the length parameter  $a^i$  is present in the notation  $q^i$  where appropriate. The momentum dependence of the fields themselves should not be expanded as our goal is to have terms of the form  $(q^i)^n(\tilde{\eta}_{{\bf q},\alpha})^m$ . If we then let partial momentum-derivatives only act on the explicite momentum-dependence in S, and neglecting the field dependence in the notation for S, the expanded action can be written as the series

$$S(\mathbf{q}) = S(0) + \frac{\partial S}{\partial q^i}(0) q^i + \frac{1}{2!} \frac{\partial^2 S}{\partial q^i \partial q^j}(0) q^i q^j + \mathcal{O}(q^3).$$
 (3.45)

Usually, the linear term cancels by symmetry of the underlying lattice. In general it is smart to here check for terms that cancel by considering any internal momentum sums that may be included in the coefficients.

Terms with products between fields  $\eta_{{m q},\alpha}$  and  $q^i$  lead to gradients of the spatially dependent fields  $\eta_{{m R},\alpha}$ , which is why the expansion in Eq. (3.45) can be called a gradient expansion. The spatially dependent fields are defined as the coefficients in the inverse Fourier transform

$$\eta_{\mathbf{q},\alpha} = \frac{1}{\sqrt{N}} \sum_{\mathbf{R}} e^{i\mathbf{q}\cdot\mathbf{R}} \eta_{\alpha}(\mathbf{R}), \tag{3.46}$$

<sup>9.</sup> This could be the spacing between hopping sites, i.e. stationary ions in an electron model.

where N is the number of terms in the sums  $\sum_{q}$  and  $\sum_{R}$ . One way of now obtaining gradients of the spatial fields is to realize that since  $q^{i}$  is small we can set

$$q^{i} \approx \sin(q^{i}) = \frac{1}{2i} \left( e^{iq^{i}} - e^{-iq^{i}} \right). \tag{3.47}$$

With this identification, all the momentum dependence in  $S(\boldsymbol{q})$  exists as phases such that the sum  $\sum_{\boldsymbol{q}} S(\boldsymbol{q})$  results in a series of Kroneckerdelta functions which we evaluate by the  $\sum_{\boldsymbol{R}}$  sums coming from the inverse Fourier transforms in Eq. (3.46). Grouping terms of displaced spatial fields, we can identify derivatives, such that

$$\eta_{\alpha}(\mathbf{R} + a\hat{e}_i) - \eta_{\alpha}(\mathbf{R}) \approx a \frac{\partial}{\partial R^i} \eta_{\alpha}(\mathbf{R}).$$
(3.48)

These identifications are justified by going to the continuum limit where  $a \rightarrow 0$ .

As an example, consider the sum

$$S = \sum_{\mathbf{q}} K_{\alpha\beta\,ij} q^i q^j \eta_{\mathbf{q},\alpha}^* \eta_{\mathbf{q},\beta}, \tag{3.49}$$

where there is an implicit summation over repeated indices  $i, j, \alpha$  and  $\beta$ . Fourier transforming the fields according to Eq. (3.46) and writing  $q^i$  as  $\sin q^i$ , we can group terms such that

$$\begin{split} S &= \frac{1}{N} \sum_{\boldsymbol{R}_{1} \boldsymbol{R}_{2}} K_{\alpha \beta \, ij} \eta_{\alpha}(\boldsymbol{R}_{1})^{*} \eta_{\beta}(\boldsymbol{R}_{2}) \frac{1}{4} \sum_{\boldsymbol{q}} \left[ e^{i \boldsymbol{q} \cdot (\boldsymbol{R}_{2} - \boldsymbol{R}_{1} + a \hat{e}_{i} - a \hat{e}_{j})} \right. \\ &+ e^{i \boldsymbol{q} \cdot (\boldsymbol{R}_{2} - \boldsymbol{R}_{1} + a \hat{e}_{j} - a \hat{e}_{i})} - e^{i \boldsymbol{q} \cdot (\boldsymbol{R}_{2} - \boldsymbol{R}_{1} + a \hat{e}_{i} + a \hat{e}_{j})} - e^{i \boldsymbol{q} \cdot (\boldsymbol{R}_{2} - \boldsymbol{R}_{1} - a \hat{e}_{i} - a \hat{e}_{j})} \right] \\ &= \sum_{\boldsymbol{R}} K_{\alpha \beta \, ij} a^{2} \frac{\partial}{\partial R_{i}} \eta_{\alpha}(\boldsymbol{R})^{*} \frac{\partial}{\partial R_{j}} \eta_{\beta}(\boldsymbol{R}). \end{split} \tag{3.50}$$

A more conventional way of converting to gradients, is to use integration by parts. Then the product rule of partial derivatives is used to obtain

$$q^{j}\eta_{\alpha}(\mathbf{R})e^{i\mathbf{q}\cdot\mathbf{R}} = i\frac{\partial\eta_{\alpha}}{\partial R^{i}}e^{i\mathbf{q}\cdot\mathbf{R}} - i\frac{\partial}{\partial R^{i}}\Big[\eta_{\alpha}(\mathbf{R})e^{i\mathbf{q}\cdot\mathbf{R}}\Big]. \tag{3.51}$$

Summing on both sides and arguing that the boundary term vanishes because  $\eta_{\alpha}(\mathbf{R}) \to 0$  as  $R^i \to \infty$ , then

$$\sum_{\boldsymbol{R}} q^{j} \eta_{\alpha}(\boldsymbol{R}) e^{i\boldsymbol{q}\cdot\boldsymbol{R}} = i \sum_{\boldsymbol{R}} e^{i\boldsymbol{q}\cdot\boldsymbol{R}} \nabla_{i} \eta_{\alpha}(\boldsymbol{R}). \tag{3.52}$$



# **Group Theory**

In this chapter we will introduce the minimum mathematical framework needed to understand the phrase "This free energy is the  $\Gamma_{5u}^-$  irreducible representation of the  $D_{4h}$  symmetry group".

A few words about notation. We will use the semicolon ';' in equations as notation for the words 'such that', e.g. when defining sets. A colon with a trailing space ': ' is used when defining maps where the symbol representing the mapping itself should be on the left while the the sets being related or how the elements of the sets are related is on the right of the colon. The colon ':' is also used as a shortcut for the words 'applied through its representation to' for when group elements are applied to vectors, where the correct representation to use for this application should be implicitly understood.

The material in this section is based on the material covered in Refs. [69] and [68], specified for the use in quantum mechanical theories of unconventional superconducting states.

## 4.1 Irreducible representations

To know what an irreducible representation (IR) is, let's start with what we mean by a reducible representation.

**Def. 4.1.** A matrix representation is **reducible** if there exists a non-trivial invariant subspace of the vector space of the representation.

The intuition is then that the vector space of the representation is reducible if a "smaller" representation is contained within it. Since there is a smaller vector space within the vector space of the original representation and this vector space is invariant, it is possible to define another representation on this smaller vector space, i.e. *reduce* the original representation. We have now used the word "invariant" a couple of times, so let's define what it means more precisely.

**Def. 4.2.** Let D(g) be a representation of the group G on the vector space V such that  $D(g) \colon V \to V$ . Then a subspace  $U \subseteq V$  is **invariant** if

$$\forall g \in G \quad u \in U \implies D(g)u \in U. \tag{4.1}$$

In other words: a vector space is invariant if it is not possible for any vector in it to escape using a representation of any group element. All representations applied to any vector in the invariant subspace must necessarily land in that same subspace from which it started.

## 4.2 BCS Hilbert Space

We define the BCS Hilbert space as the Hilbert space upon which BCS-type potentials operate. Specifically this is a reduced form of the two-state fermionic product Hilbert space  $\mathcal{H}_2=\mathcal{H}\otimes\mathcal{H}$  where  $\mathcal{H}=\mathrm{span}\{|\pmb{k},s\rangle\}$  and we only consider states that have opposite momentum. Thus this Hilbert space is given by

$$\mathcal{B} = \mathrm{span}\{|\boldsymbol{k},s_1\rangle|-\boldsymbol{k},s_2\rangle\}, \tag{4.2}$$

and the identity operator in this space can be written

$$\hat{1} = \sum_{\boldsymbol{k} s_1 s_2} |\boldsymbol{k}, s_1\rangle |-\boldsymbol{k}, s_2\rangle \langle -\boldsymbol{k}, s_2| \langle \boldsymbol{k}, s_1|.$$
(4.3)

Acting on the arbitrary vector  $|v\rangle \in \mathcal{B}$  with this identity operator, we find that in terms of this basis it can be written

$$|v\rangle = \sum_{\boldsymbol{k}\,s_1s_2} v_{s_1s_2}(\boldsymbol{k}) |\boldsymbol{k},s_1\rangle |-\boldsymbol{k},s_2\rangle, \tag{4.4}$$

where

$$v_{s_1s_2}(\boldsymbol{k}) = \langle -\boldsymbol{k}, s_2 | \langle \boldsymbol{k}, s_1 \mid v \rangle. \tag{4.5}$$

The indices  $s_1$  and  $s_2$  can take on only two values each, namely  $s_1, s_2 \in \{\uparrow, \downarrow\}$ . In total there are thus 4 different realizations of pairs,  $s_1s_2$  e.g.  $\uparrow\uparrow$  for  $v_{s_1s_2}(\boldsymbol{k})$ . Putting these different realizations of  $v_{s_1s_2}(\boldsymbol{k})$  as elements in a  $2\times 2$  matrix we get

$$v_{s_1 s_2}(\mathbf{k}) = \begin{pmatrix} v_{\uparrow \uparrow}(\mathbf{k}) & v_{\uparrow \downarrow}(\mathbf{k}) \\ v_{\downarrow \uparrow}(\mathbf{k}) & v_{\downarrow \downarrow}(\mathbf{k}) \end{pmatrix}. \tag{4.6}$$

Any  $2\times 2$  matrix can be written in the conventional basis of the 4 Pauli matrices  $\sigma^0=\mathbb{1}_{2\times 2}$ ,  $\sigma^x$ ,  $\sigma^y$ , and  $\sigma^z$ . This means that we could write the matrix in Eq. (4.6)

$$v_{s_1 s_2}(\mathbf{k}) = v_{\mathbf{k}}^0 \sigma_{s_1 s_2}^0 + v_{\mathbf{k}}^i \sigma_{s_1 s_2}^i. \tag{4.7}$$

It is however conventional to factor out a Pauli matrix  $i\sigma^y$  to the right in the expansion since this results in nice transformation properties of the coefficients as we shall see. With the spin-indices expanded in this basis it is conventional to let the function of  ${\bf k}$  that is in front of  $\sigma^0$  be called  $\psi_{\bf k}$ . The three others are conventionally denoted  $d_{{\bf k},i}$  which are components of what we call the  ${\bf d}$ -vector. Expanded in this conventional basis then  $v_{s_1s_2}({\bf k})$  takes the form

$$v_{s_1s_2}(\pmb{k}) = (\psi_{\pmb{k}}\sigma^0_{s_1s'} + d_{\pmb{k},i}\sigma^i_{s_1s'})i\sigma^y_{s's_2}, \eqno(4.8)$$

and finally the state  $|v\rangle$  can be written

$$|v\rangle = \sum_{\boldsymbol{k} \, s_1 s_2} [(\psi_{\boldsymbol{k}} \sigma^0 + \boldsymbol{d}_{\boldsymbol{k}} \cdot \boldsymbol{\sigma}) i \sigma^y]_{s_1 s_2} |\boldsymbol{k}, s_1\rangle |-\boldsymbol{k}, s_2\rangle. \tag{4.9}$$

Going one step back and writing out the different combinations of  $s_1s_2$  in  $v_{s_1s_2}(\boldsymbol{k})$  as a matrix like we did in Eq. (4.6), but now multiplying out the Pauli matrices in Eq. (4.8) we get

$$\begin{pmatrix} v_{\uparrow\uparrow}(\mathbf{k}) & v_{\uparrow\downarrow}(\mathbf{k}) \\ v_{\downarrow\uparrow}(\mathbf{k}) & v_{\downarrow\downarrow}(\mathbf{k}) \end{pmatrix} = \begin{pmatrix} -d_{\mathbf{k},x} + id_{\mathbf{k},y} & \psi_{\mathbf{k}} + d_{\mathbf{k},z} \\ -\psi_{\mathbf{k}} + d_{\mathbf{k},z} & d_{\mathbf{k},x} + id_{\mathbf{k},y} \end{pmatrix}.$$
(4.10)

<sup>1.</sup> Note that a state in  $\mathcal{B}$  that is described by a d-vector does not necessarily mean that it has d-wave symmetry, which is a symmetry of its k-space argument. It tells us that the state is a spin-triplet state.

This set of linear relations is easily inverted to yield

$$\psi_{\mathbf{k}} = \frac{1}{2}(v_{\uparrow\downarrow}(\mathbf{k}) - v_{\downarrow\uparrow}(\mathbf{k})) \tag{4.11}$$

$$d_{\boldsymbol{k},x} = \frac{1}{2}(v_{\downarrow\downarrow}(\boldsymbol{k}) - v_{\uparrow\uparrow}(\boldsymbol{k})) \tag{4.12}$$

$$d_{\boldsymbol{k},y} = -\frac{i}{2}(v_{\uparrow\uparrow}(\boldsymbol{k}) + v_{\downarrow\downarrow}(\boldsymbol{k})) \tag{4.13}$$

$$d_{\boldsymbol{k},z} = \frac{1}{2}(v_{\uparrow\downarrow}(\boldsymbol{k}) + v_{\downarrow\uparrow}(\boldsymbol{k})). \tag{4.14}$$

Since the space  $\mathcal{B}$  is fermionic we have the symmetry<sup>2</sup>

$$|\mathbf{k}, s_1\rangle|-\mathbf{k}, s_2\rangle = -|-\mathbf{k}, s_2\rangle|\mathbf{k}, s_1\rangle.$$
 (4.15)

Using this symmetry transformation on the basis vectors in the expansion of  $|v\rangle$  in Eq. (4.4), then renaming indices and finally equating coefficients term by term, we see that for the coefficients of  $|v\rangle$ , this symmetry takes the form

$$v_{s_1s_2}({\pmb k}) = -v_{s_2s_1}(-{\pmb k}). \tag{4.16}$$

## 4.3 Application of group elements

When we are talking about applying some symmetry transformation to a state, this is synonymous with applying a group element to a vector. Even more specifically, the 'applying' part means that we have some natural representation of the group on the vector space we have defined states on, and we are using the linear transformation of the representation of the group element to act on the state vector. In this thesis we will use the notation  $g:|\psi\rangle$  to refer to this procedure.

Let g be an arbitrary group element in the symmetry group G and D be a representation of G on the d-dimensional vector space V. Let V have a basis  $\{\boldsymbol{b}_i\}_{i=1}^d$ . The application of a group element to a basis vector is then defined as

$$g: \boldsymbol{b}_i = \sum_j \boldsymbol{b}_j D_{ji}(g). \tag{4.17}$$

<sup>2.</sup> We are here assuming that the state is even in (time) frequency. It is also possible to have superconducting states that are odd in frequency [70], however we will not treat that possibility here.

The application of a group element to any vector in V then is calculated by expanding the vector in the basis and applying the representation D to each basis vector separately as a linear transformation:

$$g: \boldsymbol{v} = \sum_{i} v_{i}g: \boldsymbol{b}_{i}. \tag{4.18}$$

#### 4.3.1 Active vector transformation

As g: has been defined in Eq. (4.17) it is defined in a passive perspective where the transformation is happening to the basis vectors. It is often useful and sometimes more intuitive to consider the application of g to a vector  $\mathbf{v}$  in the basis  $\{\mathbf{b}_i\}$  as an application not on the vectors themselves but on the expansion coefficients  $v_i$  of  $\mathbf{v}$  in the basis. This is the active view of the transformation. Inserting Eq. (4.17) into Eq. (4.18) we get

$$g: \boldsymbol{v} = \sum_{i} v_{i} \sum_{j} \boldsymbol{b}_{j} D_{ji}(g) = \sum_{i} v'_{i} \boldsymbol{b}_{i}, \tag{4.19}$$

where we have defined the transformed coefficients

$$v_i' = \sum_j D_{ij}(g)v_j. \tag{4.20}$$

From this calculation we see that we can consider the application of g as a transformation of the coefficients of the vector as

$$g: v_i = \sum_{j} D_{ij}(g) v_j. (4.21)$$

This defines the active transformation of a vector  $\boldsymbol{v}$  by a group element  $\boldsymbol{g}$ .

# 4.3.2 Representation on product spaces

The product space  $V\otimes V$  then has a basis  $\{{\pmb b}_i{\pmb b}_j\}_{i,j=1}^d$ . A derived representation can be constructed from D on this product space called the product representation  $D^{(D\times D)}(g)$ . This is defined through its application on the basis by

$$g: \boldsymbol{b}_{i}\boldsymbol{b}_{j} = \sum_{kl} \boldsymbol{b}_{k}\boldsymbol{b}_{l} \big[D^{(D\times D)}(g)\big]_{kl,ij} = \sum_{kl} \boldsymbol{b}_{k}\boldsymbol{b}_{l}D_{ki}(g)D_{lj}(g). \tag{4.22}$$

To apply group theory to physical problems, we need to know how the objects we are working with in the physical theory transform under group elements. The most important vector space in quantum mechanics is arguably the Hilbert space where particle states are determined by a momentum and spin quantum number. Each quantum number has its own vector space defined by the basis vectors  $|{\pmb k}\rangle$  and  $|s\rangle$  in the Dirac notation. The combination of both quantum numbers in the description of a particle state then gives a state in the product space of these vector spaces. A basis for this space is given by the vectors  $|{\pmb k},s\rangle=|{\pmb k}\rangle|s\rangle$ . Given  ${\pmb k}\in{\mathbb R}^d$  and  $s\in\{\uparrow,\downarrow\}$ , these basis vectors transform according to the product representation of the representations on each vector space, given by

$$g: |\mathbf{k}', s'\rangle = \sum_{\mathbf{k}s} |\mathbf{k}, s\rangle D_{\mathbf{k}s; \mathbf{k}'s'}^{(\mathbf{k}\times s)} = \sum_{\mathbf{k}s} |\mathbf{k}, s\rangle D_{g ss'} \delta_{\mathbf{k}, g: \mathbf{k}'}, \quad (4.23)$$

under a group element g. Here  $g\mathbf{k}'$  means application of g to the vector  $\mathbf{k}'$  through the standard representation of g in  $\mathbb{R}^d$ .  $D_{g\ ss'}$  is a representation on the spin-up spin-down vector space given by the matrix

$$D_{g\;ss'} = \sigma_{ss'}^0 \cos(\phi/2) - i\hat{\boldsymbol{u}} \cdot \boldsymbol{\sigma}_{ss'} \sin(\phi/2), \tag{4.24}$$

where  $\hat{u}$  is the rotation axis unity vector, while  $\phi$  is the angle that defines the proper rotation associated with g.  $\sigma$  is the vector notation for the 3 Pauli matrices and  $\sigma^0_{ss'} = \delta_{ss'}$  [71, 72].

In the BCS Hilbert space which we discussed in more detail in Section 4.2, the basis vectors are outer products of the momentum spin basis vectors with opposite momentum:  $\{|{\pmb k},s_1\rangle|-{\pmb k},s_2\rangle\}.$  The product representation on this vector space then transforms the basis vectors according to

$$g: |\mathbf{k}', s_1'\rangle| - \mathbf{k}', s_2'\rangle = \sum_{\mathbf{k} s_1 s_2} |\mathbf{k}, s_1\rangle| - \mathbf{k}, s_2\rangle D_{\mathbf{k} s_1 s_2; \mathbf{k}' s_1' s_2'}^{(D \times D)}(g),$$

$$(4.25)$$

where

$$D_{\mathbf{k}\,s_{1}s_{2};\mathbf{k}'\,s_{1}'s_{2}'}^{(D\times D)}(g) = D_{g\,s_{1}s_{1}'}\delta_{\mathbf{k},g:\mathbf{k}'}D_{g\,s_{2}s_{2}'}\delta_{-\mathbf{k},g:-\mathbf{k}'}. \tag{4.26}$$

Since group representations on  ${\pmb k}$  is a linear transformation then  $\delta_{-{\pmb k},g:-{\pmb k}'}=\delta_{{\pmb k},g:{\pmb k}'}$ , such that the last Kronecker delta function becomes superfluous.

## 4.3.3 Representation on $\psi$ -d functions

The coefficients of the basis expansion of a vector in the BCS Hilbert space were given in the conventional  $\psi$ -d notation in Eq. (4.9). Taking the active view of group transformations we can say that the expansion coefficients of arbitrary states in the BCS Hilbert space  $\mathcal B$  transforms like the  $v_i$  in Eq. (4.21) but where now the representation matrix D is given by the matrix  $D_{\boldsymbol k\,s_1s_2;\,\boldsymbol k'\,s_1's_2'}^{(D\times D)}$  above in Eq. (4.26). Written out then, the coefficients transform according to

$$g: v_{s_1 s_2}(\mathbf{k}) = \sum_{\mathbf{k}' \, s_1' s_2'} D_{\mathbf{k} \, s_1 s_2; \, \mathbf{k}' \, s_1' s_2'}^{(D \times D)} v_{s_1' s_2'} v_{s_1' s_2'}(\mathbf{k}'). \tag{4.27}$$

Let now  $|v\rangle$  be a state that is even in space, meaning that its expansion only consists of coefficients  $\psi({\bf k})$  in the  $\psi$ - ${\bf d}$  notation. Then we see from Eq. (4.10) that  $\psi({\bf k})$  can be written  $\psi({\bf k})=v_{\uparrow\downarrow}({\bf k})$ . The transformation properties of  $\psi({\bf k})$  are thus given by

$$g: \psi(\mathbf{k}) = g: v_{\uparrow\downarrow}(\mathbf{k}) = \sum_{\mathbf{k}' s_1' s_2'} D_{\mathbf{k}\uparrow\downarrow; \mathbf{k}' s_1' s_2'}^{(D \times D)} \psi(\mathbf{k}') (i\sigma^y)_{s_1' s_2'}$$

$$= \psi(g^{-1} : \mathbf{k}) (i\sigma^y)_{\uparrow\downarrow} = \psi(g^{-1} : \mathbf{k}). \tag{4.28}$$

In this calculation we inserted the expression of  $D_{\boldsymbol{k}\,s_1s_2;\,\boldsymbol{k}'\,s_1's_2'}^{(D\times D)}$  in Eq. (4.26) and used the equation  $D_g i \sigma^y D_g^\mathsf{T} = i \sigma^y$ , where  $D_g$  are the spin representation matrices given in Eq. (4.24).

To find the transformation properties of  ${m d}_{{m k}}$  the principle is the same as above for  $\psi({m k})$  but the calculations become more involved. We assume that the spin-momentum basis expansion of state  $|v\rangle$  consists of only odd coefficients so that  $v_{s_1s_2}({m k})={m d}_{{m k}}\cdot({m \sigma} i\sigma^y)_{s_1s_2}$ . Inserting this into the active transformation of the coefficients of  $|V\rangle$  in Eq. (4.27) and also inserting the expression for  $D_{{m k}\,s_1s_2;\,{m k}'\,s_1's_2'}^{(D\times D)}$  as we did before yields

$$\begin{split} g: v_{s_1 s_2}(\boldsymbol{k}) &= \sum_{\boldsymbol{k}' s_1' s_2'} \delta_{\boldsymbol{k}, g: \boldsymbol{k}'} D_{g \, s_1 s_1'} D_{g \, s_2 s_2} \boldsymbol{d}_{\boldsymbol{k}'} \cdot (\boldsymbol{\sigma} i \sigma^y)_{s_1' s_2'} \\ &= \sum_{\boldsymbol{s}} \left( D_g \boldsymbol{\sigma} \sigma^y D_g^\mathsf{T} \sigma^y \right)_{s_1 s} \cdot \boldsymbol{d}_{g^{-1}: \boldsymbol{k}} \ i \sigma^y_{s s_2}. \end{split} \tag{4.29}$$

Since a group transformation (aka. the linear transformation given by a group representation) can not bring a state that was even to be odd or vice versa, then the resulting state given by the transformed coefficients  $g:v_{s_1s_2}({\pmb k})$  has to remain odd, and thus they can be expanded in terms of a new  ${\pmb d}'$  such that

$$g: v_{s_1 s_2}({\pmb k}) = \sum_{{}^{\mathbf{g}}} {\pmb d}'_{g^{-1}:{\pmb k}} \cdot {\pmb \sigma}_{s_1 s} i \sigma^y_{s s_2}. \tag{4.30}$$

Having expanded both sides of the transformed coefficients with a common factor  $i\sigma^y$  to the right we can equate the remaining  $2\times 2$  spin matrices which gives an expression for  $d'_{g^{-1}:k}\cdot \boldsymbol{\sigma}$  by comparing Eq. (4.29) and Eq. (4.30). Furthermore, using the anti-commutation property  $\{\sigma^i,\sigma^j\}=2\delta_{ij}\sigma^0$  of Pauli matrices we find that

$$d'_{\boldsymbol{k},i} = \frac{1}{4} \operatorname{Tr} \left( \left\{ \sigma^{i}, \boldsymbol{d}'_{\boldsymbol{k}} \cdot \boldsymbol{\sigma} \right\} \right). \tag{4.31}$$

Inserting the expression for  $d'_{g^{-1}:k} \cdot \sigma$  and the full expression of the SU(2) spin-representation matrices  $D_g$ , which is found in Eq. (4.24), yields after some algebra

$$d'_{g^{-1}:\boldsymbol{k},i} = \frac{1}{4} \operatorname{Tr} \left( \left\{ \sigma^{i}, \boldsymbol{d}_{g^{-1}:\boldsymbol{k}} \cdot D_{g} \boldsymbol{\sigma} \sigma^{y} D_{v}^{\mathsf{T}} \sigma^{y} \right\} \right)$$

$$= R_{ij} (\hat{\boldsymbol{u}}, \phi) d_{g^{-1}:\boldsymbol{k},j}, \tag{4.32}$$

where we have defined the matrix

$$\begin{split} R_{ij}(\hat{\boldsymbol{u}},\phi) &= \delta_{ij}\cos\phi + \hat{u}_i\hat{u}_j(1-\cos\phi) - \epsilon_{ijk}\hat{u}_k\sin\phi \\ &= \begin{pmatrix} \cos\phi + \hat{u}_x^2(1-\cos\phi) & \hat{u}_x\hat{u}_y(1-\cos\phi) - \hat{u}_z\sin\phi & \hat{u}_x\hat{u}_z(1-\cos\phi) + \hat{u}_y\sin\phi \\ \hat{u}_y\hat{u}_x(1-\cos\phi) + \hat{u}_z\sin\phi & \cos\phi + \hat{u}_y^2(1-\cos\phi) & \hat{u}_y\hat{u}_z(1-\cos\phi) - \hat{u}_x\sin\phi \\ \hat{u}_z\hat{u}_x(1-\cos\phi) - \hat{u}_y\sin\phi & \hat{u}_z\hat{u}_y(1-\cos\phi) + \hat{u}_x\sin\phi & \cos\phi + \hat{u}_z^2(1-\cos\phi) \end{pmatrix}. \end{split}$$

This matrix is in fact the rotation matrix of a vector in  $\mathbb{R}^3$  by an angle  $\phi$  about a unit vector  $\hat{\boldsymbol{u}}$ . Since the coefficients of an odd state  $|v\rangle$  are fully determined by the vector  $\boldsymbol{d}$ , their transformation can be regarded just as a transformation of  $\boldsymbol{d}$  itself which thus takes the form

$$g: d_{\mathbf{k},i} = R_{ij}(\hat{\mathbf{u}}, \phi) d_{q^{-1}:\mathbf{k},j}, \tag{4.34}$$

where  $\hat{u}$  and  $\phi$  give the unit vector and angle respectively, of the proper rotation associated with g. The conclusion is thus that d transforms as a vector by the proper rotation associated with g.

# 4.3.4 Representation on ladder operators

The fermionic creation and annihilation operators  $c_{{m k},s}^{\dagger}$  and  $c_{{m k},s}$ , which we will collectively refer to as  $c_{{m k},s'}^{(\dagger)}$  are second-quantized operators that act on multi-particle states in a fermionic Fock space [68]. To properly define how a group element g transforms these operators we thus should strictly speaking first derive the representation Q(g) of g on N-particle states  $|{m k}_1,s_1\rangle\wedge\ldots\wedge|{m k}_N,s_N\rangle$  for arbitrary N and then use the relation  $g:c_{{m k},s}^{(\dagger)}=Q(g)c_{{m k},s}^{(\dagger)}Q(g)^{-1}$  to then derive their transformation properties. However, luckily, a shortcut is possible because the Q(g) representation is connected with how g acts on the single-particle basis  $\{|{m k},s\rangle\}$  through the relationship

$$Q(g)c_{\mathbf{k},s}^{(\dagger)}Q(g)^{-1} = c^{(\dagger)}(g:|\mathbf{k},s\rangle). \tag{4.35}$$

In this notation we treat the creation and annihilation operators as respectively linear and antiunitary functions of the states that they create or annihilate. If the matrix-components of the representation of g on the single-particle basis is denoted  $D_{\boldsymbol{k}s;\boldsymbol{k}'s'}^{(\boldsymbol{k}\times s)}$  then we thus have the transformation property

$$g: c_{\mathbf{k},s}^{(\dagger)} = \sum_{\mathbf{k}',s'} c_{\mathbf{k}',s'}^{(\dagger)} [D_{\mathbf{k}'s';\mathbf{k}s}^{(\mathbf{k}\times s)}]^{(*)}, \tag{4.36}$$

where the matrix-elements are complex conjugated only if we are transforming an annihilation operator.

Given that g is an element of some point group such that it can be decomposed into a rotation and a parity inversion, then we may use the representation on  $\{|k,s\rangle\}$  given in Eq. (4.23) such that

$$g: c_{\mathbf{k},s}^{(\dagger)} = \sum_{s'} c_{g:\mathbf{k},s'}^{(\dagger)} D_{g\;s's}^{(*)},$$
 (4.37)

where the matrix  $D_{g\;ss'}$  is defined in Eq. (4.24). As an example, a simple parity transformation  $g=\hat{P}$ , then transform the operators according to  $\hat{P}:c_{{m k},s}^{(\dagger)}=c_{{m k},s}^{(\dagger)}$ .

We will later need the translation transformation  $g=\pmb{R}$  as well. In real space it is defined by  $\pmb{R}:|\pmb{r},s\rangle=|\pmb{r}+\pmb{R},s\rangle$  and given by linear transformation  $L(\pmb{R})=e^{i\pmb{R}\cdot\hat{\pmb{p}}/\hbar}$  where  $\hat{\pmb{p}}$  is the momentum operator  $\hat{\pmb{p}}=\hbar\nabla/i$ . It then follows by a Fourier-transformation that the

<sup>3.</sup> The form of the linear transformation  $L({m R})$  follows directly from the multivariate Taylor expansion.

fermionic operators transform according to

$$\boldsymbol{R}: c_{\boldsymbol{k},s}^{(\dagger)} = \left(e^{-i\boldsymbol{R}\cdot\boldsymbol{k}}\right)^{(*)} c_{\boldsymbol{k},s}^{(\dagger)}. \tag{4.38}$$

Finally we discuss the transformation of time-reversal. This operation is traditionally denoted  $\Theta$  and consists of flipping the sign of time, which implies flipping the spin and momentum. Time-reversal is special in that it is an antiunitary, i.e. it transforms all linear coefficients to their complex conjugate. Any representation is because of this, often factorized into a normal linear part and a complex-conjugation operator  $\hat{K}$ . From its action on the spin-momentum basis vectors  $|k,s\rangle$ , time-reversal transforms the fermionic creation and annihilation operators according to

$$\Theta: c_{\mathbf{k},s}^{(\dagger)} = \sum_{s'} c_{-\mathbf{k},s'}^{(\dagger)} (-i\sigma_y)_{s's}. \tag{4.39}$$

Note that the matrix  $-i\sigma_y$  is not complex conjugated for the fermionic annihilation operators because of the antiunitarity of  $\Theta$ .

#### 4.4 Single-particle Hamiltonian symmetries

Any fermionic single-particle operator can be written in the spin-momentum basis as

$$\hat{H} = \sum_{\mathbf{k}\mathbf{k}'ss'} H_{\mathbf{k}\mathbf{k}'}^{ss'} c_{\mathbf{k},s}^{\dagger} c_{\mathbf{k}'s'}. \tag{4.40}$$

In this section we will employ common finite symmetry-transformations discussed in Section 4.3.4 and see how this reduces the degrees of freedom in  $\hat{H}$  so that it can be written in a more concise form. To begin we reduce the notational burden by expanding the spin-matrix elements in separate coefficients in the Pauli-matrix basis just as was done for the BCS Hilbert space vector coefficients  $v_{s_1s_2}(\boldsymbol{k})$  in Eq. (4.7) such that

$$\hat{H} = \sum_{\mathbf{k}\mathbf{k}'ss'} \left[ \xi_{\mathbf{k}\mathbf{k}'} \sigma^0 + \gamma_{\mathbf{k}\mathbf{k}'} \cdot \boldsymbol{\sigma} \right]_{ss'} c_{\mathbf{k},s}^{\dagger} c_{\mathbf{k}'s'}. \tag{4.41}$$

The transformation of a quantum mechanical operator such as the Hamiltonian is defined by  $g: \hat{H} = Q(g)\hat{H}Q(g)^{-1}$  for some many-body representation Q(g). Inserting the form of  $\hat{H}$  in Eq. (4.41) as a single-particle operator in the spin-momentum basis yields

$$g: \hat{H} = \sum_{\mathbf{k}\mathbf{k}'ss'} \left[ \xi_{\mathbf{k}\mathbf{k}'} \sigma^0 + \gamma_{\mathbf{k}\mathbf{k}'} \cdot \boldsymbol{\sigma} \right]_{ss'} (g: c_{\mathbf{k},s}^{\dagger}) (g: c_{\mathbf{k}'s'}), \quad (4.42)$$

after inserting  $Q(g)^{-1}Q(g)$  between the creation and annihilation operator. We can now use the transformation properties of these operators presented in Section 4.3.4 to derive the transformation properties of the Hamiltonian.

For a translation invariant system we need a translation invariant Hamiltonian. Using the transformation of  $c^{(\dagger)}$  in Eq. (4.38) under a translation  ${\pmb R}$ , we get that the coefficients  $\xi_{{\pmb k}{\pmb k}'}$  and  $\gamma_{{\pmb k}{\pmb k}'}$  has to satisfy the equation

$$(e^{i\mathbf{R}\cdot(\mathbf{k}'-\mathbf{k})}-1)[\xi_{\mathbf{k}\mathbf{k}'}\sigma^0+\gamma_{\mathbf{k}\mathbf{k}'}\cdot\boldsymbol{\sigma}]=0. \tag{4.43}$$

Since the sigma-matrices are linearly independent and the translation R is arbitrary, it follows that both  $\xi_{kk'}$  and  $\gamma_{kk'}$  must vanish whenever  $k \neq k'$ , i.e. these coefficients must be diagonal in k. The Hamiltonian can then be written on the reduced form

$$\hat{H} = \sum_{\mathbf{k} s s'} \left[ \xi_{\mathbf{k}} \sigma^0 + \gamma_{\mathbf{k}} \cdot \boldsymbol{\sigma} \right]_{ss'} c_{\mathbf{k},s}^{\dagger} c_{\mathbf{k},s'}. \tag{4.44}$$

In the vast majority of cases, the system is Hermitian such that the Hamiltonian is self-adjoint and has real eigenvalues. Enforcing the condition  $\hat{H}^{\dagger}=\hat{H}$  on the translationally invariant Hamiltonian in Eq. (4.44) yields the equation

$$(\boldsymbol{\xi}_{k}^{*} - \boldsymbol{\xi}_{k})\sigma^{0} + (\boldsymbol{\gamma}_{k}^{*} - \boldsymbol{\gamma}_{k}) \cdot \boldsymbol{\sigma} = 0, \tag{4.45}$$

by using the self-adjoint property of the Pauli-matrices. This implies further by the linear independence of the sigma-matrices that the coefficients  $\xi_{k}$  and  $\gamma_{k}$  are self-conjugate and hence real.

Time-reversal symmetry of the system implies that  $\hat{H}$  should be invariant under the transformation  $\Theta$ , which transforms fermionic creation and annihilation operators according to Eq. (4.39). Transforming these operators in the translationally invariant Hamiltonian in Eq. (4.44) and remembering that  $Q(\Theta)$  is an antilinear operator we get the transformed Hamiltonian

$$\Theta: \hat{H} = \sum_{\boldsymbol{k}ss'} \left[ \xi_{-\boldsymbol{k}}^* \sigma^0 - \gamma_{-\boldsymbol{k}}^* \cdot \boldsymbol{\sigma} \right]_{ss'} c_{\boldsymbol{k},s}^{\dagger} c_{\boldsymbol{k},s'}. \tag{4.46}$$

For the Hamiltonian to be invariant under time-reversal symmetry, the coefficients thus have to satisfy  $\xi_{k}=\xi_{-k}^{*}$  and  $\gamma_{k}=-\gamma_{-k}^{*}$ .

Finally we look at the transformation of the Hamiltonian under point-group elements g. Using the transformation-property of the fermionic creation and annihilation operators in Eq. (4.37), then the translation-invariant Hamiltonian in Eq. (4.44) transforms according to

$$g: \hat{H} = \sum_{\boldsymbol{k}ss'} \left[ \xi_{g^{-1}:\boldsymbol{k}} \sigma^0 + \boldsymbol{\gamma}_{g^{-1}:\boldsymbol{k}} \cdot D_g \boldsymbol{\sigma} D_g^{\dagger} \right]_{ss'} c_{\boldsymbol{k},s}^{\dagger} c_{\boldsymbol{k},s'}. \tag{4.47}$$

Using the form of the spin-rotation matrix  $\boldsymbol{D}_g$  in Eq. (4.24) we find that

$$\gamma_{g^{-1}:\mathbf{k}} \cdot D_g \boldsymbol{\sigma} D_g^{\dagger} = \gamma'_{g^{-1}:\mathbf{k}} \cdot \boldsymbol{\sigma},$$
 (4.48)

for the transformed vector

$$\gamma_{k}' = R(\hat{\boldsymbol{u}}, \phi) \gamma_{k}, \tag{4.49}$$

where  $R(\hat{\boldsymbol{u}},\phi)$  is the  $3\times 3$  matrix representation of the proper-rotation associated with g given in Eq. (4.33). Symmetry of  $\hat{H}$  under point-group transformation thus implies the symmetry relations  $\xi_{g:\boldsymbol{k}}=\xi_{\boldsymbol{k}}$  and  $R(\hat{\boldsymbol{u}},\phi)\gamma_{q^{-1}:\boldsymbol{k}}=\gamma_{\boldsymbol{k}}$  on the coefficients of the Hamiltonian.

# 4.5 Projection Operators

Let us assume that we are in a vector space V that can be divided into possibly several different IRs  $D^{(\alpha)}$  of some symmetry group G. Further, let the basis vectors of these IRs be denoted by  $\boldsymbol{b}_m^{(\alpha)}$  where m thus counts the number of basis vectors in each irrep. Then an arbitrary vector  $\boldsymbol{f} \in V$  can be written in terms of these basis vectors as

$$f = \sum_{\alpha} \sum_{m} c_m^{(\alpha)} \boldsymbol{b}_m^{(\alpha)}. \tag{4.50}$$

A projection operator can be used to extract any combination of constant  $c_m^{(\alpha)}$  multiplied by a basis vector  $\boldsymbol{b}_n^{(\alpha)}$ , where m and n can in general be different. Denoting the projection operator that picks out the mth constant multiplied by the lth basis vector in the irrep.  $\beta$  of the expansion of  $\boldsymbol{f}$ :  $P_{lm}^{(\beta)}$ , then

$$P_{lm}^{(\beta)} \boldsymbol{f} = c_m^{(\beta)} \boldsymbol{b}_l^{(\beta)}. \tag{4.51}$$

This is extremely useful in finding a basis for the IRs. To achieve this, the projection operator is defined as

$$P_{l,m}^{(\beta)} = \frac{d_{\beta}}{|G|} \sum_{g \in G} D_{lm}^{(\beta)}(g)^* g :, \tag{4.52}$$

where  $d_{\beta}$  is the dimension of irrep.  $\beta$ ,  $D_{lm}^{(\beta)}(g)$  is the lm element of the matrix representation of the group element g and finally we have used the notation g: to denote application on vectors by the relevant representation. An example is the application of g to the basis vectors  $\boldsymbol{b}_{m}^{(\alpha)}$ . Since the relevant representation of g in this case is the IR for which  $\boldsymbol{b}_{m}^{(\alpha)}$  is a basis vector, the application becomes

$$g: b_m^{(\alpha)} = \sum_n b_n^{(\alpha)} D_{nm}^{(\alpha)}(g).$$
 (4.53)

Usually, the full generality of the projection operators  $P_{l,m}^{(\beta)}$  isn't needed and it suffices to consider the diagonal projection operators  $P_{l,l}^{(\beta)} \equiv P_l^{(\beta)}$  or indeed their sum, in which case the resulting operator can be written only in terms of the irrep. characters  $\chi^{(\alpha)}(q)$  since

$$P^{(\beta)} \equiv \sum_{l} P_{l}^{(\beta)} = \frac{d_{\beta}}{|G|} \sum_{g \in G} \sum_{l} D_{ll}^{(\beta)}(g)^{*}g := \frac{d_{\beta}}{|G|} \sum_{g \in G} \chi^{(\beta)}(g)^{*}g :.$$
(4.54)

## 4.6 Symmetries of the Square Lattice

The symmetry group of the square lattice is denoted  $C_{4v}$  in the Schönflies notation. It contains 8 elements in total:

e: The identity element (do nothing),

 $C_4$ : Rotation by 90° in the positive direction (ccw),

 $C_4^{-1}$ : Rotation by 90° in the negative direction (cw),

 $C_4^2$ : Rotation by  $180^\circ$ ,

 $\sigma_x$ : Mirror about the *zy*-plane,

 $\sigma_{y}$ : Mirror about the zx-plane,

 $\sigma_{d_1}$ : Mirror about the downwards diagonal plane,

 $\sigma_{d_2}$ : Mirror about the upwards diagonal plane.

<sup>4.</sup> We are assuming the western bias of left-to-right movement here. More precisely it is the plane containing the z axis and the axis y=-x

	e	$C_4$	$C^2$	$C_4^{-1}$	Ι σ	σ	σ	σ
			$C_4$			$\sigma_y$	$\sigma_{d_1}$	$\sigma_{d_2}$
e	e	-	$C_4^2$		$\sigma_x$	$\sigma_y$	$\sigma_{d_1}$	$\sigma_{d_2}$
$C_4$	$C_4$	$C_4^2$	$C_{4}^{-1}$	e	$\sigma_{d_1}$	$\sigma_d$	$\sigma_y$	$\sigma_x$
$C_4^2$	$C_4^2$	$C_{4}^{-1}$	e	$C_4$	$\sigma_y$	$\sigma_x$	$\sigma_{d_2}$	$\sigma_{d_1}$
$C_4^{-1}$	$C_4^{-1}$	e	$C_4$	$C_4^2$	$\sigma_{d_2}$	$\sigma_{d_1}$	$\sigma_x$	$\sigma_y$
$\sigma_x$	$\sigma_x$	$\sigma_{d_2}$	$\sigma_y$	$\sigma_{d_1}$	e	$C_4^2$	$C_4^{-1}$	$C_4$
$\sigma_y$	$\sigma_y$	$\sigma_{d_1}$	$\sigma_x$	$\sigma_{d_2}$	$C_4^2$	e	$C_4$	$C_4^{-1}$
$\sigma_{d_1}$	$\sigma_{d_1}$	$\sigma_x$	$\sigma_{d_2}$	$\sigma_y^-$	$C_4$	$C_4^{-1}$	$e^{-}$	$C_4^2$
$\sigma_{d_2}$	$\sigma_{d_2}$	$\sigma_y$	$\sigma_{d_1}$	$\sigma_x$	$C_4^{-1}$	$C_4$	$C_4^2$	e

**Table 4.1:** Group multiplication table of the group  $C_{4v}$ .

This results in the group multiplication table in Table 4.1. We can check that this is correct by performing the group transformations in the top row followed by the one in the left column and seeing that this results in the group transformations where these two intersect. As an example consider the vector  $(x,y)^{\mathsf{T}}$ . Transforming this by the  $90^{\circ}$  counterclockwise rotation  $C_4$  we get  $(-y,x)^{\mathsf{T}}$ . Then mirroring this result about the yz-plane yields  $(y,x)^{\mathsf{T}}$ . We now realize that this is the same as mirroring the original vector about the axis y=x, hence  $\sigma_x C_4=\sigma_{d_2}$  as the multiplication table says.

# 4.6.1 Conjugation classes

The conjugation classes of a group is the sets of group elements that is conjugate to each other, meaning that there exists a group element g such that  $gAg^{-1}=B$  between conjugate elements A and B. Since conjugation is an equivalence relation it subdivides the group elements exactly into conjugation classes. The conjugation classes of the group  $C_{4v}$  are  $e=\{e\}, 2C_4=\{C_4,C_4^{-1}\}, C_4^2=\{C_4^2\}, 2\sigma_d=\{\sigma_{d_1},\sigma_{d_2}\}$  and  $2\sigma_v=\{\sigma_x,\sigma_y\}.$  The character  $\chi^\Gamma(g)$  of a representation  $\Gamma$  is the trace of the representation matrix  $D^\Gamma(g)$  of a certain group element g. Since the trace is cyclic, then for conjugate group elements A and B

$$\chi^{\Gamma}(B) = \operatorname{Tr}\left(D^{\Gamma}(g)D^{\Gamma}(A)D^{\Gamma}(g^{-1})\right)$$

$$= \operatorname{Tr}\left(D^{\Gamma}(g^{-1})D^{\Gamma}(g)D^{\Gamma}(A)\right)$$

$$= \operatorname{Tr}\left(D^{\Gamma}(g^{-1}g)D^{\Gamma}(A)\right) = \chi^{\Gamma}(A).$$
(4.55)

$C_{4v}$	e	$2C_4$	$C_4^2$	$2\sigma_v$	$2\sigma_d$
$\Gamma_1$	1	1 1 -1 -1 0	1	1	1
$\Gamma_2$	1	1	1	-1	-1
$\Gamma_3$	1	-1	1	1	-1
$\Gamma_4$	1	-1	1	-1	1
$\Gamma_{5}$	2	0	-2	0	0

**Table 4.2:** Character table of the group  $C_{4v}$ .

This means that the representation of all group elements in a certain conjugation class has the same character.

It is useful to list the characters of the different conjugation classes in a table of the different IRs of a group. This is because the number of conjugation classes of a finite group is the same as the number of IRs of that group. This table is known as the character table of the group. The character table of the group  $C_{4v}$  is shown in Table 4.2. This table can be derived without knowing the details of the irreducible representations but instead using character relations from basic group theory.<sup>5</sup>

## 4.6.2 Irreducible representations

The dimensionality of the irrep. can be found in Table 4.2 by looking up the first column, i.e. the column giving the character of the conjugation class  $\{e\}$ . Since the group element e maps to the identity transformation in all representations then its trace gives the dimension of the representation. From the table we see that all the IRs are 1-dimensional except for  $\Gamma_5$  which is 2 dimensional. All the 1-dimensional IR matrices are then completely determined by the character table since they are just given by the characters themselves, i.e.  $D^{\Gamma_2}(\sigma_x) = -1$ .

To find a 2-dimensional representation of  $C_{4v}$  we can imagine how a normal 2D vector  $(x,y)^\mathsf{T} \in \mathbb{R}^2$  behaves under its transformations. We take again the example of a counterclockwise rotation by  $90^\circ$  which transforms a vector

$$C_4: \begin{pmatrix} x \\ y \end{pmatrix} = \begin{pmatrix} -y \\ x \end{pmatrix} = \begin{pmatrix} 0 & -1 \\ 1 & 0 \end{pmatrix} \begin{pmatrix} x \\ y \end{pmatrix}. \tag{4.56}$$

<sup>5.</sup> Many of these relations are derived from the great orthogonality theorem which can be found e.g. in [69].

Obviously, the matrix

$$D^{(\Gamma_5)}(C_4) = \begin{pmatrix} 0 & -1\\ 1 & 0 \end{pmatrix}, \tag{4.57}$$

is the representation matrix of a two-dimensional representation of  ${\cal C}_4$ . Continuing in this way for all the group transformations yields the matrices

$$D^{(\Gamma_5)}(e) = \begin{pmatrix} 1 & 0 \\ 0 & 1 \end{pmatrix}, \qquad D^{(\Gamma_5)}(C_4) = \begin{pmatrix} 0 & -1 \\ 1 & 0 \end{pmatrix}, \quad (4.58a)$$

$$D^{(\Gamma_5)}(C_4^{-1}) = \begin{pmatrix} 0 & 1 \\ -1 & 0 \end{pmatrix}, \qquad D^{(\Gamma_5)}(C_4^2) = \begin{pmatrix} -1 & 0 \\ 0 & -1 \end{pmatrix}, \quad (4.58b)$$

$$D^{(\Gamma_5)}(\sigma_x) = \begin{pmatrix} -1 & 0 \\ 0 & 1 \end{pmatrix}, \qquad D^{(\Gamma_5)}(\sigma_y) = \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix}, \qquad (4.58c)$$

$$D^{(\Gamma_5)}(\sigma_{d_1}) = \begin{pmatrix} 0 & -1 \\ -1 & 0 \end{pmatrix}, \quad D^{(\Gamma_5)}(\sigma_{d_2}) = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}. \tag{4.58d}$$

Taking the trace of these matrices, we see that this representations characters are the same as the ones for the irrep.  $\Gamma_5$  in the character table (Table 4.2). This implies that  $\sum_g |\chi^{\Gamma_5 u}(g)|^2 = |C_{4_v}|$  which implies in turn that the representation given by the matrices in Eq. (4.58) is irreducible. Thus this is indeed the  $\Gamma_5$  IR as advertised, and we have completed the description of the representation matrices of all the IRs of  $C_{4v}$ .

# 4.6.3 Representation on odd BCS functions $d_k$

When discussing the representation of general group elements on states in the BCS Hilbert space in Section 4.3.3 we learned from Eq. (4.34) that the coefficient  $d_{\boldsymbol{k}} \in \mathbb{R}^3$  of odd states transforms by a proper rotation  $R(\hat{\boldsymbol{u}},\phi)$ . A proper rotation of a group element g is the rotation obtained when writing g as a rotation followed by an inversion P or identity. We thus obtain the representation matrices of the 3D-representation of group elements  $g \in C_{4v}$  directly from the 3D rotation matrix  $R(\hat{\boldsymbol{u}},\phi)$  by envisioning the combination of a rotation and  $C_4^2 = P$  or e that leads to g. Because of this, we denote the representation matrices of this representation R(g). As an example  $\sigma_x = C_4^2 R(\hat{\boldsymbol{x}},\pi)$ , such that the representation  $R(\sigma_x) = R(\hat{\boldsymbol{x}},\pi)$ .

Written out in its full matrix form this yields the representation matrices

$$R(C_4) = \begin{pmatrix} 0 & -1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 1 \end{pmatrix}, \quad R(C_4^{-1}) = \begin{pmatrix} 0 & 1 & 0 \\ -1 & 0 & 0 \\ 0 & 0 & 1 \end{pmatrix}, \quad (4.59a)$$

$$R(e) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix}, \quad R(C_4^2) = \begin{pmatrix} -1 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & 1 \end{pmatrix}, \quad (4.59b)$$

$$R(\sigma_x) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & -1 \end{pmatrix}, \quad R(\sigma_y) = \begin{pmatrix} -1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & -1 \end{pmatrix}, \quad (4.59c)$$

$$R(\sigma_{d_1}) = \begin{pmatrix} 0 & 1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & -1 \end{pmatrix}, \quad R(\sigma_{d_2}) = \begin{pmatrix} 0 & -1 & 0 \\ -1 & 0 & 0 \\ 0 & 0 & -1 \end{pmatrix}. \quad (4.59d)$$

## 4.7 Square Lattice Harmonics

Since  $\psi(k)$  and  $d_k$  are invariant with respect to translation by any reciprocal lattice vector  $\mathbf{Q}$ :  $\psi(\mathbf{k}+\mathbf{Q})=\psi(\mathbf{k})$ , they can be expanded in a discrete Fourier transform over the real lattice, such that

$$\psi(\mathbf{k}) = \frac{1}{\sqrt{N}} \sum_{\mathbf{R}} \psi_{\mathbf{R}} \cos \mathbf{R} \cdot \mathbf{k}, \tag{4.60}$$

and

$$d_{k} = \frac{1}{\sqrt{N}} \sum_{R} d_{R} \sin R \cdot k, \qquad (4.61)$$

where the exponential of the Fourier transform has been reduced to trigonometric functions by the parity of the functions.

We are interested in the basis vectors  $|\Gamma,q,m\rangle$  of the representations of the symmetry group  $C_{4v}$  of the 2D square lattice. In this ket notation,  $\Gamma$  gives the irrep., m enumerates the basis vectors in case the irrep. is multi-dimensional, while q gives the version of the irrep. in case the space of possible  $|v\rangle$  permits multiple versions of the same irrep. In the active view of group transformations, the question of finding the basis vectors translates to finding the basis functions of the functions  $\psi(k)$  and  $d_k$  for even and odd bases respectively. In Section 4.3.3 we

saw how these functions transformed under group transformations. In Section 4.5 we saw how the projection operators could be used to extract individual basis vectors. We will now use these operators on the Fourier expansions of  $\psi({\boldsymbol k})$  and  ${\boldsymbol d}_{\boldsymbol k}$  to extract possible basis functions given the symmetry group of the square lattice.

#### 4.7.1 Even basis functions

We remember from Eq. (4.28) that the function  $\psi(\mathbf{k})$  transforms as  $g:\psi(\mathbf{k})=\psi(g^{-1}:\mathbf{k})$  under a group transformation g. Operating on the Fourier expansion of  $\psi(\mathbf{k})$  in Eq. (4.60) with the projection operator defined in Eq. (4.52) by an arbitrary irrep.  $\Gamma$  yields

$$\begin{split} P_{l,l}^{(\Gamma)} \psi(\mathbf{k}) &= \frac{d_{\Gamma}}{8\sqrt{N}} \sum_{\mathbf{R}} \psi_{\mathbf{R}} \Big[ \big( D_{ll}^{(\Gamma)}(e) + D_{ll}^{(\Gamma)}(C_4^2) \big) \cos(\mathbf{R} \cdot \mathbf{k}) \\ &+ \big( D_{ll}^{(\Gamma)}(C_4) + D_{ll}^{(\Gamma)}(C_4^{-1}) \big) \cos(R_x k_y - R_y k_x) \\ &+ \big( D_{ll}^{(\Gamma)}(\sigma_x) + D_{ll}^{(\Gamma)}(\sigma_y) \big) \cos(R_x k_x - R_y k_y) \\ &+ \big( D_{ll}^{(\Gamma)}(\sigma_{d_1}) + D_{ll}^{(\Gamma)}(\sigma_{d_2}) \big) \cos(R_x k_y + R_y k_x). \end{split} \tag{4.62}$$

Since  ${\pmb k} \in {\mathbb R}^2$  we have in this calculation used the natural 2D representation given by the matrices in Eq. (4.58) of the group elements in  $C_{4v}$  to calculate the expressions  $g^{-1}:{\pmb k}$  inside the cosine functions. Inserting the matrix element of the different IRs of  $C_{4v}$  which we discussed in Section 4.6 we get the projected functions

$$\begin{split} P_{1,1}^{(\Gamma_1)}\psi(\boldsymbol{k}) &\propto \sum_{\boldsymbol{R}} \psi_{\boldsymbol{R}} \big[ \cos R_x k_x \cos R_y k_y + \cos R_x k_y \cos R_y k_x \big], \\ P_{1,1}^{(\Gamma_2)}\psi(\boldsymbol{k}) &\propto \sum_{\boldsymbol{R}} \psi_{\boldsymbol{R}} \big[ \sin R_x k_y \sin R_y k_x - \sin R_x k_x \sin R_y k_y \big], \\ P_{1,1}^{(\Gamma_3)}\psi(\boldsymbol{k}) &\propto \sum_{\boldsymbol{R}} \psi_{\boldsymbol{R}} \big[ \cos R_x k_x \cos R_y k_y - \cos R_x k_y \cos R_y k_x \big], \\ P_{1,1}^{(\Gamma_4)}\psi(\boldsymbol{k}) &\propto \sum_{\boldsymbol{R}} \psi_{\boldsymbol{R}} \big[ \sin R_x k_x \sin R_y k_y + \sin R_x k_y \sin R_y k_x \big], \\ P_{1,1}^{(\Gamma_5)}\psi(\boldsymbol{k}) &\propto 0, \\ P_{2,2}^{(\Gamma_5)}\psi(\boldsymbol{k}) &\propto 0. \end{split}$$

$$(4.63)$$

Since the projection operators  $P_{ll}^{(\Gamma)}$  projects onto the subspace of vectors belonging to the IR  $\Gamma$ , if we let the different basis functions of the irreps.  $\Gamma$  be denoted  $\psi^{(\Gamma),q,m}(\boldsymbol{k})$ , then we expect the projection to produce the result

$$P_{l,l}^{(\Gamma)}\psi(\mathbf{k}) = \sum_{q} c_{q,l}\psi^{(\Gamma),q,l}(\mathbf{k}). \tag{4.64}$$

Here q again enumerates the version of the basis of  $\Gamma$  possible in the space of different  $\psi({\boldsymbol k})$ , and  $c_{q,l}$  are the coefficients of  $\psi({\boldsymbol k})$  in the basis of the irrep. basis functions. Comparing Eqs. (4.64) and (4.63), we see that different sets of basis vectors can be obtained for the IRs by including different order terms in the  ${\boldsymbol R}$ -sum, i.e. different lattice neighbour sites. It is also worth noting that the IR  $\Gamma_5$  does not exist in the space of possible  $\psi({\boldsymbol k})$ , since it is an odd representation.

Including on-site, nearest neighbor and next-nearest neighbor sites in the  ${\bf R}$  sum of the projected arbitrary function  $\psi({\bf k})$  on the  $\Gamma_1$  subspace in Eq. (4.64) we see that any such function can be constructed from the three basis functions

$$\psi^{(\Gamma_1),1}(\mathbf{k}) = \frac{1}{2\pi},\tag{4.65a}$$

$$\psi^{(\Gamma_1),2}({\pmb k}) = \frac{1}{2\pi}(\cos k_x + \cos k_y), \tag{4.65b}$$

$$\psi^{(\Gamma_1),3}(\mathbf{k}) = \frac{1}{\pi} \cos k_x \cos k_y. \tag{4.65c}$$

Each of these functions give a complete basis function set of the  $\Gamma_1$  IR which can be checked by calculating how they transform under group elements  $g \in C_{4v}$ . In this case, since this is the trivial  $\Gamma_1$  representation, the functions are symmetric under all group elements g which produces the character 1 for all conjugation classes (compare with the first row in Table 4.2).

These basis functions are automatically mutually orthogonal since they belong to different IR version subspaces and their normalization coefficients have been chosen such that they are normal on the  $1^{\rm st}$  Brillouin zone, i.e.

$$\int_{-\pi}^{\pi} \int_{-\pi}^{\pi} \mathrm{d}k_x \, \mathrm{d}k_y \ \psi^{(\Gamma_1),q}({\pmb k})^* \psi^{(\Gamma_1),q'}({\pmb k}) = \delta_{qq'}. \eqno(4.66)$$

Inserting up to next-nearest neighbor sites for  ${\it R}$  in the projected functions under the remaining IRs we find the orthonormal basis functions

$$\psi^{(\Gamma_3)}(\boldsymbol{k}) = \frac{1}{2\pi}(\cos k_x - \cos k_y), \tag{4.67a}$$

$$\psi^{(\Gamma_4)}(\pmb{k}) = \frac{1}{\pi} \sin k_x \sin k_y, \tag{4.67b} \label{eq:psi_fit}$$

of the representations  $\Gamma_3$  and  $\Gamma_4$  respectively. The  $\Gamma_3$  basis is found by expansion of  ${\bf R}$  to nearest neighbor, while the one for  $\Gamma_4$  is found at the next-nearest neighbor. To get a basis vector for the representation  $\Gamma_2$  we would need to expand beyond the next-nearest neighbor site.

These basis functions are also known as square lattice harmonics. The set of square lattice harmonic functions includes the set of functions found when expanding  ${\pmb R}$  to arbitrary sites. As we have seen, they can be grouped and found through consideration of the IRs of the symmetry group of the square lattice. We have so far only considered even-in- ${\pmb k}$  basis functions. In the next section we will complete our discussion of square lattice harmonics with the inclusion of odd functions.

#### 4.7.2 Odd basis functions

Any state made out of odd basis functions is fully determined by the coefficients  $\boldsymbol{d_k}$ . These coefficients transform as  $g:\boldsymbol{d_k}=R(g)\boldsymbol{d_{g^{-1}:k}}$  under group elements  $g\in C_{4v}$ , see Eq. (4.34), where R(g) are the representation matrices in Eq. (4.59). Acting on the Fourier expansion of the arbitrary function  $\boldsymbol{d_k}$  in Eq. (4.61) with the projection operators Eq. (4.52) down on the subspace of the IR  $\Gamma_5$  of the symmetry group  $C_{4v}$  yields the results

$$P_{1,1}^{(\Gamma_5)}\boldsymbol{d_k} = \frac{\hat{\boldsymbol{z}}}{\sqrt{N}} \sum_{\boldsymbol{R}} d_{\boldsymbol{R},z} \cos R_x k_x \sin R_y k_y, \tag{4.68a}$$

$$P_{2,2}^{(\Gamma_5)} \boldsymbol{d_k} = \frac{\hat{\boldsymbol{z}}}{\sqrt{N}} \sum_{\boldsymbol{R}} d_{\boldsymbol{R},z} \sin R_x k_y \cos R_y k_y, \tag{4.68b}$$

for the two basis functions of  $\Gamma_5$ . As in the spin-singlet case we get different versions of the  $\Gamma_5$  basis vectors depending on the order of our

expansion in  $m{R}$ . Expanding to nearest neighbor sites and normalizing such that the states are orthonormal produces the basis functions

$$\boldsymbol{d}_{\boldsymbol{k}}^{(\Gamma_5),1} = -\frac{\hat{\boldsymbol{z}}}{2\pi}\sin k_y, \tag{4.69a}$$

$$\boldsymbol{d}_{\boldsymbol{k}}^{(\Gamma_5),2} = \frac{\hat{\boldsymbol{z}}}{2\pi} \sin k_x, \tag{4.69b}$$

of the two-dimensional IR  $\Gamma_5$ .

## 4.8 Decomposition of the Potential

Let at first  $\hat{V}$  be a general two-body operator that acts on an N-particle state which is a vector in  $\mathcal{H}_N = \bigotimes_{i=1}^N \mathcal{H}$ . The single particle Hilbert space  $\mathcal{H}$  in question is quantified by momentum and spin such that  $\mathcal{H} = \operatorname{span}\{|\boldsymbol{k},s\rangle\}$ . Denoting specific combinations of  $\boldsymbol{k}$  and s as a shorthand for the moment, then  $\hat{V}$  acts on basis vectors in  $\mathcal{H}_N$  as

$$\hat{V}|\alpha_1\rangle \dots |\alpha_N\rangle = \sum_{1 \leq i < j \leq N} \hat{V}_{ij}|\alpha_1\rangle \dots |\alpha_N\rangle, \tag{4.70}$$

by definition of being a two-body operator. Here  $\hat{V}_{ij}$  is an operator that only acts on the ith and jth ket. Even though  $\hat{V}$  acts on  $\mathcal{H}_N$ , because of how it can be written in terms of  $\hat{V}_{ij}$  and this only acts on two states at a time, it follows that  $\hat{V}$  is completely determined by its action on the reduced Hilbert space  $\mathcal{H}_2$ . This implies that  $\hat{V}$  is fully described by its matrix elements

$$\langle \alpha | \langle \alpha' | \hat{V} | \beta \rangle | \beta' \rangle.$$
 (4.71)

Inserting back the  $|{\pmb k},s\rangle$  notation, these matrix elements are referred to as

$$V_{\mathbf{k}_1 \mathbf{k}_2 \mathbf{k}_3 \mathbf{k}_4; s_1 s_2 s_3 s_4} = \langle \mathbf{k}_1 s_1 | \langle \mathbf{k}_2 s_2 | \hat{V} | \mathbf{k}_4 s_4 \rangle | \mathbf{k}_3 s_3 \rangle. \tag{4.72}$$

When  $\hat{V}$  is a BCS operator acting on the BCS Hilbert space described in the previous section, these matrix elements are denoted

$$V_{\mathbf{k}\mathbf{k}'; s_1 s_2 s_3 s_4} = \langle \mathbf{k}s_1 | \langle -\mathbf{k}s_2 | \hat{V} | \mathbf{k}' s_4 \rangle | -\mathbf{k}' s_3 \rangle. \tag{4.73}$$

Since  $\hat{V}$  is Hermitian, it must be diagonalizable in some basis of eigenfunctions. Barring accidental degeneracy, a basis for a d-degenerate

eigenvalue is also a basis for an IR of the symmetry group G of the Hamiltonian. In the case of accidental degeneracy then this d-dimensional vector space consists of several non-intersecting subspaces where each subspace is a basis for a (possibly different) irrep. Note that this does not mean that (barring accidental degeneracy) there exists one separate eigenvalue for each irrep. of G since there might be several different eigenvalues with different eigenspace bases but where all of them are bases for the same irrep. Regardless of these details, the connection between IRs and the eigenvalues of  $\hat{V}$  is a great help in finding the bases for which it is diagonal.

We let the basis for a  $d_{\Gamma}$ -dimensional irrep.  $\Gamma$  be denoted  $\{|\Gamma,q_{\Gamma},m\rangle\}_{m=1}^{d_{\Gamma}}$ , where  $\hat{V}$  has an eigenvalue  $V_{\Gamma,q_{\Gamma}}$  for the vectors in this basis and  $q_{\Gamma}$  is an index enumerating the different versions of bases of  $\Gamma$  that  $\hat{V}$  might have in its set of eigenspace bases. Since  $\hat{V}$  then is diagonal in this set of bases then

$$\hat{V} = \sum_{\Gamma q_{\Gamma}} V_{\Gamma, q_{\Gamma}} \sum_{m=1}^{d_{\Gamma}} |\Gamma, q_{\Gamma}, m\rangle \langle \Gamma, q_{\Gamma}, m|. \tag{4.74}$$

Because of the potential for accidental degeneracy we can not guarantee that  $V_{\Gamma,q_{\Gamma}} \neq V_{\Gamma',q_{\Gamma'}}$  for different  $\Gamma$  and  $\Gamma'$ . Inserting this expression for  $\hat{V}$  into the matrix elements in Eq. (4.73) lets us write them in terms of IR basis vectors in the momentum spin function representation:

$$V_{kk';s_{1}s_{2}s_{3}s_{4}} = \sum_{\Gamma} V_{\Gamma,q_{\Gamma}} \sum_{m=1}^{d_{\Gamma}} \Psi_{s_{1}s_{2}}^{\Gamma,q_{\Gamma}}(\mathbf{k}) \Psi_{s_{3}s_{4}}^{\Gamma,q_{\Gamma}}(-\mathbf{k}')^{\dagger}, \qquad (4.75)$$

where

$$\Psi_{s_1 s_2}^{\Gamma, q_{\Gamma}}(\mathbf{k}) = \langle \mathbf{k}, s_1 | \langle -\mathbf{k}, s_2 \mid \Gamma, q_{\Gamma}, m \rangle. \tag{4.76}$$

We can separate the set of different IR bases into bases that have vectors that transform either symmetrically or anti-symmetrically with respect to the group element of space inversion P. We call the representations of such bases even or odd representations. Even representations are those that map P to the identity operator  $\mathbbm{1}$  and as a consequence have  $\Psi^{\Gamma,q_{\Gamma},m}_{s_1s_2}(-\mathbf{k})=\Psi^{\Gamma,q_{\Gamma},m}_{s_1s_2}(\mathbf{k})$ . Writing the spin-indices in these functions in terms of Pauli matrices by using the expansion in

Eq. (4.8) and using the fermionic symmetry then even representations a have

$$\Psi_{s_1 s_2}^{a,q_a,m}(\mathbf{k}) = \psi_{\mathbf{k}}^{a,q_a,m} i \sigma_{s_1 s_2}^y. \tag{4.77}$$

Odd representations b map P to the inversion operator I such that  $\Psi^{b,q_b,m}_{s_1s_2}(-{m k})=-\Psi^{b,q_b,m}_{s_1s_2}({m k}).$  Expanding in Pauli matrices then yields

$$\Psi_{s_{1}s_{2}}^{b,q_{b},m}(\mathbf{k}) = \mathbf{d}_{\mathbf{k}}^{b,q_{b},m} \cdot (\boldsymbol{\sigma} i \sigma^{y})_{s_{1}s_{2}}. \tag{4.78}$$

Separating the sum over IRs  $\Gamma$  into sums over even (a) and odd (b) representations in the potential operator matrix elements in Eq. (4.75), we arrive at the fully expanded expression

$$\begin{split} V_{\mathbf{k}\mathbf{k}';s_{1}s_{2}s_{3}s_{4}} &= \sum_{aq_{a}} V_{a,q_{a}} \sum_{m=1}^{d_{a}} \psi_{\mathbf{k}}^{a,q_{a},m} i \sigma_{s_{1}s_{2}}^{y} \left(\psi_{-\mathbf{k}'}^{a,q_{a},m} i \sigma_{s_{3}s_{4}}^{y}\right)^{\dagger} \\ &+ \sum_{bq_{b}} V_{b,q_{b}} \sum_{m=1}^{d_{b}} \left(\mathbf{d}_{\mathbf{k}}^{b,q_{b},m} \cdot \boldsymbol{\sigma} i \sigma^{y}\right)_{s_{1}s_{2}} \left[ \left(\mathbf{d}_{-\mathbf{k}'}^{b,q_{b},m} \cdot \boldsymbol{\sigma} i \sigma^{y}\right)_{s_{3}s_{4}} \right]^{\dagger}. \end{split} \tag{4.79}$$

In this use of the dagger notation, the adjoint acts on both of the matrix indices such that  $d_{-k}^\dagger=d_k^*$  and  $\sigma_{s_1s_2}^\dagger=\sigma_{s_2s_1}^*$ .

CHAPTER 5

# **Lattice Models**

When we have a model for the free energy of a statistical mechanical system that is too complicated to calculate analytically one approach is to utilize computers and Monte-Carlo (MC) techniques. To make numerical methods more effective one common approach is to discretize continuous models down on a numerical lattice. The lattice can in principle be of any form as long as the continuum limit reproduces the original theory, however in this thesis we will exclusively focus on a square (cubic) numerical lattice due to its simplicity.

In this chapter we will introduce different aspects of discretizing a continuous free-energy model down on a square numerical lattice. If starting with a continuous model with a spatially dependent field  $f(\boldsymbol{r})$ , then the discretized model will have a corresponding field  $f_{\boldsymbol{r}}$  only defined on the numerical lattice point sites at

$$\mathbf{r} = \sum_{\mu} r_{\mu} \hat{\mu} = \sum_{\mu} a_{\mu} n_{\mu} \hat{\mu} \tag{5.1}$$

where  $a_{\mu}$  is the distance between lattice sites,  $n_{\mu} \in [0,1,\dots,N_{\mu}-1]$  and  $N_{\mu}$  is the total number of sites in the  $\mu$ -direction. The length of the numerical lattice in this direction is  $L_{\mu} = a_{\mu}N_{\mu}$ . The cubic numerical lattice is specified by  $a_{\mu} = a \ \forall \mu$  and  $\mu \in \{x,y,z\}$ . Any integrals  $\int \! \mathrm{d}^3 r \ F[f(r)]$  will in such discretization have to be replaced with sums

such that

$$\int d^3r \mapsto a^3 \sum_{r_u=a}^{Na} . \tag{5.2}$$

If we are interested in bulk properties of the model in the thermodynamic limit, then specifying realistic boundary conditions of the numerical lattice are of less importance. In this case, periodic boundary conditions are from a computational perspective a convenient choice, which are defined by the requirement that  $f_{r+L_{\mu}\hat{\mu}}=f_r$  for any direction  $\hat{\mu}$ .

# 5.1 Discretizing derivatives

In a model where fields only are defined at discrete points in space, any spatial gradient of the fields must take the form of discrete differences of the field values at these points. In a cubic grid of points with defined field-values such differences can be denoted by the forward difference operator  $\Delta_{\mu}$ , which acts on a spatially discrete function  $f_{r}$  as a forward difference in the direction of  $\hat{\mu}$  such that

$$\Delta_{\mu} f_{\mathbf{r}} = f_{\mathbf{r} + a\hat{\mu}} - f_{\mathbf{r}},\tag{5.3}$$

where a is the distance between lattice points. In a Euclidean geometry, then the natural discretization of a derivative  $\partial_{\mu}$  is  $\partial_{\mu} \mapsto \Delta_{\mu}/a$ , which reproduces the continuum derivative in the limit  $a \to 0$  with a fixed grid-size. Using an appropriate set of units, we in most cases can set a=1.

#### 5.1.1 Covariant derivatives

When discretizing continuous gauge theories, some extra care has to be taken when discretizing a covariant derivative. Because of the gauge field, the geometry is no longer naively Euclidean such that we have to rotate a field by a gauge group element to compare the field value at two spatially separate points. Given a U(1) gauge symmetry with gauge field components  $A_{\mu}(\boldsymbol{r})$ , the appropriate way of discretizing a covariant derivative is the identification [73]

$$D_{\mu}f(\mathbf{r}) = \left[\partial_{\mu} + igA_{\mu}(\mathbf{r})\right]f(\mathbf{r}) \mapsto \frac{1}{a}(f_{\mathbf{r}+a\hat{\mu}}U_{\mathbf{r},\mu} - f_{\mathbf{r}}). \tag{5.4}$$

The value of f at  $r + a\hat{\mu}$  is parallel transported back to r by the U(1) group element [74]

$$U_{r,\mu} = e^{igA_{r,\mu}},\tag{5.5}$$

where g is the coupling constant between f and the gauge field A, and

$$A_{\boldsymbol{r},\mu} \equiv \int_{\boldsymbol{r}}^{\boldsymbol{r}+a\hat{\mu}} d\boldsymbol{r} \cdot \boldsymbol{A}(\boldsymbol{r}), \tag{5.6}$$

is a link-variable, linking r to its nearest neighbors.

In the limit of  $a \to 0$  this identification reproduces the covariant derivative. Furthermore it produces terms that transform in an analogous way to the continuum version under gauge transformations such that gauge invariant terms remain invariant after discretization. In the continuous fields, a gauge transformation is defined by

$$f(\mathbf{r}) \to f(\mathbf{r})e^{i\phi(\mathbf{r})},$$

$$A_{\mu}(\mathbf{r}) \to A_{\mu}(\mathbf{r}) - \frac{1}{g}\partial_{\mu}\phi(\mathbf{r}).$$
(5.7)

Then the covariant derivative transforms as  $D_\mu f({m r}) o e^{i\phi({m r})} D_\mu f({m r})$  such that terms such as  $|D_\mu f({m r})|^2$  are invariant under gauge-transformations. Inserting the gauge transformation into the discretized field  $f_{m r}$  and the definition of the link variables  $A_{{m r},\mu}$  we see that these discretized fields transform as

$$\begin{split} f_{\boldsymbol{r}} &\to f_{\boldsymbol{r}} \, e^{i\phi_{\boldsymbol{r}}}, \\ A_{\boldsymbol{r},\mu} &\to A_{\boldsymbol{r},\mu} - \frac{1}{g} \Delta_{\mu} \phi_{\boldsymbol{r}}, \end{split} \tag{5.8}$$

where the field  $\phi_{r}$  is discretely defined on the same lattice points as  $f_{r}$ . Inserting this into the discretization of the covariant derivative on the right hand side of Eq. (5.4) we see that indeed the right hand side transforms in the same way as the left, i.e. by picking up an overall factor  $e^{i\phi_{r}}$ . This means that the discretized version of terms such as  $|D_{\mu}f(r)|^{2}$ , which were originally gauge-invariant, will remain invariant after discretization under the transformation in Eq. (5.8).

<sup>1.</sup> To show this, we see from Eq. (5.6) that  $A_{{m r},\mu} o a A_{\mu}({m r})$ , expand the exponential in  $U_{{m r},\mu}$  to first order and insert on the right hand side of Eq. (5.4).

### **5.1.2 Reduction of symmetry**

One word of caution in this connection is that the discretized gradient terms will not necessarily have all the same spatial symmetries as the originating continuous terms because the bias of the forward direction in the forward difference and the cubic structure of the numerical lattice will in general break such symmetries. The effects of the cubic symmetry of the numerical lattice can be thought of as caused by implicit lattice potentials that increases in influence towards lower temperatures and higher field strengths when the model contains an external field . Such lattice potentials can e.g. cause topological defects to have preferred positions in discretizations of theories with translational symmetry. As an example, consider again the discretization of the term  $\int\! {\rm d}^3 r \, |{\bf D} f({\bf r})|^2$  with discretized scalar field  $f_{\bf r}=\rho_{\bf r}e^{i\theta_{\bf r}}$ . The density term is rotationally symmetric  $^3$  in 3D, however the discretized version can be written

$$2\sum_{\bm{r}}\sum_{\mu}\rho_{\bm{r}}^2[1-\cos(\Delta_{\mu}\theta_{\bm{r}}+gA_{\bm{r},\mu})]. \tag{5.9}$$

From this form we can see that the term is only symmetric by rotation of  $90^{\circ}$  in the planes normal to the x, y and z directions. The discretized term contains cubic distortions when rotating in directions in-between these, hence the SO(3) rotational symmetry is broken down to the octahedral point group O.

The forward bias of the forward difference discretization scheme can also lead to breaking of symmetries that both the continuous model and the numerical lattice have in common when the scalar field consists of multiple components. Consider a density term of the form

$$\Re \left[ D_x \eta_x D_y \eta_y \right], \tag{5.10}$$

where  $\eta_x$  and  $\eta_y$  are two scalar fields that transform as components of spin such that under a  $90^\circ$   $\hat{z}$ -rotation (which is called a  $C_4$  transformation), then  $D_x \to D_y$ ,  $D_y \to -D_x$ ,  $\eta_x \to \eta_y$  and  $\eta_y \to -\eta_x$ . Inserting this into the continuous density term in Eq. (5.10) we see

<sup>2.</sup> More on this in Section 7.

<sup>3.</sup> By rotationally symmetric we mean that if we were to rotate the field configurations of f(r) and A(r) in any direction, by any amount, the term would still yield the same value.

that it remains precisely the same i.e. invariant. Now consider the discretization of this term, which reads

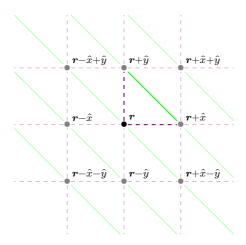
$$\begin{split} &\rho_{\boldsymbol{r}+\hat{x}}^{x}\rho_{\boldsymbol{r}+\hat{y}}^{y}\cos\left[\theta_{\boldsymbol{r}+\hat{x}}^{x}-\theta_{\boldsymbol{r}+\hat{y}}^{y}+g(A_{\boldsymbol{r},x}-A_{\boldsymbol{r},y})\right]\\ &-\rho_{\boldsymbol{r}+\hat{x}}^{x}\rho_{\boldsymbol{r}}^{y}\cos(\theta_{\boldsymbol{r}+\hat{x}}^{x}-\theta_{\boldsymbol{r}}^{y}+gA_{\boldsymbol{r},x})\\ &-\rho_{\boldsymbol{r}}^{x}\rho_{\boldsymbol{r}+\hat{y}}^{y}\cos(\theta_{\boldsymbol{r}}^{x}-\theta_{\boldsymbol{r}+\hat{y}}^{y}-gA_{\boldsymbol{r},y})\\ &+\rho_{\boldsymbol{r}}^{x}\rho_{\boldsymbol{r}}^{y}\cos(\theta_{\boldsymbol{r}}^{x}-\theta_{\boldsymbol{r}}^{y}), \end{split} \tag{5.11}$$

where we have used the notation  $\eta^a_{m r}=\rho^a_{m r}e^{i\theta^a_{m r}}$  for the discrete scalar fields. In terms of these scalar fields and link variables, a  $C_4$  transformation consists of the mappings  $\eta^x_{m r}\to\eta^y_{C_4{m r}},\eta^y_{m r}\to-\eta^x_{C_4{m r}}$  such that e.g.  $\rho^x_{{m r}+\hat{x}}\to\rho^y_{{m r}'+\hat{y}}$ . The link-variables transform as  $A_{{m r},\mu}\to A_{C_4{m r},C_4\mu}$  such that e.g.  $A_{{m r},y}\to A_{{m r}',-x}=-A_{{m r}'-\hat{x},x}$ . Using these transformations, and shifting the summation index of the external  ${m r}$ -sum, then the rotated discrete terms take the form

$$\begin{split} &-\rho_{\boldsymbol{r}-\hat{x}}^{x}\rho_{\boldsymbol{r}+\hat{y}}^{y}\cos\left[\theta_{\boldsymbol{r}-\hat{x}}^{x}-\theta_{\boldsymbol{r}+\hat{y}}^{y}-g(A_{\boldsymbol{r},y}+A_{\boldsymbol{r}-\hat{x},x})\right]\\ &+\rho_{\boldsymbol{r}}^{x}\rho_{\boldsymbol{r}+\hat{y}}^{y}\cos(\theta_{\boldsymbol{r}}^{x}-\theta_{\boldsymbol{r}+\hat{y}}^{y}-gA_{\boldsymbol{r},y})\\ &+\rho_{\boldsymbol{r}-\hat{x}}^{x}\rho_{\boldsymbol{r}}^{y}\cos(\theta_{\boldsymbol{r}-\hat{x}}^{x}-\theta_{\boldsymbol{r}}^{y}-gA_{\boldsymbol{r}-\hat{x},x})\\ &-\rho_{\boldsymbol{r}}^{x}\rho_{\boldsymbol{r}}^{y}\cos(\theta_{\boldsymbol{r}}^{x}-\theta_{\boldsymbol{r}}^{y}), \end{split} \tag{5.12}$$

which certainly is not the same as Eq. (5.11), i.e. the discretization of Eq. (5.10) is not invariant under a  $C_4$  rotation. A more immediate way of seeing the problem is to recognize that the first term in Eq. (5.11) is a next-nearest neighbor coupling on the numeric lattice that only couples sites along one diagonal, but not the other as illustrated in Figure 5.1, thus rotational symmetry is broken by the discretization.

One remedy for this kind of problem is to re-establish the broken symmetry by an average over symmetry-transformed terms. In the case of the discretization of  $\mathfrak{R}\big[D_x\eta_xD_y\eta_y\big]$  in Eq. (5.11), the  $C_4$  symmetry can thus be re-established by taking the average of Eq. (5.11), Eq. (5.12), as well as the terms obtained by transforming the discretization in Eq. (5.11) by the rotations  $C_4^2$  and  $C_4^3$ . Let  $\mathcal{F}_r$  denote the density terms in Eq. (5.11) and let  $\mathcal{T}\mathcal{F}_r$  be the terms that result when transforming  $\mathcal{F}$  by a symmetry transformation  $\mathcal{T}$ . The average of symmetry-transformed terms that re-establishes the  $C_4$  rotational symmetry can



**Figure 5.1:** Couplings between sites on a single *z*-layer of the numerical lattice from the discretization in Eq. (5.11) of the term  $\Re[D_x\eta_xD_y\eta_y]$ . On-site terms are illustrated by a point ( $\bullet$ ), while nearest neighbor and next-nearest neighbor couplings are illustrated by dashed and solid lines respectively. The couplings obtained by evaluating Eq. (5.11) at a point r are emphasized by being less transparent than the rest.

then be written

$$\begin{split} \mathcal{F}_{\pmb{r}}^{\text{sym}} &= \frac{1}{4} \sum_{\mathcal{T} \in \{1, C_4, C_4^2, C_4^3\}} \mathcal{T} \mathcal{F}_{\pmb{r}} \\ &= \frac{1}{4} \sum_{hh' = \pm 1} hh' \rho_{\pmb{r} + h\hat{x}}^x \rho_{\pmb{r} + h'\hat{y}}^y \cos \left[ \theta_{\pmb{r} + h\hat{x}}^x - \theta_{\pmb{r} + h'\hat{y}}^y + g(A_{\pmb{r}, hx} - A_{\pmb{r}, h'y}) \right]. \end{split} \tag{5.13}$$

In this expression, the next-nearest neighbour couplings are along all diagonals around the point r such that it is rotationally symmetric under  $C_4$  and thus does not explicitly break symmetries that both the original theory and the numerical lattice have in common. Such breaking of symmetries can, as we have shown above, result from the naive application of a forward difference discretization scheme when discretizing terms with multiple gradient-directions and components. In

<sup>4.</sup> The symmetric expression in Eq. (5.13) can be more easily obtained by the use of a different discretization procedure than the discretization of the covariant derivative in Eq. (5.4). Taking the average of a forward and backward difference

models with such terms, the symmetry averaged expression can then be useful in diminishing the effect of meta-stable states and faster convergence when investigating such models by means of numerical computation. Finally we would like to stress that both versions yield the same theory in the continuum limit.

### 5.2 Including an external field

The interaction between superconductors and magnetic fields is an essential aspect in the study of superconductors and thus we will need to be able to add external magnetic fields to our models in order to study this interaction. An external field is usually thought of as a constant homogeneous magnetic field in a certain direction that is a parameter of the problem rather than a variable. In other words we assume the magnetic flux to be the same everywhere and unchanging, and rather than ask what consequence the existance of a superconductor has on this field, we are interested in the effects the field has on the superconducting state. Physically this situation is relevant e.g. if a relatively small and thin sheet of superconducting material is placed in between two strong electromagnets as illustrated in Figure 5.2.

One way to introduce a constant magnetic field in a lattice model is simply to figure out what kind of vector potential  $\boldsymbol{A}(\boldsymbol{r})$  would give a constant magnetic field  $\boldsymbol{B}(\boldsymbol{r})$  through  $\boldsymbol{B} = \nabla \times \boldsymbol{A}$ , and then set the link-variables  $A_{\boldsymbol{r},\mu}$  accordingly with the only caveat being that for a lattice-model with periodic boundary conditions, then the factor  $e^{igA_{\boldsymbol{r},\mu}}$  has to satisfy periodic boundary conditions as well. This implies the condition

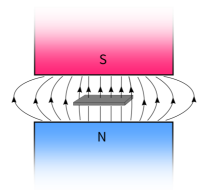
$$\forall \nu \ A_{r,\mu} = A_{r+L_{\nu}\hat{\nu},\mu} + 2\pi m_{\nu}/g,$$
 (5.14)

where  $m_{\nu} \in \mathbb{Z}$  and  $\nu$  gives a direction on the lattice.

that respects gauge-transformations we get the discretization mapping

$$D_{\mu}f(\boldsymbol{r})\mapsto (e^{igA_{\boldsymbol{r},\mu}}f_{\boldsymbol{r}+\hat{\mu}}-e^{igA_{\boldsymbol{r},-\mu}}f_{\boldsymbol{r}-\hat{\mu}})/2.$$

Applying this symmetrized covariant discretization mapping to the density term in Eq. (5.10), yields Eq. (5.13).



**Figure 5.2:** A thin sheet of superconducting material in the magnetic field produced by two magnets pointing in the same direction above and below the superconductor.

### 5.2.1 Landau Gauge

As an example, let's say we are interested in having an external field in the  $\hat{z}$ -direction with magnitude B. The vector-potential components  $A_x$  and  $A_y$  then have to satisfy the equation

$$\partial_x A_y - \partial_y A_x = B. (5.15)$$

One configuration of the vector potential, which is called the  $\mathit{Landau}$   $\mathit{gauge}$ , that satisfies this condition is  $A_y = Bx$  with the other vector potential components set to zero. Inserting this into the definition of the link-variables in Eq. (5.6) yields  $A_{r,\mu} = ar_x B \delta_{\mu,y}$ . Here x is a continuous variable while  $r_x$  is the x-component of a lattice vector. Periodic boundary conditions on the lattice implies the condition  $A_{r,y} = A_{r+L_x\hat{x},y} - 2\pi m/g$  which finally restricts the value of the field B such that the link-variables in the Landau gauge must take the form

$$A_{r,\mu} = \delta_{\mu,y} r_x \frac{2\pi m}{gL_x}, \quad m \in \mathbb{Z}, \tag{5.16}$$

where  $L_x=N_xa$  and  $N_x$  is the number of lattice sites in the x-direction. With this link-variable configuration, the field strength becomes  $B=2\pi f/ga^2$  where we have defined the filling fraction  $f=m/N_x$  which in terms of vortices gives the number of magnetic vortices pr. plaquette of the numerical lattice.

### 5.2.2 Symmetric Landau gauge

The Landau gauge has the disadvantage that it singles out a direction in the xy-plane since the vector potential is set to  ${\bf A}({\bf r})=Bx\hat{y}$  and thus only spatially dependent in the x-direction. It could because of this be argued to break a rotational symmetry of the model in the xy-plane given that Aharenov-Bohm-like effects are significant to the results. To mitigate any such concern, one can consider a symmetric gauge given by the choice  ${\bf A}({\bf r})=-{\bf r}\times B\hat{z}/2$ , which is rotationally symmetric in the xy-plane and, like the Landau gauge, produces the field  ${\bf B}=B\hat{z}$ . Inserting this choice of vector potential into the link variables yields, using implicit summation over repeated indices,  $A_{{\bf r},\mu}=\epsilon_{\mu z\nu}r_{\nu}aB/2$ . Periodic boundary conditions in this case implies two restrictions on the field value B because the vector potential varies in both the x- and y-direction. Implementing these conditions we can write the link-variables as

$$A_{\mathbf{r},\mu} = \epsilon_{\mu z\nu} r_{\nu} \frac{2\pi m}{gL_{\tau}},\tag{5.17}$$

where m is a number  $m \in \mathbb{Z}$  chosen such that there exists some  $n \in \mathbb{Z}$  such that  $mN_y = nN_x$ , i.e. m is some multiple of  $N_x/N_y$ . Then the field value is given by  $B = 2\pi f/ga^2$  for filling fraction  $f = 2m/N_x$ .

This gauge is a specification of the more general extended Landau gauge [62, 75], which is borne purely out of the assumptions of a field  $B\|\hat{z}, A(r)$  linear in r and periodic boundary conditions.

# 5.2.3 Fluctuating field

For a normal strongly type-II superconductor, the London penetration depth  $\lambda$  is much larger than the superconducting coherence length  $\xi$ . In this regime it is valid to neglect spatial fluctuations in the gauge field since any deviation around the extremal field configuration is strongly suppressed. This is called the frozen gauge approximation and makes the vector potential act only as a constraint on the value of the uniform magnetic induction given by one of the gauges presented in the above sections [62]. When the superconducting state consists of multiple components on the other hand, it becomes difficult to classify it simply in terms of type-I or type-II based solely on  $\lambda$  and  $\xi$  [76]. With multiple components, it becomes essential to fluctuate the gauge field,

because it mediates a significant indirect interaction between the components [64, 77].

Fluctuations of the gauge field imparts an energy cost on the system given in SI-units by the free energy  $^5$ 

$$F_A = \frac{1}{2\mu_0} (\nabla \times \mathbf{A})^2. \tag{5.18}$$

There are a couple of different ways of discretizing this energy for inclusion in a lattice model depending on whether one defines the link-variables compactly, i.e.  $gA_{r,\mu} \in (-\pi,\pi)$ , or non-compactly, i.e.  $gA_{r,\mu} \in (-\infty,\infty)$ . Both versions belong to the same universality class and thus produce the same results in a renormalization group sense as long as the fluctuations are sufficiently small [73]. For non-compact link-variables, we simply replace the gradient with the lattice difference operator from Eq. (5.3) divided by the lattice spacing a, and the gauge-field components by their corresponding link-variables such that

$$\partial_{\mu} \mapsto \Delta_{\mu}/a_{\mu}, 
A_{\mu}(\mathbf{r}) \mapsto A_{\mathbf{r},\mu}/a_{\mu}.$$
(5.19)

The discretized free energy pr. lattice site then becomes

$$F_{A,r} = \frac{(\mathbf{\Delta} \times \mathbf{A_r})^2}{2\mu_0 a^4} = \frac{1}{2\mu_0 a^4} \sum_{\mu} (A_{r,\mu}^{\square})^2,$$
 (5.20)

where we have defined the link-variable plaquette-sum vector, with components given by

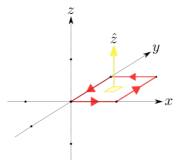
$$A_{\mathbf{r},\mu}^{\square} = \epsilon_{\mu\alpha\beta} \Delta_{\alpha} A_{\mathbf{r},\beta} = \oint_{\square_{\mu}} d\mathbf{r}' \cdot \mathbf{A}(\mathbf{r}'). \tag{5.21}$$

In the line-integral on the right hand side, the curve  $\boxdot_{\mu}$  is given by a plaquette<sup>6</sup> normal to the vector  $\hat{\mu}$ , starting at the lattice point at r and

<sup>5.</sup> One way of deriving said energy is to start with the sourceless Maxwell Lagrangian for a massless vector field  $\mathcal{L}_M=-F^{\mu\nu}F_{\mu\nu}/4\mu_0.$  In this relativistic notation  $F^{\mu\nu}=\partial^\mu A^\nu-\partial^\nu A^\mu,\,A^0=V,\,\partial_0=\partial_t/c$  and we use the metric  $g^{\mu\nu}=\mathrm{diag}(1,-1,-1,-1).$  Assuming time-independence and neglecting terms consisting only of V since they do not couple to the Higgs fields (e.g. the superconducting components) in minimal coupling, then the Lagrangian reduces to  $\mathcal{L}_M\to -(\nabla\times \mathbf{A})^2/2\mu_0$  and the free energy in Eq. (5.18) results.

<sup>6.</sup> In this context a plaquette is a square given by 4 neighboring lattice points contained in some plane.

moving along the square following the right hand rule. The integration curve given by the plaquette  $\boxdot_z$  is shown in Figure 5.3.



**Figure 5.3:** Integration path defined as  $\Box_z$  along a plaquette of the numerical lattice in the xy-plane.

To impose an external field on a system with a fluctuating field we divide the link-variables into a fluctuating part  $A_{r,\mu}^f$  with periodic boundary conditions and a constant part  $A_{r,\mu}^0$  such that  $A_{r,\mu} = A_{r,\mu}^f + A_{r,\mu}^0$ . The field is then imposed by setting the constant part such that there is a net field induction through the system, e.g. by setting it to one of the gauges in Section 5.2.1 and 5.2.2.

Chapter 6

# **Monte-Carlo Techniques**

In this chapter we discuss some techniques useful in MC simulations of systems in statistical-physics. In such systems these techniques will be used to calculate thermal averages using random numbers. Let Z denote the partition function and  ${\mathcal H}$  the Hamiltonian of the system. Then the thermal average of an observable  ${\mathcal O}$  is defined as

$$\langle \mathcal{O} \rangle = \frac{1}{Z} \sum_{\psi} \mathcal{O}(\psi) e^{-\beta \mathcal{H}(\psi)},$$
 (6.1)

where  $\psi$  denotes states of the system and we thus sum over all possible states. In the case of a quantum system, then this sum turns into a multi-dimensional integral over quantum coherent states. Now any attempt at estimating these integrals through an interpolation scheme is destined to fail because if we divide a 1-dimensional integral into M pieces and the error of the interpolation scheme scales as  $\sim M^{-\kappa}$ , then applied to a d-dimensional integral, its error will scale as  $M^{-\kappa/d}$ . What MC techniques then provides is a way of using random numbers in calculating Eq. (6.1) without actually summing over all the states. We do this by drawing random states  $\psi_i$  from a carefully selected probability distribution and using statistics to estimate how close the resulting thermal average is likely to be the true thermal average. Letting M be the number of samples, then the error scales as  $M^{-1/2}$  and is independent of the number of dimensions of the integral.

As in the case of the stationary phase approximation, the calculation of the sum in Eq. (6.1) can be made much more effective by considering which terms give large contributions. If we have a probability distribution  $\pi(\psi)$  of sampled states  $\psi$  that has peaked at states that gives large contributions to  $\langle \mathcal{O} \rangle$ , then our estimate will converge much quicker to the true value than if we were to sample states uniformly. In a sense we are interested in sampling only the important states, and hence this is called importance sampling. Let  $\{\tilde{\psi}_i\}$  be a set of states that are uniformly sampled, while  $\{\psi_i\}$  are sampled with probability distribution  $\pi(\psi_i)$ . The statistical estimator  $\langle \bar{\mathcal{O}} \rangle$  of the thermal average of the observable  $\mathcal{O}$  is then

$$\langle \bar{\mathcal{O}} \rangle = \sum_{i} \mathcal{O}(\tilde{\psi}_{i}) \frac{e^{-\beta \mathcal{H}(\tilde{\psi}_{i})}}{Z(\{\tilde{\psi}_{i}\})} = \sum_{i} \mathcal{O}(\psi_{i}) \frac{e^{-\beta \mathcal{H}(\psi_{i})}}{\pi(\psi_{i})Z(\{\psi_{i}\})}. \tag{6.2}$$

Now assuming that the state-dependence of the observable is less important than the exponential, then the largest contributions to the sum will come from states that are such that  $e^{-\beta\mathcal{H}}/Z$  is large. We thus want to pick states such that

$$\pi(\psi_i) = e^{-\beta \mathcal{H}(\psi_i)}/Z. \tag{6.3}$$

Then, given M states sampled according to this probability distribution, the statistical estimator reduces to the arithmetic average

$$\langle \bar{\mathcal{O}} \rangle = \frac{1}{M} \sum_{i} \mathcal{O}(\psi_i).$$
 (6.4)

#### 6.1 Markov-Chain Monte-Carlo method

Markov-Chain Monte-Carlo (MCMC) is a strategy of obtaining a sample of random states  $\psi_k$  where the the states are drawn sequentially in such a way that the probability  $P_k(\psi)$  of drawing a new state  $\psi_k=\psi$  is only dependent on what the last state  $\psi_{k-1}$  was. The chain developed by drawing states in this way thus has no memory of the rest of the content of the chain, except for its last link  $\psi_{k-1}$ . A chain with this property is called a Markov-Chain, hence the name.

We want the sampled states to be drawn according to the probability distribution  $\pi(\psi_k)$  discussed above. This is assured with the criteria of ergodicity and detailed balance. Ergodicity means in this context that

the states are drawn in such a way that if we were to draw infinitely many states, then we would have drawn all possible states  $\psi$  in the original sum in Eq. (6.1).

The criterion of detailed balance comes from the idea that we want the probability that a certain state is drawn to be independent of when the state is drawn in the chain. Let  $P_k(\psi)$  be the probability that  $\psi$  is drawn at the kth point in the chain. Because of the Markov-chain property, this probability is fully determined by the probability that the previous state in the chain transitions into the state  $\psi$ . Let  $\mathcal{T}(\psi' \to \psi)$  denote the probability that state  $\psi'$  transitions into state  $\psi$ , i.e. that the state  $\psi$  is drawn given a previously drawn state  $\psi'$ . Then the probability that the state drawn at the point k+1 in the chain is  $\psi$ , is given by

$$\begin{split} P_{k+1}(\psi) &= \sum_{\psi'} P(\psi_k = \psi' \ \land \ \psi' \text{ transitions to } \psi) \\ &= \sum_{\psi'} P_k(\psi') \mathcal{T}(\psi' \to \psi) \\ &= P_k(\psi) + \sum_{\psi'} \left[ P_k(\psi') \mathcal{T}(\psi' \to \psi) - P_k(\psi) \mathcal{T}(\psi \to \psi') \right], \end{split} \tag{6.5}$$

where we have used that  $\sum_{\psi'} \mathcal{T}(\psi \to \psi') = 1$  since the state must transition to some state. Now since the probability should be invariant of the point in the chain and be given by our desired probability density, we demand  $P_{k+1}(\psi) = P_k(\psi) = \pi(\psi)$  such that the sum vanishes. Because the probability density  $\mathcal{T}$  is arbitrary, the sum needs to vanish term-wise, yielding the final condition of detailed balance:

$$\pi(\psi')\mathcal{T}(\psi'\to\psi) = \pi(\psi)\mathcal{T}(\psi\to\psi').$$
 (6.6)

This states that for the process to be invariant of the point in the chain, it has to be reversible.

### 6.2 Metropolis-Hastings method

The Metropolis-Hastings (MH) method is an algorithm for drawing states in a Markov-Chain that specifies a transition probability  $\mathcal{T}(\psi \to \psi')$  between states  $\psi$  and  $\psi'$  that satisfies the detailed balance criterion. The algorithm proceeds as follows:

- 1. Given a state  $\psi_k$ , pick a new state  $\psi_p$  where the process of picking this state has an, as of now, arbitrary probability distribution denoted  $q(\psi_p \mid \psi_k)$  with the only requirement being that it leads to ergodic selection.
- 2. Accept this new proposed state  $\psi_p$ , with the probability  $\alpha(\psi_p \mid \psi_k)$ , defined as

$$\alpha(\psi_p \mid \psi_k) = \min \left\{ 1, \ \frac{\pi(\psi_p) q(\psi_k \mid \psi_p)}{\pi(\psi_k) q(\psi_p \mid \psi_k)} \right\}. \tag{6.7}$$

3. If  $\psi_p$  is accepted we set  $\psi_{k+1}=\psi_p$ . If not, then  $\psi_{k+1}=\psi_k$ . Finally return to 1. to pick the next state in the chain.

By this procedure, then the probability of transitioning between a state  $\psi$  at point k to a state  $\psi'$  at point k+1 is given by probability that the state  $\psi'$  is picked and that  $\psi'$  is accepted such that

$$\mathcal{T}(\psi \to \psi') = \alpha(\psi' \mid \psi) \, q(\psi' \mid \psi). \tag{6.8}$$

This transition probability satisfies detailed balance since inserting Eq. (6.8) and (6.7) yields

$$\pi(\psi')\mathcal{T}(\psi' \to \psi) = \min\{\pi(\psi)q(\psi' \mid \psi), \ \pi(\psi')q(\psi \mid \psi')\}$$
$$= \pi(\psi)\mathcal{T}(\psi \to \psi'). \tag{6.9}$$

#### **6.2.1 Practical considerations**

Usually, the above is a bit too general for practical implementation since we would have to calculate, or know, the probability distribution  $q(\psi' \mid \psi)$  used in picking new proposed states. If we assume q to be symmetric such that

$$q(\psi' \mid \psi) = q(\psi \mid \psi'), \tag{6.10}$$

then we don't need to calculate it explicitly since it cancels out of the equation for  $\alpha$  in Eq. (6.8).

A further simplification can be achieved by inserting the expression for  $\pi(\psi)$  in Eq. (6.3) into the q symmetric version of  $\alpha$ , which in this case reduces to

$$\alpha(\psi' \mid \psi) = \min \left\{ 1, \ e^{-\beta \left[ \mathcal{H}(\psi') - \mathcal{H}(\psi) \right]} \right\}. \tag{6.11}$$

The significance of this form is that we see the transition probability is only dependent on the difference between the energy of the updated state and the original state. If the state of the system  $\psi$  is a collection of site-dependent sub-states  $\phi(r_i)$ , e.g. how the state of an Ising chain is given by a collection of site-dependent spins, then the calculation of  $\mathcal{H}(\psi)$  has to take into account all the sites. If we update only a single site  $r_i$  of  $\psi$  to get  $\psi'$ , which we call a local MC update, then all the sites that do not have an interaction with  $oldsymbol{r}_i$  cancels out in the difference  $\mathcal{H}(\psi') - \mathcal{H}(\psi)$ . Then we only need to calculate the the difference in the sub-states that are affected by  $oldsymbol{r}_i$  to calculate the energy-difference. This is an essential property to have when creating a parallelized version of this algorithm since different parts of the lattice of sites then can be updated in an asynchronous manner without affecting each other. In other words: by simplifying to the energy difference, the update scheme becomes local which makes local MC updates grid-parallelizable.

To use pseudo-random numbers to accept a new state  $\psi'$  with probability  $\alpha$  we pick a uniformly distributed number  $r \in (0,1]$ . Then we use the fact that

$$P[r \le \alpha(\psi' \mid \psi)] = \alpha(\psi' \mid \psi), \tag{6.12}$$

so that updating the state if  $r \leq \alpha$  is equivalent to updating the state with probability  $\alpha$ . Given the form of  $\alpha$  in Eq. (6.11) then

$$r \leq \alpha(\psi' \mid \psi) \iff \ln r \leq -\beta[\mathcal{H}(\psi') - \mathcal{H}(\psi)]. \tag{6.13}$$

To update the state with probability  $\alpha$  we thus simply take the natural logarithm of r and update the state if the right hand side of Eq. (6.13) is true.

To obtain good statistics, we want, as a rule of thumb, the acceptance rate to be about 30-60% for high temperature states. The acceptance rate is defined as the number of proposed states  $\psi'$  that are accepted, divided by the total number of proposed states, and will in general be proportional to the transition probability  $\alpha(\psi'\mid\alpha)$ . This can be adjusted by changing the way new states  $\psi'$  are proposed. Let  $\psi$  be composed of site specific sub-states  $\phi(r_i)$  and let a state  $\psi'$  be proposed by

<sup>1.</sup> High temperature states refers to states that are well above any transition temperature of the system.

changing the values of the sub-state  $\phi({m r}_0)$ . Choosing values closer to the original sub-state  $\phi({m r}_o)$ , the difference  $\mathcal{H}(\psi')-\mathcal{H}(\psi)$  decreases such that  $\alpha(\psi'\mid\psi)$  in Eq. (6.11) approaches 1 and the acceptance rate increases.

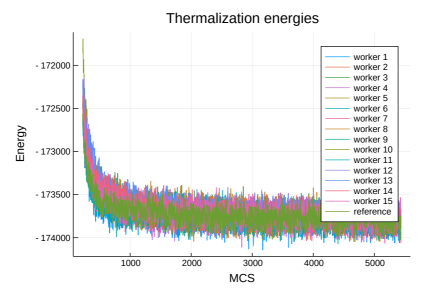
Proposing states such that the acceptance rate is very high, in this case means that the states do not change very much with each MC update. This can lead to freezing of the simulation, where the measurements do not change even after a significant number of MC updates because a large number of MC updates in a certain direction is needed to significantly change the measurements. On the other hand, too low of a acceptance rate will also freeze the simulation since then obviously states are very unlikely to change, leading to the same measurements repeatedly. Ultimately, whether the acceptance rate should be considered too high or too low, should be guided by the physics of the system since in the case that the system has reached a global minimum in the energy-landscape and has low temperature, the proper statistics is obtained by an update scheme that gives a low acceptance rate. It is not advisable to change the acceptance rate during a measurement run over decreasing temperatures as this has tended to freeze the measurements at varying temperatures leading to confusion when trying to find a transition point.

## 6.3 Thermalization procedures

Thermalization in a MCMC simulation refers to the process of discarding a number of MC updates before starting to measure the states in the Markov-chain. The reason for doing this is because the first states in the chain will usually be very unlikely states in the ensemble of states, and thus give these states an artificially high statistical weight, unless we measure long enough. That time could be very long indeed if the starting states are very unlikely, thus to get measurements in a reasonable time, these unlikely starting states are discarded.

How many states to discard is usually estimated with the help of an energy vs. Monte-Carlo sweep (MCS)<sup>2</sup> plot as shown in Figure 6.1. Since the initial state usually has a different energy than the average

<sup>2.</sup> A Monte-Carlo sweep is a term used for attempting to update all the different sites of a system once.



**Figure 6.1:** Thermalization of a  $64^3$  single-component XY-system from random initial states to a numerical temperature T=0.5. The different curves represent the energy of different realizations of the same system initialized at different random states over several MCSs. The random initial states have a relatively high energy that stabilizes to the same value for all realizations in an exponential fashion.

energy in the Markov-chain, the energy can be seen to rapidly stabilize to the average value in such a plot.

Whether the energy stabilizes from above or below will depend on what the initial state is and what the temperature of the simulation is set to. For an ab-initio state, in which the values of the sub-states are set to uniformly distributed random values in their validity range, this normally corresponds to a very high temperature and high energy state, hence thermalizing from such a state will have the energy stabilize from above. Another possibility for an ab-initio state is some kind of mean-field minimal solution of the Hamiltonian where sub-states at different sites are correlated. In this case the energy will usually be low and the thermalization energy will stabilize from below. This option has the disadvantage that if the mean field solution lies inside some local energy minimum then simulations based on it might not be able to get out and find the global minimum, as opposed to states that are

thermalized from high energy where some might find this local minimum while others fall down in the global one. In general it is best to thermalize several independent systems from different initial conditions that yield quantitatively similar results to make it less likely that the results come from a local minimum.

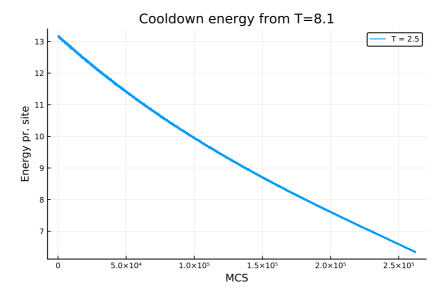
A last suggestion for an initial state of the system, is the last state of a previous simulation. In this case the thermalization will stabilize depending on the relative temperature of the two simulations. To be sure that the measurements are not correlated with the measurements of the last simulation one should discard This is a useful practice if gathering results over an extended temperature range where the systems need a large thermalization time in order to stabilize. One would then typically start measuring at a high temperature and then decrease the temperature successively in steps with a thermalization period and measure period for each step.

For systems prone to fall into local minima it was found that a more careful thermalization period analogous to the measurement procedure described above, decreased the probability of freezing into such minima. Instead of thermalizing from a high energy - high temperature state directly down to the desired temperature, which we call quenching, a cooldown period was added. During the cooldown period, the temperature was lowered stepwise from a high temperature  $T_0$  to a target temperature T with intermediate temperatures

$$T_k = \left(\frac{T}{T_0}\right)^{\frac{k}{N}} T_0. \tag{6.14}$$

The intermediate temperatures were geometrically distributed to ensure a higher density of intermediate steps at lower temperatures. At each temperature step, a fixed number of MCSs were performed such that more MCSs were done towards the lower temperature than higher. This was done because the simulations in general took longer to thermalize when the temperature decreased. An example of how the energy changed during such a thermalization period is shown in Figure 6.2.

The cooldown period was then followed by a conventional thermalization stage where the temperature was held constant at T. In most cases the energy had already stabilized at this point such that the en-



**Figure 6.2:** Energy pr. site of a  $64^3$  site model system of a p+ip chiral superconductor during the cooldown stage. The temperature is lowered as a geometric sequence and a fixed number of MCSs are done at each temperature step. Comparing with the thermalization in Figure 6.1 we see that the cooldown period gives a significantly more gradual thermalization.

ergy measurements during this extra thermalization stage typically only showed fluctuations around the mean.

# 6.4 Parallel tempering

Parallel tempering is a method of simulating multiple systems over a range of different temperatures where the systems can exchange positions with their neighbors in this temperature range according to a MH-like update step. Since all the different systems all have the same parameters except for temperature, when viewed from the perspective of a single temperature, this leads to a normal MH MCMC simulation with an occasional global update of all sites of the system whenever the current system exchanges with the system at a neighboring temperature. From the dual perspective of a single system, parallel tempering (PT) allows the system to move in temperature space.

This global updating, or movement in temperature space, has the advantage that it can prevent systems from getting stuck in local minima by allowing them to move to a higher temperature where it is easier to fluctuate to a more favorable configuration. In systems that has a jagged energy-landscape with lots of local minima this can be of great benefit and can reduce the required time it takes to measure observables with a certain accuracy by several orders of magnitude [78].

To implement parallel tempering MCMC in a temperature centric perspective, let  $\{T_i\}_{i=1}^M$  be a sorted list of M ascending temperatures, and let  $\{\lambda_i\}_{i=1}^M$  be a list of indices  $\lambda$ , that refer to replica states  $\{\psi_\lambda\}_{\lambda=1}^M$  of the system such that the replica with temperature  $T_i$  is given by  $\psi_{\lambda_i}$  and its energy is given by  $E_{\lambda_i}$ . The simulation then proceeds according to the algorithm

- 1. Perform  $\Delta t$  normal MC updates on all replica states, e.g. using the MH method.
- 2. For each replica state  $\psi_{\lambda}$  calculate the corresponding energy  $E_{\lambda}$ .
- 3. For each pair of neighboring temperatures  $T_i$  and  $T_{i+1}$  where  $T_i < T_{i+1}$ :
  - a) calculate the quantity

$$\Delta = (E_{\lambda_{i+1}} - E_{\lambda_i}) \left( \frac{1}{T_{i+1}} - \frac{1}{T_i} \right). \tag{6.15}$$

- b) Then swap the indices  $\lambda_i \leftrightarrow \lambda_{i+1}$  with probability  $\min\{1, e^{\Delta}\}$ . This can as in the MH method be done by generating a random number  $r \in [0,1)$  and then swapping indices if  $\ln r < \Delta$ .
- 4. If the replicas  $\psi_{\lambda}$  have internal knowledge of their temperatures, then distribute  $T_i$  to  $\psi_{\lambda_i}$ , for all temperatures  $T_i$ .
- 5. Sample observables and return to 1.

This algorithm is very efficiently parallelizable since the bulk of computing time will be going to doing the  $\Delta t$  MC updates which can be performed in parallel by having each replica state  $\psi_\lambda$  be assigned to a separate thread / processor. If each thread in addition keeps track of

the replicas energy at the end of the MC updates, the only information that needs to be transferred between worker processes and the process doing the PT update step is the values of the energies to the PT process, and afterwards: the set of new temperatures back to the worker processes. The PT process itself only has to calculate M-1 values and move around the indices in an array.

For the PT method to generate good statistics efficiently, some care should be taken in the distribution of the temperatures  $T_i$ . A rule of thumb is to distribute them geometrically, i.e. according to

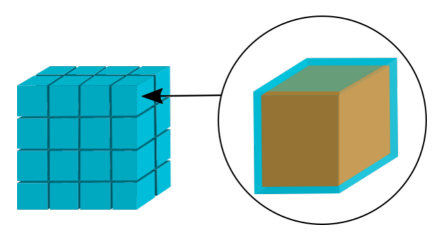
$$T_i = \left(\frac{T_M}{T_1}\right)^{\frac{k-1}{M-1}} T_1,$$
 (6.16)

with the argument that lower temperatures generally have a lower relaxation rate. With geometric distribution, the temperatures are denser towards the low end such that the acceptance rate of swaps of replicas at neighboring temperatures would in general become more flat and independent of temperature. Should the specific heat diverge at some point in the temperature range as in the case of a phase transition, then this distribution would no longer be optimal since the acceptance probability of temperature swaps is inversely proportional to  $C_v$  and thus the acceptance rate would no longer be flat.

From the perspective of an individual replica, the overall goal with the distribution of temperatures is to maximize the number of the replica moves from the lowest temperature  $T_1$ , up to the highest temperature  $T_M$  and back to the lowest temperature again. This will then maximize the number of statistically independent visits of the system at each temperature. The hope then is that a flat acceptance rate with respect to the distributed temperatures will provide a good number of round-trips. With more advanced methods such as the feedback-optimized parallel-tempering MC method the number of round-trips can be optimized [79].

#### 6.5 Grid parallelization

A simple way of utilizing multiple processor cores, or cpus on a multicpu system, is to run independent monte-carlo simulation on each processor. This is usually very efficient if a parameter of the system such as temperature is to be varied over some interval. Then each simulation



**Figure 6.3:** Illustration of the subdivision of the numerical grid into sub-cuboids. The right side illustrates that the sites in each sub-cuboid can be categorized as either internal sites, which can be updated asynchronously on different sub-cuboids and is illustrated in the figure by the internal orange cube, or sites in a border region.

could have a different value of this parameter. In this case it is recommended to also implement PT since the extra overhead is minimal and the speedup of the simulations can be significant.

Alternatively, all the simulations can run with the same value on different processes and the samples from the individual simulations can then, be combined to a super-sample. This has the advantage that the individual sampling runs can be shortened and the implementation of the parallelization is straight-forward. A drawback with this method is however that separate simulations has to be thermalized individually, such that the more the super-sample is split on different processes, the more processor time is wasted on thermalization. Additionally, if the individual simulation runs depend on some sort of freezing like how vortices freeze to the numeric lattice, then individual simulations could freeze at different angles such that they can not be combined to form good statistics.

A solution to the above issues with this simple parallelization is provided by grid-parallelization. This parallelization method is suited for simulations that consist of interacting sites either in 2D or 3D where local MC updates are to be performed. We will focus on the case of a

3D simulation where the sub-sites are organized in a numerical cubic lattice. The idea is to split the cube into different sub-cuboids as illustrated on the left side of Figure 6.3. Each sub-cuboids internal sites can then be updated in the normal MC fashion in parallel with the internal sites of other sub-cuboids. If the individual sites of the original system interacts with neighboring sites,<sup>3</sup> then it will be necessary for each subcuboid to have a border whose thickness depends on the range of the interaction. The sites in this border-region will then have an interaction with sites in the border-region of other sub-cuboids such that care has to be taken not to update a site based on the value of a neighboring site that is no longer valid. One solution to this is simply updating all the sites in the border-regions serially, i.e. update the border sites in a single sub-cuboid, communicate the updated values to the affected neighboring sub-cuboids, then move to the next sub-cuboid, etc. Or, it could be even more effective to parallelize the updates of the border sites as well by further subdividing the border-sites into different categories. As an example, one could define internal border-sites on the face of a sub-cuboid to be the border-sites that only depend on sites in the single neighboring sub-cuboid that is in the direction of the faceplane normal vector. Then all the right-facing internal border-sites can be done in parallel on all sub-cuboids, followed by all the top-facing internal border-sites, etc.

### 6.6 Reweighting

Reweighting techniques are methods of finding estimates of parameter-dependent observables at specific parameter values based on previously obtained samples of these observables from MC-simulations that have been done at parameter values independent or the reweighting parameters. Given a set of samples of some observable  $\{o_i\}_{i=1}^M$  from previous MC simulation(s), these techniques provide a set of weights  $\{w(\beta')_i\}_{i=1}^M$  that can be used to estimate the observable at a parameter value  $\beta'$  by the reweighting

$$\hat{o}(\beta') = \sum_{i} w(\beta')_{i} o_{i}. \tag{6.17}$$

<sup>3.</sup> In the case of a local MH update this can take the form of an energy-difference of the system at a proposed update-site that depends on the value of fields at neighboring sites.

This is very useful when estimating some observable over a temperature-range since then a single simulation can yield results not only for a single temperature, but for an extended region. If this region is close to a phase-transition then this can be used as a way of avoiding the critical slowing down of simulations at phase transitions by instead simulating at temperatures close to the phase-transition and then using reweighting techniques to estimate results at the phase-transition temperature. All this is possible because a simulation at a given temperature<sup>4</sup> produces a statistical distribution of energy values for the sampled states. This energy distribution will in general overlap with the energy-distribution produced when simulating at a temperature that is sufficiently close to the original. Because of this overlap, it is possible to statistically extrapolate the value of observables at the neighboring temperature.

Reweighting techniques are divided into single-histogram- and multi-histogram reweighting-techniques depending on whether they use statistical information from a single histogram or can combine histograms generated at multiple parameter values. The single-histogram techniques is used to estimate values of the observables at neighboring temperature-values and has the virtue of being comparatively simple to implement and understand. The multi-histogram techniques, on the other hand, has the advantage that the additional statistical information in general gives better estimates and is able to even give better estimates at the original temperatures the simulations were performed at, but has a more involved implementation.

In our simulations we used a Julia implementation<sup>5</sup> of the multi-histogram technique originally developed by Bojesen, a technique which they used in [80], and updated for the current Julia release by us.

## 6.6.1 Ferrenberg-Swendsen single-histogram method

To derive the Ferrenberg-Swendsen single-histogram reweighting technique, let  $o_i$  be samples of an observable  $\mathcal O$  done at sample states  $\psi_i$  of a system with Hamiltonian  $\mathcal H$  at parameter value  $\beta$ , such that  $\mathcal O(\psi_i)=$ 

<sup>4.</sup> It is possible to use reweighting techniques on other parameters of the simulation as well, as long as they are, like inverse-temperature, linear in the action.

<sup>5.</sup> This implementation is available at https://github.com/Sleort/FerrenbergSwendsenReweighting.il/tree/1.0.3\_update

 $o_i$ . Then, from out discussion of importance sampling, the estimate of the average of the observable when the states are sampled according to a probability distribution  $\pi(\psi_i)$  is given by

$$\langle \hat{\mathcal{O}} \rangle_{\beta} = \frac{\sum_{i} o_{i} e^{-\beta \mathcal{H}(\psi_{i})} / \pi(\psi_{i})}{\sum_{i} e^{-\beta \mathcal{H}(\psi_{i})} / \pi(\psi_{i})}.$$
 (6.18)

Using a simulation with importance sampling at parameter-value  $\beta$  to generate the sampled states, the probability distribution was

$$\pi(\psi_i) = \frac{e^{-\beta \mathcal{H}(\psi_i)}}{Z_\beta} = \frac{e^{-\beta \mathcal{H}(\psi_i)}}{\sum_{\psi} e^{-\beta \mathcal{H}(\psi)}},\tag{6.19}$$

where the sum in the denominator is over all possible states  $\psi$  and not sampled states. Inserting  $\pi(\psi_i)$  into  $\langle\mathcal{O}\rangle_\beta$  in Eq. (6.18), this equation reduces to the arithmetic average, however if we now imagine wanting an estimate of  $\langle\mathcal{O}\rangle_{\beta'}$  at an arbitrary parameter value  $\beta'$ , then insertion yields

$$\langle \hat{\mathcal{O}} \rangle_{\beta'} = \frac{\sum_{i} o_{i} e^{-(\beta' - \beta)\mathcal{H}(\psi_{i})}}{\sum_{i} e^{-(\beta' - \beta)\mathcal{H}(\psi_{i})}}.$$
 (6.20)

This expression then gives an estimate of the average of the observable  $\mathcal{O}$  at an arbitrary parameter value  $\beta'$  using states  $\psi_i$  that were sampled at a specific parameter value  $\beta$ . This is called the Ferrenberg-Swendsen single-histogram reweighting technique [81], and in terms of the reweighting expression in Eq. (6.17) we can read off that the weights of this technique are given by

$$w(\beta')_i = \exp\left[-\ln\left(\sum_i e^{-(\beta'-\beta)[\mathcal{H}(\psi_i)-\mathcal{H}(\psi_i)]}\right)\right]. \tag{6.21}$$

Although simple, this techniques ability to extract information about observables around the simulated parameter-value makes it extremely valuable in for instance the study of scaling relations or to accurately calculate the peak of thermodynamic variables when the MC-simulations themselves take a significant amount of computing time.

# 6.6.2 Numeric evaluation of exponential sums

The reason for introducing the extra exponential in the form of  $w(\beta')_i$  in Eq. (6.21) is because sums of exponential numbers generally are

hard to do numerically using finite-precision floating point numbers, however the logarithm of such a sum can be found using an iterative scheme. Let  $S^{(k)}$  be a sum of k exponential numbers decreasing in magnitude that presumably are too large to be stored individually such that

$$S^{(k)} = e^{a_1} + e^{a_2} + \dots + e^{a_k}, (6.22)$$

with  $a_{i+1} \leq a_i$ , is numerically hard to do. Assuming however that fractions of the numbers can be stored, then we can numerically calculate

$$\ln S^{(2)} = a_1 + \ln \left( 1 + e^{a_2 - a_1} \right). \tag{6.23}$$

Following the iteration

$$\ln S^{(k)} = \ln S^{(k-1)} + \ln \left( 1 + e^{a_k - \ln S^{(k-1)}} \right), \tag{6.24} \label{eq:6.24}$$

then  $\ln S^{(k)}$  can be found for arbitrary k without ever storing a single exponential number, only fractions of such numbers that are close to each other.

# 6.6.3 Multi-histogram Ferrenberg-Swendsen method

Let  $\{\psi_i^k\}_{i=1}^{N_k}$  be sets of states sampled at the  $N_0$  inverse temperatures  $\beta_k$  of a system with Hamiltonian  $\mathcal{H}$ . The energy of these states are then given by  $E_i^k$  and samples of an observable  $\mathcal{O}$  at these states are given by  $\mathcal{O}(\psi_i^k) = o_i^k$ . The energy-samples can then be used to construct  $N_0$  histograms

$$h_k(E) = \sum_{i=1}^{N_k} \delta_{E, E_i^k}, \tag{6.25}$$

giving the number of sampled states at a certain energy in the simulation with parameter value  $\beta_k.$  The goal is to use these histograms to estimate the density of states of the system which we for the purpose of the derivation of this method will define  $n(E) = \sum_{\psi} \delta_{E,\mathcal{H}(\psi)}.$  The essential steps in this derivation can be found in the original paper in Ref. [82], as well as Ref. [83] and [84]. With these definitions, the energetic probability distribution of the system at an inverse temperature  $\beta$  is given by

$$W(\beta, E) = n(E)e^{-\beta E}/Z_{\beta}, \tag{6.26}$$

where  $Z_{\beta}=\sum_{\psi}e^{-\beta\mathcal{H}(\psi)}$  is the partition function. Based on the sampled histograms,  $W(\beta_k,E)$  at temperature  $\beta_k$  can be estimated by  $\hat{p}_k(E)=h_k(E)/N_k$ , i.e.  $\langle\;\hat{p}_k(E)\;\rangle=W(\beta_k,E)$ . This implies that  $\langle\;h_k(E)\;\rangle=N_kW(\beta_k,E)$  and assuming for now that the samples of states  $\psi_i^k$  and  $\psi_i^k$  are statistically independent, it can be shown that

$$\langle \; h_k(E)^2 \; \rangle = N_k W(\beta_k, E) [1 + (N_k - 1) W(\beta_k, E)]. \eqno(6.27)$$

Inserting these cumulants of the histograms into the variance we get

$$\delta^2 h_k(E) = \langle \ h_k(E)^2 \ \rangle - \langle \ h_k(E) \ \rangle^2 = g_k N_k W(\beta_k, E), \eqno(6.28)$$

by assuming  $W(\beta_k,E)\ll 1$ . The factor  $g_k=1+2\tau_k$ , where  $\tau_k$  is the autocorrelation time of the samples at  $\beta_k$ , is included to generalize the result to samples where  $\psi_i^k$  and  $\psi_i^k$  are not statistically independent.

By solving Eq. (6.26) w.r.t. n(E) and inserting the estimator of  $W(\beta_k, E)$ , an estimator of the density of states is given by

$$\hat{n}_k(E) = \hat{p}_k(E) Z_{\beta_k} e^{\beta_k E}, \tag{6.29}$$

where  $Z_{\beta_k}$  is assumed known, an assumption we have to reconcile later. By the error propagation formula, then the variance of this estimator is given by

$$\delta^2 \hat{n}_k(E) = (Z_{\beta_k} e^{\beta_k E} / N_k)^2 \delta^2 h_k(E). \tag{6.30}$$

The estimator  $\hat{n}_k(E)$  is an estimator of n(E) using only a single histogram. We combine the estimators of single histograms using a weighted sum

$$\hat{n}(E) = \sum_{k} r_k \hat{n}_k(E), \tag{6.31}$$

where the coefficients  $r_k$  have to satisfy the condition  $\sum_k r_k = 1$  for the expectation value of  $\hat{n}(E)$  to give the density of states. The coefficients  $r_k$  are determined by minimizing the variance  $\delta^2 \hat{n}_k(E)$  subject to the constraint  $\sum_k r_k = 1$  using Lagrange multiplier which yields the estimator

$$\hat{n}(E) = \frac{\sum_{k=1}^{N_0} g_k^{-1} h_k(E)}{\sum_{l=1}^{N_0} N_l g_l^{-1} e^{-\beta_l E} Z_{\beta_l}^{-1}}.$$
(6.32)

The assumption that  $Z_{\beta_k}$  is known is now reconciled. Since we can write the partition function using the density of states through

$$Z_{\beta} = \sum_{\psi} e^{-\beta \mathcal{H}(\psi)} = \sum_{E} n(E)e^{-\beta E},$$
 (6.33)

then we use the density of states estimator to estimate the partition function, and use this estimate of the partition function  $\hat{Z}_{\beta_k}$  in the density of states estimator. This then creates an implicit equation for  $\hat{Z}_{\beta_k}$  that has to be solved self-consistently. Inserting the definition of  $h_k(E)$  and exchanging sums to remove the histograms, this equation takes the form

$$\hat{Z}_{\beta} = \sum_{k=1}^{N_0} \sum_{i=1}^{N_k} \frac{g_k^{-1} e^{-\beta E_i^k}}{\sum_{l=1}^{N_0} N_l g_l^{-1} e^{-\beta_l E_i^k} \hat{Z}_{\beta_l}^{-1}},$$
(6.34)

which gives  $N_0$  equations for  $N_0$  unknowns  $\hat{Z}_{\beta_k}$  when evaluated at the different  $\beta=\beta_m$ .

Solving Eq. (6.34) is usually done with the help of an iterative solution method for non-linear equations such as the Newton-Raphson method. To numerically calculate a solution, it is inconvenient to work with the full quantities  $\hat{Z}_{\beta_m}$  since these are usually extremely large. Instead it is sufficient to calculate the variables

$$L_m \equiv \ln \hat{Z}_{\beta_m} - \ln \hat{Z}_{\beta_1}, \tag{6.35}$$

since the weights in the reweighting of observables can be written in terms of them. Dividing Eq. (6.34) by  $Z_{\beta_1}$ , we get that the  $N_0-1$  equations we need to solve self consistently for the  $N_0-1$  variables  $L_m$  are given by

$$L_m = \ln \left\{ \sum_{k=1}^{N_0} \sum_{i=1}^{N_k} \frac{g_k^{-1} e^{-\beta_m E_i^k}}{\sum_{l=1}^{N_0} g_l^{-1} e^{-\beta_l E_i^k - L_l}} \right\}.$$
 (6.36)

This form has the big advantage that the overall logarithm allows us to not have to calculate the exponential sums directly, but instead only calculate logarithms of these sums. For each sum  $\sum_i e^{a_i}$  containing exponentials, which are potentially too large to be stored numerically, we simply re-exponentiate the entire sum to  $\exp \ln \sum_i e^{a_i}$  and then

use the method outlined in Section 6.6.2 to calculate  $\ln \sum_i e^{a_i}$ . Because of the overall logarithm, the exponential drops out in the last re-exponentiation such that we never have to store a single exponential number.

After finding self-consistent values for the  $N_0-1$  variables  $L_m$ , the weights  $w_i^k$  for reweighting the observable  $\mathcal O$  can be found. In terms of the density of states n(E), the thermal average of the observable is written

$$\langle \mathcal{O} \rangle_{\beta} = \frac{\sum_{E} \mathcal{O}(E) n(E) e^{-\beta E}}{\sum_{E} n(E) e^{-\beta E}}.$$
 (6.37)

Inserting the reweighting estimate of  $\hat{n}(E)$  in Eq. (6.32) for n(E) we get the reweighting estimate

$$\langle \hat{\mathcal{O}} \rangle_{\beta} = \frac{\hat{Z}_{\beta_1}}{\hat{Z}_{\beta}} \sum_{k=1}^{N_0} \sum_{i=1}^{N_k} \frac{o_i^k g_k^{-1} e^{-\beta E_i^k}}{\sum_{l=1}^{N_0} N_l g_l^{-1} e^{-\beta_l E_i^k - L_l}}, \tag{6.38}$$

where  $\hat{Z}_{\beta_1}/\hat{Z}_{\beta}$  is given by

$$\frac{\hat{Z}_{\beta}}{\hat{Z}_{\beta_1}} = \sum_{k=1}^{N_0} \sum_{i=1}^{N_k} \frac{g_k^{-1} e^{-\beta E_i^k}}{\sum_{l=1}^{N_0} N_l g_l^{-1} e^{-\beta_l E_i^k - L_l}},$$
(6.39)

through Eq. (6.34). The two equations for  $\langle \hat{\mathcal{O}} \rangle_{\beta}$  in Eqs. (6.38) and (6.39) together with the self-consistency equation in Eq. (6.36), is sufficient to describe the multi-histogram method. Notice that in these equations the histograms on which the method was derived do not figure but have been replaced by the more fundamental energy samples. This form makes the method more convenient to implement for systems with continuous energy distributions since it removes the need for a sum over all possible energies.

When calculating the exponential sums in the weights implied by Eqs. (6.38) and (6.39), numerical overflow can be avoided by first using logarithms to calculate the logarithm of a set of related un-normalized weights as before, then subtracting the maximum logarithmic value for each weight such that each weight is  $\lesssim 1$  and then using the sum of these weights to properly normalize in the end.

### 6.6.4 Initial guess

An iterative non-linear solver usually needs an initial guess at the solution. In the case of the multi-histogram method equations, a good initial guess can be provided by the single-histogram Ferrenberg-Swendsen method. Since only fractions of partition function values are needed we may set that  $\hat{Z}_{\beta_1}=1$  and use the Ferrenberg-Swendsen method based on the  $\beta_1$  energies to estimate the value  $\hat{Z}^0_{\beta_2}$  of  $\hat{Z}_{\beta_2}$  at neighboring inverse-temperature  $\beta_2$  by the formula

$$\hat{Z}_{\beta_2}^0 = \sum_{i=1}^{N_1} e^{-(\beta_2 - \beta_1)E_i^1}.$$
(6.40)

In terms of the numerically convenient variables  ${\cal L}_m$  then this first guess  ${\cal L}_2^0$  takes the form

$$L_2^0 = \ln \left[ \frac{1}{N_1} \sum_{i=1}^{N_1} e^{-(\beta_2 - \beta_1) E_i^1} \right]. \tag{6.41}$$

Continuing to estimate the partition function  $\hat{Z}_{\beta_m}$  through the single-histogram Ferrenberg-Swendsen method based on the data at  $\beta_{m-1}$ , then we may find all subsequent  $L_m^0$  by applying the iteration scheme

$$L_m^0 = L_{m-1}^0 + \ln\left[\frac{1}{N_{m-1}} \sum_{i=1}^{N_{m-1}} e^{-(\beta_m - \beta_{m-1})E_i^{m-1}}\right]. \tag{6.42}$$

	7	_
CHAPTER		

# **Vortices in superconductors**

In conventional type-I superconductors, the Meissner effect prevents any magnetic field from penetrating the superconductor when it is in the superconducting state. In a type-II superconductor, the transition between the normal- and superconducting state is more gradual than in the type-I case due to an intermediate transitional state where topological defects in the superconducting field becomes stable allowing quanta of magnetic field to pass through the material. The transitional value of the external field strength below which no magnetic field penetrates the superconductor is called  $B_{c1}$ . The upper transitional field strength above which the material stops being superconducting altogether is called  $B_{c2}$ . The state with regions of topological defects through which magnetic field quanta can penetrate, which are interspersed in a sea of superconducting state then exists between these values. It is important to note that the Meissner effect is still present in this transitional state - preventing magnetic field lines from penetrating the superconducting state, however at topological defects, the material switches to the normal state and thus allows magnetic field lines to penetrate at these points. The final continuous transition to the normal state at  $B_{c2}$  is then caused by the proliferation of vortex-loops, sending the whole material to the normal state.

The regions of normal state containing a topological defect of the superconducting state and through which magnetic field quanta can penetrate are known as superconducting vortices because they are sur-

rounded by a circulating superconducting current. This current is set up by the presence of the magnetic field and shields the rest of the superconducting condensate from its influence.

Whether a superconductor is considered type-I or type-II is conventionally given by the magnetic field penetration length  $\lambda$  and the superconducting coherence length  $\xi$  which come together to form the GL parameter  $\kappa=\lambda/\xi.$  These parameters come out of the description of the superconducting state given by the GL theory of a single-component complex field minimally coupled to a gauge field. If  $\kappa\gg 1$  then we say we have a strongly type-II superconductor, while is  $\kappa\ll 1$  the superconductor is strongly type-I. The transitional value between type-I and type-II have a theoretical mean-field value of  $\kappa=1/\sqrt{2}$ , however numerical calculations has given it the value  $\kappa=(0.76\pm0.04)/\sqrt{2}$  all within the conventional GL formalism.

In a type-II conventional superconductor without any structural defects, as we increase the field strength, we introduce more vortices into the material in order to carry the required number of magnetic field quanta. At first these vortices behave like a liquid where they mutually repel each other if they get close. As more vortices are introduced to the system, the inter-portex repulsion leads to them forming a two-dimensional lattice with equidistant lattice-spacing. Since the triangular lattice is the lattice with the highest packing fraction, i.e. the lattice that have the highest density of sites at a given lattice spacing, the lattice formed will be triangular. Such a triangular (hexagonal) lattice of single quanta vortices is known as the Abrikosov lattice since it consists of single quanta vortices which are known as Abrikosov vortices.

### 7.1 Vorticity observables

A condensate described by a complex field  $\psi$  with phase  $\theta$  can have topological defects given by discontinuities in the field  $\theta$  due to its compact nature  $(\theta \in [0,2\pi))$ . Such topological defects can be quantified by a non-zero winding-number  $N_v$  which measures how the phase  $\theta(\boldsymbol{r})$  moves around the unit circle as we change the position  $\boldsymbol{r}$  in a closed loop around the defect. These topologicals defects then lead to singularities in the field  $\nabla\theta$  which allows a nonzero value of  $\nabla\times\nabla\theta$  at

these points. Integrating over a surface S with surface normal vector  $\hat{s}$  of the system and using Stokes theorem then yields

$$\int_{S} \mathrm{d}^{2} r(\nabla \times \nabla \theta) \cdot \hat{s} = \oint_{\partial S} \nabla \theta \cdot \mathrm{d}\boldsymbol{r} = 2\pi N_{v}, \quad N_{v} \in \mathbb{Z}, \tag{7.1}$$

where  $\partial S$  is a path around the boundary of S traversed counter-clockwise. The last equality comes from the observation that  $\partial S$  is far away from the singularity such that  $\nabla \theta$  is continuous along the path and  $N_v$  thus counts the number of times the vector  $\theta$  rotates counter-clockwise back to its initial position. If there is no topological defect inside the boundary  $\partial S$  then  $\theta$  will increase as much as it decreases along the path, such that  $N_v=0$ . If a topological defect in the form of a vortex is present, then  $N_v\neq 0$  [85].  $N_v$  can then be interpreted as the total vorticity of the field  $\theta$  over the surface S. Since  $N_v$  is the total vorticity which can consist of several individual defects, then from Eq. (7.1) we see that

$$\boldsymbol{n}_v = \frac{\nabla \times \nabla \theta}{2\pi} \tag{7.2}$$

must be interpreted as a vector of local vorticity density.

If the system described above contains a gauge field that is coupled to  $\psi$ , then any meaningful observable needs to be gauge-invariant. We clearly see that the expression in Eq. (7.2) is gauge-dependent by sending  $\theta \to \theta + \phi$ . To make a gauge-invariant observable under the gauge-transformation in Eq. (5.7), we see that we need to modify the definition to

$$\boldsymbol{n}_{v} = \frac{\nabla \times (\nabla \theta + g\boldsymbol{A})}{2\pi}.$$
 (7.3)

This expression then defines  $n_v$  as a gauge invariant vector of local vorticity density of the compact field  $\theta$ .

In lattice models we want to discretize the vorticity density in Eq. (7.3) in order to effectively calculate it in MC simulations of the lattice model. In such a discrete model we have to take care to re-compactify the quantity  $\nabla \theta + g {\pmb A}$  to only be defined on some interval of length  $2\pi$ . Using the discretization mapping of  $\partial_\mu$  and  $A_\mu({\pmb r})$  from Eq. (5.19), we want  $\Delta_\mu \theta + g A_{{\pmb r},\mu} \in [-\pi,\pi)$ . Defining the operator

$$\hat{C}_{\pi} x = \operatorname{mod}(x + \pi, 2\pi) - \pi, \tag{7.4}$$

<sup>1.</sup> From vector calculus we know that for a continuously differentiable field f(r), it is the case that  $\nabla \times \nabla f = 0 \ \forall r$ .

the discretized vorticity density can be written

$$\begin{split} \boldsymbol{n}_{v,\boldsymbol{r}} &= \frac{\hat{e}_{\mu} \epsilon_{\mu\nu\lambda} \Delta_{\nu} \hat{C}_{\pi} (\Delta_{\lambda} \theta_{\boldsymbol{r}} + g A_{\boldsymbol{r},\lambda})}{2\pi a^2} \\ &= \frac{1}{2\pi a^2} \sum_{\mu} \hat{e}_{\mu} \sum_{\square_{\mu}} \hat{C}_{\pi} (\Delta_{\lambda} \theta_{\boldsymbol{r}} + g A_{\boldsymbol{r},\lambda}). \end{split} \tag{7.5}$$

Implicit summation over repeated indices is used on the first line while on the second, the components of the vector  $\boldsymbol{n}_{v,\boldsymbol{r}}$  are written as plaquettesums, which are sums of direction dependent quantities along a path  $\boldsymbol{\Box}_{\mu}$ , which is described below Eq. (5.21) and illustrated in Figure 5.3. In the plaquette-sum the directional quantity is always chosen along the path and the path is traversed according to the right hand rule with normal vector  $\hat{e}_{\mu}$  [73, 86].

If the lattice system has an external field that yields a filling fraction f, e.g. produced by one of the gauges in Section 5.2, then each plaquette-sum in Eq. (7.5) will have a contribution  $f/a^2$ . To see this assume e.g. that  $\theta_{r}$  is the same everywhere such that  $\Delta_{\lambda}\theta_{r}=0$  and insert the Landau gauge from Eq. (5.16) into the z-component of  $n_{v,r}$ . This yields  $f/a^2$ . Hence, to assure that the vortex observable yields the actual vortex quanta integer values when evaluated on a lattice with a uniform external field in the z direction with filling fraction f, we have to use the lattice function

$$n_{\boldsymbol{r}}^z = (\boldsymbol{n}_{v,\boldsymbol{r}})_z - \frac{f}{a^2}.$$
 (7.6)

If the system consists of multiple condensate components  $\psi^h = \rho^h e^{i\theta^h}$ , then we can define a separate vorticity flux density  $n_{r}^{z,h}$  for each component h by letting  $\theta \mapsto \theta^h$  in the definitions of  $n_{r}^z$  in Eqs. (7.5) - (7.6).

#### 7.2 Unconventional vortices

In a conventional superconductor, the isotropic (i.e. s-wave) nature of the superconducting state implies that stable vortices can only contain a single quanta of magnetic flux. In other words, if through some random thermal fluctuation a defect appears that contains n quanta of magnetic flux, this will soon decay into n individual vortices, which

each contain a single quanta and which we thus call single-quanta vortices. For the individual stable topological defects to contain multiple quanta of magnetic flux, the superconducting state has to be unconventional in some way. One way in which a superconductor can host stable multiple quanta vortices is if the superconducting state for some reason has an unconventional symmetry. This could e.g. be caused by unconventional (i.e. non-phononic) mechanisms of Cooper-pair formation such as van der Waahls- or spin-mediated interaction [59]. In this case multiple components might be needed in order to describe the symmetry which can result in the stabilization of vortices with double as well as fractionalized quanta [87].

A more specific example is that this can happen when a magnetic field penetrates a sample of material that is in a p+ip superconducting state, i.e. a state where the pairing function is described by two components that each have a  $k_x \pm ik_y$  dependence on the crystal momentum  ${\pmb k}$  in the continuum limit. The linear k-dependence implies a finite angular momentum of the Cooper pairs with l=1 and the phases of the components are locked by an angular momentum difference  $\Delta l=2$ . This has the consequence  $n_+=n_-+2$  on any non-trivial winding numbers  $n_+$  and  $n_-$  of the two different components which implies that if a vortex exists some place where the sub-dominant component has winding number  $n_-=0$ , then the dominant component must have  $n_+=2$  and thus the vortex must be a double quantum vortex.<sup>2</sup>

In the type of superconductor described above, the winding numbers  $n_+$  and  $n_-$  fully determine the structure of possible vortices. In the following we will use the notation  $(n_+,n_-)$  to specify these types of vortices. The possibilities for single-quanta vortices in this notation are thus the vortices (1,-1) and (-1,-3). The latter type has a higher winding number in the sub-dominant component which implies a more complex core structure and has a higher energy cost pr. vortex [88] which means that of the two, it is the (1,-1) variety that will be expected to be stable in experiments.

<sup>2.</sup> It is the winding number of the dominant component that determines the number of magnetic flux quanta that the vortex is allowed to contain because the sub-dominant component is zero in locations far away from the vortex core by the nature of being sub-dominant. Thus, it doesn't contribute to the closed loop integral in Eq. (7.1) when integrating the supercurrent in a circle around the vortex.

It is also possible to have double quanta vortices, which are either of the (2,0) or (-2,-4) variety. Again, the type of vortex with the higher winding number in the sub-dominant component exhibits a more complex core structure. For the choices of various internal parameters of such systems that we have studied, it is the (2,0) type of vortex that is associated with the lowest energy cost and thus the one that is stable [88–90].

As we have mentioned, the different types of vortices will in general have different types of core-structures even though they may permit the same number of magnetic flux quanta to penetrate. One diagnostic tool to separate different kinds of vortices is thus to observe the structure of the vortex core. Aside from plotting the actual vorticity  $n_{+}$  of the component through Eq. (7.6), this can done by e.g. plotting the amplitudes of the dominant and sub-dominant component, the phase-difference  $\theta_+ - \theta_-$  of the different components or the magnetic field in the region of the vortex core. A rendition of the essential features of plots of dominant component vorticity density  $n_{\perp}$  and phase-difference is shown in Figure 7.1 for the two vortex types (1, -1)and (2,0). We see from the figure that the double quanta vortex type (2,0) can be distinguished from the single quanta vortex by having an extended ring of vorticity density, as well as having a core region in the phase difference plot that is rotated by  $\pi/2$  radians from the asymptotic value of this phase difference. These features were used in our work to identify double and single quanta vortices in MC simulations.

#### 7.3 Ensambles of vortices

With increasing field strength, more quanta of magnetic flux will penetrate the mixed phase and thus it will contain an increasing number of vortices that form flux-lines through the system . If any structural defects are present in the system, then this leads to local suppression of the superconducting condensate such that vortices are less energetically costly, and vortices will thus be predominantly located in such regions. This is called pinning of vortices because these regions attract vortices and since their location is determined by external factors and not by the inter-vortex interactions themselves. Free vortices are mobile in response to an electric current and this leads to energy-loss and resistance in the mixed phase. Pinning regions have the effect of resist-

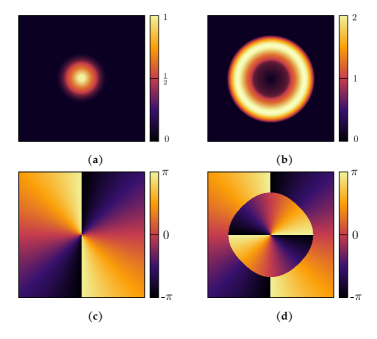


Figure 7.1: Schematic of vorticities and corresponding phase difference signature  $\theta_+ - \theta_-$  of vortices in a system with external magnetic field  $\mathbf{B} = B\hat{z}$ . a and c shows vorticity and phase-difference respectively for a singly-quantized vortex with winding number  $n_+ = 1$  and  $n_- = -1$ . b and d shows vorticity and phase difference respectively for a doubly-quantized vortex with winding number  $n_+ = 2$  and  $n_- = 0$ . The figures are directly based on the ones presented in Ref. [89].

ing such movement and thus can contribute to increasing the amount of resistance free current [91].

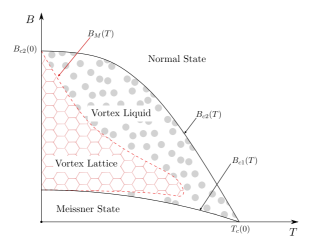
In the absence of such pinning, vortex tubes that run through the material in the direction of an external magnetic field can be ordered in a lattice according to their mutual interaction. Such a lattice is called an Abrikosov lattice or a flux line lattice since the vortex lines/tubes carry quanta of flux of the external magnetic field. The Abrikosov lattice then exists in the mixed phase of type-II superconductors and is destroyed when either the temperature or magnetic field strength is increased beyond a certain level  $B_{c2}(T)$  where the material enters the normal non-superconducting phase. This transition can be character-

ized in terms of a proliferations of unbound vortex loops which destroys the phase-coherence of the superconducting state [92].

If the interaction between the flux lines is weak compared to entropic forces such as thermal fluctuations, then fluctuations of the vortex flux lines can cause melting of the vortex lattice. In this molten state, the vortex flux lines still interact repulsively which yields an average preferred inter-vortex distance given by the balance between inter-vortex repulsion and the inclusion of the necessary number of vortices in order to carry the external magnetic field, however any directionally dependent long range correlation is lost. This corresponds to the behavior of particles in a liquid, and the molten state is thus called a vortex liquid. Such states are commonly found in high- $T_c$  superconductors such as  $YBa_2Cu_3O_{7-\delta}$ . The transition between an ordered lattice of vortices and a vortex liquid is known as a vortex lattice melting transition. This transition can also be achieved by tuning the strength of the magnetic field. This implies a magnetic field strength  $B_{c1}$ , below which the Meissner effect completely excludes all magnetic fields, and a strength  $B_M > B_{c1}$  above which the vortices behave as a liquid, i.e. without any long range correlations except that of an average distance. Finally superconductivity is destroyed at  $B_{c2} > B_M$  when a proliferation of vortex loops destroy all vortex correlations and the material enters the normal state. These different states of the vortex lattice map out a region in the B-T parameter space such as the one shown in Figure 7.2. There also exists intermediate glassy phases of vortex matter between the extremes of a completely ordered lattice and a liquid when pinning of vortices is combined with low temperature in an external magnetic field [93, 94].

The structure and behavior of a flux-line lattice in the mixed phase of a type-II superconductor is dependent on the symmetry and nature of the superconducting phase. In conventional single-component superconductors with s-wave symmetry, the vortices interact asymptotically  $^3$  through isotropic repulsion that can be modeled by a modified Bessel function of the second kid [27, 96, 97]. In a clean material, this interaction leads experimentally to a triangular (hexagonal) lattice of vortices due to this lattice symmetry having the largest packing-

<sup>3.</sup> In this context "asymptotically" means in the asymptotic limit of large separation between vortices.



**Figure 7.2:** Simplified phase diagram of the states of vortex matter in a type-II superconductor in the B-T phase space. This is based on the Lindemann criterion [95].

fraction of any two-dimensional lattice, i.e. it is the lattice that gives the highest density of vortices given a set inter-vortex distance and thus give the lowest energy configuration [98,99]. Theory predicts that also a square lattice of single quanta vortices should be possible at higher fields for  $\kappa \gtrsim 1/\sqrt{2}$  [96], however in the London-approximation, which is valid at  $\kappa \gg 1$ , the triangular symmetry is the most stable for all fields [100]. Interestingly, the vortex lattice symmetry can be incommensurate to the underlying crystal lattice structure.

In theoretical models of unconventional superconductors, such as superconductors with multiple components and non-isotropic symmetry, even more complex behavior of the mixed phase is predicted. We have already mentioned the appearance of a vortex-liquid state separate from the vortex lattice state in high- $T_c$  superconductors, which are overwhelmingly of the extreme type-II category and described by a single component unconventional d-wave symmetry. In superconductors with multiple components where each component can be modeled by a conventional London-approximation, a phase transition from the superconducting state to a superfluid state is possible in the mixed phase by melting of a composite Abrikosov vortex lattice into a state with a remaining ordered neutral mode [63].

# 7.3.1 Vortex matter in $p_x + ip_y$ -superconductors

In our work we have been specifically interested in the vortex matter of superconductors with *p*-wave symmetry. These types of superconductors are described by two components that are intrinsically coupled and give rise to unconventional composite vortices as described in Section 7.2. The stable single-quanta composite vortices, which are denoted as vortex type (1, -1) in the notation of Section 7.2, are theoretically predicted to form lattices with square symmetry [101–103]. Such symmetry has been observed in the vortex lattice of the unconventional superconductor Sr<sub>2</sub>RuO<sub>4</sub> [104–107] and is thus part of the body of evidence supporting a *p*-wave symmetry of the superconducting state of this material. The theoretical predictions are supported by numerical calculations that also show that at lower fields a triangular vortex lattice consisting of double-quanta (2,0)-vortexes is the preferred configuration [89]. Since these calculations did not account for thermal fluctuations in a convincing way, we used large-scale MC simulations to consider this effect on the vortex matter. Our results support the transition of a triangular vortex lattice consisting of doublequanta vortices to a square vortex lattice consisting of single quanta vortices at higher fields and temperatures. These results are presented in Paper II.

# 7.4 Observables of lattice symmetry

In this section we discuss two tools usable in lattice theories for considering the symmetry of vortex line lattices. These tools are based on the observables of vortex flux density discussed in Section 7.1, but in this case we are interested in measuring the structural correlations of a collection of vortex lines.

#### 7.4.1 Structure function

The structure function of a discrete cuboid system of  $N_{\mu}$  lattice sites along the  $\hat{\mu}$  direction and with local vorticity  $n_{r}^{z}$  as defined in Eq. (7.6), is defined as

$$S(\mathbf{k}_{\perp}) = \frac{1}{(fN_x N_y N_z)^2} \left\langle \left| \sum_{\mathbf{r}} a^2 n_{\mathbf{r}}^z e^{i\mathbf{k}_{\perp} \cdot \mathbf{r}_{\perp}} \right|^2 \right\rangle, \tag{7.7}$$

where a is the lattice spacing,  $r_{\perp}$  is the projected lattice vector  $r_{\perp}=r-r\cdot\hat{z}$  down on the xy-plane, and f is the filling fraction, i.e. the number of vortex quanta pr plaquette in the xy-plane. The filling fraction f relates to the inclusion of an external magnetic field in the z-direction as described in Section 7.1. This function takes a reciprocal 2D momentum vector as an argument and measures the structural correlation of the vortex lattice at this Bragg-point with normalization such that S(0)=1.

To motivate this expression, consider a continuous cuboid system with a uniform field in the z-direction with average flux density of numer of magnetic flux quanta  $\tilde{f}$  that gives rise to a lattice of vortex lines along the z-direction. Let  $n^z(r)$  be a flux density distribution of local vorticity in the z-direction such that if a vortex line with winding number  $n \in \mathbb{Z}$  goes through the point  $r_0$ , then  $\int_A \mathrm{d}^2 r \ n^z(r_0) = n$ , where A is an area that contains the vortex line. Taking the average over the z-direction keeps the value n of any vortex flux lines since they will be coherent over this dimension of the system. In contrast any contributions from vortex loops which could be the result of random thermal fluctuations will vanish in the limit of a large system size, hence

$$w(\mathbf{r}_{\perp}) = \int_0^{L_z} \mathrm{d}r_z \; n^z(\mathbf{r})/L_z, \tag{7.8}$$

filters out the vortex lines from the thermal noise. This w produces a distribution of vortex lines over the extent of the system in the xy plane. Since we are interested in structural correlations in this system we perform the 2D Fourier transform and look at its amplitude through

$$\tilde{S}(\mathbf{k}_{\perp}) = \left| \int d^2 r \ w(\mathbf{r}_{\perp}) e^{i\mathbf{k}_{\perp} \cdot \mathbf{r}_{\perp}} \right|^2.$$
 (7.9)

This function then produces a reciprocal lattice of the 2D lattice of vortex lines where Bragg-points that correspond to structural correlations have increased value. To arrive at the structure-function we need only now to take the thermal average to average over thermal fluctuations of the vortex lattice lines and normalize such that S(0)=1. To find this normalization constant, we have to calculate the integral

$$\int d^2r \, n^z(\mathbf{r}) = ?, \tag{7.10}$$

over the systems extent in the xy plane. However, from the definition of  $n^z({\bf r})$  the answer is given to us. Since  $n^z({\bf r})$  measures the flux density of vorticity in the xy-plane, then the integral is simply the total vorticity, which can be written as  $\tilde{f}L_xL_y$ , by the definition of  $\tilde{f}$ . Finally then, we arrive at the normalized dimensionless quantity

$$S(\mathbf{k}_{\perp}) = \frac{1}{(\tilde{f}L_x L_y L_z)^2} \left\langle \left| \int d^3 r \ n^z(\mathbf{r}) e^{i\mathbf{k}_{\perp} \cdot \mathbf{r}_{\perp}} \right|^2 \right\rangle. \tag{7.11}$$

Discretizing this expression through the method in Section 5, i.e. by letting  $\int\!\mathrm{d}r\mapsto a\sum_{\pmb{r}},\,n^z(\pmb{r})\mapsto n^z_{\pmb{r}}$  and  $L_\mu=aN_\nu$  then we find that the filling fraction f which is the number of vortex quanta pr. plaquette of the lattice 4 is related to  $\tilde{f}$  through  $f=a^2\tilde{f}$  and we reproduce the expression in Eq. (7.7).

As an example of how the structure function singles out specific structural correlations, consider Figure 7.3. Figure 7.3b shows a plot of the structure function for all crystal momenta  $k_{\perp}$  in the 1st Brillouin zone. The 6 yellow points surrounding the origin corresponds to correlations in the structure of the vortex lattice in these 6 directions, which implies a hexagonal lattice. The hexagonal lattice is shown directly in Fig. 7.3a, which in this case is a hexagonal lattice of double-quanta vortices. Choosing specific points in the plot of Fig. 7.3b and plotting the structure functions value at different values of a parameters of the system, e.g. temperature, is a common method for evaluating different phases of the structure of the vortex lattice for example for measuring when the vortex lattice melts into a vortex liquid [63, 64, 108–110].

# 7.4.2 Angular histogram

Building on the idea of measuring a specific point in the structure function to signify a structural transition we developed an angular histogram approach that is robust towards rotations of the vortex lattice. We found this to be important in measuring the transition from a hexagonal to a square vortex lattice since the angular symmetry of the model allowed the hexagonal vortex lattice to freeze in various direc-

<sup>4.</sup> We note that by the definitions here, f is a dimensionless quantity, while  $\tilde{f}$  has dimension inverse length square.

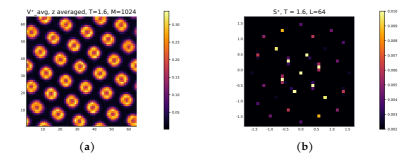
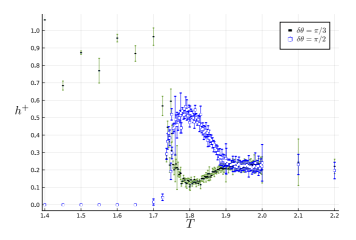


Figure 7.3: Plots of vorticity of the +-component of a p+ip superconductor system. Figure a shows a plot of the real space vorticity which corresponds to a thermal average of a discretized version of  $w(\mathbf{r}_{\perp})$  from Eq. (7.9). Fig. b shows the corresponding structure function which shows a creat hexagonal structure of the vortex line lattice.

tions. To combat this rotation, we built a histogram of the angular distance between peaks in the structure function over several MC steps. For a hexagonal lattice, such a histogram is peaked at the bin containing the angular distance  $\pi/3$ , while a square lattice would be peaked at  $\pi/2$ . Plotting these bins of angular distance over various temperatures, we were able to measure the transition from the square to the hexagonal lattice as seen in Figure 7.4.



**Figure 7.4:** Plot of the bins of angular distance  $\delta\theta=\pi/3$  and  $\delta\theta=\pi/2$ , as a function of simulation temperature.

The histogram was constructed algorithmically by creating a set of angular distances between peaks of the structure function for each MC step. We first found the radius where the peaks were located by searching the average structure function over the entire MC series, within a specified radius interval for the radius  $\rho_m$  that produced the largest value of the discretization of the integral

$$\int_0^{2\pi} d\theta \ S(\rho, \theta), \tag{7.12}$$

where  $S(\rho,\theta)$  is the structure function given in polar coordinates about the Bragg-point  ${\bf k}_\perp=0$ . The entire series of MC data was then blocked into sections containing  $\Delta \tau$  numbers of individual MC measurements of the structure function. Each such interval of structure function measurements were then averaged over to yields separate averaged measurements of the structure function. This blocking is absolutely necessary in order to reveal the hidden vortex lattice from the noise. From each block t, a structure function ring  $S^t(\theta)$  was then created by selecting the highest value of the blocked structure function over a ribbon centered at radius  $\rho_m$  such that

$$S^t(\theta) = \max_{\rho_m - \delta \rho \leq \rho \leq \rho_m + \delta \rho} \{ S^t(\rho, \theta) \}. \tag{7.13}$$

A collection of n peaks  $P^t=\{\theta^p\}$  was then found for each block by finding the highest possible  $S_m$  such that  $S^t(\theta)$  crossed the line  $S_m$  n times. From this set of peaks, then all possible distances between these peak positions were constructed by

$$\Theta^t = \{\delta\theta_{ij} = |\theta^p_i - \theta^p_j| \quad | \quad i \neq j, \ \theta^p_i, \theta^p_j \in P^t\}. \tag{7.14}$$

Let  $\Theta=\bigcup_t \Theta^t$  be the union of all block sets of mutual angular peak distances. The final histogram h was then constructed based on all of the distances in  $\Theta$ . Let the bin in this histogram of the interval  $[0,2\pi)$  that contains the angular distance  $\delta\theta$  be denoted  $\Delta\delta\theta$  such that

$$\Delta \delta \theta = [\delta \theta - \delta \theta_{-}, \delta \theta + \delta \theta_{+}), \tag{7.15}$$

for some non-negative  $\delta\theta_-$  and  $\delta\theta_+$ . The value of the histogram  $h(\Delta\delta\theta)$  at this bin was then calculated by

$$h(\Delta \delta \theta) = \frac{1}{|\Delta \delta \theta| |\Theta|} \sum_{\delta \theta' \in \Theta} \delta_{\delta \theta' \in \Delta \delta \theta}, \tag{7.16}$$

where  $|\Delta\delta\theta|$  is the size of bin  $\Delta\delta\theta$ ,  $|\Theta|$  is the number of mutual distances  $\delta\theta'$  in  $\Theta$  and  $\delta_{\delta\theta'\in\Delta\delta\theta}$  is the Kronecker delta function defined as

$$\delta_{\delta\theta'\in\Delta\delta\theta} = \begin{cases} 1 &: \delta\theta' \in \Delta\delta\theta \\ 0 &: \delta\theta' \notin \Delta\delta\theta \end{cases} . \tag{7.17}$$

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# **Outlook**

In this thesis we have given an introduction to some of the fundamental techniques employed in our theoretical investigations into the nature of unconventional superconductivity with p-wave symmetry. These investigations resulted in three papers.

In Paper I we used a group-theoretical approach to motivate the form of the effective interaction potential between electrons whose low energy excitations could be described in terms of Cooper-pairs with p-wave symmetry. This potential was then used as a basis for deriving the effective free energy for such a superconductor when it was influenced by explicite spin-orbit interaction. We found that the effective free energy had the expected form given by its group-theoretical constraints, but that previous assumptions about its coefficients needed revision because of the effect of spin-orbit coupling.

In Paper II we used large-scale Monte-Carlo simulations to investigate the vortex matter of a p-wave superconductor with a free energy similar to the one derived in Paper I. We found a transition between a square vortex-lattice of single-quanta vortices to an hexagonal vortex lattice consisting of double quanta vortices as the temperature was lowered in a finite field parallel to the crystallographic c-axis.

In Paper III we investigated this same superconductor when exposed to zero external magnetic field and found an Ising phase transition in the neutral sector of the theory. This transition did not separate from the phase-transition of the charged sector in contrast to other mod-

els of two-component superconductors. The reason for the connection between the charged and neutral modes seemed to be because of their group-theoretical nature as components of a single irreducible representation. This identification implies equal stiffness to both components and an explicit coupling through mixed gradient and mixed-component terms in the effective free energy.

Looking towards the future, it now seems less likely that the unconventional superconductor  $\mathrm{Sr_2RuO_4}$  should be theoretically modeled as a p+ip superconductor in spite of the strong evidence for its spontaneous time-reversal-symmetry breaking nature [111–113] and the good agreement between theory and experiment for the qualitative nature of its vortex lattices [89, 106]. Rather, the prevailing view based on the current evidence has shifted to suggesting a degeneracy between a  $d_{x^2-y^2}$  and a  $g_{xy(x^2-y^2)}$  superconducting state [114, 115]. It might in this regard be interesting to use Monte-Carlo simulations to investigate the vortex lattice behavior of such a superconductor and see how it matches with experiments on  $\mathrm{Sr_2RuO_4}$ .

In the exploration of such a novel symmetry state it might be beneficial to utilize more modern forms of Monte-Carlo analysis such as those offered by the advances in machine-learning to yield convincing results in an efficient manner [116–118].

Building on the results of Paper I, it would be of interest to investigate numerically how the kinetic dimensionless phenomenological parameters in the Ginzburg-Landau model depend on the microscopic parameters, especially spin-orbit coupling strength and spin-orbit coupling spin *z*-component.

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