Development of a high-performance gyrokinetic turbulence code for the edge and scrape-off layer of magnetic confinement fusion devices

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Abstract

Understanding the role of turbulence in the edge and scrape-off layer (SOL) of magnetic confinement fusion devices is key to predict their heat exhaust and energy confinement time. Turbulence is commonly studied with computer simulations based on the gyrokinetic equations, a five-dimensional, nonlinear, integro-partial differential system that describes the evolution of the distribution of particles in phase space.

Performing gyrokinetic simulations of edge and SOL plasmas is particularly difficult. First, the plasma density and temperature is low and the turbulent fluctuations are of the same size as the background plasma. This requires a so-called full-$f$ treatment of the gyrokinetic equations. Second, the magnetic geometry of the edge and SOL is complex. It contains regions of open magnetic flux surfaces, X-points and, in stellarator fusion experiments, even regions where field lines are oriented stochastically. This renders flux-coordinates, commonly used in gyrokinetic turbulence simulations of the plasma core, ineffective.

To tackle these issues, we present a new gyrokinetic turbulence code, GENE-X, in this work. GENE-X implements a full-$f$ version of the gyrokinetic system that includes electromagnetic effects and can handle turbulence with large fluctuation amplitudes. Furthermore, GENE-X can perform simulations in realistic magnetic geometries by implementing the so-called flux-coordinate independent approach. In this work, we detail the equations implemented in the code, the coordinate system, the discretisation scheme and careful numerical and physical verifications. In addition, we present the first application of GENE-X to ASDEX Upgrade (AUG). The simulations of AUG are performed with realistic and reduced electron-to-ion mass ratio, with and without collisions over a time of 1ms, studying profile evolution. We compare the results to previous simulations with the Braginskii fluid code GRILLIX and experimental measurements. Furthermore, we discuss the influence of the electron-to-ion mass ratio and collisions on the profiles and the heat exhaust.
Publications

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1. Introduction

1.1. Motivation

The climate of our planet is changing – with dramatic effects. Glaciers melt, sea levels rise and, as a consequence, the natural habitat of many species is destroyed [1]. It is scientific consensus that this trend cannot be explained by natural climate variability and is caused by the anthropogenic emission of greenhouse gases [2]. To limit global warming and its tremendous effects, it is therefore vital to reduce the emission of greenhouse gases by using and developing new, environmentally friendly, energy sources.

Inspired by the energy production of stars, fusion energy offers a unique pathway to achieve that goal. Energy produced by fusion power plants would be CO$_2$-free, safe and sustainable for generations to come. Consequently, developing net energy-producing fusion experiments, as a first step towards a fusion power plant, has been a dream since the 1950s.

1.1.1. Nuclear fusion

Many stars, like our sun, produce energy by fusing hydrogen via the Bethe-Weizäcker cycle or the proton-proton chain [3]. These reactions have a low cross section at temperatures and densities producible in the laboratory. Therefore, they are not suited for fusion reactors on earth. In the most promising fusion experiments today, hydrogen is replaced by its isotopes deuterium and tritium (D-T). The D-T fusion process has the highest known cross section at experimentally relevant parameters and is therefore the easiest to access [4]. The products of the fusion reaction have a slightly lower mass than the inputs. This difference in mass is converted into kinetic energy by the phenomenon known as mass defect. During the D-T fusion process, visualised in Fig. 1.1, a neutron with 14.1 MeV energy and a helium atom with 3.5 MeV energy is released.

![Diagram of D-T fusion reaction]

Figure 1.1.: Figure displaying the fusion reaction of deuterium and tritium to helium and a neutron. The reaction releases a total of 17.6 MeV energy.
However, making atoms fuse is not simple. Due to the Coulomb force, nuclei repel each other. To make fusion reactions possible, the two nuclei need to get close enough such that quantum tunnelling through the Coulomb barrier is likely to happen. This can either be achieved by high temperatures or high pressures. In these conditions, the atoms are ionised and form the state of a plasma. Part of the energy, produced in the fusion process, stays within the plasma and makes, in theory, a self-sustained fusion reaction possible. To obtain such an energy producing, also called burning, plasma, the product of its temperature $T$, density $n$ and energy confinement time $\tau_E$ must exceed the threshold $nT\tau_E > 3 \cdot 10^{21}\text{KeVsm}^{-3}$.

\[ (1.1) \]

1.1.2. Confinement and plasma stability

In stars the plasma is confined with the help of gravity. Temperatures and densities are large and energy confinement times long. As a result, the fusion reaction in stars sustains itself and produces energy. While different confinement strategies exist on earth, the most promising experiments use magnetic fields to confine the plasma. Due to the Lorentz force, charged particles are trapped to a spiral like motion around magnetic field lines. To avoid particle losses at the end of field lines, they are bend into a toroidal shape. The two most promising concepts for fusion power plants are the tokamak and the stellarator [6]. In tokamaks densities around $10^{19}/\text{m}^3 - 10^{20}/\text{m}^3$, Temperatures of $1\text{KeV} - 40\text{KeV}$ and energy confinement times of $\tau_E = 0.1\text{s} - 1\text{s}$ can be achieved [5, 7].

In first approximation, the plasma and its interaction with the magnetic field $B$, in these fusion devices, can be described as a fluid within magnetohydrodynamics (MHD). The equilibrium state of the plasma is classified by the force balance [8]

\[ j \times B = c\nabla p, \]  

(1.2)

between the Lorentz force $j \times B$, caused by the plasma current $j$, and the plasma pressure $p$. The variable $c$ denotes the speed of light. In axisymmetric devices, like the tokamak, every magnetic field line generates a magnetic flux surface. On a magnetic flux surface the pressure is constant and particles stream freely along magnetic field lines. An MHD equilibrium for the tokamak ASDEX Upgrade (AUG) is visualised in Fig. 1.2.

1.1.3. Transport and turbulence

The confinement of the plasma is limited by radial transport perpendicular to magnetic flux surfaces, leading to particle and energy loss. The first component of radial transport is caused by diffusive processes due to collisions in the plasma. However, the radial transport measured in experiments cannot be explained by collisional transport alone. The second component is given by transport due to turbulent fluctuations within the plasma and commonly referred to as anomalous transport [9, 7]. The turbulence is driven by microinstabilities that are caused by density and temperature fluctuations. They grow by extracting energy from gradients
within the plasma. The turbulence transports the energy from the microstabilities, in a cascade like way, to different spatial scales, where it is dissipated.

1.1.4. The edge and scrape-off layer

Inevitably, the plasma comes into contact with the main wall in the boundary region of the fusion device. To avoid damage, the plasma needs to be cooled by orders of magnitude before it hits the wall. Consequently, the boundary region of magnetic confinement fusion devices is characterised by steep density and temperature gradients, driving microinstabilities and turbulence.

In the fusion community, the boundary region is conventionally divided into two parts. The first is the so-called scrape-off layer (SOL). In the SOL magnetic field lines are open and the plasma is in contact with the device wall. The second is the so-called edge of the device. It represents the region just before the last closed flux surface. For visualisation, the different regions are shown at the example of AUG in Fig. 1.2.

In recent years, it has become apparent that understanding turbulent transport in the edge and SOL is important for the energy confinement time and the feasibility of fusion energy. The reason is given in the following two subsections.
1.1.5. Understanding the L-H transition

To achieve long energy confinement times, most fusion experiments operate in high confinement regimes, like the H-mode [11]. These operating modes are characterised by a transport barrier in the edge that limits turbulence and, as a consequence, increases the energy confinement time. The goal of the upcoming fusion experiment ITER [12], to achieve a burning plasma and produce more than ten times the energy injected, is based on the H-mode [13]. Measurements at different experiments have shown that the transport barrier is accompanied by a drop in the radial electric field just before the separatrix [14]. This drop induces a poloidal rotation of the plasma through the $E \times B$ drift that shears turbulent vortices and, consequently, suppresses turbulent transport [14]. As a result of the transport barrier, the temperature and density in plasma core increase and the gradients, close to the edge of the device, are steepened.

Over the last decades, tremendous efforts have been undertaken to understand the formation of the transport barrier and the shift into the high confinement regime, also referred to as the L-H transition [15]. Experimentally, the transition into H-Mode is triggered when the plasma heating power exceeds a critical value [11]. There exist many different models that try to explain the L-H transition. A popular choice, that can explain some of the experimentally measured features of the L-H transition, are predator-prey models [16, 17, 18]. We explain them at the example of [17]. The turbulence takes the position of the prey. With increasing heating power the turbulence increases. After it has reached a certain threshold, it excites zonal flows, the predator, that regulate the turbulence. As soon as the drive of the zonal flows becomes stronger than their damping, they grow exponentially and annihilate the turbulence. At the same time a steep pressure profile with a poloidal flow builds that takes over the role of the zonal flow in damping the turbulence. The result is a quiescent H-mode with a steep pressure profile, a large poloidal flow, no zonal flow and no turbulence.

Some of these results have been reproduced with fluid turbulence simulations [19, 20, 21]. Additionally, simulations with the gyrokinetic turbulence code XGC1 showed that effects due to ion-orbit losses, that are not resolved within fluid codes, are important in the L-H transition [22]. The scientific evidence suggests that turbulence plays an integral part in the L-H transition and predictive, first principle modelling is needed to understand it.

1.1.6. Limiting the particle and heat exhaust of fusion devices

To avoid damage to the fusion device by hot particles hitting the wall, limiting the particle and heat exhaust poses an important problem. Particles are transported over the separatrix and stream along magnetic field lines onto the wall. This generates heat fluxes up to $500\text{MW}/\text{m}^2$ in present tokamaks and is expected to reach $1\text{GW}/\text{m}^2$ in ITER [23]. To avoid damage to the wall, the heat flux must be reduced to $10\text{MW}/\text{m}^2$ normal to the plates for actively cooled materials in steady-state operation [23]. The heat flux onto the divertor is commonly reduced by using shallow incident angles between the magnetic field lines and the divertor plates [24, 25], which effectively distributes the heat flux onto a wider area. The mechanism
that describes the width of the scrape-off layer heat flux, commonly called $\lambda_q$, has not been understood and predictions of $\lambda_q$ yield vastly different results for different plasma models. At the example of ITER, predictions from scaling laws constructed out of measurements of present experiments suggest $\lambda_q \approx 1$mm [26]. Calculations performed with the transport code SOLPS yield $\lambda_q = 3.6$mm [27] and recent studies with the gyrokinetic turbulence code XGC1 showed that electron turbulence can lead to a broadening of $\lambda_q$ to 5.9mm [28]. The heat flux onto the divertor is also reduced by inserting a neutral gas into the SOL. The neutral gas gets ionised from the particles in the plasma. Free electrons recombine with the ions and emit gamma radiation. This provides an efficient mechanism to transport energy out of the plasma before it hits the wall. It is even possible to radiate all the excess energy of the plasma before it comes into contact with the wall. This state is also known as detachment [29]. In order to understand and accurately describe the particle and heat exhaust of a fusion experiment, all processes involved and their interrelation need to be understood. This includes the turbulent transport of heat from the core over the separatrix into the SOL, the dynamics of the plasma in the SOL itself and its interaction with the wall.

1.1.7. Turbulence simulation

As we have established so far, turbulence plays integral part in the feasibility of fusion energy. Due to the limited number of analytical results, numerical methods are employed to study turbulence in fusion experiments.

High fidelity simulations of turbulent fusion plasmas are based on the gyrokinetic equations, a set of nonlinear, integro-partial differential equations that describe the evolution of the distribution of particles in a five-dimensional phase space [30]. The first gyrokinetic simulations were performed in the 80s and 90s based on many simplifying assumptions [31, 32, 33, 34, 35]. Around the turn of the millennium, the simulations became more realistic and were able to provide quantitative insights into the properties of plasma turbulence in the core of fusion devices [36, 37, 38, 39, 40, 41, 42, 43, 44, 45].

Expanding the simulations from the core to the edge and SOL is difficult. In the boundary region the gradients are steep and the temperature and density is low. As a result, the amplitude of the turbulent fluctuations is comparable to the plasma background and a so-called full-$f$ treatment of the equations is necessary. Furthermore, the geometry of the magnetic equilibrium is complex. We explain this challenge in more detail in the next subsection.

Due to the high computational demand of gyrokinetic simulations and the increased collisionality in the edge and SOL, fluid and gyrofluid models are often used [46, 47, 48, 49, 50, 51]. They assume that collisions move the velocity distribution of particles close to a Maxwellian that can be described with a few moments like density and temperature. They are either based on the drift reduced Braginskii equations [52, 53] or on gyrofluid models [54]. However, kinetic effects are known to be important in the edge and SOL [55, 56, 57]. Consequently, several different projects have been launched to simulate gyrokinetic turbulence in the edge and SOL, namely XGC [58], COGENT [59], GKEYLL [60], GENE [61], GYSELA [62, 63] and PICLS [64]. The codes are based on different numerical approaches with their respective advantages and
disadvantages. The different approaches are split into three categories.

**The particle-in-cell approach**

The first gyrokinetic plasma simulations were based on the so-called particle-in-cell (PIC) approach [31, 32, 33]. Therein, a sample of so-called markers are integrated along the characteristics of the gyrokinetic Vlasov equation, and the electromagnetic fields are solved on a three-dimensional grid in position space. Particle-in-cell codes are straightforward to implement and have shown good scalability on modern supercomputers [28]. Due to the probabilistic nature of these algorithms, the results contain statistical noise which has shown to be problematic for certain applications [65, 66, 67]. From the gyrokinetic edge and SOL codes presented, XGC and PICLS implement the PIC method. To resolve the full distribution function in SOL plasmas and keep the numerical noise at a manageable level, many markers are needed which may require larger computational resources than comparable continuum codes.

**The Eulerian approach**

In continuum or Eulerian schemes the gyrokinetic Vlasov-Maxwell system is discretised on a five-dimensional position-velocity space grid. They were first introduced around the year 2000 [68, 39, 38, 56, 40] and use a variety of numerical methods available for partial differential equations. This includes finite difference, finite volume, finite element and discontinuous Galerkin methods with explicit and implicit time stepping schemes. Compared to PIC codes, continuum codes are free of numerical noise at the expense of introducing a velocity space grid. From the gyrokinetic edge and SOL codes mentioned above, COGENT, GKEYLL and GENE implement a continuum method. GKEYLL uses an energy conservative discontinuous Galerkin discretisation, COGENT discretises the gyrokinetic system with a high order finite volume method and GENE features a mixed shock stable finite-volume, finite difference discretisation scheme.

**The semi-Lagrangian approach**

In the semi-Lagrangian approach the gyrokinetic Vlasov-Maxwell system is, similar to Eulerian schemes, discretised on a five-dimensional position-velocity space grid. Therefore, they are also free of numerical noise. Semi-Lagrangian schemes are commonly divided into three categories: (i) The (standard) backward [69], (ii) the forward [70] and (iii) the conservative [71] semi-Lagrangian method. We explain them at the example of the backward method. The distribution function is evolved by following the characteristics of the gyrokinetic Vlasov equation from a given grid point, on the five-dimensional position-velocity space grid, backwards in time for one timestep. The new value of the distribution function at the grid point is the interpolated value at the foot of that characteristic curve, using the values of the distribution function at the previous timestep [69]. Semi-Lagrangian schemes can allow for larger timesteps than comparable Eulerian schemes, which makes them, next to fusion applications, also attractive for numerical weather predictions and atmospheric models [72]. The gyrokinetic edge and SOL code GYSELA implements the semi-Lagrangian approach.
1.1.8. Handling X-point geometries

Especially challenging for turbulence simulations in the edge and SOL is the complex magnetic geometry. There are regions of open and closed magnetic field lines, and, in stellarator fusion experiments, even regions, where field lines are oriented stochastically. Many gyrokinetic codes present in the community implement so-called flux-coordinate systems [73]. While these allow for efficient simulations in regions with regular magnetic flux surfaces, they become ill-defined at the separatrix or in stochastic regions [74]. To allow for predictive edge and SOL simulations, coordinate systems free of singularities that can treat realistic magnetic geometries must be used. XGC1 has demonstrated that PIC codes can perform simulations in X-point geometries by using a cylindrical coordinate system. PIC codes follow the trajectories of marker particles in continuous real space and therefore parallel structures in the plasma are well represented. However, coordinate systems agnostic of the magnetic field are suboptimal for continuum codes. They use a five-dimensional phase space grid and a fine toroidal resolution is needed to represent structures that are aligned to magnetic field lines. This increases the cost of the simulation significantly because more grid points and, in case of explicit time stepping, a smaller timestep is needed. To allow for efficient simulations in X-point geometry with continuum codes, locally field-aligned methods based on the flux-coordinate independent approach (FCI) [75, 76] have been developed and tested in fluid codes [48, 77, 51, 78]. Within the fluid community, they have shown to be effective for studying edge and SOL turbulence. Recently, the use of FCI has enabled the simulation of turbulence in the diverted geometry of AUG [79, 80]. In the gyrokinetic community, locally field-aligned methods have been implemented in COGENT [81] and used to perform five-dimensional simulations in a geometry with one X-point. Further, field-aligned interpolation has been used to accelerate semi-Lagrangian codes [82, 83].

1.2. Overview

Predictive, first principle modelling of turbulence is essential to understand the plasma properties in the boundary of fusion devices. To this end, we present a new gyrokinetic turbulence code that is tailored towards edge and SOL simulations in this work. The code is based on the continuum approach to resolve the full distribution in regions in the SOL where the plasma density is low. We put a special focus on handling the geometric complexities. For this reason, we employ a locally field-aligned coordinate system based on the FCI approach. By using the FCI approach, we are not limited by the magnetic geometry and are able to perform simulations in geometries with a single and even multiple X-points. This will enable the investigation of advanced divertor concepts [84, 85] within gyrokinetic theory in the future. Furthermore, the coordinate system is well-defined in stochastic regions, and a generalisation of the code to stellarator geometries is straightforward. The implementation follows the example of the Braginskii fluid code GRILLIX [74, 48, 86] and uses unstructured, locally Cartesian grids. These grids allow for a high flexibility regarding the geometry while ensuring good numerical properties and a high computational efficiency [87]. The discretisation follows
the example of the GENE code [38, 45] and uses explicit time stepping with a mixed order finite difference scheme. The numerical solution of the gyrokinetic Vlasov-Maxwell system on modern supercomputers is resource intensive. To maximize the performance of the code and prepare it for the heterogeneous architectures of future exascale machines, we put a special emphasis on software engineering and performance. Based on its scientific ancestor, GENE, we call the code GENE-X.

The major scientific contributions of this work are threefold. First, next to GENE-X, only XGC1 and COGENT can perform gyrokinetic turbulence simulations in X-point geometry. To ensure good scientific practice, it is vital to have multiple codes, using different numerical techniques, that can reproduce each others findings. Second, we demonstrate that the FCI approach together with the presented numerical scheme works well with a gyrokinetic model. Third, we present gyrokinetic turbulence simulations of the edge and SOL of ASDEX Upgrade that include a magnetic X-Point.

This work is structured as follows. In Chapter 2, we give an introduction into gyrokinetic theory and derive the full-f, electromagnetic model implemented in the code. We continue with the rigorous derivation of the FCI coordinate system and the metric coefficients in Chapter 3. In Chapter 4, we discuss the normalisation of the equations and present the finite difference discretisation scheme implemented. In Chapter 5, we detail the software engineering of the GENE-X code and demonstrate the performance of the software design. We present thorough verification benchmarks in different magnetic geometries, including a full order analysis with the method of manufactured solutions, in Chapter 6. We finish this work with presenting electromagnetic, gyrokinetic simulations of the edge and SOL of ASDEX Upgrade in Chapter 7.
2. Gyrokinetic theory

We start the discussion of gyrokinetic theory at the example of a single particle with mass $m$ and charge $q$ that moves in a homogeneous magnetic field with field strength $B$. We denote the unit vector of the magnetic field with $\mathbf{b} = \mathbf{B}/B$. Due to the Lorentz force, the particle gyrates around a magnetic field line with the cyclotron frequency

$$\Omega = \frac{qB}{mc}. \quad (2.1)$$

The speed of the particle on the gyroorbit, $v_\perp = |\mathbf{v} - (\mathbf{b} \cdot \mathbf{v})\mathbf{b}|$, is constant and the magnetic moment of the particle,

$$\mu = \frac{mv_\perp^2}{2B}, \quad (2.2)$$

is conserved. Therefore, the constant gyromotion can be neglected. In a fusion device the conservation of the magnetic moment is violated. First, the magnetic field is non-uniform. Second, the plasma produces density and current fluctuations which produce electromagnetic fields themselves. In the following, we denote the perturbed electrostatic potential with $\phi_1$ and the perturbed electromagnetic vector potential with $\mathbf{A}_1$. These fields alter the trajectory of particles via the Lorentz force, which ultimately leads to turbulence.

The plasma in a fusion device is a collection of particles of different species, like electrons, ions or impurities. The phase space distribution of all particles of a given species, denoted by $\sigma$, is described with their distribution function $f_\sigma$. The time evolution of the distribution function is given by the Vlasov equation that is coupled to Maxwell’s equations to provide the electromagnetic fields $\phi_1$ and $\mathbf{A}_1$. This system contains a multitude of physical effects, including the full gyroorbits of particles, Debye shielding and Langmuir waves. While these are interesting phenomena, they act on significantly smaller time scales than turbulence. Numerical solutions of the full Vlasov-Maxwell system at the scale of medium size fusion experiments and on turbulence time scales are not even possible on the largest supercomputers available today. Consequently, it is necessary to derive a simplified model that removes the fast timescales from the system and makes the simulation possible. This is the goal of gyrokinetic theory.

In gyrokinetic theory the goal is achieved by introducing a new set of phase space coordinates, called gyrocentre coordinates. They include a generalised magnetic moment $\mu_{gy}$ that is an exact invariant of the motion of the particle. In addition, the dynamics associated to the fast gyromotion of the particle, called $\theta_{gy}$, are decoupled from the other gyrocentre coordinates [88]. This removes the fast timescales from the system and reduces the dimensionality from six to five.
The reduction from the full kinetic system to the gyrokinetic subset is controlled with so-called ordering parameters. They express the spatial and temporal scales of different phenomena and are used to control their size. In the following, we define the ordering parameters used in standard gyrokinetic theory [88].

1. We call the length scale of perturbations in the magnetic field $L_B = B/|\nabla B|$ and define the ordering parameter $\epsilon_B = \rho_T/L_B$, where
   \[
   \rho_T = \frac{m c v_{T,\perp}}{|q|B}
   \]
   is the gyroradius and $v_{T,\perp}$ is the perpendicular velocity of a thermal ion. We assume that the length scale of the magnetic fluctuations is much larger than the gyroradius of a thermal ion, i.e. $\epsilon_B \ll 1$.

2. We define ordering parameter for electrostatic perturbations as $\epsilon_\delta = k_{\perp}\rho_T q\phi_1/T_i$, where $k_{\perp}$ represents the wavenumber of the electrostatic fluctuation perpendicular to the magnetic field and $T_i$ the ion temperature. We assume that $\epsilon_\delta \ll 1$ and that fluctuations of the magnetic vector potential $A_1$ are of the order $\phi_1 c/v_T$.

3. We call the wavevector of fluctuations parallel to the background magnetic field $k_{||}$. In strongly magnetised plasmas parallel perturbations act on larger scales than perpendicular ones. We reflect this property by assuming that $\epsilon_\parallel = k_{||}/k_{\perp} \ll 1$.

4. We call the characteristic frequency of fluctuations in the plasma $\omega$ and assume that the frequency is much smaller than the ion cyclotron frequency $\Omega_i$, i.e. $\epsilon_\omega = \omega/\Omega_i \ll 1$.

For the derivation of the gyrokinetic model, different orderings between the four different parameters can be used. In this work, we employ the standard ordering used by gyrokinetic codes in the community [36, 37, 38, 40, 41, 42, 43, 44, 45]. It is assumed that fluctuations in the background magnetic field are smaller than fluctuations in the electrostatic potential, i.e. $\epsilon_B \sim \epsilon_\delta^2$. Further, it is assumed that $\epsilon_\parallel \sim \epsilon_\delta$ and $\epsilon_\omega \sim \epsilon_\delta$. The projects mentioned above study plasma turbulence in the core of fusion devices. At the point of writing it is an area of active research whether this ordering and the resulting gyrokinetic theory is also applicable in the edge and SOL. Different studies with hybrid and fully kinetic models indicate that the standard gyrokinetic model is still useful [89, 90, 91, 92].

In the remainder of this chapter, we present the gyrokinetic Vlasov-Maxwell system that is implemented in the GENE-X code. As gyrokinetic theory is fundamentally linked to coordinate transformations on manifolds, we start by discussing the basics of Hamiltonian mechanics. We proceed by giving an overview of the gyrokinetic reduction and present the gyrokinetic Vlasov-Maxwell system. Finally, we simplify the system to a self-consistent subset, suitable for the first implementation in the GENE-X code and discuss conservation laws.
2.1. Hamiltonian mechanics

The plasma in a fusion device is formally described by a manifold which we call \( \mathcal{M} \). This manifold carries a differentiable structure inherited from differentiable coordinate charts \( (\varphi, U) \) or a collection of coordinate charts \( (\varphi_a, U_a) \), called an atlas\(^1\). Charts assign a coordinate \( q \in V = \varphi(U) \) to every point on the manifold \( \mathcal{M} \). What is commonly known as a coordinate map is the inverse of the chart \( \varphi^{-1} : V \to \mathcal{M} \), assigning a point on the manifold to every coordinate. As the charts are diffeomorphic, the coordinate map is always well-defined. In the application of fusion physics the manifold is embedded in \( \mathbb{R}^3 \) and a natural Cartesian coordinate system exists. We call the Cartesian coordinates of the three dimensional manifold \( q = (x_c, y_c, z_c) \in V \). Particles in the plasma have a velocity. The velocity space at position \( q \) is modelled by the tangent space of the manifold \( T_q\mathcal{M} \). The inverse of the coordinate chart \( \varphi^{-1} \) provides a natural basis for the tangent space \( T_q\mathcal{M} \) by

\[
e_i := \frac{\partial \varphi^{-1}}{\partial q_i}.
\] (2.4)

The basis of the cotangent space \( T^*_q\mathcal{M} \), denoted by \( e^i = dq_i \), is defined by the condition

\[
e^i \cdot e_j = \delta_j^i.
\] (2.5)

We denote the Cartesian coordinates in velocity space, induced by the Cartesian coordinates in real space, with \( (v_x, v_y, v_z) \). In the Lagrangian formulation of classical mechanics the state of a particle is described by its position on the manifold \( q \in \mathcal{M} \) and its generalised velocity \( \dot{q} \in T_q\mathcal{M} \). The equations of motion are determined by the so-called Lagrangian

\[
L : (q, \dot{q}, t) \mapsto L(q, \dot{q}, t)
\] (2.6)

through the Euler-Lagrange equations

\[
\frac{d}{dt} \frac{\partial L}{\partial \dot{q}_i} = \frac{\partial L}{\partial q_i}.
\] (2.7)

In the Lagrange formalism arbitrary coordinate transformations of the coordinates \( q \) are possible with a transformation to a different coordinate chart. In Hamiltonian mechanics the generalised velocity \( \dot{q} \) is replaced by the generalised momentum

\[
p_i := \left( \frac{\partial L}{\partial \dot{q}_i} \right)(q, \dot{q}, t).
\] (2.8)

The generalised momentum \( p \) is a cotangent vector and contained in the cotangent space \( T^*_q\mathcal{M} \). The vector bundle of the cotangent spaces at all the points in the manifold \( \mathcal{M} \) defines the phase space. We denote it with \( \mathcal{P} := T^*\mathcal{M} \). The phase space is a six-dimensional, differentiable manifold and different charts can be used to define coordinate systems. The phase space \( \mathcal{P} \) carries a symplectic structure given by the closed, non-degenerate differential

\(^1\)We construct an atlas of \( \mathcal{M} \) in Chapter 3 when we introduce a locally field-aligned coordinate system.
The 2-form $\omega^2$ can be interpreted as a skew-symmetric scalar product on $TP$. Like a normal scalar product, Eq. (2.9) introduces a natural isomorphism between differential forms and vector fields on the manifold. A vector $\xi \in TP$ is mapped to a differential form via [93, pp. 203–204]

$$I : \xi \mapsto \omega^1 = (\eta \mapsto \omega^2(\eta, \xi)) .$$

(2.10)

In the coordinate system $(p, q)$ the isomorphism $I$ can be represented as a simple matrix multiplication with [93, p. 203]

$$I = \begin{pmatrix} 0 & 1_3 \\ -1_3 & 0 \end{pmatrix} .$$

(2.11)

A coordinate transformation $\tau$ on the phase space $P$ is called canonical if it preserves the 2-form $\omega^2$ [93, p. 206]. The volume element of the phase space is given by $\omega^2 \wedge \omega^2 \wedge \omega^2$ and therefore canonical transformations preserve the volume element in phase space. The Hamiltonian $H$ is a function on the phase space. With the inverse of the isomorphism $I$, a vector field can be associated to the Hamiltonian via $I^{-1} dH$. In the coordinates $(p, q)$ the Hamiltonian vector field reads [93, p. 203]

$$I^{-1} dH = \begin{pmatrix} \frac{\partial H}{\partial p} \\ -\frac{\partial H}{\partial q} \end{pmatrix} .$$

(2.12)

The Hamiltonian flow $g^\lambda_H : P \to P$ is defined as a 1-parameter group of diffeomorphisms on the manifold $P$ such that the rate of change of the flow, contained in the tangent space $TP$, is given by the Hamiltonian vector field, i.e. [93, p. 204]

$$\frac{d}{d\lambda} \bigg|_{\lambda=0} g^\lambda_H = I^{-1} dH .$$

(2.13)

In the coordinates $(p, q)$, Eq. (2.13) gives rise to Hamilton’s equations [93, p. 204]

$$\dot{q}_i = \frac{\partial H}{\partial p_i},$$

$$\dot{p}_i = -\frac{\partial H}{\partial q_i} .$$

(2.14)

They describe the trajectories of the Hamiltonian flow in phase space. The Hamiltonian orbit parameter $\lambda$ can be interpreted as the time. The Poisson bracket $\{f, h\}$ of two functions on the manifold $P$ is defined as the derivative of the function $f$ in the direction of the Hamiltonian
flow of $h$ \cite[pp. 214–215]{93}
\[
\{f, h\} := \left. \frac{d}{d\lambda} \right|_{\lambda=0} f(g_h^\lambda) \\
= df(I^{-1} dh) \\
= \omega^2(I^{-1} dh, I^{-1} df)
\] (2.15)

and yields the well-known form in the coordinates $(p, q)$ \cite[p. 215]{93}
\[
\{f, h\} = \sum_{i=1}^{3} \frac{\partial f}{\partial q_i} \frac{\partial h}{\partial p_i} - \frac{\partial f}{\partial p_i} \frac{\partial h}{\partial q_i}.
\] (2.16)

In simple words, Eq. (2.15) shows that the rate of change of a function along the Hamiltonian flow is given by the Poisson bracket. It can be shown that a transformation is canonical if and only if it preserves the structure of the Poisson bracket \cite[p. 216]{93}. After this formal discussion of Hamiltonian mechanics on symplectic manifolds, we introduce the Lagrangian 1-form that connects Hamiltonian to Lagrangian mechanics and is used in the presentation of the gyrokinetic model in the next section. In the following, the phase space is extended by the Hamiltonian orbit parameter $\lambda$ \cite{30}. The Lagrangian 1-form reads
\[
\omega^1 := p \, dq - H \, d\lambda.
\] (2.17)

The part $p \, dq$ is often referred to as the symplectic part and $H \, d\lambda$ as the Hamiltonian part of the 1-form. The symplectic 2-form $\omega^2$ is obtained by taking the exterior derivative $\omega^2 = d(p \, dq)$ of the symplectic part of $\omega^1$. Using Eq. (2.15), $\omega_1$ gives rise to the Poisson bracket and therefore Hamiltonian mechanics. Further, the 1-form $\omega^1$ allows for a straightforward connection to Lagrangian mechanics. The integral over $\omega^1$ along a trajectory $\gamma$ in the extended phase space starting from $(p_0, q_0, \lambda_0)$ and going to $(p_1, q_1, \lambda_1)$ has an extremum under variations of $\gamma$. The extremum is given by the integral curves of Hamilton’s equations \cite[pp. 243–244]{93}. The connection to the Lagrangian, as the Legendre transformation of the Hamiltonian, is given by
\[
\int_\gamma p \, dq - H \, d\lambda = \int_{\lambda_0}^{\lambda_1} (p \, q - H) \, d\lambda = \int_{\lambda_0}^{\lambda_1} L \, d\lambda.
\] (2.18)

In the following sections, we give an overview of the derivation of the gyrokinetic equations. The starting point is the form $\omega^1$ of a charged particle in an electromagnetic field. The coordinate transformation into gyrocentre coordinates is non-canonical and therefore results in a non-canonical expression for the symplectic part of $\omega^1$ and the Poisson bracket \cite{30}.

### 2.2. Common coordinate systems for fusion plasmas

Before we start with the derivation of the gyrokinetic Vlasov-Maxwell system, we introduce different coordinate systems that are more suited to describe fusion plasmas than the
In magnetic confinement fusion devices the shape of the plasma is formed by the magnetic field. Therefore, it is convenient to use a cylindrical and cylindrical-toroidal coordinate system that reflects its structure. The cylindrical coordinate system is given by

\begin{align*}
R &= \sqrt{x_c^2 + y_c^2} \quad (2.19) \\
\varphi &= \text{atan2}(y_c, x_c) \quad (2.20) \\
Z &= z_c \quad (2.21)
\end{align*}

and the cylindrical-toroidal coordinate system by [73]

\begin{align*}
r &= \sqrt{(\sqrt{x_c^2 + y_c^2} - R_0)^2 + z_c^2} \quad (2.22) \\
\theta &= \text{atan2}(z_c, \sqrt{x_c^2 + y_c^2} - R_0) \quad (2.23) \\
\varphi &= \text{atan2}(y_c, x_c), \quad (2.24)
\end{align*}

where \( R_0 \) denotes the major radius of the torus. As the plasma is magnetised, charged particles follow magnetic field lines. Therefore, it is useful to also introduce different coordinates in velocity space. Given an orthonormal basis \((b_1, b_2, b)\) in the tangent space \(T_qM\) at the point \(q \in M\) and the vector \(v \in T_qM\), we define the coordinates \((v_1, v_2, v_{\parallel})\) by the coordinate transformation

\begin{align*}
v_1 &= b_1 \cdot v \\
v_2 &= b_2 \cdot v \\
v_{\parallel} &= b \cdot v \quad (2.25)
\end{align*}

and the coordinates \((v_{\parallel}, \mu, \theta)\) by the coordinate transformation

\begin{align*}
\mu &= \frac{m_\sigma (v_1^2 + v_2^2)}{2B} \quad (2.28) \\
\theta &= \text{atan2}(v_2, v_1). \quad (2.29)
\end{align*}

The new coordinates in velocity space can be interpreted in a simple way. The coordinate \(v_{\parallel}\) denotes the velocity of a charged particle along the field line, \(\mu\) the magnetic moment and \(\theta\) the angle of the gyromotion. The kinetic energy of the particle is given by \(E = \mu B + \frac{1}{2} m_\sigma v_{\parallel}^2\).

Further, we define the gyromotion vectors

\begin{align*}
\rho(\theta) &= b_1 \cos \theta - b_2 \sin \theta \quad (2.30) \\
\bot(\theta) &= -b_1 \sin \theta - b_2 \cos \theta \quad (2.31)
\end{align*}

forming a second orthonormal basis set \((\bot(\theta), \rho(\theta), b)\)\(^\dagger\). As the magnetic field in general changes for every point \(q \in M\), so do the coordinates in velocity spaces.

\(^\dagger\)The set \((b_1, b_2, b)\) is also known as the fixed frame and \((\bot(\theta), \rho(\theta), b)\) as the rotating frame [94].
2.3. Derivation of the gyrokinetic Vlasov-Maxwell system

The derivation of the gyrocentre coordinates starts with a charged particle moving in the background magnetic field $B$ under the influence of time dependent perturbed electromagnetic fields $A_1$ and $\phi_1$. In the presence of time dependent electromagnetic fields, the energy $k$ of the particle is not conserved and changes according to the additional equation of motion [30]

$$\frac{dk}{dt} = q_\sigma \left( \frac{\partial \phi_1}{\partial t} - v \cdot \frac{\partial A_1}{\partial t} \right).$$

(2.32)

To account for the change of energy, the phase space is expanded by the canonically conjugate pair $(t,k)$, where $k$ is constrained to the conserved energy in the time independent case [30]. This allows the formulation of Hamilton’s equations of motion with time dependent electromagnetic fields [30].

### 2.3.1. The gyrokinetic Lagrangian 1-form

The 1-form $\omega^1$, introduced in Eq. (2.17), describing the trajectory of the charged particle in the extended phase space coordinates $(R, \varphi, Z, t, v_\parallel, \mu, \theta, k, \lambda)$ reads

$$\omega^1 = \left( \frac{q_\sigma}{c} A(R) + \epsilon_\delta \frac{q_\sigma}{c} A_1(R, t) + m_\sigma v \right) \cdot dR - k \, dt - H \, d\lambda,$$

(2.33)

with the Hamiltonian

$$H = \frac{1}{2} m_\sigma v^2 + \epsilon_\delta q_\sigma \phi_1(R, t) - k.$$

(2.34)

The physical motion of the particle takes place on the subspace $H = 0$, and Hamilton’s equation of motion for the time, $\dot{t} = 1$, can be used to replace the Hamiltonian orbit parameter $\lambda$ with the time $t$ [30].

Both the symplectic and Hamiltonian part of the Lagrangian 1-form depend on the velocity and therefore have a dependence on the gyroangle $\theta$. The symplectic part of $\omega^1$ depends on the perturbed magnetic vector potential $A_1$ and the Hamiltonian part on the perturbed electrostatic potential $\phi_1$. The derivation of the gyrokinetic 1-form comprises of two steps. First, the perturbed fields $A_1$ and $\phi_1$ are removed from $\omega^1$ and a transformation into so-called guiding-centre coordinates $(R_{gc}, \varphi_{gc}, Z_{gc}, v_{\parallel gc}, \mu_{gc}, \theta_{gc})$ is performed. The guiding-centre transformation removes the fast gyromotion time scales that are associated with the magnetic background $A$. In a second step, the perturbed fields $A_1$ and $\phi_1$ are added to the guiding-centre 1-form again and a transformation into gyrocentre coordinates $(R_{gy}, \varphi_{gy}, Z_{gy}, v_{\parallel gy}, \mu_{gy}, \theta_{gy})$ is performed. The gyrocentre transformation removes the fast gyromotion time scales caused by the perturbed electromagnetic fields. In this work, we employ the so-called symplectic gyrocentre model [95, 30]. The symplectic gyrocentre model retains explicit time derivatives in the Poisson bracket and has been shown to be well suited for gyrokinetic continuum simulations [96, 97]. In contrast, so-called Hamiltonian gyrocentre models [95, 30] are often used in particle-in-cell codes [43]. They have the advantage that they are free of explicit time derivatives of the perturbed electromagnetic vector po-
potential, but can lead to the so-called Ampère cancellation problem in gyrokinetic turbulence simulations [98, 99, 100].

We start with the discussion of the guiding-centre coordinates. For brevity, we state and interpret the result of the transformation. For an explicit calculation we refer to [101, 102, 103, 88]. In guiding-centre coordinates, the form \( \omega^1 \) reads

\[
\omega^1_{gc} = \left( \frac{1}{\epsilon_B} \frac{q_\sigma}{c} A(R_{gc}) + \epsilon_B m_\sigma v_{\| gc} b(R_{gc}) - \epsilon_B \frac{m_\sigma c}{q_\sigma} \mu_{gc} R^* \right) \cdot \partial R_{gc} + \epsilon_B \frac{m_\sigma c}{q_\sigma} \mu_{gc} \partial \theta_{gc} - k \, dt - H_{gc} \, d\lambda + \mathcal{O}(\epsilon_B^2),
\]

(2.35)

where the expression \( R^* = \nabla b_1 \cdot b_2 + (b \cdot \nabla \times b) b/2 \) and the Hamiltonian

\[
H_{gc} = \frac{1}{2} m_\sigma v_{\| gc}^2 + \mu_{gc} B(R_{gc}) - k
\]

(2.36)

were used. As a result of the guiding-centre transformation, the symplectic and Hamiltonian part of \( \omega^1 \) are independent of the gyroangle \( \theta_{gc} \). The Poisson bracket of the system is calculated from the symplectic part of \( \omega^1 \) by deriving the matrix representation of the isomorphism \( I \), its inverse \( I^{-1} \) and using Eq. (2.15). It reads [30, 88]

\[
\{ f, g \}_{gc} = \frac{1}{\epsilon_B} \frac{q_\sigma}{m_\sigma c} \left( \frac{\partial f}{\partial \theta_{gc}} \frac{\partial g}{\partial \mu_{gc}} - \frac{\partial f}{\partial \mu_{gc}} \frac{\partial g}{\partial \theta_{gc}} \right) + \frac{B^*_{gc}}{\epsilon_B B^*_{\| gc}} \cdot \left( \nabla^*_{gc} \frac{\partial g}{\partial v_{\| gc}} - \frac{\partial f}{\partial v_{\| gc}} \nabla^*_{gc} g \right)
\]

\[
- \epsilon_B \frac{c b}{q_\sigma B^*_{\| gc}} \cdot \left( \nabla^*_{gc} f \times \nabla^*_{gc} g \right)
\]

\[
+ \left( \frac{\partial f}{\partial k} \frac{\partial g}{\partial t} - \frac{\partial f}{\partial t} \frac{\partial g}{\partial k} \right),
\]

(2.37)

where the definitions

\[
\nabla^*_{gc} = \nabla + R^* \frac{\partial}{\partial \theta_{gc}}
\]

(2.38)

\[
B^*_{gc} = B + \epsilon_B \frac{m_\sigma c}{q_\sigma} v_{\| gc} \nabla \times b - \epsilon_B^2 \frac{m_\sigma c^2}{q_\sigma^2} \mu_{gc} \nabla \times R^*
\]

(2.39)

were used. The parallel component of \( B^*_{gc} \) is called \( B^*_{\| gc} = b \cdot B^*_{\| gc} \). The Jacobian of the transformation from laboratory coordinates \( (R, \varphi, \mu, v_{\|}, \theta) \) into guiding-centre coordinates \( (R_{gc}, \varphi_{gc}, Z_{gc}, v_{\| gc}, \mu_{gc}, \theta_{gc}) \) is given by \( J_{gc} = m_\sigma B^*_{\| gc} \). Although the guiding-centre coordinates are already quite involved, they have a straightforward interpretation: The relation between the particle and its guiding-centre position is given by \( R = R_{gc} + \rho_{gc}(R_{gc}, \mu_{gc}, \theta_{gc}) \) with the guiding-centre displacement

\[
\rho_{gc}(R_{gc}, \mu_{gc}, \theta_{gc}) = \frac{m_\sigma c}{q_\sigma B(R_{gc})} \sqrt{\frac{2 \mu_{gy} B(R_{gy})}{m} \rho(\theta_{gc}, R_{gc}) + \mathcal{O}(\epsilon_B)}.
\]

(2.40)

We only present the guiding-centre displacement \( \rho_{gc} \) up to order \( \epsilon_B \). For the full expression

\[\text{The guiding-centre Jacobian is often also given as } J_{gc} = m_\sigma B^*_{\| gc}. \] The difference originates from the use of \( p_{\|} \) instead of \( v_{\|} \) in the guiding-centre coordinates.
we refer to [88, Appendix A]. The guiding-centre position can be interpreted as the centre of the gyromotion of the particle given the non-uniform magnetic field. We continue with the discussion of gyrocentre coordinates. To perform the transformation, the perturbed electromagnetic fields are added to $\omega^1$ again. The 1-form is transformed into

$$\omega^1_{gc} \to \omega^1_{gc} + \frac{q_\sigma}{c} A_1(R_{gc} + \rho_{gc}, t) \cdot d(R_{gc} + \rho_{gc}) - q_\sigma \phi_1(R_{gc} + \rho_{gc}, t) d\lambda$$

(2.41)

and regains an explicit dependence on the fast gyroangle $\theta_{gc}$ through the guiding-centre displacement vector $\rho_{gc}$. Before we present the 1-form in gyrocentre coordinates, we introduce further notation that is used throughout this chapter. We define the well-known gyroaverages and orthogonal components of the electromagnetic vector potential by

$$\psi_1(R_{gy}, v_{\| gy}, \mu_{gy}, t) = \phi_1(R_{gy} + \rho_{gy}, t)$$

$$A_{1\|}(R_{gy}, t) := b(R_{gy}) \cdot A_1(R_{gy}, t)$$

$$A_{1\perp}(R_{gy}, t) := A_1(R_{gy}, t) - A_{1\|}(R_{gy}, t)b,$$

(2.42)

and the fluctuating part of a function $g$, called $\tilde{g}$, by $\tilde{g} := g - G[g]$. We define the parallel and orthogonal components of the electromagnetic vector potential by

$$A_{1\|}(R_{gy}, t) := b(R_{gy}) \cdot A_1(R_{gy}, t)$$

$$A_{1\perp}(R_{gy}, t) := A_1(R_{gy}, t) - A_{1\|}(R_{gy}, t)b,$$

(2.43)

and we introduce the effective electrostatic potential

$$\psi_1(R_{gy}, v_{\| gy}, \mu_{gy}, \theta_{gy}, t) = \phi_1(R_{gy} + \rho_{gy}, t)$$

$$- \sqrt{\frac{2\mu_{gy} B(R_{gy})}{mc^2}} \perp(\theta_{gy}, R_{gy} + \rho_{gy}) \cdot A_{1\perp}(R_{gy} + \rho_{gy}, t).$$

(2.44)

The transformation to gyrocentre coordinates and, in the process, the elimination of the fast gyroangle in the electromagnetic fields is done with the help of Lie transformations [104]. Methods free of Lie transformations based on variational averaging have been used for the derivation as well [105]. Similar to the gyrocentre reduction, we state the results. For a detailed derivation we refer to [106, 30, 88]. In gyrocentre coordinates, the 1-form reads

$$\omega^1_{gy} = \left(\frac{q_\sigma}{c} (A(R_{gy}) + \epsilon_3 A_1(R_{gy} + \rho_{gy})) + m_\sigma v_{\| gy} b(R_{gy}) - \frac{m_\sigma c}{q_\sigma} \mu R^* + \frac{m_\sigma c}{q_\sigma} \mu_{gy} d\theta_{gy} - k dt - H_{gy} d\lambda + O(\epsilon_3^2)\right) \cdot dR_{gy}$$

(2.45)

with the gyrocentre Hamiltonian [30]

$$H_{gy} = H_{gc} + \epsilon_3 q_\sigma G[\psi_1] + \epsilon_3^2 \left(\frac{q_\sigma^2}{2mc^2} \left(G \left[|A_1(R_{gy} + \rho_{gy}, t)|^2\right] - G \left[A_{1\|}(R_{gy} + \rho_{gy}, t)\right]^2\right) - \frac{q_\sigma^2}{2B(R_{gy})^2} \frac{\partial G}{\partial \mu_{gy}} \cdot b(R_{gy}) \cdot \nabla \tilde{\psi}_1 \times \int \nabla \tilde{\psi}_1 d\theta_{gy}\right)$$

$$+ G \left[A_{1\perp}(R_{gy} + \rho_{gy}, t)\right] \cdot \frac{q_\sigma}{B} b \times \nabla G[\psi_1] + O(\epsilon_3^3).$$

(2.46)
Analogous to Eq. (2.37), the Poisson bracket in gyrocentre coordinates is obtained by deriving the matrix representation of the isomorphism $I$, calculating $I^{-1}$ and using Eq. (2.15). It reads \[ (2.47) \]
\[
\{f, g\}_{gy} = \frac{q_s}{m_c} \left( \frac{\partial f}{\partial \theta_{gy}} \frac{\partial g}{\partial \mu_{gy}} - \frac{\partial f}{\partial \mu_{gy}} \frac{\partial g}{\partial \theta_{gy}} \right) + \frac{B^*_{\parallel}}{\mu_{gy}} \cdot \left( \nabla_{gy}^* f \frac{\partial g}{\partial v_{\parallel|gy}} - \frac{\partial f}{\partial v_{\parallel|gy}} \nabla_{gy}^* g \right) - \epsilon \frac{\partial f}{\partial \mu_{gy}} \frac{\partial g}{\partial \theta_{gy}} \frac{\partial H_{gy}}{\partial \mu_{gy}} \cdot \left( \nabla_{gy}^* f \times \nabla_{gy}^* g \right) + \left( \frac{\partial f}{\partial \mu_{gy}} \frac{\partial g}{\partial \theta_{gy}} - \frac{\partial f}{\partial \theta_{gy}} \frac{\partial g}{\partial \mu_{gy}} \right),
\]
where the definitions \[ (2.48) \]
\[
\nabla_{gy}^* f = \nabla_{gc}^* f - \epsilon \frac{q_s}{c} \left( \frac{\partial G[A_1(R_{gy} + \rho_{gc}, t)]}{\partial t} \frac{\partial f}{\partial \mu_{gy}} - \epsilon \delta q_s \frac{\partial G[A_1(R_{gy} + \rho_{gc}, t)]}{\partial \mu_{gy}} \frac{\partial f}{\partial \theta_{gy}} \right)
\]
and \[ \[ (2.49) \]
\[
\frac{\partial f}{\partial \mu_{gy}} = \frac{\partial f}{\partial \mu_{gy}} - \epsilon \delta \frac{q_s}{c} \left( \frac{\partial G[A_1(R_{gy} + \rho_{gc}, t)]}{\partial t} \frac{\partial f}{\partial \mu_{gy}} - \epsilon \delta \frac{\partial G[A_1(R_{gy} + \rho_{gc}, t)]}{\partial \mu_{gy}} \frac{\partial f}{\partial \theta_{gy}} \right)
\]
were used. With Eqs. (2.47) and (2.48), Hamilton’s equations of motion in gyrocentre coordinates read \[ (2.50) \]
\[
\dot{R}_{gy} = \frac{B^*_{\parallel}}{\mu_{gy}} \frac{\partial H_{gy}}{\partial v_{\parallel|gy}} + \epsilon \frac{q_s}{c} \nabla_{gy}^* f \cdot \left( \nabla_{gy}^* f \times \nabla_{gy}^* g \right) + \epsilon \frac{q_s}{c} \delta \frac{\partial G[A_1(R_{gy} + \rho_{gc}, t)]}{\partial t} \frac{\partial f}{\partial \theta_{gy}}
\]
\[
\dot{v}_{\parallel|gy} = - \frac{B^*_{\parallel}}{\mu_{gy}} \cdot \left( \nabla_{gy}^* f \cdot \left( \nabla_{gy}^* f \times \nabla_{gy}^* g \right) + \epsilon \frac{q_s}{c} \delta \frac{\partial G[A_1(R_{gy} + \rho_{gc}, t)]}{\partial t} \frac{\partial f}{\partial \theta_{gy}} \right)
\]
\[
\dot{\mu}_{gy} = 0
\]
\[
\dot{t} = 1.
\]
For brevity, we omitted the equations of motion for $\theta$ and $k$. As required at the beginning of the section, the equations of motion for $R_{gy}, v_{\parallel|gy}$ and $\mu_{gy}$ are independent of the fast gyroangle $\theta_{gy}$. With the gyrokinetic Lagrangian 1-form the dynamics of charged particles in the plasma is specified. We proceed by deriving the gyrokinetic Vlasov-Maxwell system to obtain equations for the distribution function as well as the perturbed electromagnetic fields. From now on, we drop the $gy$ annotation on the gyrocentre coordinates and assume that we work in gyrocentre coordinates unless explicitly specified otherwise. In addition, we omit the ordering parameters $\epsilon$ and $\epsilon_B$.

### 2.3.2. The Vlasov equation in gyrocentre coordinates

The particle follows a trajectory through the eight-dimensional extended phase space. The path is denoted with $\gamma : \lambda \mapsto \gamma(\lambda) = (R(\lambda), t(\lambda), v_{\parallel}(\lambda), \mu(\lambda), \theta(\lambda), k(\lambda))$ such that $\gamma(0)$ represents the initial state of the particle. According to Liouville’s theorem in Hamiltonian
mechanics, the distribution function in extended phase space, denoted by $F_\sigma$, is constant along the trajectories $\gamma$ of the system, i.e.

$$F_\sigma(\gamma(\lambda)) = F_\sigma(\gamma(0)).$$  \hspace{1cm} (2.56)

Differentiating Eq. (2.56) with respect to the orbit parameter yields the Vlasov equation in extended phase space

$$\frac{dF_\sigma}{dt} = \dot{R} \cdot \nabla F_\sigma + i \frac{dF_\sigma}{dt} + \dot{v}_\| \frac{\partial F_\sigma}{\partial v_\|} + \dot{\mu} \frac{\partial F_\sigma}{\partial \mu} + \dot{\theta} \frac{\partial F_\sigma}{\partial \theta} + k \frac{\partial F_\sigma}{\partial k} = 0. \hspace{1cm} (2.57)$$

As mentioned in the beginning, the physical motion of the particle takes place on the hypersurface $H = 0$. The extended phase space is reduced to six dimensions by projecting it onto the hypersurface defined by $H = 0$ and using Hamilton’s equations of motion for $t$ to replace the orbit parameter $\lambda$. To meet these physical requirements, the extended distribution function $F_\sigma$ is related to the six-dimensional, physical distribution function $f_\sigma$ via

$$F_\sigma = f_\sigma \delta(H(k)), \quad \text{where} \quad \delta \text{is the Dirac delta function} \ [30].$$

The Vlasov equation on the six-dimensional hypersurface is obtained by integrating Eq. (2.57) over $k$. It reads

$$\frac{df_\sigma}{dt} = \frac{\partial f_\sigma}{\partial t} + \dot{R} \cdot \nabla f_\sigma + \dot{v}_\| \frac{\partial f_\sigma}{\partial v_\|} + \dot{\mu} \frac{\partial f_\sigma}{\partial \mu} + \dot{\theta} \frac{\partial f_\sigma}{\partial \theta} = 0. \hspace{1cm} (2.58)$$

Similar to the last section, the particle distribution function $f_\sigma$ can be split into a gyroaveraged part $G[f_\sigma]$ and a fluctuating part $\tilde{f}_\sigma$ such that $f_\sigma = G[f_\sigma] + \tilde{f}_\sigma$. By construction of the gyrocentre coordinates, both parts evolve independently in time according to [105]

$$\frac{\partial G[f_\sigma]}{\partial t} + \dot{R} \cdot \nabla G[f_\sigma] + \dot{v}_\| \frac{\partial G[f_\sigma]}{\partial v_\|} + \dot{\mu} \frac{\partial G[f_\sigma]}{\partial \mu} = 0 \hspace{1cm} (2.59)$$

$$\frac{\partial \tilde{f}_\sigma}{\partial t} + \dot{R} \cdot \nabla \tilde{f}_\sigma + \dot{v}_\| \frac{\partial \tilde{f}_\sigma}{\partial v_\|} + \dot{\mu} \frac{\partial \tilde{f}_\sigma}{\partial \mu} + \dot{\theta} \frac{\partial \tilde{f}_\sigma}{\partial \theta} = 0. \hspace{1cm} (2.60)$$

In gyrokinetic theory the fluctuating part $\tilde{f}_\sigma$ is discarded and only the gyroaveraged part $G[f_\sigma]$ is retained. This removes the fast time scales associated with the gyroangle $\theta$ from the system and, therefore, reduces the phase space from six to five dimensions. The equation of motion for the gyroaveraged distribution function is known as the gyrokinetic Vlasov equation. In slight abuse of notation we denote the gyroaveraged distribution function $G[f_\sigma]$ with $f_\sigma$ and the gyroaveraged extended distribution function $G[F_\sigma]$ with $F_\sigma$ for the remainder of this work. The gyrokinetic Vlasov equation reads

$$\frac{\partial f_\sigma}{\partial t} + \dot{R} \cdot \nabla f_\sigma + \dot{v}_\| \frac{\partial f_\sigma}{\partial v_\|} = 0, \hspace{1cm} (2.61)$$

where the time derivative of the gyrocentre trajectories $\dot{\gamma}$ are given by Hamilton’s equation of motion presented in Eqs. (2.52) and (2.53).
2.3.3. The gyrokinetic action functional

To solve the dynamics of the plasma self consistently, additional equations for the perturbed electromagnetic fields $\phi_1$ and $A_1$ are needed. They are commonly derived via variational methods from the total gyrokinetic action functional of the plasma. We start with the particle Lagrangian, which is obtained from $\omega^1$ via Eq. (2.18). Expressed in gyrocentre coordinates, the particle Lagrangian reads

$$L(\gamma, \dot{\gamma}, \lambda) = \left( \frac{q_\sigma}{c} (A + A_1) + m_\sigma v_\parallel b - \frac{m_\sigma c}{q_\sigma} \mu R^* \right) \cdot \dot{R} + \frac{m_\sigma c}{q_\sigma} \mu \dot{\theta} - k_t - H. \quad (2.62)$$

From Eq. (2.62) the total gyrokinetic Lagrangian can be derived [107, 106]. It yields

$$L(\gamma, \dot{\gamma}, \phi_1, A_1, \lambda) = \sum_\sigma \int F_\sigma(\gamma(0), t) L(\gamma(0), \dot{\gamma}(0), t) d\Omega + \int \frac{E^2 - B_{\text{tot}}^2}{8\pi} dV dt, \quad (2.63)$$

where we used the volume element of the extended phase space $d\Omega = dW dV dt$, the volume element of velocity space $dV = 2\pi J_{\text{gy}}/m^3 \sigma d\mu$, the gyrocentre Jacobian $J_{\text{gy}} = J_{\text{gc}} + \frac{m^2_\sigma G}{b \cdot \nabla \times A_1}$, the volume element of real space $dV$, the electric field $E = -\nabla \phi_1$ and the magnetic field $B_{\text{tot}} = \nabla \times (A + A_1)$. In the following, we use cylindrical coordinates $(R, \varphi, Z)$ in real space such that $dV = R dR d\varphi dZ$. The gyrokinetic action functional is defined as the time integral of the total gyrokinetic Lagrangian

$$I(\gamma, \phi_1, A_1, \lambda) := \int_0^\lambda L(\gamma(\lambda), \dot{\gamma}(\lambda), \phi_1(\gamma(\lambda), \lambda), A_1(\gamma(\lambda), \lambda)) d\lambda. \quad (2.64)$$

The equations of motion for the particle in gyrocentre coordinates as well as the gyrokinetic field equations can be derived from the action functional with variational calculus [106, 88, 30].

To perform the calculation, we define the functional derivative. Let $I : V \to \mathbb{R}$ be a functional defined on a suitable vector space $V$. The functional derivative $\delta I : V \to \mathbb{R}$ for $\zeta \in V$ is the functional defined by

$$\delta I(\psi; \zeta) := \lim_{\epsilon \to 0} \frac{I(\psi + \epsilon \zeta) - I(\psi)}{\epsilon}. \quad (2.65)$$

By requiring $\delta I(\phi_1; \delta \phi_1) = 0$ the gyrokinetic Poisson equation is derived, by requiring $\delta I(A_1; \delta A_1) = 0$ the gyrokinetic Ampère’s law is derived and by requiring $\delta I(\gamma; \delta \gamma) = 0$ the Euler-Lagrange equations for the particle Lagrangian

$$\frac{d}{dt} \frac{\partial L}{\partial \dot{\gamma}_i} = \frac{\partial L}{\partial \gamma_i} \quad (2.66)$$

are derived, which are equivalent to Hamilton’s equation of motion presented in Eqs. (2.52), (2.53), (2.54) and (2.55). For simplicity, we perform the variation with respect to $\phi_1$ and $A_1$ after we have simplified the model. For a derivation of the field equations of the full system, represented by Eq. (2.64), we refer to [30, 88].
2.3.4. Simplification of the gyrokinetic Vlasov-Maxwell system

In order to focus on the development of a gyrokinetic turbulence code targeted for the edge and SOL, it is beneficial to further simplify the gyrokinetic Vlasov-Maxwell system. The physics in the edge and SOL is not only described by the gyrokinetic interaction of the plasma with the electromagnetic fields, but also by collisions and neutral physics, which are not covered by gyrokinetic theory. If needed, the approximations made can be lifted step by step in the future. We think that it is very important to understand the nature and origin of the approximations made. Therefore, we explain and justify the simplifications in this section in detail. In order to satisfy precise conservation laws with self-consistent equations, we simplify the system such that the Hamiltonian structure of the equations is retained. We start by neglecting all orthogonal fluctuations of the electromagnetic vector potential \( A_1 \), i.e.

\[
A_{1\perp} = 0. \tag{2.67}
\]

Orthogonal fluctuations in the electromagnetic vector potential give rise to parallel fluctuations in the magnetic field. They have shown to be important in the simulations of kinetic ballooning modes in the core of fusion devices [108]. We focus on the implementation of parallel electromagnetic fluctuations that already model finite \( \beta \) effects and decrease the frequency of kinetic shear Alfvén waves, which has a positive effect on the timestep of the simulation and is discussed further in Section 6.4. With this simplification the electromagnetic vector potential is solely given by its parallel component

\[
A_1 = A_{1\parallel} b. \tag{2.68}
\]

We neglect the \( \mathcal{O}(\epsilon B) \) term \( R^* \). Further, we employ a long wavelength limit \( k_\perp \rho \ll 1 \) [109] simplifying the gyrocentre displacement

\[
g(R + \rho) \approx g(R) + \rho \cdot (\nabla_\perp g)(R). \tag{2.69}
\]

With this simplification gyroaverages of functions of the form \( G[g(R + \rho)] \) reduce to the function \( g \) itself. Terms of the form \( G[\bar{g}] \) vanish. With the long wavelength approximation and the vanishing of \( A_\perp \), the second order part of the Hamiltonian, \( H_2 \), is greatly simplified and reads [109]

\[
H_2 \approx -\frac{q_a^2}{2B(R)} \frac{\partial G}{\partial \mu} \left[ \bar{\psi}^2 \right] \\
\approx -\frac{m_e c^2}{2B^2} \left| \nabla_\perp \phi_1 \right|^2. \tag{2.70}
\]

The long wavelength limit is often referred to as the drift-kinetic limit of the gyrokinetic equations [57, 59, 60]. Although most gyroaverage operations vanish, the second order, finite-Larmor radius effect in Eq. (2.70) is retained. It gives rise to the polarisation that is used to calculate the electrostatic potential \( \phi_1 \) in the gyrokinetic Poisson equation. The
simplified Hamiltonian reads \( H = H_0 + H_1 + H_2 \) with
\[
H_0 = \frac{1}{2} m_\sigma v_\parallel^2 + \mu B - k
\]
(2.71)
\[
H_1 = q_\sigma \phi_1
\]
(2.72)
\[
H_2 = -\frac{m_\sigma e^2}{2B^2} |\nabla \phi_1|^2
\]
(2.73)
and the simplified Poisson bracket in gyrocentre coordinates is given by
\[
\{ f, g \} = \frac{q_\sigma}{m_\sigma c} \left( \frac{\partial f}{\partial \theta} \frac{\partial g}{\partial \mu} - \frac{\partial f}{\partial \mu} \frac{\partial g}{\partial \theta} \right) + \frac{B^*}{m_\sigma B^\parallel} \cdot \left( \nabla f \frac{\partial g}{\partial v_\parallel} - \frac{\partial f}{\partial v_\parallel} \nabla g \right)
\]
\[
- \frac{q_\sigma}{m_\sigma c} \frac{\partial A^\parallel}{\partial t} \left( \frac{\partial f}{\partial k} \frac{\partial g}{\partial v_\parallel} - \frac{\partial f}{\partial v_\parallel} \frac{\partial g}{\partial k} \right)
\]
\[
- \frac{cb}{q_\sigma B^*} \cdot (\nabla f \times \nabla g) + \frac{\partial f}{\partial t} \frac{\partial g}{\partial k} - \frac{\partial f}{\partial k} \frac{\partial g}{\partial t},
\]
(2.74)
with the guiding-centre magnetic field
\[
B^* = B + \frac{m_\sigma e}{q_\sigma} v_\parallel \nabla \times b + \nabla \times (A^\parallel b).
\]
(2.75)
In this form \( B^* \) depends on \( A^\parallel \). Next, we simplify the free field part of the Lagrangian. We employ the quasi-neutrality approximation and neglect the \( E^2 \) term in Eq. (2.63) [106]. We linearise the polarisation by assuming that \( H_2 \) acts on the initial distribution \( F_{0\sigma} \) [110, 106].

The gyrokinetic Lagrangian with the described simplifications is given by
\[
\mathcal{L} = \sum_\sigma \int \left( \left( \frac{q_\sigma}{c} (A + A^\parallel b) + m_\sigma v_\parallel b \right) \cdot \dot{R} \right.
\]
\[
+ \frac{m_\sigma c}{q_\sigma} \mu \dot{t} - k \dot{t} - \frac{1}{2} m_\sigma v_\parallel^2 - \mu B - q_\sigma \phi_1 + k \right) F_{0\sigma} d\Omega
\]
\[
+ \int \left( \sum_\sigma \frac{m_\sigma e^2}{2B^2} |\nabla \phi_1|^2 F_{0\sigma} - \frac{1}{8\pi} |\nabla \times (A^\parallel b)|^2 \right) dV dt.
\]
(2.76)

Hamilton’s equations of motion read
\[
\dot{R} = \frac{\vec{B}^*}{B^\parallel} + \frac{c}{q_\sigma B^\parallel} \vec{b} \times (\mu \nabla B + q_\sigma \nabla \phi_1)
\]
(2.77)
\[
\dot{v}_\parallel = -\frac{\vec{B}^*}{m_\sigma B^\parallel} \cdot (\mu \nabla B + q_\sigma \nabla \phi_1) - \frac{q_\sigma}{m_\sigma c} \frac{\partial A^\parallel}{\partial t},
\]
(2.78)
where we have omitted the second order term, generated by \( H_2 \), to ensure energetic consistency due to the linearisation of the quasi-neutrality equation [110]. Finally, we assume that
the magnetic field created by $A_{1\parallel}$ is orthogonal to $b$ by simplifying

$$
\nabla \times (A_{1\parallel} b) = \nabla A_{1\parallel} \times b + A_{1\parallel} \nabla \times b
\approx \nabla A_{1\parallel} \times b.
$$

(2.79)

This approximation cannot be made on the Lagrangian entirely. The field part of the Lagrangian can be simplified

$$
\frac{1}{8\pi} |\nabla \times (A_{1\parallel} b)|^2 \approx \frac{1}{8\pi} |\nabla_{\perp} A_{1\parallel}|^2.
$$

Nevertheless, the modification of the symplectic part is not straightforward. Therefore, we perform the simplification on the Poisson bracket of the system, where $B^\ast$, presented in Eq. (2.75), is modified to

$$
B^\ast \approx B + m_\sigma v_{\parallel} c \frac{e}{q_\sigma} \nabla \times b + \nabla A_{1\parallel} \times b.
$$

(2.80)

This simplification removes the $A_{1\parallel}$ dependence of $B^\ast$ because $b \cdot (\nabla A_{1\parallel} \times b) = 0$. Consequently, $J_{\parallel v} = J_{v\parallel}$ and the velocity space volume element, $dW$, is independent of $A_{1\parallel}$, which decouples the gyrokinetic field equations.

### 2.3.5. The gyrokinetic field equations

The quasi-neutrality equation is derived by requiring that the variation of the action functional with respect to $\phi_1$ vanishes. The variation reads

$$
\delta I(\phi_1; \delta \phi_1) = - \sum_\sigma q_\sigma \int F_\sigma \delta \phi_1 \, d\Omega + \sum_\sigma \int \frac{m_\sigma c^2}{B^2} F_{0\sigma} \nabla_\perp \phi_1 \cdot \nabla_\perp \delta \phi_1 \frac{2\pi}{m_\sigma} B^\ast \parallel \delta \phi_1 \, dv_{\parallel} \, d\mu \, dV \, dt
\approx 0 \forall \delta \phi_1.
$$

(2.81)

This is also known as the weak formulation of the quasi-neutrality equation and can be used directly in a finite element discretisation scheme [111]. We are interested in a finite difference discretisation and derive a strong formulation of the quasi-neutrality equation. We use the product rule to modify the integrand

$$
\frac{F_{0\sigma}}{B^2} B^\ast |\nabla_\perp \phi_1 \cdot \nabla_\perp \delta \phi_1| = \nabla \cdot \left( \delta \phi_1 \frac{F_{0\sigma}}{B^2} B^\ast \nabla_\perp \phi_1 \right) - \nabla \cdot \left( \frac{F_{0\sigma}}{B^2} B^\ast \nabla_\perp \phi_1 \right) \delta \phi_1
$$

(2.82)

and apply the divergence integral theorem, where we assume that the boundary integral vanishes

$$
- \sum_\sigma q_\sigma \int F_\sigma \delta \phi_1 \, d\Omega - \sum_\sigma \int \frac{1}{B^\ast} \nabla \cdot \left( \frac{m_\sigma c^2}{B^2} F_{0\sigma} B^\ast \nabla_\perp \phi_1 \right) \delta \phi_1 \, d\Omega = 0 \forall \delta \phi_1.
$$

(2.83)

By applying the fundamental lemma of variational calculus, we obtain the strong form of the quasi-neutrality equation

$$
- \nabla \cdot \left( \sum_\sigma \frac{m_\sigma c^2}{B^2} n_{0\sigma} \nabla \phi_1 \right) = \sum_\sigma q_\sigma \int f_\sigma \, dW,
$$

(2.84)
with the initial density \( n_{0\sigma} = \int f_{0\sigma} \, dW \). Similar to the quasi-neutrality equation, we derive Ampère’s law by requiring that the variation of the action functional with respect to \( A_{1\|} \) vanishes

\[
\delta I(A_{1\|}; \delta A_{1\|}) = \sum_{\sigma} \frac{q_{\sigma}}{c} \int v_{\|} F_{\sigma} \delta A_{1\|} \, d\Omega - \frac{1}{4\pi} \int \nabla_{\perp} A_{1\|} \cdot \nabla_{\perp} \delta A_{1\|} \, dV \, dt = 0 \forall \delta A_{1\|}. \tag{2.85}
\]

After applying the divergence theorem and requiring that boundary terms vanish, we obtain Ampère’s law

\[
- \nabla_{\perp} \cdot \nabla_{\perp} A_{1\|} = 4\pi \sum_{\sigma} \frac{q_{\sigma}}{c} \int f_{\sigma} v_{\|} \, dW, \tag{2.86}
\]

where we have used the definition \( \nabla_{\perp} := \nabla \cdot \nabla_{\perp} \). In the symplectic formulation of the gyrokinetic Vlasov equation, presented above, explicit time derivatives of \( A_{1\|} \) appear. To allow for a straightforward temporal discretisation in Section 4.2, we derive a generalised Ohm’s law by taking the time derivative of Eq. (2.86) and solving for \( \frac{\partial A_{1\|}}{\partial t} \) directly

\[
- \nabla_{\perp} \frac{\partial A_{1\|}}{\partial t} = 4\pi \sum_{\sigma} \frac{q_{\sigma}}{c} \int \frac{\partial f_{\sigma}}{\partial t} v_{\|} \, dW. \tag{2.87}
\]

We write the Vlasov equation as

\[
\frac{\partial f_{\sigma}}{\partial t} = \left( \frac{\partial f_{\sigma}}{\partial t} \right)^{\ast} + \frac{q_{\sigma}}{m_{\sigma} c} \frac{\partial f_{\sigma}}{\partial v_{\parallel}} \frac{\partial A_{1\|}}{\partial t}, \tag{2.88}
\]

where \( \left( \frac{\partial f_{\sigma}}{\partial t} \right)^{\ast} \) denotes the part excluding explicit time derivatives. Inserting Eq. (2.88) into Eq. (2.87) yields the generalised Ohm’s law

\[
- \left( \nabla_{\perp} + 4\pi \sum_{\sigma} \frac{q_{\sigma}^2}{m_{\sigma} c^2} \right) \frac{\partial f_{\sigma}}{\partial v_{\parallel}} v_{\|} \, dW \right) \frac{\partial A_{1\|}}{\partial t} = 4\pi \sum_{\sigma} \frac{q_{\sigma}}{c} \int \left( \frac{\partial f_{\sigma}}{\partial t} \right)^{\ast} v_{\|} \, dW. \tag{2.89}
\]

Eqs. (2.61), (2.77), (2.78), (2.80), (2.84), (2.86) and (2.89) describe the system implemented in the GENE-X code.

### 2.3.6. Conservation laws

The gyrokinetic Vlasov-Maxwell system, presented in the last section, fulfils multiple conservation laws. On the one hand, conservation laws are important to ensure numerical stability, on the other hand, they can serve as a method of verification of the numerical scheme. In this section, we discuss the conservation of the particle number, momentum and energy of the gyrokinetic system. As we performed the approximation in Eq. (2.75) on the Poisson bracket and not on the Lagrangian of the system, we proof the conservation by direct calculation. We start with the energy

\[
E = E_{H} - E_{\phi} + E_{A}, \tag{2.90}
\]
with the contributions

\[ E_H = \sum_{\sigma} \int H_\sigma f_\sigma \, dV \, dW = \sum_{\sigma} \int \left( \frac{1}{2} m_\sigma v_{\parallel}^2 + \mu B + q_\sigma \phi \right) f_\sigma \, dV \, dW \]  
(2.91)

\[ E_A = \int \frac{1}{8\pi} |\nabla_\perp A_{\parallel}|^2 \, dV \]  
(2.92)

\[ E_\phi = \int \frac{m_\sigma e^2}{2B^2} n_{0\sigma} |\nabla_\perp \phi_1|^2 \, dV. \]  
(2.93)

We assume that the domain is periodic and boundary terms vanish. The time derivative of the Hamiltonian part of the energy reads

\[ \frac{\partial E_H}{\partial t} = \sum_{\sigma} \int \left( \frac{\partial H_\sigma}{\partial t} f_\sigma + H_\sigma \frac{\partial f_\sigma}{\partial t} \right) \, dV \, dW \]

\[ = \sum_{\sigma} \int \left[ q_\sigma \frac{\partial \phi_1}{\partial t} f_\sigma + H_\sigma \left( \{ f_\sigma, H_\sigma \}^* + \frac{q_\sigma}{m_\sigma c} \frac{\partial A_{\parallel}}{\partial v_{\parallel}} \right) \right] \, dV \, dW \]

\[ = \sum_{\sigma} \int \left[ q_\sigma \frac{\partial \phi_1}{\partial t} f_\sigma - \frac{q_\sigma}{c} \frac{\partial A_{\parallel}}{\partial t} v_{\parallel} f_\sigma \right] \, dV \, dW, \]  
(2.94)

where we used the integral property of the Poisson bracket

\[ \int H_\sigma \{ f_\sigma, H_\sigma \}^* \, dV \, dW = 0, \]  
(2.95)

proved in Appendix A.4, and partial integration in \( v_{\parallel} \). The Poisson bracket \( \{ , \}^* \) is given by Eq. (2.74) without the \( \partial A_{\parallel}/\partial t \) term. We continue with the energy contribution of the electromagnetic vector potential. The rate of change of the electromagnetic energy is simplified with Ampère’s law to

\[ \frac{\partial E_A}{\partial t} = \int \frac{1}{8\pi} \frac{\partial}{\partial t} |\nabla_\perp A_{\parallel}|^2 \, dV = \int \frac{1}{4\pi} \nabla_\perp A_{\parallel} \cdot \nabla_\perp \frac{\partial A_{\parallel}}{\partial t} \, dV \]

\[ = - \int \frac{1}{4\pi} \Delta_\perp A_{\parallel} \frac{\partial A_{\parallel}}{\partial t} \, dV = \sum_{\sigma} \frac{q_\sigma}{c} \int \frac{\partial A_{\parallel}}{\partial t} v_{\parallel} f_\sigma \, dV \, dW, \]  
(2.96)

and the rate of change of the electrostatic energy is simplified with the quasi-neutrality equation to

\[ \frac{\partial E_\phi}{\partial t} = \sum_{\sigma} \int \frac{m_\sigma e^2}{2B^2} n_{0\sigma} \frac{\partial}{\partial t} |\nabla_\perp \phi_1|^2 \, dV = \sum_{\sigma} \int \frac{m_\sigma e^2}{B^2} n_{0\sigma} \nabla_\perp \phi_1 \cdot \nabla_\perp \frac{\partial \phi_1}{\partial t} \, dV \]

\[ = - \sum_{\sigma} \int \nabla \cdot \left( \frac{m_\sigma e^2}{2B^2} n_{0\sigma} \nabla \phi_1 \right) \frac{\partial \phi_1}{\partial t} \, dV = \sum_{\sigma} \int q_\sigma \frac{\partial \phi_1}{\partial t} f_\sigma \, dV \, dW. \]  
(2.97)

Combining Eqs. (2.94), (2.96) and (2.97) yields

\[ \frac{\partial E}{\partial t} = \frac{\partial E_H}{\partial t} - \frac{\partial E_\phi}{\partial t} + \frac{\partial E_A}{\partial t} = 0. \]  
(2.98)
The expression for the energy is further simplified by using the weak form of the quasi-neutrality equation, presented in Eq. (2.81), with \( \delta \phi_1 = \phi_1 \)

\[
E = \sum_{\sigma} \int \left( \frac{1}{2} m_\sigma v_\parallel^2 + \mu B + \frac{1}{2} q_\sigma \phi_1 \right) f_\sigma \, dV \, dW + \int \frac{1}{8 \pi} |\nabla_\perp A_{1\parallel}|^2 dV .
\]  

(2.99)

Liouville’s theorem states that the characteristics of the system satisfy the conservation law \([30, 42]\)

\[
\nabla \cdot \left( \mathcal{J}_{gy} \dot{R} \right) + \frac{\partial}{\partial v_\parallel} \left( \mathcal{J}_{gy} v_\parallel \right) = 0 .
\]  

(2.100)

This property can be used to write the gyrokinetic Vlasov equation in the form of a conservation law

\[
\frac{\partial (\mathcal{J}_{gy} f_\sigma)}{\partial t} + \nabla \cdot \left( \mathcal{J}_{gy} f_\sigma \dot{R} \right) + \frac{\partial}{\partial v_\parallel} \left( \mathcal{J}_{gy} f_\sigma v_\parallel \right) = 0 .
\]  

(2.101)

Integrating Eq. (2.101) over phase space and assuming that boundary terms vanish yields the conservation of the number of particles

\[
n_\sigma = \int f_\sigma \, dV \, dW .
\]  

(2.102)

Finally, we discuss the conservation of canonical toroidal momentum. For a complete discussion of momentum conservation of the gyrokinetic Vlasov-Maxwell system we refer to \([110, 112]\). The canonical toroidal momentum of a particle is defined as

\[
p_\varphi := \frac{\partial L}{\partial \dot{\varphi}},
\]  

(2.103)

where \( \varphi \) denotes the toroidal angle in a cylindrical coordinate system. We use the canonical toroidal momentum in Section 4.5 to construct an initial condition for the simulations. The canonical toroidal momentum of a particle is conserved under the assumption of an axisymmetric equilibrium \( B = B(R,Z) \) and a time independent electromagnetic vectorpotential \( A_{1\parallel} \). In the following, we assume that \( A_{1\parallel} = 0 \). The conservation of the canonical toroidal momentum follows directly from the Euler-Lagrange equations and the independence of the particle Lagrangian \( L \) of \( \varphi \), i.e.

\[
\frac{\partial L}{\partial \dot{\varphi}} = 0 .
\]  

(2.104)

Using \( R = R e_R + Z e_Z \) and \( \dot{R} = \dot{R} e_R + R \dot{\varphi} e_\varphi + \dot{Z} e_Z \), the canonical toroidal momentum reads

\[
p_\varphi = \left( \frac{q_\sigma}{c} A + m_\sigma v_\parallel b \right) Re_\varphi
\]

\[
= \frac{q_\sigma}{c} R e_\varphi + m_\sigma v_\parallel b_\varphi R .
\]  

(2.105)
We simplify the expression by using the poloidal flux function defined as†

\[
\psi_p := \frac{1}{2\pi} \int_{D_R(0)} B \cdot dF = \frac{1}{2\pi} \int_{D_R(0)} \nabla \times A \cdot dF = \frac{1}{2\pi} \int_{\partial D_R(0)} A \cdot dl = RA_{\varphi},
\]

where the axisymmetry of the electromagnetic vector potential and Stokes theorem were used. The set \(D_R(0)\) denotes a disk of radius \(R\) with the origin at the magnetic axis in the \(R,Z\) plane. With the poloidal flux function, the canonical toroidal momentum simplifies to

\[
p_{\varphi} = \frac{q_{\sigma}}{c} \psi_p + m_{\sigma} v_{||} b_{\varphi} R.
\]

The conserved canonical toroidal momentum of the total gyrokinetic system reads [112]

\[
P_{\varphi} := \sum_{\sigma} \int \left( \frac{q_{\sigma}}{c} \psi_p + m_{\sigma} v_{||} b_{\varphi} R \right) f_{\sigma} dV dW.
\]

### 2.4. Conclusion and outlook

In this chapter, we presented an overview of gyrokinetic theory and introduced the gyrokinetic model implemented in the GENE-X code. The model presented is consistent, evolves the full distribution function and fulfils multiple exact conservation laws. We put a special emphasis on a precise presentation of the gyrokinetic theory and a rigorous simplification from the full second order theory to the model implemented. This makes the generalisation of the model straightforward in the future.

The model implemented uses a long wavelength limit, a linearisation of the polarisation density and neglects perpendicular electromagnetic fluctuations. For the application of GENE-X to the edge and SOL, the linearisation of the polarisation density should be lifted first. Next to a nonlinear quasi-neutrality equation, this generates additional terms in the gyrokinetic Vlasov equation as the second order part of the Hamiltonian \(H_2\), generating the polarisation, acts on the full distribution function \(f_{\sigma}\) and not the linearisation \(f_{0\sigma}\) [110, 106]. Furthermore, the implementation of gyroaverages and, consequently, further second order terms would be interesting.

Next to the gyrokinetic plasma model, other physical effects like collisions and neutral physics are important in the edge and SOL. In Chapter 7, we present first simulations that use a simplified Bhatnagar-Gross-Krook (BGK) collision operator [113]. The implementation of more sophisticated collision operators, like a full-\(f\) Lennard-Bernstein/Dogherty operator [114, 115], is work in progress [116]. To study the effect of neutrals on gyrokinetic edge and SOL turbulence, implementing the simple fluid neutral model used in the GRILLIX code [80] is a good starting point for future developments.

†Note that the poloidal flux function defined here contains a factor of \(1/(2\pi)\), as it appears in the Grad-Shafranov equation. It is also common to define the poloidal flux without the factor of \(1/(2\pi)\), e.g. in [8].
3. Coordinate systems

Due to the strong magnetic field in fusion devices, structures in the plasma are homogeneous along and heterogeneous perpendicular to magnetic field lines. This property is also referred to as the flute-like character of the plasma and produces small gradients parallel to magnetic field lines. It can be exploited in turbulence simulations by choosing a coordinate system in which parallel gradients resolve to a partial derivative in one of the coordinate variables. This enables the use of a reduced resolution in the corresponding direction, sufficient to resolve the small gradients, and spares computational resources.

For predictive edge and SOL simulations it is further important that the chosen coordinate system is well-defined in realistic magnetic geometries, including the separatrix and one or even multiple X-points.

In this chapter, we discuss coordinate systems suitable for turbulence simulations in magnetic confinement fusion devices. We discuss the advantages and disadvantages of established flux-coordinate systems and introduce the locally field-aligned coordinate system based on the flux-coordinate independent approach [76], implemented in the GENE-X code. This coordinate system is free of coordinate singularities while still being aligned to the magnetic field. It can handle general fusion device geometries including multiple X-points and stochastic regions.

3.1. Flux coordinate systems

In the gyrokinetic community so-called flux-coordinate systems are common. In these magnetic field lines are straight and one coordinate, we call it $\zeta$, can be chosen such that the desired property

$$\mathbf{b} \cdot \nabla f \propto \frac{\partial f}{\partial \zeta}$$

(3.1)

is fulfilled. We start the discussion of flux-coordinate systems from the cylindrical-toroidal coordinates introduced in Chapter 2. For simplicity, we assume that the magnetic field is axisymmetric and given as a function of $(r, \theta, \varphi)$. The first coordinate $\psi$ of the flux-coordinate system is a label for the flux surfaces. It can for example be realised with the toroidal flux function, the poloidal flux function, the volume of a flux surface or the pressure on a flux surface. [73]. Consider a magnetic field line on a given flux surface. The rotational transform angle $\iota$ of the magnetic field line is defined as the poloidal angle $\theta$ travelled by a field line

---

1The coordinate transformations presented in this chapter affect the gyrocentre position $\mathbf{R}$ and not the velocity space.
after one poloidal transit [73]. It can be expressed as [73]

$$
\zeta := \frac{\ell}{2\pi} := \frac{d\psi_p}{d\psi_t},
$$

(3.2)

where $\psi_p$ represents the poloidal flux function, defined in Eq. (2.106), and

$$
\psi_t := \frac{1}{2\pi} \int_{A_t} B \cdot dF
$$

(3.3)

the toroidal flux function. The set $A_t$ is given by the part of a $\varphi = \text{const}$ surface that lies within the magnetic flux surface under consideration [73]. The $q$ factor is defined as the reciprocal of $\zeta$, i.e.

$$
q := \frac{1}{\zeta}.
$$

(3.4)

The two remaining coordinates of the flux-coordinate system are chosen such that magnetic field lines are described by straight lines on a flux surface. Such a coordinate system can be constructed by replacing either the toroidal angle or the poloidal angle with a new flux-coordinate, or both at the same time [73]

$$
\varphi \rightarrow \varphi_f
$$

(3.5)

$$
\theta \rightarrow \theta_f.
$$

(3.6)

The choice of $\varphi_f$ and $\theta_f$ is not unique and many different flux coordinate systems can be constructed. Often used are for example the coordinates implemented in the PEST code [117] and the coordinate systems described by Hamada [118] and Boozer [119]. For the further discussion, we assume generic flux angles $\theta_f$ and $\varphi_f$. The flux-coordinate system can be modified such that the magnetic field is represented as a Clebsch form

$$
B = \nabla \psi \times \nabla \nu,
$$

(3.7)

with the new coordinate [73]

$$
\nu = \frac{1}{2\pi} \frac{\partial \psi_t}{\partial \psi} \varphi_f - \frac{1}{2\pi} \frac{\partial \psi_p}{\partial \psi} \varphi_f.
$$

(3.8)

In the Clebsch form $\psi$ and $\nu$ identify a magnetic field line on a magnetic flux surface. The position on a magnetic field line is then provided by the additional coordinate $\zeta$. As $\nabla \psi \cdot b = 0$ and $\nabla \nu \cdot b = 0$, the magnetic field vector $b$ is proportional to $\nabla \zeta$ and parallel derivatives resolve to partial derivatives in $\zeta$. In tokamaks it is common to use the poloidal flux function as a flux surface label such that [73]

$$
\psi = \psi_p
$$

(3.9)

$$
\nu = q\theta_f - \varphi_f
$$

(3.10)

$$
\zeta = \theta_f,
$$

(3.11)
and in stellarators it is common to use the toroidal flux function as a flux surface label such that [73]

\[ \psi = \frac{\psi_k}{2\pi} \tag{3.12} \]
\[ \nu = \theta_f - \nu \varphi_f \tag{3.13} \]
\[ \zeta = \varphi_f . \tag{3.14} \]

Nevertheless, these coordinate systems have two major drawbacks making them unsuitable for the simulation of edge and SOL turbulence. We explain the first problem at the example of the coordinates given in Eqs. (3.9), (3.10) and (3.11). The term \( q \theta_f \) in the coordinate \( \nu \) increases with the poloidal angle \( \theta_f \). This introduces a distortion in the grid for large poloidal angles and large magnetic shear [120]. This problem can be partially solved by using the so-called shifted-metric approach [120]. The manifold \( M \) of the plasma, introduced in Chapter 2, is divided over the variable \( \theta_f \) into a family of overlapping sectors, labelled by \( \{ \theta_k \} \). Within each sector the coordinate transformation is given by

\[ \psi = \frac{\psi_p}{2\pi} \tag{3.15} \]
\[ \nu = q(\theta_f - \theta_k) - \varphi_f \tag{3.16} \]
\[ \zeta = (\theta_f - \theta_k) . \tag{3.17} \]

If the number of sectors is chosen equal to the number of grid points in \( \theta_f \), the grid in \( \psi, \nu \) is orthogonal. Second, flux-coordinates are generally ill-defined at the separatrix. The reason is the following. In flux coordinates magnetic field lines are straight and their equation can be written as [73]

\[ \theta_f = \nu \varphi_f + \theta_0 . \tag{3.18} \]

This implies that the flux-coordinates need to fulfil the relation

\[ \frac{d\theta_f}{d\varphi_f} = \ast . \tag{3.19} \]

At the X-point the toroidal transform vanishes and the angle \( \theta_f \) reduces to a constant \( \theta_0 \)

\[ \theta_f = \theta_0 . \tag{3.20} \]

Hence, the coordinate transformation into flux-coordinates cannot be bijective [74] and is ill-defined. To visualise the behaviour of flux coordinates close to the separatrix, we present the lines of constant \( \psi \) and \( \theta_f \) of the PEST coordinate system in Fig. 3.1. In summary, it is not possible to use flux coordinate systems to simulate turbulence across the separatrix.
3.2. A locally field-aligned coordinate system

Including the separatrix and handling large magnetic shear is important for edge and SOL plasmas. Therefore, we introduce a locally field-aligned coordinate system following the flux-coordinate independent (FCI) approach [76] in this section. This coordinate system retains the property that parallel gradients can be expressed as a partial derivative in one of the coordinate variables while being able to handle the separatrix and large shear. The coordinate system is similar to the one used in the GRILLIX code [86]. We introduce the coordinate transformation and express the gyrokinetic Vlasov-Maxwell system in the new coordinates.

3.2.1. Coordinate transformation

The results presented in this subsection have been published in [87]. For the completeness of the presentation the inverse coordinate transformation, presented in Eq. (3.27), has been added.

Consider a family of poloidal planes located at the angles $\varphi_k$. We define the locally field-aligned coordinate system in a region $[\varphi_k - \Delta \varphi, \varphi_k + \Delta \varphi]$ around the poloidal plane. For linear fusion devices or calculations with a large aspect ratio assumption, the toroidal angle $\varphi$ is interpreted as a Cartesian coordinate labelling poloidal planes. For every $k$ we define a different coordinate system. Hence, the coordinate system is local to every poloidal plane.
We choose $\Delta \varphi$ large enough that the coordinate systems for different poloidal planes overlap and the collection of all these coordinate systems, or charts respectively, form an atlas of the manifold $\mathcal{M}$. In the following, we assume without loss of generality that the poloidal plane is located at $\varphi_k = 0$. It is always possible to rotate the cylindrical $(R, \varphi, Z)$ coordinate system such that $\varphi_k = 0$ is fulfilled. To derive the coordinate transformation, we use a Cartesian reference frame $(x_c, y_c, z_c)$ on the poloidal plane such that $x_c$ points in the direction of the cylindrical $R$ coordinate, $z_c$ points in the direction of the cylindrical $Z$ coordinate and $y_c$ out of the poloidal plane. The magnetic field line is a curve $\gamma : [0, 1] \to \mathcal{M}$ on the configuration space manifold $\mathcal{M}$ and defined by the differential equation

$$\frac{d\gamma}{d\lambda} = b \circ \gamma. \quad (3.21)$$

Let $\gamma_{\text{cart}} = (\gamma_{x_c}, \gamma_{y_c}, \gamma_{z_c})$ be the curve of the magnetic field line expressed in the Cartesian reference frame. Eq. (3.21) reads

$$\frac{d\gamma_{x_c}}{d\lambda} = b_{x_c} \circ \gamma_{\text{cart}} \quad (3.22) \quad \frac{d\gamma_{y_c}}{d\lambda} = b_{y_c} \circ \gamma_{\text{cart}} \quad (3.23) \quad \frac{d\gamma_{z_c}}{d\lambda} = b_{z_c} \circ \gamma_{\text{cart}}. \quad (3.24)$$

We define the locally field-aligned coordinate system $(x, y, z)$ by giving the coordinate transformation $\tau$ from the locally field-aligned coordinate system to the Cartesian reference frame

$$\tau : \begin{pmatrix} x \\ y \\ z \end{pmatrix} \mapsto \begin{pmatrix} x_c \\ y_c \\ z_c \end{pmatrix} = \begin{pmatrix} x + \int_0^y b_{x_c}(\gamma_{\text{cart}}(u)) \, du \\ \int_0^y b_{y_c}(\gamma_{\text{cart}}(u)) \, du \\ z + \int_0^y b_{z_c}(\gamma_{\text{cart}}(u)) \, du \end{pmatrix}. \quad (3.25)$$

Next, we construct the inverse $\tau^{-1}$. The function $\gamma_{y_c}$ assigns the $y_c$ coordinate of the magnetic field line to its orbit parameter $\lambda$ and is invertible in a region around the poloidal plane. In this region the coordinate transformation can be inverted by substituting the orbit parameter $u$, in the integral, with the function $\gamma_{y_c}^{-1}(u)$ and using the derivative rule of the inverse function

$$\frac{d\gamma_{y_c}^{-1}}{du}(u) = \frac{1}{\frac{d\gamma_{y_c}}{du}(\gamma_{y_c}^{-1}(u))} = \frac{1}{b_{y_c}(\gamma_{y_c}^{-1}(u))}. \quad (3.26)$$

The inverse reads

$$\tau^{-1} : \begin{pmatrix} x_c \\ y_c \\ z_c \end{pmatrix} \mapsto \begin{pmatrix} x \\ y \\ z \end{pmatrix} = \begin{pmatrix} x_c - \int_0^{y_c} \frac{b_{x_c}(\gamma_{\text{cart}}(\gamma_{y_c}^{-1}(u)))}{b_{y_c}(\gamma_{\text{cart}}(\gamma_{y_c}^{-1}(u)))} \, du \\ \int_0^{y_c} \frac{b_{y_c}(\gamma_{\text{cart}}(\gamma_{y_c}^{-1}(u)))}{b_{y_c}(\gamma_{\text{cart}}(\gamma_{y_c}^{-1}(u)))} \, du \\ z_c - \int_0^{y_c} \frac{b_{z_c}(\gamma_{\text{cart}}(\gamma_{y_c}^{-1}(u)))}{b_{y_c}(\gamma_{\text{cart}}(\gamma_{y_c}^{-1}(u)))} \, du \end{pmatrix}. \quad (3.27)$$

The new coordinate system can be interpreted in a simple way. The $y$ coordinate corresponds to the length of the magnetic field line starting at the point $(x_c, y_c = 0, z_c)$. The $x$ and $z$
coordinates correspond to the $x_c$ and $y_c$ coordinates of the point the field line is originating from. On the poloidal plane, i.e. $y_c = \varphi = y = 0$, the locally field-aligned coordinates coincide with the Cartesian reference frame. The Jacobian matrix of the coordinate transformation evaluated at $y = 0$ reads

$$J(x, 0, z) = \begin{pmatrix} \frac{\partial \tau}{\partial x} (x, 0, z), \frac{\partial \tau}{\partial y} (x, 0, z), \frac{\partial \tau}{\partial z} (x, 0, z) \end{pmatrix} = \begin{pmatrix} 1 & b_{x_c} & 0 \\ 0 & b_{y_c} & 0 \\ 0 & b_{z_c} & 1 \end{pmatrix}, \quad (3.28)$$

and its inverse is given by

$$J^{-1}(x_c, 0, z_c) = \begin{pmatrix} \frac{\partial \tau^{-1}}{\partial x_c} (x_c, 0, z_c), \frac{\partial \tau^{-1}}{\partial y_c} (x_c, 0, z_c), \frac{\partial \tau^{-1}}{\partial z_c} (x_c, 0, z_c) \end{pmatrix} = \begin{pmatrix} 1 & -\frac{b_{x_c}}{b_{y_c}} & 0 \\ 0 & \frac{1}{b_{y_c}} & 0 \\ 0 & -\frac{b_{z_c}}{b_{y_c}} & 1 \end{pmatrix}. \quad (3.29)$$

For clarity, we omit the dependence of $b_{x_c}$, $b_{y_c}$, and $b_{z_c}$ on $x$ and $z$. The basis of the tangent space at $y_c = 0$ is given by the columns of the Jacobian matrix

$$e_x = \begin{pmatrix} 1 \\ 0 \\ 0 \end{pmatrix}, \quad e_y = \begin{pmatrix} b_{x_c} \\ b_{y_c} \\ 0 \end{pmatrix}, \quad e_z = \begin{pmatrix} 0 \\ 0 \\ 1 \end{pmatrix}. \quad (3.30)$$

The basis is non-orthogonal. The basis vector $e_y$ is equal to the unit vector of the magnetic field. The coordinate system is depicted in Fig. 3.2. We continue by introducing the metric tensor and the dual basis of the coordinate system to set the stage for the derivation of the differential operators.
3.2.2. Metric coefficients

The results presented in this subsection have been published in [87]. For the completeness of the presentation the inverse metric, presented in Eq. (3.33), and a discussion about the orthogonal projection operator, presented in Eqs. (3.41)–(3.44), have been added.

The components of the metric tensor read

\[ g_{ij} = J^T J \] (3.31)

\[ = \begin{pmatrix} 1 & b_{xc} & 0 \\ b_{xc} & 1 & b_{ze} \\ 0 & b_{ze} & 1 \end{pmatrix} , \] (3.32)

the inverse reads

\[ g^{ij} = \frac{1}{b_{yc}^2} \begin{pmatrix} b_{xc}^2 + b_{yc}^2 & -b_{xc} & b_{xc} b_{ze} \\ -b_{xc} & 1 & -b_{ze} \\ b_{xc} b_{ze} & -b_{ze} & b_{ze}^2 + b_{yc}^2 \end{pmatrix} \] (3.33)

and the Jacobian determinant is given by

\[ J_{fc1} := \sqrt{\det (g_{ij})} = b_{yc} . \] (3.34)

As gradients correspond to 1-forms, they are contained in the dual of the tangent space. The basis of the dual space can be expressed as

\[ e^i = g^{ij} e_j , \] (3.35)

where the basis vectors \( e_i \) of the tangent space, presented in Eq. (3.30), were used. Evaluating Eq. (3.35) yields

\[ e^x = \begin{pmatrix} 1, \ -\frac{b_{xc}}{b_{yc}}, \ 0 \end{pmatrix} \] (3.36)

\[ e^y = \begin{pmatrix} 0, \ \frac{1}{b_{yc}}, \ 0 \end{pmatrix} \] (3.37)

\[ e^z = \begin{pmatrix} 0, \ -\frac{b_{ze}}{b_{yc}}, \ 1 \end{pmatrix} . \] (3.38)

We derive the basis expansion of the covariant \( b \) vector

\[ b = b_i e^i . \]

Remembering that \( b = e_y \), due to the alignment of the \( y \) coordinate line with the magnetic field line, and using the relation between reciprocal sets of basis vectors \( e_i = g_{ij} e^j \), we find that

\[ b = e_y = g_{2ij} e^j = b_{xc} e^x + e^y + b_{ze} e^z . \] (3.39)
This expression provides us with covariant components of the magnetic unit vector in the chosen coordinate system

\[ (b_i) = (b_{xc}, 1, b_{zc}) . \]  

Finally, we derive two new basis vectors \( e_{1\perp} \) and \( e_{2\perp} \) that form an orthogonal basis system \((e_y, e_{1\perp}, e_{2\perp})\) with the magnetic field unit vector \( e_y \). By using the Gram-Schmidt process on the basis vectors \((e_y, e_z, e_x)\) and rescaling the third output vector of the Gram-Schmidt process, we obtain

\[ e_{1\perp} = \begin{pmatrix} -b_{xz} / b_{zc} \\ -b_{zc} / b_{yc} \\ 1 - b_{zc}^2 \end{pmatrix} \quad e_{2\perp} = \begin{pmatrix} 1 \\ -b_{yc} / b_{zc} \\ 0 \end{pmatrix} . \]  

We use Eq. (3.41) to construct a projection operator into the two-dimensional plane orthogonal to \( e_y \). Given a vector \( v = v^x e_x + v^y e_y + v^z e_z \) in the locally field-aligned coordinate system. We perform a change of basis into \((e_y, e_{1\perp}, e_{2\perp})\)

\[ v = \begin{pmatrix} v^y + b_{xc} v^x + b_{yc} v^z \\ v^z + b_{zc} b_{yc} / (1 - b_{zc}^2) v^x \\ v^x - b_{xc}^2 v^z / (1 - b_{zc}^2) \end{pmatrix} e_{1\perp} + \begin{pmatrix} v^x \\ v^z \\ v^z \end{pmatrix} e_{2\perp} \]  

and calculate the orthogonal projection \( v_\perp \) by setting the component parallel to \( e_y \) to zero. Reverting the change of basis to \((e_x, e_y, e_z)\) yields \( v_\perp \) expressed in the locally field-aligned coordinate system

\[ v_\perp = v^x e_x - (b_{xc} v^x + b_{zc} v^z) e_y + v^z e_z . \]  

Formally, we define the orthogonal projection operator via

\[ P_\perp : v \rightarrow v^x e_x - (b_{xc} v^x + b_{zc} v^z) e_y + v^z e_z . \]  

With these we are set to derive differential operators in the new coordinate system.

### 3.2.3. Differential operators

The results presented in this subsection have been published in [87]. For the completeness of the presentation a discussion about the operator \( \nabla \cdot (h \nabla \perp f) \), presented in Eqs. (3.56)–(3.60), has been added.

There are four types of differential operators in the gyrokinetic Vlasov-Maxwell system which need to be expressed in the locally field-aligned coordinate system

\[ b \cdot \nabla f, \quad (\nabla \times b) \cdot \nabla f, \quad (b \times \nabla h) \cdot \nabla f \quad \text{and} \quad \nabla \cdot (h \nabla \perp f) , \]  

where \( b = e_y \) and \( f, h \) are arbitrary scalar functions. The first operation corresponds to the
parallel derivative and is given by

\[ b \cdot \nabla f = e_y \cdot e^i \frac{\partial f}{\partial x^i} = \frac{\partial f}{\partial y}. \]  

(3.46)

Eq. (3.46) shows that parallel derivatives are given by derivatives in the \( y \) coordinate. Due to the flute-like property of the plasma, mentioned in the first section of this chapter, it is, similar to flux-coordinates, possible to use a much coarser resolution in toroidal than in poloidal direction. This saves computational resources. For the second and third operator we express the curl and cross product in the new coordinate system. The curl of the magnetic field unit vector \( b \) is given by

\[ \nabla \times b = \frac{1}{J_{eci}} \sum_{k=1}^{3} \epsilon_{ijk} \left( \frac{\partial b_j}{\partial x^i} - \frac{\partial b_i}{\partial x^j} \right) e_k, \]

\[ = \frac{1}{b_{yc}} \left[ \frac{\partial b_z}{\partial y} e_x - \frac{\partial b_x}{\partial y} e_z + \left( \frac{\partial b_x}{\partial z} - \frac{\partial b_z}{\partial x} \right) e_y \right], \]  

(3.47)

where the covariant components \( b_i \), presented in Eq. (3.40), were used. The gradient of a function \( f \) in a curvilinear coordinate system can be expressed as

\[ \nabla f = \frac{\partial f}{\partial x^i} e^i. \]  

(3.48)

We obtain the second operation by combining Equations (3.47) and (3.48) and considering that \( e_i \cdot e^j = \delta^j_i \)

\[ (\nabla \times b) \cdot \nabla f = \frac{1}{b_{yc}} \left[ \frac{\partial b_z}{\partial y} \frac{\partial f}{\partial x} - \frac{\partial b_x}{\partial y} \frac{\partial f}{\partial z} + \left( \frac{\partial b_x}{\partial z} - \frac{\partial b_z}{\partial x} \right) \frac{\partial f}{\partial y} \right]. \]  

(3.49)

The third operation is a triple vector product of the magnetic unit vector and two gradients of scalar functions. The gradients of the functions \( h \) and \( f \) are given by

\[ \nabla h = \frac{\partial h}{\partial x^i} e^i \quad \nabla f = \frac{\partial f}{\partial x^j} e^j. \]  

(3.50)

The vector product in the curvilinear coordinate system takes the form

\[ A \times B = \frac{1}{J_{eci}} \sum_{k=1}^{3} \epsilon_{ijk} \left( A_i B_j - A_j B_i \right) e_k \]  

(3.51)

By combining Eqs. (3.39), (3.50) and (3.51), we obtain the vector product of the magnetic field unit vector with the gradient of \( h \)

\[ b \times \nabla h = \frac{1}{b_{yc}} \left[ \left( \frac{\partial h}{\partial z} - b_z \frac{\partial h}{\partial y} \right) e_x + \left( b_y \frac{\partial h}{\partial y} - \frac{\partial h}{\partial x} \right) e_z \right. \]

\[ + \left. \left( b_x \frac{\partial h}{\partial x} - b_y \frac{\partial h}{\partial z} \right) e_y \right]. \]  

(3.52)
Taking the dot product of the last expression with the gradient of the function $f$ yields the third operation

$$\mathbf{b} \times \nabla h \cdot \nabla f = \frac{1}{b_{yc}} \left[ \left( \frac{\partial h}{\partial z} - b_{zc} \frac{\partial h}{\partial x} \right) \frac{\partial f}{\partial x} + \left( b_{x} \frac{\partial h}{\partial y} - b_{y} \frac{\partial h}{\partial x} \right) \frac{\partial f}{\partial z} \right] + \left( b_{zc} \frac{\partial h}{\partial x} - b_{x} \frac{\partial h}{\partial y} \right) \frac{\partial f}{\partial y}.$$  (3.53)

We continue by rewriting Eq. (3.53) in terms of canonical Poisson brackets

$$\{h, f\}_{x,y} = \frac{\partial h}{\partial x} \frac{\partial f}{\partial y} - \frac{\partial h}{\partial y} \frac{\partial f}{\partial x}. \quad (3.54)$$

This rewriting is used in the discretisation, presented in Chapter 4, where we employ the Arakawa scheme for discretising Poisson brackets [121]

$$\mathbf{b} \times \nabla h \cdot \nabla f = \frac{b_{x} b_{yc}}{b_{yc}} \left( \frac{\partial h}{\partial y} \frac{\partial f}{\partial x} - \frac{\partial h}{\partial x} \frac{\partial f}{\partial y} \right) + \frac{b_{z} b_{yc}}{b_{yc}} \left( \frac{\partial h}{\partial x} \frac{\partial f}{\partial z} - \frac{\partial h}{\partial z} \frac{\partial f}{\partial x} \right) + \frac{1}{b_{yc}} \left( \frac{\partial h}{\partial y} \frac{\partial f}{\partial x} - \frac{\partial h}{\partial x} \frac{\partial f}{\partial y} \right) + \frac{1}{b_{yc}} \{h, f\}_{y,z} \cdot (3.55)$$

The fourth operation is the divergence of a scalar function times the orthogonal gradient. The orthogonal gradient of the scalar function $f$ is defined with the orthogonal projection operator $\mathcal{P}_\perp$, presented in Eq. (3.44), as

$$\nabla_\perp f \cdot v := \nabla f \cdot \mathcal{P}_\perp(v) \quad (3.56)$$

and yields in the locally field-aligned coordinate system

$$\nabla_\perp = e^x \left( \frac{\partial}{\partial x} - b_{x} \frac{\partial}{\partial y} \right) + e^z \left( \frac{\partial}{\partial z} - b_{z} \frac{\partial}{\partial y} \right). \quad (3.57)$$

The divergence of a vector field $v$ in general curvilinear coordinate system reads

$$\nabla \cdot v = \frac{1}{J_{ki}} \sum_{i=1}^{3} \sum_{j=1}^{3} \frac{\partial}{\partial x^i} (g^{ij} v_i J_{ki}). \quad (3.58)$$

Combining Eqs. (3.57) and (3.58) yields the fourth operation in locally field-aligned coordinates

$$\nabla \cdot (h \nabla_\perp) = \frac{1}{J_{ki}} \sum_{i=1}^{3} \sum_{j=1}^{3} \frac{\partial}{\partial x^i} (g^{ij} h \nabla_\perp J_{ki}). \quad (3.59)$$

In Eqs. (3.32) and (3.57) the metric tensor and the perpendicular gradient are evaluated on the poloidal plane at $y = 0$. As Eq. (3.59) contains $y$ derivatives of the metric tensor and the perpendicular gradient, the full expressions of both are needed. As a result of the
y derivatives, the field equations on different poloidal planes are coupled and need to be solved at once. To save computational resources, we decouple the field equations on different poloidal planes by assuming that

$$ \nabla \cdot (h \nabla \perp f) \approx \left( \frac{1}{J_{\text{cyl}}} \frac{\partial}{\partial x} \left( J_{\text{cyl}} h \frac{\partial f}{\partial x} \right) + \frac{\partial}{\partial z} \left( h \frac{\partial f}{\partial z} \right) \right). $$

(3.60)

In simulations of linear fusion devices or with a large aspect ratio assumption, the toroidal angle $\varphi$ is interpreted as a Cartesian coordinate labelling poloidal planes and the Jacobian $J_{\text{cyl}}$ is given by $J_{\text{cyl}} = 1$. In toroidal geometries it reads $J_{\text{cyl}} = x$. This approximation can be performed on the field part of the Lagrangian by replacing the orthogonal gradient with the gradient in cylindrical coordinates

$$ \nabla \perp \approx \nabla_{R,Z} := e_R \frac{\partial}{\partial R} + e_Z \frac{\partial}{\partial Z}. $$

(3.61)

By performing the approximation on the Lagrangian, the gyrokinetic Vlasov-Maxwell systems remains consistent. In summary, Eqs. (3.46), (3.49), (3.53) and (3.59) provide an explicit form of the operations in the gyrokinetic Vlasov-Maxwell system in the locally field aligned coordinate system. In GENE-X we replace Eq. (3.59) with Eq. (3.60).

### 3.2.4. Explicit form of the Vlasov-Maxwell system

The gyrokinetic Vlasov equation given by Eqs. (2.61), (2.77), (2.78) and (2.80) reads in the new coordinates

$$ \frac{\partial f}{\partial t} + v_{\parallel} \frac{B^*}{B_{\parallel}} \cdot \nabla f_{\sigma} + \frac{c}{q_{\sigma} B_{\parallel}} b \times (\mu \nabla B + q_{\sigma} \nabla \phi_1) \cdot \nabla f_{\sigma} - \frac{B^*}{m_{\sigma} B_{\parallel}} \cdot (\mu \nabla B + q_{\sigma} \nabla \phi_1) \frac{\partial f_{\sigma}}{\partial v_{\parallel}} $$

$$ - \frac{q_{\sigma}}{m_{\sigma} c} \frac{\partial A_{\parallel}}{\partial t} \frac{\partial f_{\sigma}}{\partial v_{\parallel}} = 0 $$

(3.62)

with the components

\begin{align*}
(a) &= v_{\parallel} \frac{B}{B_{\parallel}} \frac{\partial f_{\sigma}}{\partial y} + m_{\sigma} v_y^2 \frac{c}{q_{\sigma} B_{\parallel}} \frac{1}{b_y} \left( \frac{\partial b_{y_1}}{\partial y} \frac{\partial f_{\sigma}}{\partial y} - \frac{\partial b_{y_1}}{\partial y} \frac{\partial f_{\sigma}}{\partial x} + \frac{b_{x_1}}{b_y} \frac{\partial b_{x_1}}{\partial y} \frac{\partial f_{\sigma}}{\partial z} \right) \\
&+ \frac{v_{\parallel}}{B_{\parallel}} \left( \frac{\partial A_{\parallel}}{\partial y} \frac{\partial f_{\sigma}}{\partial y} - \frac{\partial A_{\parallel}}{\partial y} \frac{\partial f_{\sigma}}{\partial z} + b_{x_1} \left( \frac{\partial A_{\parallel}}{\partial y} \frac{\partial f_{\sigma}}{\partial x} - \frac{\partial A_{\parallel}}{\partial y} \frac{\partial f_{\sigma}}{\partial y} \right) \right) \\
&+ \frac{1}{b_y} \left( \frac{\partial A_{\parallel}}{\partial x} \frac{\partial f_{\sigma}}{\partial z} - \frac{\partial A_{\parallel}}{\partial z} \frac{\partial f_{\sigma}}{\partial x} \right). \quad (3.63)
\end{align*}
Ampère's law reads
\[ \nabla \times \mathbf{B} = \mu_0 \mathbf{j}. \]

In this subsection, we show how to perform parallel derivatives in practice, it is necessary to trace the position of magnetic field lines from one poloidal plane to another. In this subsection, we show how to perform

\[ (b) = \frac{c}{q_0 B_z} \left[ \mu \left( \frac{b_{xz}}{b_{yc}} \left( \frac{\partial B}{\partial y} \right) \frac{\partial f_\sigma}{\partial x} - \frac{\partial B}{\partial z} \frac{\partial f_\sigma}{\partial y} \right) + \frac{b_{xz}}{b_{yc}} \left( \frac{\partial B}{\partial y} \right) \frac{\partial f_\sigma}{\partial x} - \frac{\partial B}{\partial z} \frac{\partial f_\sigma}{\partial y} + \frac{1}{b_{yc}} \left( \frac{\partial \phi_1 f_\sigma}{\partial y} - \frac{\partial \phi_1 f_\sigma}{\partial z} \right) \right] \]

and

\[ (c) = \frac{1}{m_\sigma B_z^2} \left[ \mu \left( \frac{b_{xyc}}{b_{yc}} \left( \frac{\partial A_{\parallel}}{\partial y} \right) - \frac{\partial A_{\parallel}}{\partial z} \right) + \frac{b_{xyc}}{b_{yc}} \left( \frac{\partial A_{\parallel}}{\partial y} \right) - \frac{\partial A_{\parallel}}{\partial z} \right) \]

and

\[ \frac{q_0}{\mu_0 C c} \left( \frac{\partial A_{\parallel}}{\partial y} \right) - \frac{\partial A_{\parallel}}{\partial z} \right) \]

\[ \frac{q_0}{\mu_0 C c} \left( \frac{\partial A_{\parallel}}{\partial y} \right) - \frac{\partial A_{\parallel}}{\partial z} \right) \]

and

\[ \frac{q_0}{\mu_0 C c} \left( \frac{\partial A_{\parallel}}{\partial y} \right) - \frac{\partial A_{\parallel}}{\partial z} \right) \]

In the new coordinate system, employing Eq. (3.60), the quasi-neutrality equations reads

\[ - \left( \frac{1}{j_{cly}} \frac{\partial}{\partial x} \left( j_{cly} \sum_\sigma m_\sigma c^2 n_{0\sigma} \frac{\partial}{\partial x} \right) + \frac{\partial}{\partial z} \left( \sum_\sigma m_\sigma c^2 n_{0\sigma} \frac{\partial}{\partial z} \right) \right) \phi_1 = \sum_\sigma q_\sigma \int f_\sigma dW, \]

\[ \text{(3.66)} \]

Ampère's law reads

\[ - \left( \frac{1}{j_{cly}} \frac{\partial}{\partial x} \left( j_{cly} \frac{\partial}{\partial x} \right) + \frac{\partial^2}{\partial z^2} \right) A_{\parallel} = 4\pi \sum_\sigma q_\sigma \int f_\sigma v_{\parallel} dW \]

\[ \text{(3.67)} \]

and the generalised Ohm's law reads

\[ - \left( \frac{1}{j_{cly}} \frac{\partial}{\partial x} \left( j_{cly} \frac{\partial}{\partial x} \right) + \frac{\partial^2}{\partial z^2} \right) f_\sigma \left( \frac{\partial f_\sigma}{\partial t} \right) v_{\parallel} dW \]

\[ \text{(3.68)} \]

3.2.5. Field line tracing

To calculate parallel y derivatives in practice, it is necessary to trace the position of magnetic field lines from one poloidal plane to another. In this subsection, we show how to perform
the field line tracing in the cylindrical coordinate system. We call the curve of the field line expressed in cylindrical coordinates
\[ \gamma_{\text{cyl}} = (\gamma_R, \gamma_\phi, \gamma_Z) . \]
In cylindrical coordinates, Eq. (3.21) reads
\[
\begin{align*}
\frac{d\gamma_R}{d\lambda} &= b_R \circ \gamma_{\text{cyl}} \quad (3.69) \\
\frac{d\gamma_\phi}{d\lambda} &= b_\phi \circ \gamma_{\text{cyl}} \quad (3.70) \\
\frac{d\gamma_Z}{d\lambda} &= b_Z \circ \gamma_{\text{cyl}} . \quad (3.71)
\end{align*}
\]
The length of the curve \( \gamma_{\text{cyl}} \) starting at the point \( \gamma_{\text{cyl}}(0) = R_0 = (R_0, \phi_0, Z_0) \) is given by
\[
l : R_0 \mapsto \int_0^a |b(\gamma_{\text{cyl}}(\lambda))|d\lambda = a , \quad (3.72)
\]
which demonstrates that the orbit parameter \( \lambda \) of the curve labels the length of the field line.
To follow the position of a magnetic field line, starting at \( R_0 \), around the fusion device, we parametrise the curve of the magnetic field line with its poloidal angle instead of its length.
We define the function that assigns the poloidal angle of the field line to its length as
\[
s : \phi \mapsto \int_0^a |b(\gamma_{\text{cyl}}(\lambda))|d\lambda = a = \gamma_{\phi}^{-1}(\phi) , \quad (3.73)
\]
where \( a \) is chosen such that \( \gamma_{\phi}(a) = \phi \). The function \( s \) can be simplified
\[
s(\phi) = \int_0^a |b(\gamma_{\text{cyl}}(\lambda))|d\lambda \\
= \int_0^{\gamma_{\phi}^{-1}(\phi)} 1 d\lambda \\
= \int_{\phi_0}^{\phi} \frac{d\gamma_{\phi}^{-1}(u)}{du} du \\
= \int_{\phi_0}^{\phi} \frac{1}{b_\phi(\gamma_{\text{cyl}}(\gamma_{\phi}^{-1}(u)))} du , \quad (3.74)
\]
where we have used Eq. (3.26). By taking the derivative of Eq. (3.74), we obtain an expression for the derivative of the field line length
\[
\left( \frac{ds}{d\phi} \right)(\phi) = \frac{1}{b_\phi(\gamma_{\text{cyl}}(\gamma_{\phi}^{-1}(\phi)))} . \quad (3.75)
\]
Similar to the function \( s(\phi) \), parametrising the field line length by the toroidal angle \( \phi \), it is possible to derive functions \( Z(\phi) \) and \( R(\phi) \). They parametrise the \( R \) and \( Z \) coordinates the
magnetic field line visits by the poloidal angle \( \varphi \). Their derivatives read

\[
\begin{align*}
\left( \frac{dR}{d\varphi} \right) (\varphi) &= \frac{b_R(\gamma_{\text{cyl}}(\gamma_{\varphi}^{-1}(\varphi)))}{b_{\varphi}(\gamma_{\text{cyl}}(\gamma_{\varphi}^{-1}(\varphi)))} \\
\left( \frac{dZ}{d\varphi} \right) (\varphi) &= \frac{b_Z(\gamma_{\text{cyl}}(\gamma_{\varphi}^{-1}(\varphi)))}{b_{\varphi}(\gamma_{\text{cyl}}(\gamma_{\varphi}^{-1}(\varphi)))} \\
\left( \frac{d\varphi}{d\varphi} \right) (\varphi) &= 1.
\end{align*}
\]  

(3.76) (3.77) (3.78)

Eqs. (3.75), (3.76), (3.77) and (3.78) provide the position of the magnetic field line and its length in cylindrical coordinates given a starting position of the field line \( R_0 \) and a poloidal angle \( \varphi \) travelled. We solve the system numerically with an explicit Runge-Kutta method of order eight called **DOP853** [122, II. 5]. The integration is implemented in the library **PARALLAX**. For more details on **PARALLAX** we refer to Chapter 5.

### 3.3. Conclusion and outlook

When performing turbulence simulations of the edge and SOL, it is important to use a coordinate system that (i) is aligned to the magnetic field and (ii) is well-defined for realistic magnetic geometries, including the separatrix and multiple X-points. Flux-coordinate systems, which have been successfully used in fusion theory for decades, suffer from a coordinate singularity at the separatrix that renders them inapplicable for predictive edge and SOL simulations.

In this chapter, we presented a locally field-aligned coordinate system that is based on the FCI approach. This coordinate system is not restricted by the magnetic geometry and allows for the simulation in geometries with single and even multiple X-points. Furthermore, the coordinate system is still aligned to the magnetic field such that derivatives along the magnetic field line are represented as a derivative in the coordinate \( y \). This respects the flute-like structure of plasma turbulence and allows to reduce the toroidal resolution needed for simulations.

As the coordinate system is not limited by the magnetic geometry, it would be interesting to generalise the code to 3D geometries in the future. This would enable the exploration of turbulence in the SOL of stellarator fusion experiments as well as stochastic regions and magnetic islands.

---

†In the code, the toroidal component of the magnetic field is given with respect to the normalised basis vector \( e_{\varphi c} = e_\varphi / |e_\varphi| = e_\varphi / J_{\text{cyl}} \). Therefore, the right-hand side of Eqs. (3.75), (3.76) and (3.77) is multiplied with an additional factor of \( J_{\text{cyl}} \) and \( b_{\varphi} \) is replaced with \( b_{\varphi c} \).
4. Numerical scheme

For a stable, long-time integration of the gyrokinetic Vlasov-Maxwell system it is important to select a numerical scheme that reflects the properties of the equations. In this chapter, we present the numerical scheme that is used in the GENE-X code. We start with the derivation of a dimensionless form of the gyrokinetic Vlasov-Maxwell system, appropriate for numerical discretisation. Next, we discuss which discretisation strategies are suited for the locally field aligned coordinate system, presented in the last chapter, and detail the discretisation scheme implemented in the GENE-X code. Finally, we discuss numerical diffusion and boundary as well as initial conditions for the simulations.

4.1. Normalisation

For the numerical implementation it is useful to renormalise the equations. With the normalisation different physical quantities can be scaled to numerical values of order $O(1)$. This is beneficial because floating point numbers are most dense around 1 and therefore smaller differences between numbers can be resolved. Furthermore, it is possible to choose different normalisations for different species. This is for example useful for the representation of the distribution function in velocity space. Due to their lower weight, electrons move faster than ions. By adjusting the normalisation of the velocity coordinate of the electron species, it is possible to use the same velocity space grid for ions and electrons.

The following paragraph has been published in [87]. As the gyrokinetic model has been generalised to include electromagnetic effects, the normalisation of the parallel component of the electromagnetic vector potential has been added.

We use the normalisation from the GENE code [45]. We denote reference quantities with a ref subscript and renormalised quantities with a hat, i.e. for the mass $m_\sigma = \hat{m}_\sigma m_{\text{ref}}$. The reference values, defining the scales of the simulation, are given in Tab. 4.1. With the reference quantities from Tab. 4.1, we normalise the gyrocentre coordinates as

$$x = \hat{x} L_{\text{ref}}, \quad z = \hat{z} L_{\text{ref}}, \quad \varphi = \hat{\varphi},$$

where $T_{0\sigma}(\psi_0) = T_{\text{ref}}(\hat{T}_{0\sigma}(\psi_0))$ denotes the initial plasma temperature at a given flux surface $\psi_0$ and $v_{T_\sigma}(\psi_0) = \sqrt{2 T_{0\sigma}(\psi_0)/m_\sigma} = c_{\text{ref}} \hat{v}_{T_\sigma}(\psi_0)$ the corresponding thermal velocity. Similarly, $n_{0\sigma}(\psi_0) = n_{\text{ref}} \hat{n}_{0\sigma}(\psi_0)$ denotes the initial plasma density. Finally, we normalise the distribution function, the electrostatic potential and the parallel component of the electromagnetic

$$v|| = \hat{v}_{||} v_{T_\sigma}(\psi_0), \quad \mu = \hat{\mu} \frac{T_{0\sigma}(\psi_0)}{B_{\text{ref}}}, \quad t = \frac{L_{\text{ref}}}{c_{\text{ref}}},$$

where $T_{0\sigma}(\psi_0) = T_{\text{ref}}(\hat{T}_{0\sigma}(\psi_0))$ denotes the initial plasma temperature at a given flux surface $\psi_0$ and $v_{T_\sigma}(\psi_0) = \sqrt{2 T_{0\sigma}(\psi_0)/m_\sigma} = c_{\text{ref}} \hat{v}_{T_\sigma}(\psi_0)$ the corresponding thermal velocity. Similarly, $n_{0\sigma}(\psi_0) = n_{\text{ref}} \hat{n}_{0\sigma}(\psi_0)$ denotes the initial plasma density. Finally, we normalise the distribution function, the electrostatic potential and the parallel component of the electromagnetic
Elementary charge \( e \)  
\( \) s.t. \( q_\sigma = \hat{q}_\sigma e \)  
Reference mass \( m_{\text{ref}} \)  
\( \) s.t. \( m_\sigma = \bar{m}_\sigma m_{\text{ref}} \)  
Reference temperature \( T_{\text{ref}} \)  
Reference length \( L_{\text{ref}} \)  
Reference magnetic field strength \( B_{\text{ref}} \)  
Reference density \( n_{\text{ref}} \)  
Reference velocity \( c_{\text{ref}} \)  
Reference gyrofrequency \( \Omega_{\text{ref}} \)  
Reference gyroradius \( \rho_{\text{ref}} \)  
Reference \( \beta \) \( \beta_{\text{ref}} = 8\pi n_{\text{ref}} T_{\text{ref}}/B_{\text{ref}}^2 \)  

Table 4.1.: Elementary reference values for mass, temperature, length, magnetic field strength and density and derived reference values for velocity, gyrofrequency, gyroradius and thermal to magnetic pressure ratio. This table has been published in [87]. To account for the electromagnetic model, the reference \( \beta \) has been added.

vector potential according to

\[
\begin{align*}
  f_\sigma &= \frac{n_0(\psi_0)}{v_{T_\sigma}(\psi_0)^3} \hat{f}_\sigma \\
  \phi_1 &= \frac{T_{\text{ref}}}{e} \hat{\phi}_1 \\
  A_1\parallel &= \rho_{\text{ref}} B_{\text{ref}} \hat{A}_{1\parallel}.
\end{align*}
\]

4.1.1. Normalisation of the gyrokinetic Vlasov-Maxwell system

We normalise the gyrokinetic Vlasov-Maxwell system in parts. We start with the normalisation of \( B^* \)

\[
B^* = B + \frac{m_\sigma v_\parallel c}{q_\sigma} \nabla \times b + \nabla A_1\parallel \times b
\]

\[
= B_{\text{ref}} \left( \hat{B} + \sqrt{2m_\sigma T_{\text{ref}}(\psi_0)} \frac{\hat{v}_\parallel}{q_\sigma} L_{\text{ref}} \hat{\nabla} \times b + \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \hat{\nabla} \hat{A}_{1\parallel} \times b \right)
\]

\[
= B_{\text{ref}} \hat{B}^*.
\]

As a corollary of Eq. (4.5), the parallel component of \( B^* \) is normalised in the same way, i.e. \( B^*\parallel = B_{\text{ref}} \hat{B}^*\parallel \). The partial derivatives of the distribution function \( f_\sigma \) are normalised as

\[
\begin{align*}
  \frac{\partial f_\sigma}{\partial t} &= \frac{n_0(\psi_0)}{v_{T_\sigma}(\psi_0)} \frac{c_{\text{ref}}}{L_{\text{ref}}} \frac{\partial \hat{f}_\sigma}{\partial t} \\
  \nabla f_\sigma &= \frac{n_0(\psi_0)}{v_{T_\sigma}(\psi_0)} \frac{1}{L_{\text{ref}}} \hat{\nabla} \hat{f}_\sigma \\
  \frac{\partial f_\sigma}{\partial v_\parallel} &= \frac{n_0(\psi_0)}{v_{T_\sigma}(\psi_0)} \frac{1}{v_{T_\sigma}(\psi_0)} \frac{\partial \hat{f}_\sigma}{\partial \hat{v}_\parallel}.
\end{align*}
\]
We proceed by normalising the equations of motion of the gyrocentres. The change of the gyrocentre position reads

\[
\dot{R} = \frac{v_\parallel B^*}{B^\parallel} + \frac{c b}{q_\sigma B^\parallel} \times (\mu \nabla B + q_\sigma \nabla \phi_1)
\]

\[
= c_{\text{ref}} \left( \sqrt{\frac{2T_{0\sigma}(\psi_0)}{m_\sigma}} \dot{\psi} + \frac{\rho_{\text{ref}} b}{L_{\text{ref}} q_\sigma B^*} \times \left( \hat{T}_{0\sigma}(\psi_0) \hat{\mu} \nabla \hat{B} + \hat{q}_\sigma \nabla \hat{\phi}_1 \right) \right),
\tag{4.9}
\]

and the change of the parallel velocity reads

\[
\dot{v}_\parallel = -\frac{B^*}{m_\sigma B^\parallel} \cdot (\mu \nabla B + q_\sigma \nabla \phi_1) - \frac{q_\sigma}{m_\sigma c} \frac{\partial A_1}{\partial t}
\]

\[
= -\frac{c_{\text{ref}}^2 B^*}{L_{\text{ref}} m_\sigma B^\parallel} \cdot \left( \hat{T}_{0\sigma}(\psi_0) \hat{\mu} \nabla \hat{B} + \hat{q}_\sigma \nabla \hat{\phi}_1 \right) - \frac{c_{\text{ref}}^2 q_\sigma}{L_{\text{ref}} m_\sigma} \frac{\partial \hat{A}_1}{\partial t}.
\tag{4.10}
\]

By collecting the normalised parts and dividing by the factor \(c_{\text{ref}}/L_{\text{ref}} n_{0\sigma}(\psi_0)/\sqrt{\psi_0}(\psi_0)\) we obtain the normalised Vlasov equation

\[
\frac{\partial \hat{f}_\sigma}{\partial t} + \sqrt{\frac{2T_{0\sigma}(\psi_0)}{m_\sigma}} \frac{\hat{B}^*}{B^\parallel} \cdot \nabla \hat{f}_\sigma + \frac{\rho_{\text{ref}} b}{L_{\text{ref}} q_\sigma B^*} \times \left( \hat{T}_{0\sigma}(\psi_0) \hat{\mu} \nabla \hat{B} + \hat{q}_\sigma \nabla \hat{\phi}_1 \right) \nabla \hat{f}_\sigma
\]

\[
= \frac{\hat{B}^*}{\sqrt{2m_\sigma T_{0\sigma}(\psi_0)B^\parallel}} \cdot \left( \hat{T}_{0\sigma}(\psi_0) \hat{\mu} \nabla \hat{B} + \hat{q}_\sigma \nabla \hat{\phi}_1 \right) \frac{\partial \hat{f}_\sigma}{\partial \phi_\parallel} - \frac{\hat{q}_\sigma}{\sqrt{2m_\sigma T_{0\sigma}(\psi_0)}} \frac{\partial \hat{A}_1}{\partial \phi_\parallel} \frac{\partial \hat{f}_\sigma}{\partial \phi_\parallel}
\]

\[
= 0.
\tag{4.11}
\]

We continue with the normalisation of the quasi-neutrality equation, given in Eq. (2.84). We start with the right-hand side

\[
\sum_\sigma q_\sigma \int f_\sigma \, dW = \sum_\sigma q_\sigma \int_0^\infty \int_{-\infty}^\infty 2\pi \frac{B^\parallel}{m_\sigma} f_\parallel \, dv_\parallel \, d\mu
\]

\[
= en_{\text{ref}} \sum_\sigma \hat{n}_{0\sigma}(\psi_0) \hat{q}_\sigma \int_0^\infty \int_{-\infty}^\infty \pi \hat{B}^\parallel \hat{f}_\sigma \, dv_\parallel \, d\hat{\mu}.
\tag{4.13}
\]

The left-hand side reads

\[
- \nabla \cdot \left( \sum_\sigma \frac{m_\sigma c n_{0\sigma}}{B^2} \nabla_\perp \phi \right) = -en_{\text{ref}} \left( \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right)^2 \nabla \cdot \left( \sum_\sigma \frac{\hat{m}_\sigma \hat{n}_{0\sigma}}{B^2} \nabla_\perp \hat{\phi}_1 \right).
\tag{4.14}
\]

Combining both sides and multiplying by \((n_{\text{ref}}e)^{-1}\) yields the normalised quasi-neutrality equation

\[
- \left( \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right)^2 \nabla \cdot \left( \sum_\sigma \frac{\hat{m}_\sigma \hat{n}_{0\sigma}}{B^2} \nabla_\perp \hat{\phi}_1 \right) = \sum_\sigma \hat{n}_{0\sigma}(\psi_0) \hat{q}_\sigma \int_0^\infty \int_{-\infty}^\infty \pi \hat{B}^\parallel \hat{f}_\sigma \, dv_\parallel \, d\hat{\mu}.
\tag{4.15}
\]
The normalisation of Ampère’s law is similar to the quasi-neutrality equation. We start with the right-hand side

$$4\pi \sum_{\sigma} \frac{q_\sigma}{c} \int f_\sigma v_\parallel dW = 4\pi \sum_{\sigma} \frac{q_\sigma}{c} \int_0^\infty \int_{-\infty}^\infty \frac{B^*_\parallel}{m_\sigma} f_\sigma v_\parallel d\nu_\parallel d\mu
$$

$$= \frac{B_{ref}}{\rho_{ref}} \frac{\beta_{ref}}{2} \sum_{\sigma} \hat{n}_{0\sigma}(\psi_0) \hat{q}_\sigma \sqrt{2T_{0\sigma}(\psi_0)}/m_\sigma \int_0^\infty \int_{-\infty}^\infty \pi B^*_\parallel \hat{f}_\sigma \nu_\parallel d\nu_\parallel d\mu.$$

The left-hand side reads

$$- \Delta_\perp A_1|| = - \frac{\rho_{ref} B_{ref}}{L^2_{ref}} \hat{\Delta}_\perp \hat{A}_1||. \quad (4.16)$$

Combining Eqs. (4.16), (4.17) and dividing by $B_{ref}/\rho_{ref}$ yields the normalised Ampère’s law

$$- \left( \frac{\rho_{ref}}{L_{ref}} \right)^2 \Delta_\perp \hat{A}_1|| = - \frac{\beta_{ref}}{2} \sum_{\sigma} \hat{n}_{0\sigma}(\psi_0) \hat{q}_\sigma \sqrt{2T_{0\sigma}(\psi_0)}/m_\sigma \int_0^\infty \int_{-\infty}^\infty \pi B^*_\parallel \hat{f}_\sigma \nu_\parallel d\nu_\parallel d\mu.$$

The normalised form of the generalised Ohm’s law can either be derived explicitly or by taking the normalised time derivative of Eq. (4.18). It reads

$$- \left( \frac{\rho_{ref}}{L_{ref}} \right)^2 \Delta_\perp \hat{A}_1|| = \frac{\beta_{ref}}{2} \sum_{\sigma} \hat{n}_{0\sigma}(\psi_0) \hat{q}_\sigma \sqrt{2T_{0\sigma}(\psi_0)}/m_\sigma \int_0^\infty \int_{-\infty}^\infty \pi B^*_\parallel \hat{f}_\sigma \nu_\parallel d\nu_\parallel d\mu
$$

$$= \frac{\beta_{ref}}{2} \sum_{\sigma} \hat{n}_{0\sigma}(\psi_0) \hat{q}_\sigma \sqrt{2T_{0\sigma}(\psi_0)}/m_\sigma \int_0^\infty \int_{-\infty}^\infty \pi B^*_\parallel \left( \frac{\partial \hat{f}_\sigma}{\partial t} \right)^* \hat{\nu}_\parallel d\nu_\parallel d\mu. \quad (4.19)$$

The explicit form of the normalised gyrokinetic Vlasov-Maxwell system, expressed in the locally field-aligned coordinate system, is given in Appendix A.2.

### 4.1.2. Normalisation of the local and canonical Maxwellian

In order to initialise the simulation or set radial boundary conditions, it is useful to have a normalised Maxwellian distribution function at our disposal. We denote the Maxwellian distribution function with $F_{0\sigma}$ and normalise it in the same way as $f_\sigma$. It reads as a function of the flux surface label $\psi$, $\hat{\nu}_\parallel$ and $\hat{\mu}$

$$\hat{F}_{0\sigma}(\psi, \hat{\nu}_\parallel, \hat{\mu}) = \frac{\hat{n}_{0\sigma}(\psi)}{(\pi \hat{T}_{0\sigma}(\psi))^{3/2}} e^{-\frac{\hat{\nu}_\parallel^2 + \hat{\mu}^2}{\hat{T}_{0\sigma}(\psi)}}, \quad (4.20)$$

where the normalised temperature and density profiles $\hat{T}_{0\sigma}(\psi) = T_{0\sigma}(\psi)/T_{0\sigma}(\psi_0)$ and $\hat{n}_{0\sigma}(\psi) = n_{0\sigma}(\psi)/n_{0\sigma}(\psi_0)$ were used. Next to the local Maxwellian, we make use of a so-called canonical Maxwellian. The canonical Maxwellian is defined in terms of constants of motion and will be discussed in detail in Section 4.5. It uses the canonical toroidal momentum $p_{\phi}$ as a replacement for the flux surface label $\psi$. The new modified radial coordinate $\psi_C$ is defined
\[
\psi_C = \frac{c}{q_\sigma} p_\varphi \\
= \psi_\rho + \frac{c}{q_\sigma} m_\sigma v_\parallel b_\varphi R,
\]

and normalised as

\[
\psi_C = \psi_\rho + \rho_{\text{ref}} \sqrt{2 m_\sigma T_0 (\psi_0)} \dot{q}_\sigma \dot{v}_\parallel b_\varphi \hat{R}.
\]

### 4.2. Discretisation

We use the method of lines to construct the discretisation scheme. The gyrokinetic Vlasov-Maxwell system can be written in the form

\[
\frac{\partial f}{\partial t} = L[f],
\]

where \(L\) is a nonlinear, integro-partial differential operator acting on \(f\)\(^1\). In the method of lines the operator \(L\) acting on phase space is discretised first. This yields a system of ordinary differential equations

\[
\frac{d(f_i)}{dt} = L_D[(f_i)],
\]

where \((f_i)\) corresponds to a vector containing the values of the distribution function at the phase space grid points \(i\), and \(L_D\) denotes the discretised differential operator. The initial value problem in Eq. (4.24) can then be solved with a method of choice. In the following, we discuss different spatial and temporal discretisation strategies.

#### 4.2.1. Spatial discretisation

There are multiple different strategies to construct a spatial discretisation of an integro-partial differential operator. Most schemes fall in the class of finite difference, finite volume or finite element schemes. There are also schemes in between these classes. A good example are schemes developed in the area of geometric discretisations and mimetic finite differences \([123, 124, 125]\). Therein, differential operators are discretised based on the preservation of mathematical properties of the continuous system. Further, spectral methods based on Fourier decomposition in real space \([68, 44]\) or an expansion in orthogonal Hermite and Laguerre polynomials in velocity space \([126, 127, 128]\) are often used to exploit symmetries of magnetically confined fusion plasmas.

**Spatial discretisation of the gyrokinetic Vlasov equation**

We discuss different spatial discretisation schemes in the language of differential forms to explicitly clarify the structure and properties of the quantities and operators involved. As

\(^1\)In this form the function \(f\) is a multi-dimensional function with a component for each species and the nonlinear, integro-partial differential operator \(L\) represents the whole gyrokinetic Vlasov-Maxwell system including the quasi-neutrality equation, Ampère’s law and the generalised Ohm’s law.
Figure 4.1.: Figure displaying the overlap of flux boxes originating from two poloidal planes at $\varphi = 0$ and $\varphi = \pi/2$. In a conservative finite volume or finite element discretisation, fluxes are exchanged between different boxes. In the locally field-aligned coordinate system, the flux boxes overlap in a non-trivial way demonstrated on a staggered poloidal plane at $\varphi = \pi/4$.

discussed in Chapter 2, the gyrokinetic Vlasov equation can be written in advective and conservative form. On the continuous level both formulations are equivalent. Which one is chosen makes a difference for the discretisation. Let $\Lambda^n(\mathcal{P})$ be the space of the differential $n$-forms on the phase space $\mathcal{P}$. In the conservative form of the Vlasov equation the distribution function corresponds to a six-form on the manifold, $f^{(6)} \in \Lambda^6(\mathcal{P})$, that can be integrated to obtain the particle number in a given volume. The Vlasov equation can be written as

$$\frac{\partial f^{(6)}}{\partial t} = -d\zeta^{(5)}$$

(4.25)

$$\zeta^{(5)} = \iota_u f^{(6)},$$

(4.26)

where the five-form $\zeta^{(5)}$ corresponds to the five-dimensional flux out of a six-dimensional unit volume and $\iota_u$ denotes the interior product. The vector field $u$ represents the drift velocities and is itself a nonlinear function of $f$. This form of the Vlasov equation is well suited for finite volume or finite element discretisation schemes because the distribution function corresponds to volume averages and $\zeta^{(5)}$ to fluxes. In the locally field-aligned coordinate system it is difficult to correctly calculate fluxes because flux boxes from different poloidal planes do not align. They overlap in a non-trivial way and sophisticated book keeping would be needed to correctly handle the fluxes at the intersection points and ensure conservation. Exemplarily, the overlap of flux boxes is demonstrated in Fig. 4.1.

In the advective form the distribution function corresponds to a scalar function or zero form
Figure 4.2.: Figure displaying the grid in the $x$ and $z$ coordinates at the example of AUG, used for simulations in Chapter 7. The grid is unstructured, such that the boundary of fusion devices can be modelled, and Cartesian in the neighbourhood of every point. For a clear presentation only every 20th grid point is shown.

\[ f^{(0)} \in \Lambda^0(P). \] The Vlasov equation can be written as

\[
\frac{\partial f^{(0)}}{\partial t} = -i_u \zeta^{(1)} \]

\[ \zeta^{(1)} = df^{(0)}, \]

where $\zeta^{(1)}$ is a one form corresponding naturally to the gradient of $f$. The advective form of the Vlasov equation is well suited for a finite difference discretisation because gradients correspond to differences of scalar function values.

Due to the non-conformal overlap of flux boxes, we discretise the advective form of the Vlasov equation with a finite difference discretisation scheme. As the coordinate system is defined in regions around a family of poloidal planes, we choose a Cartesian grid in the toroidal angle $\varphi$. In the coordinates $x$ and $z$ in the poloidal plane we use an unstructured grid that is Cartesian in the neighbourhood of every point. The grid is constructed such that $\Delta x = \Delta z$. We label each grid point by a one-dimensional index $i$ and store the $x$ and $z$ coordinates in one-dimensional arrays $x[i]$ and $z[i]$. This makes the implementation flexible because one can remove points from and add points to the grid and thus model the complex shape of the fusion device. Requiring that the grid is Cartesian in the neighbourhood of every point allows us to use efficient numerical schemes for Cartesian grids [87]. The grid is visualised at the example of an AUG equilibrium in Fig 4.2. In the velocity space coordinate $v_\parallel$ we use a Cartesian grid. We discretise the first derivatives in $x$, $z$ and $v_\parallel$ direction, apart from the
nonlinear Poisson brackets

\[
\{ \dot{A}_{1||}, \dot{f}_\sigma \}_{z,x} = \frac{\partial \dot{A}_{1||}}{\partial z} \frac{\partial \dot{f}_\sigma}{\partial x} - \frac{\partial \dot{A}_{1||}}{\partial x} \frac{\partial \dot{f}_\sigma}{\partial z},
\]

(4.29)

\[
\{ \phi_1, \dot{f}_\sigma \}_{z,x} = \frac{\partial \phi_1}{\partial z} \frac{\partial \dot{f}_\sigma}{\partial x} - \frac{\partial \phi_1}{\partial x} \frac{\partial \dot{f}_\sigma}{\partial z},
\]

(4.30)

with a fourth order centred finite difference scheme

\[
\left( \frac{\partial \dot{f}_\sigma}{\partial x} \right)(\hat{x}_i) = \frac{1}{12 \Delta x} \left( \dot{f}_\sigma(\hat{x}_{i-2}) - 8 \dot{f}_\sigma(\hat{x}_{i-1}) + 8 \dot{f}_\sigma(\hat{x}_{i+1}) - \dot{f}_\sigma(\hat{x}_{i+2}) \right) + O((\Delta x)^4).
\]

(4.31)

For the terms in Eqs. (4.29) and (4.30), we use the second order scheme introduced by Arakawa [121]

\[
\{ \hat{\phi}_1, \dot{f}_\sigma \}_{z,x}(\hat{x}_i, \hat{z}_j) =
\]

\[
-\frac{1}{12(\Delta x)^2} \left[ \left( \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j-1}) + \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j+1}) - \dot{f}_\sigma(\hat{x}_i, \hat{z}_{j+1}) - \dot{f}_\sigma(\hat{x}_i, \hat{z}_{j-1}) \right) \right. \]

\[
\left. \left( \hat{\phi}_1(\hat{x}_{i+1}, \hat{z}_j) - \hat{\phi}_1(\hat{x}_i, \hat{z}_j) \right) + \left( \dot{f}_\sigma(\hat{x}_{i-1}, \hat{z}_{j-1}) + \dot{f}_\sigma(\hat{x}_{i-1}, \hat{z}_{j+1}) - \dot{f}_\sigma(\hat{x}_i, \hat{z}_{j+1}) - \dot{f}_\sigma(\hat{x}_i, \hat{z}_{j-1}) \right) \right]
\]

\[
\left( \hat{\phi}_1(\hat{x}_i, \hat{z}_{j-1}) - \hat{\phi}_1(\hat{x}_i, \hat{z}_{j+1}) \right) + \left( \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j-1}) + \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j+1}) - \dot{f}_\sigma(\hat{x}_{i-1}, \hat{z}_{j-1}) - \dot{f}_\sigma(\hat{x}_{i-1}, \hat{z}_{j+1}) \right)
\]

\[
\left( \hat{\phi}_1(\hat{x}_{i-1}, \hat{z}_{j-1}) - \hat{\phi}_1(\hat{x}_{i-1}, \hat{z}_{j+1}) \right) + \left( \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j-1}) - \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j+1}) \right) \left( \hat{\phi}_1(\hat{x}_{i+1}, \hat{z}_{j+1}) - \hat{\phi}_1(\hat{x}_{i+1}, \hat{z}_{j-1}) \right)
\]

\[
+ \left( \hat{\phi}_1(\hat{x}_{i-1}, \hat{z}_{j-1}) - \hat{\phi}_1(\hat{x}_{i-1}, \hat{z}_{j+1}) \right) \left( \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j-1}) - \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j+1}) \right)
\]

\[
\left( \hat{\phi}_1(\hat{x}_{i+1}, \hat{z}_{j-1}) - \hat{\phi}_1(\hat{x}_{i+1}, \hat{z}_{j+1}) \right) \left( \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j-1}) - \dot{f}_\sigma(\hat{x}_{i+1}, \hat{z}_{j+1}) \right)
\]

\[
+ O((\Delta x)^2).
\]

(4.32)

The \( y \) direction follows magnetic field lines originating on a poloidal plane. We define the grid in the \( y \) direction by the intersection points of magnetic field lines, originating from the \( x \), \( z \) grid points, with neighbouring poloidal planes. The intersection points in general do not fall on a grid point. We use bicubic interpolation to approximate the value of the distribution function on the intersection points. The intersection of the magnetic field line with neighbouring poloidal planes and the bicubic interpolation is visualised in Fig. 4.3. The intersection points are calculated during the initialisation step of GENE-X by solving Eqs. (3.75), (3.76), (3.77) and (3.78). As the length of the magnetic field line differs between different positions on the poloidal plane, the grid in \( y \) direction is non-uniform. Similar to Eq. (4.31), we discretise the first derivatives in the \( y \) direction with a fourth order centred finite difference scheme. We construct the scheme on the non-uniform \( y \) grid with the help of Lagrange polynomials. Let \((\hat{y}_{-2}, \hat{y}_{-1}, \hat{y}_0, \hat{y}_1, \hat{y}_2)\) describe the grid points in \( y \) direction local to a poloidal plane and \((\dot{\sigma}(\hat{x}_i, \hat{y}_{-2}, \hat{z}_j), \dot{\sigma}(\hat{x}_i, \hat{y}_{-1}, \hat{z}_j), \dot{\sigma}(\hat{x}_i, \hat{y}_0, \hat{z}_j), \dot{\sigma}(\hat{x}_i, \hat{y}_1, \hat{z}_j), \dot{\sigma}(\hat{x}_i, \hat{y}_2, \hat{z}_j))\) the corresponding interpolated values of the distribution function. The polynomial interpo-
Figure 4.3.: Figure displaying a magnetic field line originating at the poloidal plane at $\varphi = 0$ and hitting different poloidal planes at $\varphi \in \{-\pi/2, -\pi/4, \pi/4, \pi/2\}$. The points where the field line intersects a poloidal plane are denoted by red dots. The value of the distribution function is obtained via bicubic interpolation around the intersection point. The area included in the interpolation on every poloidal plane is shaded in blue.

The distribution of the distribution function along the field line is given by

$$\hat{f}_{\sigma, \text{int}}(\hat{x}_i, \hat{y}, \hat{z}_j) = \sum_{k=-2}^{2} \hat{f}_\sigma(\hat{x}_i, \hat{y}_k, \hat{z}_j) \ell_k(\hat{y}), \quad (4.33)$$

where $\ell_k$ denotes the Lagrange polynomial

$$\ell_k(\hat{y}) = \prod_{-2 \leq m \leq 2, m \neq k} \frac{\hat{y} - \hat{y}_m}{\hat{y}_k - \hat{y}_m}. \quad (4.34)$$

The first derivative in $\hat{y}$ direction is discretised as

$$\left( \frac{\partial \hat{f}_\sigma}{\partial \hat{y}} \right)(\hat{x}_i, 0, \hat{z}_j) = \sum_{k=-2}^{2} \hat{f}_\sigma(\hat{x}_i, \hat{y}_k, \hat{z}_j) \left( \frac{d\ell_k}{d\hat{y}} \right)(0) + O\left( (\Delta \hat{y})^4 \right) + O\left( \frac{(\Delta \hat{x})^3}{\Delta \hat{y}} \right) + O\left( \frac{(\Delta \hat{z})^3}{\Delta \hat{y}} \right). \quad (4.35)$$

The derivative of the Lagrange polynomials can be evaluated analytically and stored in an array for every grid point in $\hat{x}$, $\hat{z}$ at initialisation of the code such that the evaluation of Eq. (4.35) is fast during time stepping. The explicit form of the derivatives of the Lagrange polynomials is given in Appendix A.1.
Spatial discretisation of the field equations

The quasi-neutrality equation, Ampère’s law and the generalised Ohm’s law can be written in the general form

\[
\hat{\xi} \left( \frac{1}{\hat{J}_{\text{cyl}}} \frac{\partial}{\partial \hat{x}} \left( \hat{J}_{\text{cyl}} \frac{\partial \hat{\phi}_1}{\partial \hat{x}} \right) + \frac{\partial}{\partial \hat{z}} \left( \hat{c} \frac{\partial \hat{\phi}_1}{\partial \hat{z}} \right) \right) + \hat{\lambda} \hat{\phi}_1 = \hat{b},
\]

with the coefficients \( \xi, c, \lambda \) and \( b \). The latter of the two coefficients consist out of velocity space integrals over the distribution function. The velocity space integrals are discretised as

\[
\int \hat{f}_\sigma \, d\hat{v}_\parallel \, d\hat{\mu} \approx \sum_{l=1}^{N_{\parallel}} w_l \sum_{m=1}^{N_{\mu}} w_m \hat{f}_\sigma(\hat{v}_\parallel l, \hat{\mu}_m),
\]

with the weights \( w_l \) and \( w_m \) for the \( \hat{v}_\parallel \) and \( \hat{\mu} \) integration respectively. For the weights and the \( \mu \) grid different options exist. In the \( \hat{v}_\parallel \) direction the grid is, as discussed above, Cartesian and integration weights corresponding to trapezoidal and Simpson integration are available. The trapezoidal weights read

\[
w_l = \frac{\Delta \hat{v}_\parallel}{2} (1, 2, \ldots, 2, 1)
\]

and the Simpson weights read [129, Section 4.1]

\[
w_l = \frac{\Delta \hat{v}_\parallel}{48} (17, 59, 43, 49, 48, \ldots, 48, 49, 43, 59, 17).
\]

In the \( \hat{\mu} \) direction also two options are available. The first option is a quadratic grid in \( \hat{\mu} \)

\[
\hat{\mu}_m = \left( m - 1 \right) + \frac{1}{2} \frac{L_{\hat{\mu}}}{N_{\hat{\mu}}} ^2,
\]

with trapezoidal weights \( w_m \) according to

\[
w_m = \begin{cases} 
\frac{1}{2} \left( \hat{\mu}_2 - \hat{\mu}_1 \right) & \text{for } m = 1 \\
\frac{1}{2} \left( \hat{\mu}_N - \hat{\mu}_{N-1} \right) & \text{for } m = N_{\hat{\mu}} \\
\frac{1}{2} \left( \hat{\mu}_{m+1} - \hat{\mu}_{m-1} \right) & \text{otherwise}
\end{cases}
\]

The second option are grid points and weights implementing Gaussian quadrature on the interval \([0, \infty)\). As there is no differentiation in \( \hat{\mu} \), the grid can be optimized regarding numerical integration. The grid nodes \( \hat{\mu}_m \) are formed by the roots of the Laguerre polynomials \( L_m \) and the integration weights are given by [130]

\[
w_m = \frac{\hat{\mu}_m e^{\hat{\mu}_m}}{(m+1)^2 L_{m+1}(\hat{\mu}_m)^2}.
\]

The default options are Simpson quadrature in \( \hat{v}_\parallel \) and Gauss-Laguerre quadrature in \( \hat{\mu} \). Equipped with a discrete expression for \( \lambda \) and \( b \), we discretise the left-hand side of Eq. (4.36)
with second order centred finite differences

\[
-\xi \left( \frac{1}{J_{cyl}} \frac{\partial}{\partial \hat{z}} \left( \hat{J}_{cyl} \frac{\partial \hat{\phi}_1}{\partial \hat{z}} \right) + \frac{\partial}{\partial \hat{x}} \left( \hat{c} \frac{\partial \hat{\phi}_1}{\partial \hat{x}} \right) \right) + \lambda \hat{\phi}_1 =
\]

\[
- \frac{\xi (\hat{x}_i, \hat{z}_j)}{J_{cyl,i} \Delta \hat{x}} \left[ \frac{\hat{J}_{cyl,i+1} \hat{c} (\hat{x}_{i+1}, \hat{z}_j) + \hat{J}_{cyl,i} \hat{c} (\hat{x}_i, \hat{z}_j)}{2} \right] \left( \frac{\hat{\phi}_1 (\hat{x}_{i+1}, \hat{z}_j) - \hat{\phi}_1 (\hat{x}_i, \hat{z}_j)}{\Delta \hat{x}} \right)
\]

\[
- \frac{\xi (\hat{x}_i, \hat{z}_j)}{J_{cyl,i} \Delta \hat{z}} \left[ \frac{\hat{J}_{cyl,i} \hat{c} (\hat{x}_i, \hat{z}_{j+1}) + \hat{J}_{cyl,i} \hat{c} (\hat{x}_i, \hat{z}_j)}{2} \right] \left( \frac{\hat{\phi}_1 (\hat{x}_i, \hat{z}_{j+1}) - \hat{\phi}_1 (\hat{x}_i, \hat{z}_j)}{\Delta \hat{z}} \right)
\]

\[
+ \lambda (\hat{x}_i, \hat{z}_i) \hat{\phi}_1 (\hat{x}_i, \hat{z}_i)
\]

\[
+ O \left( (\Delta \hat{x})^2 \right) + O \left( (\Delta \hat{z})^2 \right)
\]

(4.43)

Due to the elliptic operator and the Arakawa scheme, the spatial discretisation is of order two.

### 4.2.2. Temporal discretisation

We solve the remaining initial value problem, presented in Eq. (4.24), with an explicit Runge-Kutta method of order four [122, pp. 132–140] specified by the Butcher tableau

<table>
<thead>
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</table>

(4.44)

According to [131], the fourth order Runge-Kutta method combined with centred finite differences of fourth order, as introduced above, is a stable and efficient scheme to solve multidimensional linear advection equations. The timestep remains constant throughout the simulation. The size of the timestep is typically limited by either the Courant-Friedrichs-Lewy condition of the parallel electron advection or the frequency of kinetic shear Alfvén waves [132, 133]. The timestep restriction due to the parallel electron advection can be estimated as [134, Section II 1.4]

\[
\Delta \hat{t} \lesssim 2.82 \sqrt{\frac{m_e}{2T_0(\psi_0)}} \frac{\Delta \hat{y}_{\min}}{v_{\| \max}}
\]

(4.45)
4.3. Numerical diffusion

The discretisation scheme presented in the last section is almost free of numerical diffusion. This is desirable because it mimics the property that the gyrokinetic Vlasov-Maxwell system is purely advective and free of diffusion on the continuous level. However, numerical diffusion is useful to stabilise the simulations and remove spurious effects caused by the discretisation scheme. As mentioned above, the chosen numerical scheme can integrate multi-dimensional linear advection problems stably and efficiently. Nevertheless, the gyrokinetic Vlasov-Maxwell system solved by the GENE-X code is significantly more complicated than a multi-dimensional linear advection problem and involves bicubic interpolation as well as the evaluation of integrals. Instabilities are typically caused by modes with a high wavenumber such that the scale of their oscillation is close to the spacing of the numerical grid. High wavenumber modes can be damped with the help of a diffusion operator. We explain the effect of the diffusion operator without loss of generality at the example of the parallel $y$ direction. A general diffusion operator of order $n$, in the direction $y$ and with strength $\epsilon$ is given by \[ D_n = (-i)^n \epsilon (\Delta \hat{y})^n \frac{\partial^n}{\partial \hat{y}^n}. \] (4.46)

Applying the diffusion operator $D_n$ to a Fourier mode in $y$ yields

\[ D_n e^{ik\hat{y}} = -\epsilon k^n e^{ik\hat{y}}. \] (4.47)

As can be seen from Eq. (4.47), the diffusion operator reproduces the mode with a negative prefactor resulting in effective damping. The damping strength $\epsilon k^n_y$ is stronger for high $k_y$ and weaker for low $k_y$ modes. Therefore, the high frequency waves are efficiently damped while low frequency waves remain relatively unperturbed. The higher the derivative in the diffusion operator, the better the damping of high frequency and the preservation of low frequency modes. In numerical simulations a compromise between the order of the derivative and the cost of the evaluation must be found. With a centred finite difference scheme the evaluation of the second, fourth, sixth and eighth derivative to second order accuracy requires a stencil of size three, five, seven and nine respectively. As explained in the last section, we use centred finite difference stencils of order four with a stencil size of five to evaluate the first derivative. Similarly, we use a five-point stencil, comprising the fourth derivative evaluated up to second order accuracy, for the numerical diffusion operator.

Unstable, high wavenumber modes appear in all spatial and in the $v_y$ direction. In the $x, z$ direction the main driver of grid scale oscillations is caused by Runge’s phenomenon due to the bicubic interpolation used to calculate parallel derivatives. We use the two-dimensional
The biharmonic operator discretised as \[136\]

\[
\epsilon_{\hat{x}, \hat{z}} \Delta \hat{x}^4 \Delta \hat{z}^4 \hat{f}_\sigma = \epsilon_{\hat{x}, \hat{z}} \left( 
\hat{f}_\sigma (\hat{x}_{i-2}, \hat{z}_j) + \hat{f}_\sigma (\hat{x}_i, \hat{z}_{j-2}) + \hat{f}_\sigma (\hat{x}_{i+2}, \hat{z}_j) + 2 \left( \hat{f}_\sigma (\hat{x}_{i-1}, \hat{z}_{j-1}) + \hat{f}_\sigma (\hat{x}_{i-1}, \hat{z}_{j+1}) + \hat{f}_\sigma (\hat{x}_{i+1}, \hat{z}_{j-1}) + \hat{f}_\sigma (\hat{x}_{i+1}, \hat{z}_{j+1}) \right) \right. 
- 8 \left( \hat{f}_\sigma (\hat{x}_{i-1}, \hat{z}_j) + \hat{f}_\sigma (\hat{x}_{i+1}, \hat{z}_j) + \hat{f}_\sigma (\hat{x}_i, \hat{z}_{j-1}) + \hat{f}_\sigma (\hat{x}_i, \hat{z}_{j+1}) \right) 
+ 20 \hat{f}_\sigma (\hat{x}_i, \hat{z}_j) \right) + O(\Delta \hat{x}^2) \tag{4.48}
\]

to smooth the interpolation. For the diffusion coefficient \(\epsilon_{\hat{x}, \hat{z}}\) we use values between \(\epsilon_{\hat{x}, \hat{z}} = 1\) and \(\epsilon_{\hat{x}, \hat{z}} = 5\) in typical simulations.

In the parallel direction the discretisation scheme destabilises high wavenumber modes that are stable in the non-discrete system \[135\]. The diffusion operator in the \(y\) direction is discretised, similarly to Eq. (4.35), as

\[
\left( \frac{\partial^4 \hat{f}_\sigma}{\partial \tilde{y}^4} \right) (\hat{x}_i, 0, \hat{z}_j) = \sum_{k=-2}^{2} \hat{f}_\sigma (\hat{x}_i, \tilde{y}_k, \hat{z}_j) \left( \frac{d^4 \hat{f}_\sigma}{d \tilde{y}^4} \right) (0) 
+ O \left( (\Delta \tilde{y})^3 \right) + O \left( \left( \frac{\Delta \hat{x}}{\Delta \tilde{y}} \right)^3 \right). \tag{4.49}
\]

The explicit form of the derivatives of the Lagrange polynomials is given in Appendix A.1. Due to the non-uniform grid in \(y\) direction, the grid spacing is not multiplied to the operator as in Eq. (4.48). Therefore, the diffusion coefficient must be adapted when the number of poloidal planes is increased. In typical simulations with 32 poloidal planes we choose values between \(\epsilon_{\hat{y}} = 5 \cdot 10^{-3}\) and \(\epsilon_{\hat{y}} = 1 \cdot 10^{-2}\).

Finally, we introduce numerical diffusion in the \(v_\parallel\) direction. Solutions of the Vlasov equation develop filament structures superimposed over a smooth part of the distribution function over time \[137\]. Similar to studying Fourier modes on periodic domains, the mode structure on the domain \((-\infty, \infty)\) in \(v_\parallel\) can be studied by expanding the distribution function in Hermite polynomials. The filamentation process causes excitations of higher and higher mode numbers. When the largest mode in the discrete system gets excited, the truncation acts as a reflecting boundary and the perturbation travels back towards lower mode numbers \[138\]. As a result, recurrence occurs and the distribution function is restored to a state close to the initial condition. Recurrence can be prevented by introducing an entropy producing, diffusive term that prevents the filamentation of the velocity space to scales that are not resolved by the grid resolution \[135\]. We discretise the numerical diffusion operator in \(v_\parallel\) with centred finite differences as

\[
\epsilon_{\hat{v}_\parallel} \Delta \hat{v}_\parallel^4 \left( \frac{\partial \hat{f}_\sigma}{\partial \hat{v}_\parallel} \right) (\hat{v}_\parallel) = \epsilon_{\hat{v}_\parallel} \left( \hat{f}_\sigma (\hat{v}_\parallel | t-2) - 4 \hat{f}_\sigma (\hat{v}_\parallel | t-1) + 6 \hat{f}_\sigma (\hat{v}_\parallel | t) - 4 \hat{f}_\sigma (\hat{v}_\parallel | t+1) + \hat{f}_\sigma (\hat{v}_\parallel | t+2) \right) 
+ O \left( (\Delta \hat{v}_\parallel)^2 \right). \tag{4.50}
\]

According to the analysis presented in \[135\], we typically choose a diffusion coefficient of \(\epsilon_{\hat{v}_\parallel} = 0.0125\).
4.4. Boundary conditions

For the simulations presented in this work we use Dirichlet boundary conditions. We set the distribution function to zero on the \(v_\parallel\) boundary and equal to the initial distribution \(f_{0\sigma}\) on the \(x,z\) boundary. In the \(y\) direction we apply the same Dirichlet boundary condition when a magnetic field line hits the \(x,z\) boundary in between two planes. In \(\mu\) there is no differentiation and therefore no boundary conditions is needed. We apply the Dirichlet boundary condition for both the advective inflow and outflow through the boundary. As the Vlasov equation is advective, only a boundary condition for the inflow is needed. To avoid spurious effects due to the boundary condition for the advective outflow, we include the option to apply diffusion in a small buffer zone around the \(x,z\) boundary\(^1\). The diffusion operator is discretised with a second order finite difference scheme

\[
-\epsilon_{\text{buffer}}(\hat{x}_i, \hat{z}_j) \left( \Delta \hat{x}_i, \Delta \hat{z}_j \right) \left( \hat{f}_\sigma(\hat{x}_{i-1}, \hat{z}_j) + \hat{f}_\sigma(\hat{x}_{i+1}, \hat{z}_j) + \hat{f}_\sigma(\hat{x}_i, \hat{z}_{j-1}) + \hat{f}_\sigma(\hat{x}_i, \hat{z}_{j+1}) - 4\hat{f}_\sigma(\hat{x}_i, \hat{z}_j) \right) + O(\Delta \hat{x}_i^2) .
\]

(4.51)

We set the strength of the diffusion in the buffer zone as large as the timestep of the simulation permits. The electrostatic potential \(\phi_1\) as well as the parallel component of the magnetic vector potential \(A_{1\parallel}\) are set to zero on the \(x,z\) boundary. This corresponds to a grounded, perfectly conducting wall.

4.5. Initial conditions

In order to study the behaviour and evolution of turbulence, a suitable initial condition for the simulation must be selected. The initial condition should fulfil two properties. First, it should be close to experimental input parameters, which are usually present in the form of density and temperature profiles for different species. Second, it should be an equilibrium of the gyrokinetic Vlasov-Maxwell system. If the initial condition fulfils both properties, instabilities can be triggered by adding a small perturbation to the initial condition. These instabilities grow exponentially and develop turbulence. The property that the initial condition without the added perturbation is an exact equilibrium of the gyrokinetic Vlasov-Maxwell system is crucial for the development of turbulence. If this property is not met, dynamics generated from the initial condition are present from the start and might hide, alter or even prevent the generation of turbulence in the system.

We generate the initial condition with the help of constants of motion of the particle trajectories. The kinetic energy of a particle, \(E = 1/2m_\sigma v_\parallel^2 + \mu B\), and the magnetic moment \(\mu\) are conserved. The conservation of \(\mu\) can be broken, if additional terms like a collision operator are added to the gyrokinetic Vlasov-Maxwell system. The presence of additional conserved quantities is dependent on the magnetic geometry used. In the following, we discuss suitable

\(^1\)By adding a diffusion term, the Vlasov equation turns from a hyperbolic to a parabolic problem. For the parabolic problem Dirichlet boundary conditions can be set consistently for the in- and outflow boundary.
initial conditions for different geometries. The first is a slab geometry, where the magnetic field is given by
\[
\hat{B} = \hat{B} e_\varphi
\] (4.52)
and the second is a circular geometry with concentric flux surfaces, where the magnetic field is given by
\[
\hat{B} = \hat{B} \left( e_\varphi + \frac{\hat{r}}{q(\hat{r})} e_\theta \right).
\] (4.53)
The function \(q(\hat{r})\) denotes the \(q\) profile. For both the slab and circular geometry we use a large aspect ratio assumption such that \(\varphi\) represents a Cartesian coordinate labelling poloidal planes. In slab and circular geometry the gradient and curvature drift of a particle are confined to a flux surface. Therefore, the poloidal flux function \(\psi_p\) is a constant of motion. The poloidal flux function is converted into a flux surface label \(\psi\) by normalising it with the value of the poloidal flux function at the separatrix \(\psi_{sep}\) and the magnetic axis \(\psi_{ax}\). In addition, we take the square root of the normalised result
\[
\psi(\hat{\psi}_p) := \begin{cases} 
\sqrt{\frac{\hat{\psi}_p - \hat{\psi}_{ax}}{\hat{\psi}_{sep} - \hat{\psi}_{ax}}} & \text{for } \hat{\psi}_p > 0 \\
0 & \text{for } \hat{\psi} \leq 0
\end{cases}.
\] (4.54)
The case distinction is needed to account for negative values of the normalised flux surface label. In slab and circular geometry we initialise the simulation with local Maxwellians with given normalised temperature and density profiles \(\hat{T}_{\sigma}\) and \(\hat{n}_{\sigma}\), defined in Section 4.1.2,
\[
\hat{F}_{0\sigma}(\psi(\hat{\psi}_p), \hat{v}_\parallel, \mu) = \frac{\hat{n}_{\sigma}(\psi(\hat{\psi}_p))}{\pi \hat{T}_{\sigma}(\psi(\hat{\psi}_p))^{3/2}} e^{-\frac{\hat{\psi}_p^2 + \mu \hat{B}}{\hat{T}_{\sigma}(\psi(\hat{\psi}_p))}}.
\] (4.55)
For toroidal geometries with a background magnetic field \(\hat{B} \propto 1/R = 1/(\hat{R}_0 + \hat{r}\cos(\theta))\) the \(\nabla \hat{B}\) drift points out of a flux surface and, consequently, the flux surface label is no constant of motion anymore. For the following discussion we assume that the equilibrium is axisymmetric, that the model contains no electromagnetic fluctuations and that the initial condition is free of an electrostatic potential. We discuss the implications of the assumptions at the end of this section. Under the assumption of axisymmetry and the absence of an electromagnetic vector potential, the canonical toroidal momentum, defined in Section 2.3.6, is conserved. Following [139, 140, 42], we define a modified poloidal flux function with the help of the canonical toroidal momentum. The corrected flux function is given by Eq. (4.21) and in normalised units by Eq. (4.22). The so-called canonical Maxwellian is obtained by replacing the poloidal flux function \(\hat{\psi}_p\) with the corrected poloidal flux function \(\hat{\psi}_C\), i.e.
\[
\hat{F}_{C\sigma}(\hat{\psi}_C, \hat{v}_\parallel, \mu).
\] (4.56)
The canonical Maxwellian is a function of constants of motion and therefore conserved. Due to the \(\sqrt{2m_\sigma}\) dependence of Eq. (4.22), the correction to the local Maxwellian is by a factor
of 60 larger for deuterium than for electrons. The corrected flux function $\psi_C$ has an explicit dependence on $v_\parallel$ and therefore introduces a mean flow, similar to the Pfirsch-Schlüter current [8], in parallel direction to the magnetic field. This flow has been investigated in detail in previous studies [139, 140] and has been mentioned to be problematic for the onset and the properties of turbulence. To overcome the flow, a correction to $\psi_C$ has been defined that is a constant of motion for passing particles. It reads [139, 140]

$$\psi_{C,s} = \psi_C + \frac{m_e c}{q_e} R_0 \bar{v}_\parallel,$$  \hspace{1cm} (4.57)

with

$$\bar{v}_\parallel = \text{sign}(v_\parallel) \sqrt{2(E - \mu B_{\text{max}})} \mathcal{H}(E - \mu B_{\text{max}}),$$ \hspace{1cm} (4.58)

and in normalised units

$$\hat{\psi}_{C,s} = \hat{\psi}_C + \frac{\rho_{ref}}{L_{ref}} \sqrt{2 \hat{m}_e \hat{T}_{0e}}(\psi_0) \text{sign}(\hat{v}_\parallel) \sqrt{\hat{E} - \hat{\mu B}_{\text{max}}} \mathcal{H}(\hat{E} - \hat{\mu B}_{\text{max}}).$$ \hspace{1cm} (4.59)

In the following, we call the canonical Maxwellian constructed with $\hat{\psi}_{C,s}$ instead of $\hat{\psi}_C$ shifted canonical Maxwellian. To visualise the difference between the local Maxwellian, canonical Maxwellian and shifted canonical Maxwellian, we display the ion distribution function for all three Maxwellians, for the cyclone base case setup described in Section 6.5, in Fig. 4.4. The distribution function is shown for the first $\hat{\mu} = 0.044$ grid point and at the outboard mid-plane at $\psi = 0.62$. The shape of the canonical Maxwellian is different from the local Maxwellian. Due to the sign function in Eq. (4.58), the shifted canonical Maxwellian is non-differentiable close to $\hat{v}_\parallel = 0$. This can cause stability and convergence problems based on the discretisation scheme chosen. The density at the outboard mid-plane, resulting from an initialisation of the distribution function with a given density profile $\hat{n}_{p\sigma}$, is shown for the different Maxwellians in Fig. 4.5.

Initialising ions and electrons with canonical Maxwellians results in good conservation of the initial condition if the assumptions above are met. Particularly, the electrostatic potential and the induced electromagnetic vector potential need to vanish. It has also been shown that initialising the ions with a canonical Maxwellian works well in the limit of an adiabatic electron response [139].

With both kinetic ions and electrons, the situation changes. It is not possible to initialise both ions and electrons with canonical Maxwellians and achieve exact quasi-neutrality at the same time. As mentioned above, electrons are lighter than ions and therefore their canonical Maxwellian is close to a local Maxwellian. As shown in Fig. 4.5, this results in different density profiles which generate a non-zero charge density. Further, the canonical Maxwellian is not constant on magnetic flux surfaces due to the explicit dependency on $b_\psi$. The variation of the canonical Maxwellian on a flux surface is again different for ions and electrons. We have tried different approaches to solve this problem.

1. The canonical Maxwellian for electrons is close to a local Maxwellian. Therefore, it is reasonable to initialise the ions with a canonical Maxwellian, calculate the resulting ion
Figure 4.4.: Figure displaying a slice of the ion distribution initialised with a local Maxwellian, a canonical Maxwellian and a shifted canonical Maxwellian for the cyclone base case configuration described in Section 6.5. The distribution function is shown for the first grid point in $\mu = 0.044$ and at the outboard mid-plane at $\psi = 0.62$. Due to the sign function in Eq. (4.58), the shifted canonical Maxwellian is non-differentiable close to $v_\parallel = 0$.

Figure 4.5.: Figure displaying the ion density profile calculated from an ion distribution function initialised with a local Maxwellian, a canonical Maxwellian and a shifted canonical Maxwellian for the cyclone base configuration described in Section 6.5. The density of both the canonical and shifted canonical Maxwellian differ from the local Maxwellian. The shifted canonical Maxwellian is closer to the local Maxwellian than the canonical Maxwellian.
density and initialise the electrons with a local Maxwellian and the given ion density. While this initial state is quasi-neutral per construction, we find that the conservation of the electron distribution is so bad that quickly after initialisation large charge densities and, consequently, a radial electric field is generated.

2. In the second approach, ions and electrons are initialised with a canonical Maxwellian. As this state is not quasi-neutral, we calculate the ion density $n_i$ and electron density $n_e$ after loading the distribution function and correct the electron distribution function by the factor $n_i/n_e$. This ensures exact quasi-neutrality of the initial condition because ion and electron densities are equal. Nevertheless, the correction breaks the conservation of the initial electron distribution. The correction factor $n_i/n_e$ is a function of the coordinates $R, Z$, which are no constants of motion. If the correction factor $n_i/n_e$ is small enough, the charge density generated by the electron dynamics is small and turbulence can develop freely. We find that this is not the case for realistic profiles. The reason is simple. As shown in Fig. 4.4 and Fig. 4.5, the distribution function of the ions is significantly different from a local Maxwellian. Consequently, the ion density is different from the initial profile and different from the electron density. This results in a large difference between ion and electron density and a large correction factor $n_i/n_e$, which effectively breaks the conservation.

3. The third approach proceeds similarly to the second one with the only difference that the shifted instead of the standard canonical Maxwellian is used. Consequently, the ion distribution function is closer to a Maxwellian and the resulting ion density profile is closer to the profile generated by a local Maxwellian. As a result, ion and electron density are approximately equal and the correction factor $n_i/n_e$ is smaller than in the second approach. This is the standard initial condition implemented in the GENE-X code for toroidal equilibria. It works for a lot of profiles and geometries and allows for the stable simulation of turbulence. Nevertheless, the solution is still dependent on the right parameters.

The above analysis holds true in the limit of vanishing electromagnetic vector potential $A_{1∥}$. As soon as electromagnetic fluctuations are activated, the situation changes. First, the canonical toroidal momentum and the canonical Maxwellian are modified and obtain a dependence on $A_{1∥}$. Let us call the electromagnetic vector potential entering the canonical Maxwellian $A_{1∥init}$. As discussed above, the canonical Maxwellian introduces a mean flow, which leads to an electromagnetic vector potential $A_{1∥generated}$. The canonical Maxwellian is only conserved if $A_{1∥init} = A_{1∥generated}$, which is in general not fulfilled. Therefore, the initial condition implemented in the GENE-X code for toroidal equilibria is only stable for small $\beta_{ref}$. In addition, the conservation is broken in non-axisymmetric equilibria.
4.6. Conclusion and outlook

In this chapter, we presented the numerical scheme implemented in the GENE-X code. The numerical scheme was based on centred finite differences and motivated by the use of the locally field-aligned coordinate system, presented in Chapter 3. The chosen numerical scheme contains little numerical diffusion and therefore respects the property that the gyrokinetic Vlasov-Maxwell system is free of diffusion on the continuous level. The total scheme is of order two, although multiple operators are implemented to fourth order accuracy. Furthermore, we explained why it is difficult to construct a stable initial condition for the full-f gyrokinetic Vlasov-Maxwell system and detailed the initial condition for our simulations.

In the future, three upgrades to the numerical scheme would be advantageous. First, the elliptic operators and the nonlinear Poisson brackets should be discretised with numerical schemes that are fourth order accurate. Consequently, this would render the discretisation of the complete gyrokinetic Vlasov-Maxwell system accurate to fourth order. Second, the scheme in its current form does neither preserve the number of particles, energy or momentum to machine precision nor the positivity of the distribution function. It would be interesting to construct a conservative numerical scheme within the framework of mimetic finite differences that preserves the particle number and if possible even momentum and energy. For the inclusion of collisions into the simulations it is necessary to take roots of the moments of the distribution function. This is only well-defined if the moments remain positive. It will be important to explore schemes that preserve the positivity of the distribution function in the future. Third, it would be desirable to develop and implement a numerical solver that can calculate an initial condition for the simulations that is stable to machine precision.
5. Implementation

Our goal is to create a scientific code that is both fast and extensible. These two requisites contradict each other to a certain degree. For a code being extensible data encapsulation, well-structured interfaces and code abstraction is necessary. For execution speed it is best to work on raw data such that capabilities of modern hardware, like vectorisation, can be used efficiently. In the following chapter, we present and discuss our code design and prove that it provides excellent performance while being extensible. We start by presenting the combined project structure with the GRILLIX code. Next, we discuss the software engineering of GENE-X and introduce the data structures and algorithms, called operators. In addition, we discuss the structure of the time stepping and the main program. We validate the performance of the implementation with the help of the roofline model [141] and the measurement of strong [142] as well as weak scaling [143] up to 20480 CPU cores.

5.1. Project structure

As detailed in the last two chapters, GENE-X uses the FCI approach and unstructured, locally Cartesian grids. Consequently, it shares a lot of technology with the Braginskii fluid code GRILLIX. To strengthen the collaboration between both codes, we created a library based on GRILLIX, called PARALLAX, that contains the functionality to generate the grid for different magnetic equilibria, trace magnetic field lines using dop853, perform bicubic polynomial interpolation and solve elliptic equations within poloidal planes. The library is now used by both GRILLIX and GENE-X and allows us to join forces and reduce maintenance work in both codes.

5.2. Software engineering

GENE-X is written in Fortran 2008. We decided to choose Fortran 2008 over C++ because it combines modern software engineering concepts, like object orientation, with powerful array handling capabilities.

5.2.1. Data structures

We store the data of the distribution function as a five-dimensional and the data of the electromagnetic fields as two-dimensional contiguous arrays\(^1\). As all performance demand-
ing operations are performed on this data, it is crucial to store it in contiguous memory and such allow for vectorisation, cache line optimisation and optimisations by the compiler. The five-dimensional array is stored in the `data_storage_5d_t` class and the two-dimensional arrays in the `data_storage_2d_t` class respectively. The data is distributed over different Message Passing Interface (MPI) [144] processes. The exchange of ghost cells, necessary for the calculation of stencil operations, is encapsulated in the classes. The MPI ghost exchange is implemented via non-blocking communication and is performed automatically by calling `start_exchange` and `finish_exchange`. The arrays themselves are not encapsulated and a pointer is retrieved by calling `get_pointer`. These design decisions allow for the efficient implementation of operators acting on the fields. The array storing the data is either allocated on CPU or GPU. CPU allocation is implemented in the child class `data_storage_5d_cpu_t` and GPU allocation in `data_storage_5d_gpu_t`. GPU and CPU implementations are instantiated and chosen at runtime. The design of the data structures is depicted as an Unified Modelling Language (UML) diagram in Fig. 5.1.

5.2.2. Operators

The calculation of the right-hand side of the Vlasov equation, the solution of the field equations as well as diagnostics are implemented as operators acting on raw data. The interface of the operators is defined in a base operator class. Different realisations of the operator are implemented as children of the base class. A good example is the operator evaluating the part \((\partial f_\sigma / \partial t)^*\), independent of explicit time derivatives, of the right-hand side of the Vlasov equation, called `op_rhs_vlasov_eq_static_t` in GENE-X. The class `op_rhs_vlasov_eq_static_t` defines the interface of the operator and the children `op_rhs_vlasov_eq_static_cpu_t` or `op_rhs_vlasov_eq_static_gpu_t` contain CPU or GPU implementations of the operator. This way of selecting different execution schemes is known as the strategy design pattern [145, pp. 315–323]. The operators receive contiguous pointers to raw data as input. As different operators require different inputs, each operator is implemented as its own class. Next to the operator for evaluating the right-hand side, there are for example operators to obtain scalar and multi-dimensional moments of the distribution function called `op_mom_0d_t` and `op_mom_2d_t`. The design of the operators is depicted as UML diagrams in Fig. 5.2.

5.2.3. The state vector and time stepping

After having introduced the data structures and operators, we describe how they interact in the code. The state of the electromagnetic model, often called state vector, is defined by the distribution function, the electrostatic potential and the parallel component of the electromagnetic vector potential. They are encapsulated in the `state_vector_t` class. With that also the raw data, accessible by the `get_pointer` method of the data storage types, is encapsulated. The operators need access to the raw data. This is achieved in an encapsulated way by passing the operator as a `visitor` [145, pp. 331–344] to the state vector by the `apply` instead of three-dimensional arrays consisting out of the unstructured grid and \(\varphi\).
Figure 5.1.: UML diagram representing the design of the data structures in GENE-X. The data for the distribution function, the electrostatic potential and the parallel component of the electromagnetic vector potential are stored in 2d and 5d arrays in the classes `data_storage_5d_t` and `data_storage_2d_t`. The MPI ghost exchange is encapsulated and initiated with `start_exchange` and `finish_exchange`. The data is stored either on CPU or GPU.
Implementations of the operators in GENEX differ by target, e.g., CPU or GPU, or by the numerical scheme used.

Figure 5.2: UML diagrams representing the design of the operators in GENEX. The operators are organized in classes that perform computations on contiguous pointers to raw data. The parent class is abstract and defines the interface of the operation. The children provide different implementations of the operator. Implementations differ by target, e.g., CPU or GPU, or by the numerical scheme used.
Within the `apply` method the operator instance has access to all private data in the state vector and can perform operations on the raw data. This design is depicted as an UML diagram in Fig. 5.3. With this software structure it is simple to extend the state vector by introducing more fields or extend operators acting on the state vector. The interface of the operators is likely to change in the future when additional phenomena, like the perpendicular component of the electromagnetic vector potential, is included in the model. As the operators are implemented as their own class, further fields and effects can be added by modifying the operator locally.

![UML diagram](image)

**Figure 5.3.:** UML diagram representing the structure of the state vector. The state vector contains multiple instances of the data storage types to store the electrostatic potential, the electromagnetic vector potential, the distribution function or other fields. Operators are applied to the state vector by passing them as a visitor to the `apply` method. This ensures that the raw data, stored in the data structure, remains in the state vector and is not accessible from the outside.

In order to evolve the physical system, we need to apply operators to the state vector in a way specified by the time stepping scheme. For simulations of different models and geometries, it might be beneficial to have multiple time stepping schemes at our disposal. The interface of the time stepping is defined in the abstract `time_step_t` class. It contains one or more different state vectors, depending on the specific time stepping scheme implemented. A time step is calculated by calling `step`, diagnostics are written by calling `diagnose` and the state vector is saved to disk as a checkpoint by calling `save`. Our workhorse time stepping scheme is, as presented in Section 4.2, the fourth order Runge-Kutta method implemented as a child of `time_step_t` in `rk4_t`. The time stepping is controlled from the main program `genex`. The time stepping design is once again depicted as an UML diagram in Fig. 5.4.

### 5.2.4. Continuous integration

Next to the actual software design, automating and optimising the development process is an important issue for us. We employ the software engineering practice called *Continuous Integration* (CI) [146] in the development process.

Due to the modular software design, presented in the last subsections, the code consists of many independent and flexible units with well-defined interfaces which are simple to test. We test the functionality of the interfaces with the help of the Fortran unit test library `pFUnit`[147]. `pFUnit` allows to automatically execute tests for a different amount of MPI processes and OpenMP threads making the unit tests comprehensive.
Figure 5.4.: UML diagram representing the structure of the main program. The time step is handled by the `time_step_t` class. `time_step_t` is responsible for storing one or multiple instances of the `state_vector_t` representing the state of the simulation. It evolves the simulation one time step by calling `step`, it writes diagnostics by calling `diagnose` and it saves the current state of the simulation as a checkpoint to disk by calling `save`.

The build process is automatized with CMake and supports the GNU as well as Intel Fortran compiler. Upon integration of new implementations in the master branch, the code is built automatically with both the GNU and Intel compiler for debug and release configurations. Furthermore, the unit tests and fast integration tests are run for all configurations. New functionality can only be incorporated into the master branch when all tests succeed.

### 5.3. Parallelisation and performance

The implementation is parallelised with a hybrid approach using both OpenMP [148] and MPI [144]. Currently, no GPU implementation of the operators and data structures is available. The reason for choosing a hybrid approach is twofold. First, modern supercomputers are heterogeneous. CPU based machines usually consist out of compute nodes with one or multiple sockets each. The nodes themselves are connected with a fast network. With a hybrid approach it is straightforward to adjust the parallelisation to the hardware needs. OpenMP is well suited to parallelise tasks within a node because it makes use of the shared memory architecture. MPI is used to exchange information over the network connecting the nodes. Second, by separating intra- and inter-node parallelisation it is easier to optimise for the
use of accelerators in the future. For instance, OpenMP can be replaced with OpenACC [149] or a specific CUDA [150] implementation can be written for the datastructures and operators introduced in Section 5.2.

We use OpenMP to parallelise over the $x$, $z$ direction within the poloidal plane. Therefore, no inter node communication is necessary during the solution of the elliptic field equations. We employ a domain decomposition with MPI over the $v_{\parallel}$, $\mu$, $\varphi$ and species dimensions.

All benchmarks are performed on the supercomputer Cobra of the Max-Planck Computing and Data Facility. All compute nodes of Cobra feature two Intel Xeon Gold 6148 processors with 20 cores and 27.5Mib L3 Cache each. The software is compiled with the Intel Fortran Compiler Version 19.1.1.

5.3.1. Node level performance

We analyse the node level performance with the roofline model [141]. The goal of the roofline model is to calculate the maximal attainable rate of floating-point operations per second (Flop/s) for a given program. Before we discuss the model, we introduce the terminology needed. The arithmetic intensity $I$ is the quotient of floating-point operations per bytes loaded. We explain the term at the example of the saxpy operation

$$\text{do } i = 1, N$$
$$y(i) = y(i) + sa * x(i)$$
$$\text{enddo} .$$

The arithmetic intensity is $I = \frac{2}{24} = \frac{1}{12} \approx 0.083$ because in each iteration there are two floating-point operations, two eight-byte loads and one eight-byte store. Further, we call the maximal memory bandwidth of the hardware $B$ and the hardware limit of the rate of floating-point operations $F$. The theoretically achievable flop rate, or roofline, of a program is then given by

$$R = \min \{B \cdot I, F\} . \quad (5.1)$$

In order to apply the roofline model to our implementation, we measure the maximum memory bandwidth with a stream benchmark provided by the likwid [151, 152] toolkit\(^\dagger\). The measured maximum main memory bandwidth is $B = 150\text{Gb/s}$ and the maximum L3 Cache memory bandwidth is $B = 660\text{Gb/s}$. For the hardware limit of the rate of floating-point operations we use the specifications by Intel. The maximum rate of scalar addition operations is $F = 190\text{GFlop/s}$ and of vectorised addition operations is $F = 1400\text{GFlop/s}$. All our operators are bound by memory bandwidth such that the hardware limit of the rate of floating-point operations is not of high significance. For the saxpy example the roofline rate of floating-point operations evaluates to $R = 12.5\text{GFlop/s}$.

We perform the roofline analysis for the operators consuming major parts of the computation time in GENE-X. The operators are given in Tab. 5.1. We select a problem size with 677000 grid points in the poloidal plane, $n_\varphi = 1$, $n_{v_{\parallel}} = 16$, $n_\mu = 8$ that reflects a typical load for

\(^{\dagger}\)We found that the theoretical memory bandwidth specified by Intel for the Intel Xeon Gold 6148 dual socket system is not achievable in real applications.
<table>
<thead>
<tr>
<th>Operator</th>
<th>Arithmetic intensity</th>
<th>Performance / (GFlop/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>op_saxpy_cpu_t</td>
<td>0.083</td>
<td>11.9</td>
</tr>
<tr>
<td>op_lin_comb_cpu_t</td>
<td>0.13</td>
<td>14.8</td>
</tr>
<tr>
<td>op_diag_mom_0d_cpu_t</td>
<td>0.10</td>
<td>121</td>
</tr>
<tr>
<td>op_diag_mom_2d_cpu_t</td>
<td>0.12</td>
<td>163</td>
</tr>
<tr>
<td>op_rhs_vlasov_eq_static_cpu_t</td>
<td>0.092</td>
<td>28.3</td>
</tr>
<tr>
<td>op_rhs_vlasov_eq_dynamic_cpu_t</td>
<td>0.19</td>
<td>51.9</td>
</tr>
</tbody>
</table>

Table 5.1.: Table displaying the arithmetic intensity and the measured performance of operators consuming major parts of the computation time in GENE-X. op_saxpy_cpu_t and op_lin_comb_cpu_t are used to combine the stages of the RK4 time stepping scheme. op_diag_mom_0d_cpu_t and op_diag_mom_2d_cpu_t represent diagnostic operators with scalar and two-dimensional output. Finally, op_rhs_vlasov_eq_static_cpu_t and op_rhs_vlasov_eq_dynamic_cpu_t evaluate the right-hand side of the Vlasov equation.

We measure the arithmetic intensity as well as the flop rate of the operations with Intel Advisor 2020. Intel Advisor reports the arithmetic intensity of the L1 Cache. That means all loads and stores from and to L1 cache are counted in the calculation of the arithmetic intensity. The result of the roofline analysis is shown in Fig. 5.5. The op_saxpy_cpu_t and op_lin_comb_cpu_t operators perform close to the main memory bandwidth roofline. This is expected and a nice validation of the model because there is no possibility for cache reuse. The diagnostic operators op_diag_mom_0d_cpu_t and op_diag_mom_2d_cpu_t fall above the L3 cache roofline. As they load the distribution function once to calculate six different moments a high amount of cache reuse is expected. The operators evaluating the right-hand side of the gyrokinetic Vlasov equation, op_rhs_vlasov_eq_static_cpu_t and op_rhs_vlasov_eq_dynamic_cpu_t, fall in between the DRAM and L3 cache roofline indicating a decent amount of cache reuse. It is difficult to estimate the maximal theoretical performance of these more sophisticated operators because the cache reuse is unknown.

### 5.3.2. Strong scaling

After discussing the intra-node performance, we proceed by testing the MPI scaling of the implementation. We start with the strong scaling that was first described by Amdahl [142]. In strong scaling a problem with a fixed size is selected and the speedup of the execution time is measured as a function of the number of compute nodes [153]. We use the AUG example, presented in Chapter 7, with a resolution of $\Delta x = \Delta z = 1.91 \cdot 10^{-4}$ mm resulting in 223000 grid points in the poloidal plane, $n_\nu = 24$, $n_{v_1} = 64$ and $n_\mu = 16$. The resolution has been reduced such that we are able to fit the simulation onto 1 node. The memory usage with 1 node is approximately 700GB exhausting the available 768GB of main memory on the fat partition on Cobra. The total number of grid points is equal to $11.0 \cdot 10^9$. We start with 1 node and scale up to 512 nodes. We first fill the $\mu$ and species communicator with nodes as

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†The number of grid points includes a factor of two from the two species.
minimal MPI communication is needed in these directions. After they are filled, we use the \( \varphi \) and \( v_\parallel \) communicator that require a ghost exchange via MPI. We measure the wall clock time per time step averaged over 100 iterations. For the run with only 1 node, we average the execution time over 50 timesteps to be able to run the benchmark within the time limit of 24h imposed by our supercomputing centre. The time of the first iteration is discarded due to the presence of memory allocations and initialisations. The results are shown in Fig. 5.6. The times are given in Appendix B.1. We define the efficiency of the strong scaling by

\[
\eta := \log_2 \left( \frac{t_n}{t_{2n}} \right)
\]

(5.2)

where \( t_n \) and \( t_{2n} \) denote the time per timestep with \( n \) nodes and \( 2n \) nodes respectively. We measure an average reduction of execution time by a factor of 1.9 upon doubling the number of nodes. This corresponds to an efficiency of \( \eta = 0.93 \), demonstrating excellent strong scaling.

### 5.3.3. Weak scaling

We continue with the weak scaling that was first described by Gustafson [143] as a re-evaluation of Amdahl’s work. In weak scaling the number of compute nodes is increased while keeping the problem size per node constant [153]. This results in larger problems the more nodes are chosen. Similar to the strong scaling, we select the problem based on the AUG simulation presented in Chapter 7. As we have more memory per node available than in the strong scaling, we choose a higher resolution of \( \Delta x = \Delta z = 9.54 \cdot 10^{-4} \text{mm} \) resulting in 892000 points in the poloidal plane. We start the scaling measurement with 1 node and...
Number of Nodes

Speedup

Measured scaling $\mu$

Measured scaling $\sigma$

Measured scaling $\varphi$

Measured scaling $v_{||}$

Scaling $\eta=1$

Scaling $\eta=0.93$

Figure 5.6.: Figure displaying the strong scaling analysis of the GENE-X code over different MPI communicators. For reference, the ideal scaling with an efficiency of $\eta = 1$ and the scaling with an efficiency of $\eta = 0.93$ is shown. On average, GENE-X shows an efficiency of $\eta = 0.93$, demonstrating excellent strong scaling.

$n_x = 2$, $n_{v_{||}} = 8$, $n_{\mu} = 2$ and fill the MPI communicators one after another up to 512 nodes and $n_x = 16$, $n_{v_{||}} = 64$ and $n_{\mu} = 16$. The number of grid points is equal to $57.1 \cdot 10^6$ per node. Again, we measure the time per timestep by averaging the execution time over 100 timesteps. The results are shown in Fig. 5.7. We observe that the time per timestep is almost constant from 1 to 512 nodes, demonstrating excellent weak scaling.

5.4. Conclusion and outlook

We started this chapter with the goal to create a scientific code that is fast and extensible. We achieved both by implementing a software design that is focused around data structures and operators, acting on raw data. Both are encapsulated in classes and implementations, specialised to different compute architectures, can be provided. We demonstrated that the implemented software design is fast by validating the intra-node performance against the roofline model and the inter-node performance by measuring the strong and weak scaling from 1 to 512 compute nodes. The strong scaling showed an efficiency of 93% and the weak scaling was nearly ideal.

Many supercomputers today feature accelerated compute nodes with GPUs [154]. To make use of these architectures, it will be important to exploit the modular code structure of GENE-X and provide a GPU implementation of the data structures and operators in the future.
Figure 5.7.: Figure displaying the weak scaling analysis of the GENE-X code. The measured time per timestep is almost constant from 1 to 512 nodes, demonstrating excellent weak scaling.
6. Verification

With the help of good software engineering practices, presented in Chapter 5, we ensure that the implementation of the numerical scheme is correct. The next step is to show that the numerical scheme converges and returns approximate solutions of the gyrokinetic Vlasov-Maxwell system.

To this end, we present different tests of the GENE-X code in this chapter. We start with purely numerical tests and analyse the accuracy of the field line tracing, which is central for calculating derivatives in the locally field-aligned coordinate system, and perform a convergence order analysis of the whole gyrokinetic Vlasov-Maxwell system with the Method of Manufactured Solutions (MMS) [155]. Next, we test different physical properties of the system. Each of the physical tests probes a different aspect of the code. We start with testing the linear regime of the gyrokinetic system in the limit of no electromagnetic fluctuations, i.e. $A_1 = 0$, by calculating and comparing ion temperature gradient growth rates in screw pinch geometry against analytical predictions. This test is well suited to compare the performance of the locally field-aligned coordinate system to standard flux-coordinates. We test the nonlinear phase by checking the conservation of energy during a turbulence simulation. We proceed with testing the electromagnetic implementation by measuring and comparing the frequency and damping rate of Alfvén waves in slab geometry. Finally, we perform a well studied electromagnetic Cyclone Base Case (CBC) benchmark [156] that tests the full gyrokinetic Vlasov-Maxwell system in toroidal geometry.

6.1. Accuracy of the field line tracing

As explained in Chapter 3, the length of a magnetic field line as well as the intersection points with neighbouring poloidal planes is calculated by solving a system of ordinary differential equations. We use the algorithm dop853 for solving the system. dop853 is an embedded Runge-Kutta method of order eight that uses a fifth order error estimator with a third order correction [122, II. 10]. The local approximation error is controlled by providing relative and absolute error tolerances. It is kept roughly below $\text{rtol} \times \text{abs(sol)} + \text{atol}$ [122, Appendix], where $\text{rtol}$ and $\text{atol}$ represent the relative and absolute error tolerance and $\text{sol}$ the value of the solution. In the following, we test the convergence and precision of the integrator for our application.

We choose a circular geometry with the magnetic field specified in Eq. (4.53) and a large aspect ratio assumption such that $\varphi$ represents a Cartesian coordinate labelling poloidal planes. The $q$ profile is given by

$$q(\hat{r}) = q_0 + \hat{s} \cdot \hat{r}.$$  \hspace{1cm} (6.1)
For this geometry the system of equations presented in Eqs. (3.75), (3.76), (3.77) and (3.78) can be solved analytically. The position and the length of a magnetic field line starting at $(\hat{r}_0, \theta_0, \varphi_0)$, parametrised as a function of the poloidal angle $\varphi$, read

$$\hat{r}(\varphi) = \hat{r}_0$$  \hspace{1cm} (6.2)

$$\theta(\varphi) = \theta_0 + \frac{(\varphi - \varphi_0)}{q(\hat{r})}$$  \hspace{1cm} (6.3)

$$\hat{s}(\varphi) = \sqrt{1 + \left(\frac{\hat{r}}{q(\hat{r})}\right)^2} (\varphi - \varphi_0)$$  \hspace{1cm} (6.4)

First, we analyse the accuracy of the position of the field line. We compare the coordinate $\hat{z} = \hat{Z} = \hat{R}_0 + \hat{r} \sin(\theta)$, obtained from the numerical solution with dop853, against the analytical solution for the angles $\varphi \in \{\pi/8, \pi, 100\pi\}$, corresponding to $1/16$, $1/2$ and $50$ toroidal turns. For the $z$ coordinate the absolute error

$$\epsilon = |\hat{z}_{\text{dop}} - \hat{z}|$$  \hspace{1cm} (6.5)

is measured. We set the relative tolerance of dop853 to machine precision and scale the absolute tolerance. The results are shown in the upper image in Fig. 6.1. For $\varphi = \pi/8$ the field line tracing is precise to machine precision for every tolerance. For larger trace angles we observe a convergence with decreasing tolerance. The measured error is always larger than the provided tolerance. This is expected because the tolerance limits the local and not the global approximation error of the numerical solution.

Next, we analyse the accuracy of the calculation of the field line length. As the field line length increases with increasing trace angle $\varphi$, we measure the relative numerical error

$$\epsilon = \frac{|\hat{s}_{\text{dop}} - \hat{s}|}{|\hat{s}|}$$  \hspace{1cm} (6.6)

for the angles $\varphi \in \{\pi/8, \pi, 100\pi\}$. We set the absolute tolerance to machine precision and scale the relative tolerance. The results are shown in the lower image in Fig. 6.1. We observe a similar behaviour as for the $z$ coordinate. For $\varphi = \pi/8$ the numerical integration is exact to machine precision. For larger trace angles we observe convergence. In the simulations presented in this work, we use 16 or more poloidal planes. The trace angle required to calculate parallel derivatives with the fourth order scheme, presented in Chapter 4, is equal to $\pi/8$ or smaller. Therefore, the accuracy of the tracing close to machine precision.
Figure 6.1.: Figure displaying the accuracy of the field line tracing with the dop853 algorithm. We test three different trace angles $\varphi \in \{\pi/8, \pi, 100\pi\}$. Above the absolute error of the $z$ coordinate of the field line and below the relative error of the field line length is shown. With increasing tolerance the field line tracing converges. For clarity, the convergence is highlighted with a dashed line.
6.2. Method of manufactured solutions

In this section, we present a full convergence order analysis of the numerical scheme in slab, circular and toroidal geometry with MMS. The explanation and introduction into MMS have been published in [87]. In this section, the results published in [87] are generalised from the electrostatic to the electromagnetic model, detailed in Chapter 2, with the improved numerical scheme described in Chapter 4.

The gyrokinetic Vlasov-Maxwell system is a complicated integro-partial differential system of equations and exact solutions are only known for special cases. Therefore, it is not possible to comprehensively check the convergence of the numerical solution by comparing it against exact solutions. Nevertheless, it is possible to add extra terms to the Vlasov-Maxwell system such that it obtains any solution. This is known as the method of manufactured solutions (MMS) [155]. In order to explain it in more detail, we rewrite the gyrokinetic Vlasov-Maxwell system in a different form. Consider the integro-partial differential operators $L(\phi)$, $N(f)$ and $A(f)$ such that

\begin{align}
    L(\phi_1, A_{1||})[f] &= 0 \\
    N(f)[\phi_1] &= 0 \\
    A(f)[A_{1||}] &= 0
\end{align}

represents the gyrokinetic Vlasov-Maxwell system. $L$ represents the Vlasov operator which depends on the electrostatic potential and the parallel component of the electromagnetic vector potential, $N$ represents the operator that forms the quasi-neutrality equation and $A$ the operator that forms Ampère’s law. In the method of manufactured solutions a modified system of equations

\begin{align}
    L(\phi_1, A_{1||})[f] &= S_f \\
    N(f)[\phi_1] &= S_{\phi_1} \\
    A(f)[A_{1||}] &= S_{A_{1||}}
\end{align}

is solved, where the sources $S_f$, $S_{\phi_1}$ and $S_{A_{1||}}$ are included in the right-hand side of the equations such that the given functions $f_{MMS}$, $\phi_{1MMS}$ and $A_{1||MMS}$ are a solution to the system. The functions $f_{MMS}$, $\phi_{1MMS}$ and $A_{1||MMS}$ can be chosen arbitrarily. The sources $S_{\phi_1}$, $S_f$ and $S_{A_{1||}}$ are calculated by plugging the solutions into Eqs. (6.7), (6.8) and (6.9). As the solutions are known this is a straightforward process and can be done with a computer algebra system. We use Mathematica [157]. The functions $f_{MMS}$, $\phi_{1MMS}$ and $A_{1||MMS}$ are called the manufactured solutions.

Numerical solutions of the modified Vlasov-Maxwell system, presented in Eqs. (6.10), (6.11) and (6.12), can be compared to the exact solutions $f_{MMS}$, $\phi_{1MMS}$ and $A_{1||MMS}$. This makes it possible to calculate relative errors of the numerical solution and to study the order of convergence of the numerical scheme.
As manufactured solutions we choose the functions

\[
\begin{align*}
    f_{\text{MMS}}(\hat{r}, \varphi, \theta, \hat{\nu}, \hat{\mu}, \hat{t}) &:= \cos^2(\hat{\omega} \hat{t}) \cos^2(n_{\text{tor}} \varphi) \sin^2(n_{\text{pol}} \theta) \\
    &\times \sin^2\left(\pi n_{\text{rad}} (\hat{r} - \hat{r}_{\text{min}})/(\hat{r}_{\text{max}} - \hat{r}_{\text{min}})\right) e^{-\left(\hat{v}_\parallel^2 + \hat{\mu} B\right)} , \\
    A_{1\,\parallel\text{MMS}}(\hat{r}, \varphi, \theta, \hat{t}) &:= \phi_{1\,\text{MMS}}(\hat{r}, \varphi, \theta, \hat{t}) := 2 \cos^2(\hat{\omega} \hat{t}) \cos(n_{\text{tor}} \varphi) \sin(2n_{\text{pol}} \theta) \\
    &\times \sin\left(2\pi n_{\text{rad}} (\hat{r} - \hat{r}_{\text{min}})/(\hat{r}_{\text{max}} - \hat{r}_{\text{min}})\right),
\end{align*}
\]

representing a mode. We run the verification in three different magnetic geometries testing different aspects of the implementation. For the first test we choose a slab geometry with the magnetic field specified in Eq. (4.52). In slab geometry multiple terms in the Vlasov equations like the curvature or \(\nabla B\) drift vanish. Furthermore, the magnetic field is uniform and points in the \(\varphi\) direction. Therefore, the magnetic field is automatically aligned to the coordinate system and the locally field-aligned coordinate system coincides with the intrinsic coordinate system of the manifold. This renders the field line tracing trivial and thus enables us to test the implementation without complications arising from the locally field-aligned coordinate system. For the second test we choose a circular geometry with the magnetic field specified in Eq. (4.53) and a constant \(q\) profile \(q(\hat{r}) = q_0\). In the circular geometry the magnetic field lines are twisted and the field line tracing as well as the interpolation is tested. For both the slab and circular geometry we use a large aspect ratio assumption such that \(\varphi\) represents a Cartesian coordinate labelling poloidal planes. Third, we test a toroidal magnetic equilibrium that incorporates all terms in the gyrokinetic Vlasov equation. The magnetic field is given by

\[
\hat{B} = \frac{\hat{B}}{R_0 + \hat{r} \cos(\theta)} \left( e_\varphi + \frac{\hat{r}}{q(\hat{r})} e_\theta \right),
\]

with the \(q\) profile

\[
q(\hat{r}) = q_0 + \alpha \left(\frac{\hat{r}}{\hat{a}}\right)^2,
\]

minor radius \(a\) and major radius \(R_0\). We run the MMS analysis with the parameters given in Table 6.1 with four different resolutions presented in Table 6.2. The radial box size is limited to \(\hat{r}_{\text{min}} = 0\) and \(\hat{r}_{\text{max}} = 1\) in slab geometry and \(\hat{r}_{\text{min}} = 0.1\) and \(\hat{r}_{\text{max}} = 0.5\) in both circular and toroidal geometry. In velocity space \((\hat{v}_\parallel, \hat{\mu})\), we choose a domain size of \([-3,3] \times [0,9]\) and in the toroidal angle \(\varphi\) of \([0,2\pi]\). We calculate one period of the manufactured solution up to \(\hat{t} = 1\). We analyse the results by calculating the relative \(L_2\) and \(L_\infty\) error of the numerical solution given by \(\|f - f_{\text{MMS}}\|_p/\|f_{\text{MMS}}\|_p\) where \(p \in \{2, \infty\}\). The results are shown in Fig. 6.2.

We observe that the numerical scheme converges to second order for all three geometries. Initially, the convergence is faster than second order because the linear advection terms in the
\[ N_\varphi \quad \Delta \hat{r} = \Delta \hat{z} \quad N_{v \parallel} \quad N_\mu \quad \Delta \hat{t} \]

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<td>0.003125</td>
<td>80</td>
<td>40</td>
<td>0.00125</td>
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</table>

Table 6.2.: Table displaying the different resolutions used for the method of manufactured solutions.

Gyrokinetic Vlasov equation are discretised with a fourth order scheme. For higher resolutions the numerical error introduced by the second order discretisation of the field equations and the nonlinear Poisson brackets starts to dominate, and we observe the expected second order convergence.

6.3. Screw pinch studies

After having demonstrated the convergence of the numerical scheme, we verify if the code reproduces the correct physical results. The starting point is a physical test where we measure growth rates of an ion temperature gradient mode in a screw pinch geometry. This section has been published in [87].

We use a setup similar to the work in [158, 83]. We model the screw pinch with the circular geometry, described in the last section, with a constant \( q \) profile \( q(r) = q_0 \). We simulate two different cases. In the first case we choose \( q_0 \to \infty \), corresponding to a slab like geometry\(^1\), and in the second case we choose a finite \( q \). This allows us to compare the performance of ordinary field-aligned coordinates, in the slab case, to the locally field-aligned coordinate system with interpolation for the finite \( q \) case. We choose the reference length parameters \( L_{\text{ref}} = 7.74 \cdot 10^{-1} \) m, \( B_{\text{ref}} = 1 \) T, \( m_{\text{ref}} = 1 \) u and \( T_{\text{ref}} = 1 \) keV such that the ion gyroradius \( \rho_{\text{ref}} = 3.23 \cdot 10^{-3} \) m and \( L_{\text{ref}}/\rho_{\text{ref}} = 239 \). The simulation box ranges from \( \hat{r}_{\text{min}} = 4.17 \cdot 10^{-3} \) to \( \hat{r}_{\text{max}} = 6.05 \cdot 10^{-2} \) spanning over \( 14.5 \rho_{\text{ref}} \). In velocity space \((\hat{v}_{\parallel}, \hat{\mu})\) we choose a domain size of \([-4, 4] \times [0, 16]\) and in the toroidal angle \( \varphi \) of \([0, 2\pi]\). We use the trapezoidal integration weights and the quadratic grid in \( \mu \), described in Section 4.2, and no hyperdiffusion as well as no buffer zone. We use a realistic mass ratio with \( m_e = 1/3600 m_i \).

We initialise the simulation with local Maxwellians, presented in Eq. (4.20), and given temperature \( T_{0e} \) and density profiles \( n_{0e} \). Maxwellians are an exact equilibrium of the gyrokinetic Vlasov equation in the given geometry and thus provide a stable initial condition. In order to study growth rates, we initialise a mode as a perturbation of the Maxwellian

\[
\hat{f}_\sigma(\hat{t} = 0, \hat{r}, \theta, \varphi, \hat{v}_{\parallel}, \hat{\mu}) = \hat{F}_{0\sigma}(\hat{r}, \theta, \varphi, \hat{v}_{\parallel}, \hat{\mu}) \\
\times \left( 1 + \epsilon \exp \left( -\left( \frac{\hat{r} - \hat{r}_{\text{mid}}}{\delta \hat{r}_{\text{mode}}} \right)^2 \right) \cos (m\theta + n\varphi) \right). 
\]

We choose the profiles \( \hat{T}_{0i}, \hat{T}_{0e}, \hat{n}_{0i} \) and \( \hat{n}_{0e} \), appearing in the Maxwellian distribution function,

\(^1\)Numerically we implement \( q_0 \to \infty \) by setting \( q_0 = 10^8 \).
Figure 6.2.: Figure displaying the relative $L_2$ (top) and $L_{\infty}$ (bottom) error of the numerical solution in the MMS test as a function of the resolution. The resolution is represented by the number of poloidal planes. The complete resolutions are specified in Tab. 6.2. The test is performed for a slab, circular and toroidal geometry. For reference a dashed line representing second order convergence is drawn in black. The numerical scheme converges to second order for all geometries.
according to

\[ P(\hat{r}) = C_P \exp \left( -\kappa_p \delta \hat{r}_p \tanh \left( \frac{\hat{r} - \hat{r}_{\text{mid}}}{\delta \hat{r}_p} \right) \right). \]  

(6.17)

The parameters for the profile and the initial condition are given in Table 6.3. The initial density profile is chosen to be constant by setting \( \kappa_n = 0 \). The profiles are depicted in Figure 6.3.

In order to compare the growth rates to analytical predictions, we use an approximate dispersion relation derived in [83, Appendix]. The dispersion relation has been derived with the assumption of adiabatic electron response and large aspect ratio, i.e. \( r/(qR_0) \to 0 \). Both approximations are valid for the parameters chosen and hence we expect the numerical results to be close to the prediction of the dispersion relation. In order to present the dispersion relation, we need to define a single Fourier mode of \( \hat{\phi}_1 \):

\[ \hat{\phi}_1 = \hat{\phi}_{1,m,n,\omega} \exp \left( i(m\theta + n\varphi - \hat{\omega}t) \right). \]  

(6.18)

| \( \epsilon \)   | \( 1.00 \cdot 10^{-4} \) |
| \( \hat{r}_{\text{mid}} \) | \( 3.05 \cdot 10^{-2} \) |
| \( \delta \hat{r}_{\text{mode}} \) | \( 5.90 \cdot 10^{-3} \) |
| \( C_{n_i} = C_{n_e} \) | \( 1.65 \cdot 10^1 \) |
| \( C_{T_i} = C_{T_e} \) | \( 1.00 \) |
| \( \delta \hat{r}_{T_i} = \delta \hat{r}_{T_e} \) | \( 6.05 \cdot 10^{-3} \) |
| \( \delta \hat{r}_{n_i} = \delta \hat{r}_{n_e} \) | \( 1.21 \cdot 10^{-2} \) |
| \( \kappa_{T_i} = \kappa_{T_e} \) | \( 6.62 \cdot 10^1 \) |
| \( \kappa_{n_i} = \kappa_{n_e} \) | \( 0 \) |

Table 6.3.: Table defining the parameters for the screw pinch profile for ions and electrons specified in Equation (6.17). This table has been published in [87].
The dispersion relation reads [83, Appendix]

\[ \frac{L_{\text{ref}}}{\rho_{\text{ref}}} \left[ \frac{\partial^2}{\partial \hat{r}^2} + \left( \frac{1}{\hat{r}} + \frac{1}{\hat{n}_0} \frac{\partial}{\partial \hat{r}} \right) \frac{\partial}{\partial \hat{r}} - \frac{m^2}{\hat{r}^2} \right] \hat{\phi}_{1,m,n,\omega}(\hat{r}) = \]

\[ \left\{ - \frac{1}{T_{0i}} (1 + z Z(z)) - \frac{1}{k_{\|} B_0 L_{\text{ref}}} \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right. \]

\[ \left. \left[ Z(z) \left( \frac{1}{\hat{n}_0} \frac{\partial}{\partial \hat{r}} - \frac{1}{2 T_{0i}} \frac{\partial}{\partial \hat{r}} \right) + z (1 + z Z(z)) \frac{1}{T_{0i}} \frac{\partial}{\partial \hat{r}} \right] \right\} \]

\times \hat{\phi}_{1,m,n,\omega}(\hat{r}), \quad (6.19) \]

with \( z = \hat{\omega}/\hat{k}^* \), \( \hat{k}^* = k_{\|} \sqrt{2 T_{0i}} \) and the parallel wavenumber \( k_{\|} = (m/q + n)b_\phi \). Furthermore, we used the plasma dispersion function

\[ Z(u) = \frac{1}{\sqrt{\pi}} \int_{-\infty}^{\infty} \exp(-x^2) \frac{dx}{x - u} = i \sqrt{\pi} \exp(-u^2)(1 + \text{erf}(iu)) \]

\[ \text{erf}(x) = \frac{2}{\sqrt{\pi}} \int_{0}^{x} \exp(-t^2) \, dt. \quad (6.20) \]

The dispersion relation is a nonlinear, differential eigenvalue problem that has to be solved for the radial electrostatic profile \( \hat{\phi}_{1,m,n,\omega} \) and the complex eigenvalue \( \hat{\omega} \) simultaneously. As we are not aware of methods to solve the equation analytically, we employ a numerical method. First, we discretise the differential operators to obtain an ordinary nonlinear eigenvalue problem. Similar to the numerical scheme presented in Chapter 4, we choose fourth order centred finite differences to discretise the \( r \) derivative. At the boundary, we implement Dirichlet boundary conditions and set \( \hat{\phi}_{1,m,n,\omega}(\hat{r}_{\min}) = \hat{\phi}_{1,m,n,\omega}(\hat{r}_{\max}) = 0 \). We choose 128 points for the discretisation.

The remaining nonlinear eigenvalue problem can be solved by calculating the roots of the characteristic polynomial. We use a determinant free method based on the Newton iteration described in [159, pp. 28–29]. As an initial guess for the Newton iteration we choose \( \omega_{\text{init}} = -2 + 4i \). The calculation is done with Mathematica [157]. The results are presented in Appendix B.2.

The dispersion relation and thus the eigenvalues \( \hat{\omega} \) are a function of the poloidal mode number \( m \) and the parallel wavenumber \( k_{\|} \). Hence, the eigenvalues \( \hat{\omega} \) do not depend on \( q \) explicitly. For a given poloidal mode number \( m \), we can select two toroidal mode numbers \( n_{\text{slab}} \) and \( n_{\text{screw}} \), for the \( q = \infty \) and \( q \neq \infty \) case respectively, such that the parallel wavenumbers and, consequently, the growth rates are equal. For the slab case the poloidal magnetic field vanishes, and the parallel wavenumber reads \( k_{\|,\text{slab}} = n_{\text{slab}} \). For the \( q \neq \infty \) case we choose the toroidal mode number \( n_{\text{screw}} \) such that \( k_{\|,\text{screw}} = k_{\|,\text{slab}} \), i.e.

\[ n_{\text{screw}} = k_{\|,\text{screw}} - \frac{b_\phi m}{r} = n_{\text{slab}} - \frac{m}{q}. \quad (6.21) \]
Table 6.4.: Table displaying the different initial mode configurations for the simulation. On the left, the mode numbers for the \( q = \infty \) case and on the right, the mode numbers for the \( q \neq \infty \) case are shown. The poloidal and toroidal mode numbers are constructed such that the parallel mode number \( k_\parallel \) is equal to one in all cases. This table has been published in [87].

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<th>( q )</th>
<th>( m )</th>
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<th>( \hat{k}_\parallel )</th>
<th>( q )</th>
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<td>( \infty )</td>
<td>10</td>
<td>1</td>
<td>1</td>
<td>10/3</td>
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<td>( \infty )</td>
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<td>1</td>
<td>1</td>
<td>15/3</td>
<td>15</td>
<td>-2</td>
<td>1</td>
</tr>
<tr>
<td>( \infty )</td>
<td>20</td>
<td>1</td>
<td>1</td>
<td>20/3</td>
<td>20</td>
<td>-2</td>
<td>1</td>
</tr>
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</table>

Having two setups with the same growth rates allows us to compare the performance of field-aligned coordinates in the \( q = \infty \) case to the locally field-aligned coordinates in the \( q \neq \infty \) case.

The \( q \) factors and mode numbers chosen for the simulation are shown in Table 6.4. They are selected such that a broad range of modes and \( q \) factors are tested.

### 6.3.1. Growth rates

We measure the growth rate from the \( L_2 \) norm of the electrostatic potential. The \( L_2 \) norm is calculated numerically with the trapezoidal quadrature rule similar to the charge density in the quasi-neutrality equation. We determine the growth rate by performing a linear fit of \( \log ||\hat{\phi}_1|| \) between the times \( \hat{t} = 1 \) and \( \hat{t} = 3.5 \). This is shown exemplarily in Fig. 6.4.

We use a resolution of \( \Delta \hat{x} = \Delta \hat{z} = 6.46 \cdot 10^{-4} \), corresponding to \( 0.2 \rho_{\text{ref}} \), \( n_\varphi = 16 \), \( n_{v_\parallel} = 256 \), \( n_\mu = 16 \) and a timestep of \( \Delta t = 1.4 \cdot 10^{-3} t_{\text{ref}} = 3.5 \text{ns} \). The timestep is constant throughout the simulation and limited by the parallel electron streaming. We study the convergence of the simulation at the example of the \( q = 15/3 \) case. Upon doubling the resolution in each dimension, \( \Delta \hat{x} \rightarrow 0.5 \Delta \hat{x}, \Delta \hat{z} \rightarrow 0.5 \Delta \hat{z}, n_\varphi \rightarrow 2n_\varphi, n_{v_\parallel} \rightarrow 2n_{v_\parallel} \) and \( n_\mu \rightarrow 2n_\mu \), the growth rate of the \( q = 15/3 \) mode changes less than 1%. Therefore, we assume a 1% error on the growth rates obtained from our simulations.

The growth rates obtained from the simulation as well as the solution of the approximate dispersion relation are shown in Fig. 6.5. The simulations for \( q = \infty \) and \( q \neq \infty \) agree well with each other within the 1% error margin, as predicted by the dispersion relation. Furthermore, the numerical results are close to the solution of the dispersion relation. They agree within the 1% numerical error margin for the poloidal mode numbers \( m = 15 \) and \( m = 20 \). For the poloidal mode numbers \( m = 5 \) and \( m = 10 \) the measured growth rates are slightly higher than growth rates obtained from the dispersion relation. As discussed above, the dispersion relation approximates the growth rates and thus exact agreement is not expected.

### 6.3.2. Energy and particle conservation

In order to study the behaviour of the nonlinear phase of the simulation, we investigate the conservation of the particle number and energy during the onset of turbulence. We choose
Figure 6.4.: Figure displaying the $L_2$ norm of the electrostatic potential as a function of time. The growth rate is obtained by performing a linear fit of $\log \| \phi_1 \|_2$ between the times $\hat{t} = 1$ and $\hat{t} = 3.5$. The fit is drawn in a blue dashed line. Shown are the results from the simulation with $q = 15/3$ and $m = 15$. This figure has been published in [87].

Figure 6.5.: Figure displaying the growth rates obtained from the approximate dispersion relation and the simulations with different $q$ factors and mode numbers specified in Table 6.4. Vertical lines through the markers represent the error of the simulation. The measured growth rates are close to the prediction of the dispersion relation. This figure has been published in [87].
the same screw pinch setup as presented in the last section and run the \( q = 15/3 \) case. We use a resolution of \( \Delta \hat{x} = \Delta \hat{z} = 3.23 \cdot 10^{-4} \), corresponding to \( 0.1 \rho_{\text{ref}}, n_{\varphi} = 32, n_{v_{\parallel}} = 64 \) and \( n_{\mu} = 16 \). We run the simulation until \( \hat{t} = 10 \), when the turbulence starts to hit the wall. The chosen Dirichlet boundary conditions allow an energy flux through the domain boundaries, which becomes effective when eddies hit the wall. As an example, the ion density is depicted in the linear and nonlinear phase in Fig. 6.6. The energy of the ions and electrons is shown in Fig. 6.7. The number of particles is conserved up to a precision of \( 10^{-7} \). The energy is conserved up to a precision of \( 10^{-8} \).† In the linear phase the conservation is improved. As soon as the nonlinear phase starts energy gets transferred between electrons and ions.

### 6.4. Alfven waves in slab geometry

We continue with the physical verification of the electromagnetic part of the model. In this section, we compare the frequency and damping rate of Alfven waves in slab geometry between numerical simulation with GENE-X and analytical estimates, following the work [132, 97].

#### 6.4.1. Derivation of the dispersion relation

We start by deriving a dispersion relation that provides the analytical estimates for the frequencies and damping rates of the Alfven waves. The dispersion relation is similar to the one used in [132, 97]. We perform the derivation explicitly in order to control the assumptions made and obtain the correct equation in our normalisation. Electrons are, due to their light weight, more mobile than ions and the main generator of electromagnetic fields. Therefore, we assume that the ions are stationary, i.e. the frequency of the Alfven waves fulfills \( \omega_{\text{Alfven}} \gg k_{\parallel} v_{T_{i}} \). Further, we employ a \( \delta f \) splitting such that \( \hat{f}_{e} = \delta \hat{f}_{e} + \hat{f}_{M} \) with

\[
\hat{f}_{M}(\hat{v}_{||}) = \frac{1}{\sqrt{\pi}} e^{-\hat{v}_{||}^{2}},
\]

(6.22)

We neglect all nonlinear terms. We choose two spatial dimensions \( x \) and \( y \). In compliance with the locally field aligned coordinate system presented in Chapter 3, \( y \) labels the direction parallel to the magnetic field. The normalisation is chosen, as introduced in Section 4.1, such that \( \hat{m}_{e} = 1 \) and \( \hat{q}_{e} = -1 \). The normalised gyrokinetic Vlasov equation for the electrons, only taking the parallel dynamics into account, reads

\[
\frac{\partial \delta \hat{f}_{e}}{\partial \hat{t}} = -\sqrt{\frac{2}{\hat{m}_{e}}} \left( \frac{\partial }{\partial \hat{y}} \left( \frac{\partial \delta \hat{f}_{e}}{\partial \hat{y}} + \hat{q}_{e} \left( \frac{\partial \hat{\phi}_{1}}{\partial \hat{y}} + \frac{\partial \hat{A}_{1\parallel}}{\partial \hat{y}} \right) \right) \hat{F}_{M} \right),
\]

(6.23)

†The energy conservation of \( 10^{-8} \), presented here, is significantly better than the result published in [87]. The reason is that the factor \( 1/2 \) in front of the \( q_{e} \phi_{1} \) term in Eq. (2.99) was missing in the energy diagnostic of the GENE-X code. This resulted in an overestimation of the electrostatic energy contribution and, consequently, an incorrect energy measurement.
Figure 6.6.: Figure displaying the ion density in the poloidal plane located at $\varphi = \pi$ in the linear phase at $t = 3.5$ and the nonlinear phase at $t = 6.5$ of a screw pinch simulation with $q = 15/3$. The simulation is initialised with the temperature and density profile shown in Fig. 6.3 and perturbed with an unstable $m = 15$, $n = -2$ mode. This figure has been published in [87]. For a unified presentation, the color bars and font sizes have been adapted.
Figure 6.7.: Figure displaying the normalised energy $E(t)/E(0)$ for electrons and ions of a simulation in screw pinch geometry with $q = 15/3$. The simulation is initialised with the temperature profile shown in Figure 6.3 and perturbed with an unstable $m = 15$, $n = −2$ mode. The simulation develops turbulence at approximately $\hat{t} = 4$. The total energy is conserved to a precision of $10^{-8}$. This figure has been published in a similar form in [87]. The origin of the difference is explained in the footnote on page 86.

with the corresponding field equations

$$- \left( \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right)^2 \frac{\partial^2 \hat{\phi}_1}{\partial \hat{x}^2} = \hat{q}_e \int_{-\infty}^{\infty} \delta \hat{f}_e \, d\hat{v}_\parallel \tag{6.24}$$

$$- \left( \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right)^2 \frac{\partial^2 \hat{A}_\parallel}{\partial \hat{x}^2} = \hat{q}_{\text{ref}} \sqrt{\frac{2}{m_e}} \int_{-\infty}^{\infty} \hat{v}_\parallel \delta \hat{f}_e \, d\hat{v}_\parallel. \tag{6.25}$$

Similar to Eq. (6.18), we switch to a Fourier representation of a single mode

$$\delta \hat{f}_e (\hat{x}, \hat{y}, \hat{v}_\parallel, \hat{t}) = \delta \hat{f}_{k_x, k_y} (\hat{x}, \hat{v}_\parallel) e^{i(k_x \hat{x} + k_y \hat{y})} \tag{6.26}$$

$$\hat{\phi}_1 (\hat{x}, \hat{y}, \hat{t}) = \hat{\phi}_{k_x, k_y} (\hat{t}) e^{i(k_x \hat{x} + k_y \hat{y})} \tag{6.27}$$

$$\hat{A}_\parallel (\hat{x}, \hat{y}, \hat{t}) = \hat{A}_{k_x} (\hat{y}, \hat{t}) e^{i k_x \hat{x}}. \tag{6.28}$$

Inserting Eqs. (6.26), (6.27) and (6.28) into Eqs. (6.23), (6.24) and (6.25) gives the linear system

$$-i \omega \delta \hat{f}_e = - \sqrt{\frac{2}{m_e}} \hat{v}_\parallel \left( i \hat{k}_y \delta \hat{f}_e - \hat{q}_e \left( i \hat{k}_y \hat{\phi}_1 + \frac{\partial \hat{A}_\parallel}{\partial \hat{t}} \right) \right) \hat{F}_M \tag{6.29}$$

$$\left( \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right)^2 \hat{k}_x^2 \hat{\phi}_1 = \hat{q}_e \int_{-\infty}^{\infty} \delta \hat{f}_e \, d\hat{v}_\parallel \tag{6.30}$$

$$\left( \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right)^2 \hat{k}_z^2 \frac{\partial \hat{A}_\parallel}{\partial \hat{t}} = i \omega \hat{q}_{\text{ref}} \frac{1}{2} \sqrt{\frac{2}{m_e}} \int_{-\infty}^{\infty} \hat{v}_\parallel \delta \hat{f}_e \, d\hat{v}_\parallel. \tag{6.31}$$

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We define \( \bar{\omega} = \sqrt{m_e/2\omega /k_y} \) and rewrite Eq. (6.29)

\[
\delta \dot{f}_e = -\frac{i \hat{v}_\| \hat{q}_e (i \frac{\partial A_\|}{\partial t} - \hat{\phi}_1)}{\hat{v}_\| - \bar{\omega}} F_M.
\]  

(6.32)

Inserting Eq. (6.32) into Eq. (6.30) yields

\[
\hat{\phi}_1 = \left( \frac{L_{\text{ref}}}{\rho_{\text{ref}}} \right)^2 \frac{1}{k_z^2} \left( \int_{-\infty}^{\infty} \frac{v}{\bar{\omega}} F_M(v) \, dv \right) \left( i \frac{\partial A_\|}{\partial t} - \hat{\phi}_1 \right)
\]

\[
= \left( \frac{L_{\text{ref}}}{\rho_{\text{ref}}} \right)^2 \frac{1}{k_z^2} \left( 1 + \bar{\omega} Z(\bar{\omega}) \right) \left( i \frac{\partial A_\|}{\partial t} - \hat{\phi}_1 \right),
\]

(6.33)

where we have used the plasma dispersion function and Eq. (A.19). Similarly, we insert Eq. (6.32) into Eq. (6.31)

\[
\frac{\partial A_\|}{\partial t} = -i \omega \left( \frac{L_{\text{ref}}}{\rho_{\text{ref}}} \right)^2 \frac{1}{k_z^2} \frac{2}{\bar{m}_e} \left( \int_{-\infty}^{\infty} \frac{v^2}{\bar{\omega}} F_M(v) \, dv \right) \left( i \frac{\partial A_\|}{\partial t} - \hat{\phi}_1 \right)
\]

\[
= -i \omega \left( \frac{L_{\text{ref}}}{\rho_{\text{ref}}} \right)^2 \frac{1}{k_z^2} \frac{2}{\bar{m}_e} \left( \bar{\omega} (1 + \bar{\omega} Z(\bar{\omega})) \right) \left( i \frac{\partial A_\|}{\partial t} - \hat{\phi}_1 \right),
\]

(6.34)

where we used Eq. (A.20). Subtracting Eq. (6.33) from \( i \hat{k}_y \) times Eq. (6.34) gives the final dispersion relation

\[
\hat{k}_y^2 \left( \frac{L_{\text{ref}}}{\rho_{\text{ref}}} \right)^2 + (1 + \bar{\omega} Z(\bar{\omega})) \left( 1 - \frac{\beta_{\text{ref}} \bar{\omega}^2}{\bar{m}_e} \right) = 0.
\]

(6.35)

We solve the dispersion relation numerically with the `FindRoot` function of Mathematica [157]. We choose \( \bar{\omega} = 1 - 0.1i \) as the starting value of the iterative algorithm. The results for different mode numbers and plasma beta are given in Appendix B.4.

### 6.4.2. Numerical simulation

We continue with describing the setup of the numerical simulation with GENE-X. We choose the reference parameters \( L_{\text{ref}} = 1.0 \text{m}, \) \( T_{\text{ref}} = 1.0 \text{KeV} \) and \( B_{\text{ref}} = 3.231 \times 10^{-3} \text{T} \) such that \( \rho_{\text{ref}} = L_{\text{ref}} = 1 \text{m} \). We choose a realistic mass ratio with \( \hat{m}_i = 1 \) and \( \hat{m}_e = 1/3600 \). We simulate in slab geometry and use the magnetic field described in Eq. (4.52). In slab geometry, the toroidal angle \( \varphi \) is interpreted as a Cartesian coordinate and coincides with the locally field-aligned \( y \) coordinate. We choose a box size of \( (\hat{x}, \hat{y}, \hat{z}, \hat{v}_\|, \hat{\mu}) \in [0, 2\pi] \times [0, 2\pi] \times [0, 2\pi] \times [-3, 3] \times [0, 9] \). The initial condition is given by

\[
\hat{f}_i(\hat{t} = 0, \hat{x}, \hat{y}, \hat{z}, \hat{v}_\|, \hat{\mu} = \pi^{-3/2} e^{-i \hat{v} \hat{\mu}})
\]

\[
\hat{f}_e(\hat{t} = 0, \hat{x}, \hat{y}, \hat{z}, \hat{v}_\|, \hat{\mu} = \pi^{-3/2} e^{-i \hat{v} \hat{\mu}} \cdot \left( 1 + \epsilon \cos (\hat{k}_y \hat{y}) \sin (\hat{k}_x \hat{x}) \right).
\]

We choose the strength of the mode perturbation as \( \epsilon = 10^{-2} \). We simulate with a constant parallel mode number \( \hat{k}_y = 1 \). The orthogonal mode number is varied from \( \hat{k}_x \in \)
Figure 6.8.: Figure displaying the time trace of the logarithm of the $L_2$ norm of the electrostatic potential for $\beta_{\text{ref}}/\hat{m}_e = 0.2$ and $\hat{k}_x = 1/4$. The red crosses indicate local maxima and the green crosses local minima. The damping rate is determined by a linear fit through the local maxima, drawn as a red dashed line.

{1/128, 1/64, 1/32, 1/16, 1/8, 1/4, 1/2, 1}. The distribution function is constant along $z$ and constant on the boundary. For the electromagnetic fields we choose periodic boundary conditions in $z$ and Dirichlet boundary conditions in $x$. We set the electromagnetic fields to zero for $x = 0$ and $x = 2\pi$. Due to the periodic boundary condition, there is no dynamic in the $z$ direction and the system is effectively two dimensional, as in Eq. (6.35). With the mode perturbation in $x$ we set the perpendicular mode in the electrostatic field $\hat{\phi}_1$. We use a resolution of $\Delta\hat{x} = \Delta\hat{z} = 0.05$, $n_\varphi = 32$, $n_{\hat{\phi}_1} = 64$, $n_{\hat{\mu}} = 16$ and a constant timestep of $\Delta\hat{t} = 1 \cdot 10^{-3}$. We use no hyperdiffusion and no buffer zone.

We measure the frequency and damping rate of the Alfven wave at the $L_2$ norm of the electrostatic potential. An example of the time trace of the $L_2$ norm is shown in Fig. 6.8. To determine the damping rate and frequency, we search for local minima and maxima of the oscillatory time trace. We define a local maximum as a point $\hat{t}_i$ such that $\|\hat{\phi}_1\|_2(\hat{t}_i) > \|\hat{\phi}_1\|_2(\hat{t}_j) \forall j \in \{i-2, i-1, i+1, i+2\}$, i.e. $\|\hat{\phi}_1\|_2(\hat{t}_i)$ has to be larger than all of its four nearest neighbours. In a similar fashion, we define a local minimum by the condition $\|\hat{\phi}_1\|_2(\hat{t}_i) < \|\hat{\phi}_1\|_2(\hat{t}_j) \forall j \in \{i-2, i-1, i+1, i+2\}$ such that the $L_2$ norm is smaller than its four nearest neighbours. The matched local maxima and minima of the example are shown in Fig. 6.8 as crosses. We determine the growth rate of the oscillation as the slope of the linear fit through the local maxima. For the kinetic shear Alfven waves the growth rate is always negative and the waves are damped via Landau damping. Let $\{\hat{t}_{\text{min},i} | i \in \{1, \ldots N_{\text{min}}\}\}$ label the position of the minima. We determine the frequency of the oscillation as

$$\Re \hat{\omega} \approx \frac{1}{N_{\text{min}} - 1} \sum_{i=1}^{N_{\text{min}} - 1} \frac{\pi}{\hat{t}_{\text{min},i+1} - \hat{t}_{\text{min},i}}.$$  

(6.38)
6.4.3. Results

The results from the numerical simulation with GENE-X as well as from the solution of the dispersion relation are shown in Fig. 6.9. The numerical results agree well with the prediction of the dispersion relation for all plasma beta and mode numbers tested. For low perpendicular mode numbers and large plasma beta the damping of the Alfvén waves is very weak. In that case the induced electric field $\partial \hat{A}_1 / \partial \hat{t}$ is determined by the two integrals in Eq. (2.89). These integrals need to be discretised consistently such that numerical errors arising from the discretisation cancel and the physics is correctly resolved [97, Appendix. A]. This is also known as the Ampère cancellation problem [98, 99, 100]. As shown by the results, the consistent discretisation scheme used in GENE-X does not suffer from the cancellation problem.

6.5. Cyclone base case

We close this chapter by performing a well-studied CBC benchmark [156] that probes the full electromagnetic model in toroidal geometry. Following [156], we measure the growth rate and frequency of different toroidal modes at a fixed plasma $\beta$. We choose the reference parameters $L_{\text{ref}} = 1.67\, \text{m}$, $B_{\text{ref}} = 2.0\, \text{T}$, $m_{\text{ref}} = 1\, \text{u}$, $T_{\text{ref}} = 2.14\, \text{keV}$ and $n_{\text{ref}} = 4.66 \cdot 10^{19} \, \text{m}^{-3}$ such that the plasma $\beta$ is given by $\beta_{\text{ref}} = 1\%$. The reference ion gyroradius reads $\rho_{\text{ref}} = 3.34 \cdot 10^{-3} \, \text{m}$. The minor radius is given by $a = 0.60\, \text{m} = 0.36L_{\text{ref}}$. We choose a radial extent of $\hat{r}_{\text{min}} = 3.60 \cdot 10^{-2} = 0.1\hat{a}$ to $\hat{r}_{\text{max}} = 3.24 \cdot 10^{-1} = 0.9\hat{a}$. The magnetic field is given by the toroidal geometry described in Eq. (6.14) with the $q$ profile

$$q(\hat{r}) = 0.86 - 0.16 \left( \frac{\hat{r}}{\hat{a}} \right) + 2.52 \left( \frac{\hat{r}}{\hat{a}} \right)^2.$$  \hspace{1cm} (6.39)

In velocity space $(\hat{v}_\parallel, \hat{\mu})$ we choose a domain size of $[-4, 4] \times [0, 16]$ and in the toroidal angle $\varphi$ of $[0, 2\pi)$. We use a realistic mass ratio of $m_e = 1/3600m_i$. Similar to the screw pinch benchmark, we initialise the simulation with local Maxwellians, presented in Eq. (4.20), with the profiles $\hat{T}_{0i}$, $\hat{T}_{0e}$, $\hat{n}_{0i}$ and $\hat{n}_{0e}$, chosen according to Eq. (6.17). The parameters for the profiles are given in Tab. 6.5 and the profiles are shown in Fig. 6.10. This test poses challenges for GENE-X. As discussed in Section 4.5, the initial condition constructed with local Maxwellians is unstable and prevents us from studying the growth rate and frequency

$$\begin{array}{l|l}
\hat{r}_{\text{mid}} & 5.00 \cdot 10^{-1} \\
C_{ni} = C_{ne} & 1.00 \\
C_{Ti} = C_{Te} & 1.00 \\
\delta \hat{T}_{Ti} = \delta \hat{T}_{Te} & 3.00 \cdot 10^{-1} \\
\delta \hat{n}_{ni} = \delta \hat{n}_{ne} & 3.00 \cdot 10^{-1} \\
\kappa_{Ti} = \kappa_{Te} & 2.49 \\
\kappa_{ni} = \kappa_{ne} & 8.00 \cdot 10^{-1} \\
\end{array}$$

Table 6.5.: Table defining the parameters for the CBC profile for ions and electrons specified in Equation (6.17).
Figure 6.9.: Figure displaying the frequency $\text{Re} \hat{\omega}$ and the damping rate $-\text{Im} \hat{\omega}$ for different perpendicular mode numbers and different $\hat{\beta}_{\text{ref}}/\hat{m}_e$. The results obtained with the numerical calculation from GENE-X agree well with the results from the dispersion relation for all parameters tested.
of specific toroidal modes. Therefore, we remove the dynamics of the initial condition from the system by evaluating the right-hand side of the gyrokinetic Vlasov-Maxwell system on the initial condition and subtracting the result every timestep. Using this filtering process is similar to using the assumption that the $\nabla B$ drift of the background vanishes, which is usually made in $\delta f$ codes [139]. To trigger the growth of unstable modes, we perturb the initial distribution function with a sample $x$ from the uniform distribution on the interval $[0, 10^{-6}]$ according to

$$\hat{f}_\sigma \rightarrow \hat{f}_\sigma + x.$$  \hspace{1cm} (6.40)

We select modes with a fixed toroidal mode number by using a toroidal Fourier filter on the electrostatic potential and the parallel component of the electromagnetic vector potential. The dominant unstable modes in the prescribed system are either ion temperature gradient (ITG) or trapped electron modes (TEM) [156]. They are aligned to the magnetic field and obtain part of their toroidal structure through the poloidal rotation of the magnetic field. Using the FCI approach, the parallel dynamics is resolved via field line tracing and a high resolution within the poloidal plane. The toroidal resolution is usually sparse and of the order of $n_\varphi = 16$. This makes toroidal Fourier filtering difficult because the highest resolvable mode number is $n_\varphi/2^\dagger$. To filter higher toroidal mode numbers, the number of poloidal planes needs to be increased. The computational demand of the simulations increases quadratically with

\[^\dagger\text{At this point we would like to emphasise that only the Fourier filter and not the code itself is limited to modes with a mode number } n_\varphi/2. \text{ In the simulation ITGs and TEMs of significantly higher mode numbers are resolved. The reason for this is that the modes are field-aligned and most of the toroidal structure of the mode is generated by the poloidal rotation of the magnetic field. This rotation and the parallel structure of the modes along the field line is well resolved because the coordinate system is aligned to the magnetic field. For instance, in a simulation with } n_\varphi = 20 \text{ and a toroidal Fourier filter of } n = 5 \text{ we obtain the mode } n = 15. \text{ The mode } n = 15 \text{ is not covered by the filter because the number of poloidal planes is too low. It is still present in the system due to the high parallel resolution of FCI and grows faster than the mode } n = 5.\]
\( n_\varphi \) because, in addition to the increase in number of grid points, the timestep decreases proportional to the increase in \( n_\varphi \). Therefore, we restrict the analysis to the toroidal mode numbers \( n = 5, 10, 15 \). As discussed in Chapter 2, we solve the gyrokinetic Vlasov-Maxwell system in a long wavelength limit. The modes \( n = 5, 10, 15 \) correspond to perpendicular mode numbers \( k_\perp = nq(a/2)/(a/2) \) of \( k_\perp \rho_{\text{ref}} = 0.178, 0.157, 0.235 \). These are significantly smaller than \( k_\perp \rho_{\text{ref}} = 1 \) such that a good agreement between the literature and the simulations performed with GENE-X can be expected.

We measure the growth rate and the frequency from the maximum of the parallel component of the electromagnetic vector potential on the poloidal plane located at \( \varphi = \pi \). Similar to the screw pinch benchmark, we determine the growth rate by performing a linear fit of \( \log \max_{\hat{x},\hat{z}}(\hat{A}_\parallel(x, z, \varphi = \pi)) \). We calculate the frequency by subtracting the fit from the signal, measuring the position of two minima of the oscillations \( \{\hat{t}_{\text{min},1}, \hat{t}_{\text{min},2}\} \) and using \( \text{Re} \hat{\omega} = 2\pi/(\hat{t}_{\text{min},2} - \hat{t}_{\text{min},1}) \). As GENE-X is a nonlinear code and the frequency is small in relation to the growth rate, we are in general not able to measure the position of more than two minima before the nonlinear phase starts.

**Growth rates and frequencies**

We use a resolution of \( \Delta \hat{x} = \Delta \hat{z} = 3.34 \cdot 10^{-3} \) corresponding to \( 1.67\rho_{\text{ref}}, n_{v_\parallel} = 48 \) and \( n_\mu = 12 \). For simulations with the \( n = 5 \) toroidal filter we use \( n_\varphi = 32 \), for the \( n = 10 \) toroidal filter we use \( n_\varphi = 48 \) and for the \( n = 15 \) toroidal filter we use \( n_\varphi = 64 \). The timestep is constant throughout the simulation and reads \( \Delta t = 1 \cdot 10^{-3}t_{\text{ref}}, \Delta t = 0.5 \cdot 10^{-3}t_{\text{ref}} \) and \( \Delta t = 0.25 \cdot 10^{-3}t_{\text{ref}} \) for the three different toroidal filters.

We study the convergence of the simulation at the example of the simulations with the \( n = 10 \) and \( n = 15 \) toroidal filters. A higher toroidal mode number leads to a higher poloidal mode number and therefore requires a higher resolution within the poloidal plane. We increase the resolution of the two simulations to \( \Delta \hat{x} \rightarrow 0.7\Delta \hat{x}, \Delta \hat{z} \rightarrow 0.7\Delta \hat{z} \) and \( n_\varphi \rightarrow 2n_\varphi \) and measure the growth rate and frequency. We do not observe significant changes in the frequency upon increasing the resolution. The error of the frequency is limited by the measurement error of the minima of the oscillations. We observe significant changes in the growth rate. The growth rate for the \( n = 10 \) simulation increases by 10% from 0.186 to 0.205 and the growth rate of the \( n = 15 \) simulation by 30% from 0.271 to 0.356. Therefore, we assume an error of 10% on the growth rate of the \( n = 5 \) and \( n = 10 \) simulations and an error of 30% on the growth rate of the \( n = 15 \) simulation. To obtain more precise results for the \( n = 15 \) simulation, a higher resolution is needed, which is expensive due to the specific properties of the locally field-aligned coordinate system discussed above. The measured growth rates and frequencies are shown in Fig. 6.11. Both the growth rates and frequencies agree well with the reference results obtained with the GENE code [156].

Exemplarily, a cross section of the electrostatic potential and the parallel component of the electromagnetic vector potential for the simulations with a Fourier filter of the mode \( n = 10 \) are shown in Fig. 6.12. The structure of the \( n = 10 \) mode is clearly visible in both fields.
Figure 6.11.: Figure displaying the measured growth rate (top) and frequency (bottom) of modes with a different toroidal mode number. For reference, the results obtained with the GENE code [156] are drawn as a dashed line. The measured growth rates and frequencies agree well.
Figure 6.12.: Figure displaying the electrostatic potential (top) and the parallel component of the electromagnetic vector potential (bottom) on the poloidal located at $\varphi = \pi$ at the time $t = 80t_{\text{ref}}$ for the simulation with a toroidal Fourier filter of $n = 10$. 
6.6. Conclusion and outlook

In this chapter, we presented thorough tests of the GENE-X code. The test were designed to comprehensively probe different parts of the numerical scheme and their implementation in the code. We started by performing precise numerical tests to demonstrate that the chosen numerical scheme, in combination with the locally field-aligned coordinate system, is capable of solving the gyrokinetic Vlasov-Maxwell system. This included the measurement of the accuracy of the field line tracing and a comprehensive convergence order analysis with MMS. We demonstrated that the numerical scheme converges to second order accuracy. We proceeded with physically motivated benchmarks and probed the electrostatic part of the model by comparing growth rates in a screw pinch geometry against analytical predictions and tested the nonlinear regime by studying the conservation of energy. We tested the electromagnetic implementation by calculating the damping rates and frequencies of Alfén waves in slab geometry. Finally, we tested the whole gyrokinetic Vlasov-Maxwell system in toroidal geometry by performing a well studied CBC benchmark. GENE-X passed all tests.

In the future, it would be beneficial to perform further tests that probe the nonlinear regime. A suitable candidate would be a nonlinear version of the CBC benchmark [160]. Furthermore, it would be interesting to study the damping of zonal flows with the Rosenbluth-Hinton test [161, 160].
7. Simulations of the edge and scrape-off layer of ASDEX Upgrade

In this chapter, we demonstrate the full capability of the GENE-X code by performing simulations of the edge and SOL of ASDEX Upgrade. The simulations are performed in the diverted X-point geometry of the L-mode discharge #36190.

We present the results from three simulations at different physical parameters. The first simulation uses a reduced electron-to-ion mass ratio of $\mu = m_e/m_i = 1/400$, the second simulation a realistic electron-to-deuterium mass ratio of $\mu = 1/3600$ and the third simulation a realistic mass ratio and a collision operator. This allows us to study the effect of the mass ratio and collisions on edge and SOL turbulence. The simulations presented are performed over a timescale of 1ms until profile saturation. This chapter is structured as follows. First, we describe the setup of the simulation and discuss the parameters used. We proceed by presenting the final plasma density and temperature profiles and the radial electric field. The results are compared to Braginskii fluid [52] simulations, performed with the GRILLIX code [79, 80], and experimental measurements. Finally, we look at the power balance of the simulations and study the $E \times B$ heat flux over the separatrix and the parallel heat flux in the SOL. In particular, we measure the parallel heat flux onto the divertor and the SOL fall-off length $\lambda_q$ by performing an Eich fit [26] of the parallel heat flux profile.

7.1. Simulation setup

We use the magnetic equilibrium of the AUG discharge #36190 at the time $t = 3.3s$ for the simulations. The simulations are performed in the edge and SOL ranging from the flux surface $\psi = 0.9$ to $\psi = 1.05$. We use the canonical Maxwellian described in Section 4.5 to initialise the simulations. The initial condition is constructed to allow for comparison with previous Braginskii fluid simulations with GRILLIX [79, 80]. The profiles for electrons and ions, entering the canonical Maxwellian, are equal. The form of the profiles is given by the function

$$P(\psi) = \begin{cases} P_{\text{max}} & \text{if } \psi \leq \psi_{\text{min}} \\ c_1 \sin(c_2 \psi + c_3) + c_4 & \text{if } \psi_{\text{min}} < \psi < \psi_{\text{max}} \\ P_{\text{min}} & \text{else} \end{cases}$$

(7.1)
Figure 7.1.: Figure displaying the initial temperature and density profiles for the simulations. Ion and electron temperatures are equal. The functional form of the profile is a sine function described in Eq. (7.1). The measured initial profile, presented here, differs slightly from the analytical form, described in Eq. (7.1), because a canonical Maxwellian is used.

with the coefficients

\[ c_1 = \frac{(P_{\text{max}} - P_{\text{min}})}{2} \quad (7.2) \]
\[ c_2 = \frac{\pi}{(\psi_{\text{max}} - \psi_{\text{min}})} \quad (7.3) \]
\[ c_3 = \frac{\pi}{2} - c_2 \psi_{\text{min}} \quad (7.4) \]
\[ c_4 = \frac{(P_{\text{max}} + P_{\text{min}})}{2} \quad (7.5) \]

For both the temperature and density profile, we choose \( \psi_{\text{min}} = 0.92 \) and \( \psi_{\text{max}} = 0.995 \). In order to obtain a stable initial condition, it is important that the distribution function is constant in regions with \( \psi > 1 \) at the start of the simulation. On the core boundary \( \psi_{\text{min}} = 0.90 \) we choose a temperature of \( T_{\text{max}} = 350\text{eV} \) and a density of \( n_{\text{max}} = 2 \cdot 10^{19}\text{m}^{-3} \), coinciding with the experimental measurements of the ion temperature and density at AUG and the GRILLIX simulations. On the SOL boundary \( \psi_{\text{max}} = 1.05 \) we choose a density of \( n_{\text{min}} = 0.2 \cdot 10^{19}\text{m}^{-3} \) and a temperature of \( T_{\text{min}} = 35\text{eV} \). The density is selected according to experimental measurements. The temperature boundary value of 35eV is motivated by the available velocity space resolution. In gyrokinetic simulations the velocity information about the distribution function is retained. In the plasma core, the temperature is by orders of magnitude larger than in the SOL. Consequently, the dynamics of fast particles in the plasma core and slow particles in the SOL need to be resolved on the velocity space grid. In order to reduce the computational demand of the simulations, we limit the factor between the temperature at the core boundary and the SOL boundary to ten. To trigger the growth of unstable modes, we perturb the initial condition, according to Eq. (6.40), with a sample \( x \) from...
the uniform distribution on the interval \([0, 10^{-3}]\). The plasma profiles after initialisation with the canonical Maxwellian are depicted in Fig. 7.1. The initial ion density of the simulation in the poloidal plane is shown in Fig. 7.2.

As explained in Section 4.4, we use Dirichlet boundary conditions in the simulations. The boundary value of the distribution function in real space is equal to the initial condition and remains constant throughout the simulation. This implies that the temperature and density on the core boundary are fixed to 350eV and \(2.0 \cdot 10^{19} m^{-3}\), and the core boundary functions as heat and particle source. On the SOL boundary the temperature and density is fixed to 35eV and \(0.2 \cdot 10^{19} m^{-3}\), and the boundary functions as a heat and particle sink. By fixing the temperature at the SOL boundary to 35eV, we ensure that the temperature in the SOL does not reach values significantly below 35eV, and the dynamics is well resolved within the available velocity space resolution.

We choose the reference length \(L_{\text{ref}} = 1.65\text{m}\), corresponding to the major radius, the reference magnetic field \(B_{\text{ref}} = 2.5T\), corresponding to the on-axis magnetic field, \(m_{\text{ref}} = 1u\), corresponding to the mass of a hydrogen ion and \(T_{\text{ref}} = 0.050\text{keV}\), corresponding to the ion temperature in the near SOL. The reference gyroradius reads \(\rho_{\text{ref}} = 2.89 \cdot 10^{-4}\text{m}\) and \(L_{\text{ref}}/\rho_{\text{ref}} = 5.71 \cdot 10^3\). We use a resolution of \(\Delta x = \Delta z = 1.43\text{mm}\) in the poloidal plane which corresponds to roughly 1.9 ion gyroradii at a (separatrix) temperature of 100eV and a (outboard mid-plane) magnetic field of 2T. As the temperature and magnetic field vary over the device, so does the resolution in terms of local gyroradii. In velocity space \((\hat{v}_\parallel, \hat{\mu})\) we use different resolutions for the simulations with and without collisions. For the conservation of energy, particles and momentum in the simulations with collision operator, it is vital to increase the domain size in velocity space such that larger parts of the tail of distribution function are included. In the simulations without collisions, we use a resolution of \(n_{v_\parallel} = 72\) and \(n_\mu = 16\) and a domain size of \([-7, 7] \times [0, 49]\). In the simulations with collisions, we use a resolution of \(n_{v_\parallel} = 80\) and \(n_\mu = 16\) and a domain size of \([-8, 8] \times [0, 81]\). In the toroidal angle \(\varphi\), we use a resolution of \(n_\varphi = 24\) and a domain size of \([0, 2\pi]\), covering the whole torus. To study the convergence of the numerical results, we performed simulations at an in plane resolution of \(\Delta \hat{x} \rightarrow 0.7\Delta \hat{x}, \Delta \hat{z} \rightarrow 0.7\Delta \hat{z}\). We did not observe significant changes of the results.

The timestep is \(\Delta t = 6 \cdot 10^{-4} t_{\text{ref}} = 14.3\text{ns}\) and constant throughout the simulation. The simulation is run for 42 time units corresponding to 1ms of plasma evolution.

Close to the boundary and before the divertor plates we apply diffusion in a buffer zone. We set the parameter controlling the strength of the diffusion, introduced in Section 4.2, to \(\epsilon_{\text{buffer}} = 200\), which is close to the maximum value allowed by the CFL condition and the chosen timestep. Furthermore, we use hyperdiffusion in the dimensions \(x, y, z\) and \(v_\parallel\). The parameters controlling the strength of the hyperdiffusion, introduced in Section 4.2, are chosen as \(\epsilon_{\hat{x}, \hat{z}} = 5, \epsilon_{\hat{y}} = 0.06\) and \(\epsilon_{\hat{v}_\parallel} = 0.025\). The strength of the hyperdiffusion in \(x, y, z\) is estimated via trial and error. We use the estimation presented in [135] for the value of the hyperdiffusion in \(v_\parallel\). It is worth discussing the computational cost of the simulations. The simulations presented in this work were performed on the supercomputer Cobra of the Max-
Figure 7.2.: Figure displaying the ion density of the initial state of the simulation on the poloidal plane at $\varphi = \pi$. The separatrix is drawn in green, the device wall in blue and the divertor legs in grey. The distribution function is initialised according to Eq. (7.1) and constant in the SOL.
Planck Computing and Data Facility (MPCDF) and the A3 partition of the supercomputer Marconi of Cineca. Both systems feature compute nodes with Intel SkyLake Processors with 40 and 48 cores respectively. The cost for one simulation at the resolution presented is $2.5 \times 10^6$ CPU hours. We used 256 nodes for the simulations which required a runtime of approximately 9 days. Doubling the resolution within the poloidal plane quadruples the computational cost because the resolution is increased uniformly in the $x$ and $z$ direction. This means that a resolution of $\Delta x = \Delta z = 7.53 \times 10^{-4}$ mm, corresponding to resolving one thermal ion Larmor radius on the separatrix, increases the computational demand by a factor of roughly 3.6. Of further interest is a higher resolution in velocity space to reduce the temperature in the SOL. Upon decreasing the temperature by a factor of 2, the resolution requirement in $v_\parallel$ and $\mu$ increases by a factor of $\sqrt{2}$, and the total computational cost increases by a factor of $\sqrt{2^2} = 2$. Doubling the resolution in the $\phi$ direction leads to a quadrupling of the computational cost because the number of points increase and the timestep needs to be reduced at the same time. This means that an increase of the number of poloidal planes from $n_\phi = 24$ to $n_\phi = 32$ leads to increase in the computational demand by roughly 1.8.

Collision model

For the simulations with collisions, we use a simple collision model based on the Bhatnagar-Gross-Krook (BGK) collision operator [113]. The collision operator supports multi-species collisions and reads

$$\hat{C}_{\alpha\beta} \hat{f}_\alpha = \hat{\nu}_{\alpha\beta} \left( \frac{\hat{B}}{B_\parallel} \hat{M}_{\alpha\beta} - \hat{f}_\alpha \right), \quad (7.6)$$

$$\hat{M}_{\alpha\beta} = \frac{\hat{n}_\alpha}{(\pi T_{\alpha\beta})^{3/2}} \exp \left( -\frac{(v_\parallel - \hat{u}_{\alpha\beta})^2 + \hat{\mu} B}{T_{\alpha\beta}} \right), \quad (7.7)$$

where $\alpha, \beta$ denote the different species and $\hat{n}_\alpha$ the current plasma density. The coefficients $\hat{u}_{\alpha\beta}, T_{\alpha\beta}$ are chosen such that the collision operator conserves the particle number, parallel momentum, energy and Boltzmann’s $H$-Theorem on the continuous level [162]. In addition, the collision frequency $\hat{\nu}_{\alpha\beta}$ is chosen such that the temperature relaxation rates match the rates of the Boltzmann collision operator [162]. They read

$$\hat{u}_{\alpha\beta} = \frac{m_\alpha \hat{u}_\alpha + \sqrt{m_\beta m_\alpha} \hat{u}_\beta}{m_\alpha + m_\beta} \quad (7.8)$$

$$\hat{T}_{\alpha\beta} = \frac{\hat{T}_\alpha + \hat{T}_\beta}{2} + \frac{\hat{m}_\alpha \hat{m}_\beta}{3(m_\alpha + m_\beta)} \left( \frac{\hat{u}_\alpha}{\sqrt{m_\alpha}} - \frac{\hat{u}_\beta}{\sqrt{m_\beta}} \right)^2 \quad (7.9)$$

$$\hat{\nu}_{\alpha\beta} = \frac{n_{\text{ref}} L_{\text{ref}}}{T_{\text{ref}}^2} \frac{16\hat{n}_\beta \sqrt{2\pi \hat{m}_\alpha \hat{m}_\beta} (Z_\alpha Z_\beta e)^2}{3 \left( \hat{m}_\alpha \hat{T}_\beta + m_\beta \hat{T}_\alpha \right)^{3/2}} \ln \Lambda. \quad (7.10)$$

The Coulomb logarithm $\ln \Lambda$ is calculated at the beginning of the simulation according to model number four described in [163] and varies over the computational domain. For more
details on the collision operator in GENE-X we refer to [116]. The numerical scheme, presented in Section 4.2, does not preserve the positivity of the distribution function. The temperature \( \hat{T} \) can become negative and cause the collision operator to become imaginary. As the temperature and density get closer to zero, the collision frequency \( \hat{\nu} \) diverges. This requires the timestep of the simulation to go to zero. To prevent these problems, we use a density floor in the collision operator of \( n = 0.1 \cdot 10^{19} \text{m}^{-3} \) for the ion and electron density and \( T = 21 \text{eV} \) for the ion and electron temperature.

7.2. Plasma profiles and fluctuation amplitudes

We start the analysis of the AUG simulations by looking at the plasma density and the electrostatic potential of the final state at \( t = 1 \text{ms} \). We present the analysis at the example of the simulation with realistic mass ratio. The plasma density is shown in Fig. 7.3 and the electrostatic potential in Fig. 7.4. Compared to the initial condition, shown in Fig. 7.2, plasma density has been transported out radially and populates the whole confined region. Turbulent structures are dominant on the low field side and most prominent in the confined region. Turbulent streamers start close to the core boundary and end at the separatrix. The density in the SOL is significantly lower than in the confined region. On the outboard mid-plane (OMP) small density blobs are visible that have passed the separatrix and entered the SOL. The electrostatic potential in the confined region is dominated by zonal flows [164]. There is a rapid drop in the electrostatic potential close to the separatrix giving rise to a radial electric field that generates a poloidal rotation of the plasma along the separatrix. In all simulations there is little dynamics in the private flux region. This finding is consistent with the simulations of the discharge performed with GRILLIX. There are two possible explanations for the findings. First, the private flux region is cold and the Larmor radius of thermal ions small. Therefore, the in-plane resolution could not be high enough to resolve the small scale dynamics in the private flux region. A second explanation could be the absence of physical effects related to neutral gases in the simulation. Recent simulations performed with GRILLIX demonstrated turbulent structures in the private flux region when a neutral gas model was added [80].

In the following, we analyse different plasma profiles at the OMP. The profile data at the OMP is measured by interpolating the results along a line that starts at the magnetic axis and is orthogonal to the magnetic flux surfaces via cubic spline interpolation. The data for the density and temperature profiles is taken at \( t = 1 \text{ms} \) and averaged over all 24 poloidal planes. The fluctuation amplitude shown is calculated as the standard deviation of the profiles over the 24 poloidal planes. The average and standard deviation of the radial electric field is calculated including the last 100\( \mu \text{s} \) of the simulation. This averages over the fluctuations in the radial electric fields due to oscillations of the geodesic acoustic mode (GAM) [165].
Figure 7.3.: Figure displaying the ion density of the final state of the simulation on the poloidal plane at \( \varphi = \pi \). Similar to Fig. 7.2, the separatrix, wall and divertor is shown. Compared to the initial condition presented in Fig. 7.2, plasma density populates the whole confined region, and the dynamics is dominated by turbulence. In the SOL on the low field side density blobs are visible that have been transported over the separatrix.
Figure 7.4.: Figure displaying the electrostatic potential of the final state of the simulation on the poloidal plane at $\phi = \pi$. Similar to Fig. 7.2, the separatrix, wall and divertor is shown. In the confined region the dynamics is dominated by zonal flows that change their structure over time. Upon crossing the separatrix there is a significant drop in the electrostatic potential giving rise to a radial electric field.
7.2.1. Density

The density profiles are shown in Fig. 7.5. The measured average value of the density profile is similar for all three simulations performed. We observe a significantly larger fluctuation amplitude in the simulation with reduced mass ratio indicating a stronger turbulent drive. The density in the far SOL of the GENE-X simulations is limited by the boundary condition. The shape of density profiles is similar to the shape of density profile obtained with GRILLIX without neutral physics. The slope of the density profiles is shallower. The density in the SOL in the GRILLIX simulations is lower. This is because the density in the GRILLIX simulations is not fixed by the boundary condition. The shape of the measured density profile of the experiment differs significantly. GRILLIX simulations with neutrals reproduce the experimental measurement quite well. This could be a hint that neutrals are important for the formation of the density profile and should be included in gyrokinetic edge and SOL simulations.

7.2.2. Temperature

Following [52], we define the temperature of the species $\sigma$ as the kinetic energy of the plasma in a coordinate system in which the parallel flow $u_\sigma = 0$, i.e.

$$T_\sigma := \frac{2}{3n_\sigma} \int_0^\infty \int_{-\infty}^\infty \left( \frac{1}{2} m_\sigma \left( v_\parallel - \frac{u_\sigma}{n_\sigma} \right)^2 + \mu B \right) f_\sigma \, dW,$$  

(7.11)
Figure 7.6.: Figure displaying the mean ion temperature profiles measured at the outboard mid-plane of the three different GENE-X simulations performed. In addition, the fluctuation amplitude is drawn as a shaded area around the mean. For comparison, the ion temperature profile of GRILLIX simulations with and without neutrals is shown.

Figure 7.7.: Figure displaying the mean electron temperature profiles measured at the outboard mid-plane of the three different GENE-X simulations performed. In addition, the fluctuation amplitude is drawn as a shaded area around the mean. For comparison, the electron temperature profile of GRILLIX simulations with and without neutrals is shown.
where the expression for the density

\[ n_\sigma := \int_0^\infty \int_{-\infty}^{\infty} f_\sigma \, dW \]  

and parallel flow

\[ u_\sigma := \int_0^\infty \int_{-\infty}^{\infty} v_\parallel f_\sigma \, dW \]  

were used. The ion temperature profiles are shown in Fig. 7.6 and the electron temperature profiles in Fig. 7.7. We start with the analysis of the ion temperature profiles. In the confined region the ion temperature profiles of the simulations with realistic mass ratio and the GRILLIX simulation without neutrals agree well. The density of the simulation with reduced mass ratio is slightly increased. Similiar to the density profile, we observe that the fluctuation amplitude of the simulation with reduced mass ratio is increased. In the SOL we observe differences in the profiles. In the simulations with realistic mass ratio the temperature is slightly reduced. Collisions do not seem to have an impact on the ion temperature. Nevertheless, the ion temperature of all three simulations is in the range of 100eV–250eV and significantly larger than the temperature floor of 35eV. We proceed with analysing the electron temperature profile. In contrast to the ion temperature, none of the measured electron temperature profiles agree with the GRILLIX simulations. The profiles of the simulations without collision are close to each other. Again, the temperature of the simulation with reduced mass ratio is slightly increased and the fluctuation amplitude is larger. The electron temperature profile with collisions differs significantly from the simulations without collisions. It is significantly lower and reaches values close to the temperature boundary value of 35eV in the SOL. Its slope agrees well with the experimental measurement. Comparing the velocity space of the simulations with and without collisions, we observe that the electrons are efficiently de-trapped. The perpendicular kinetic energy is transferred to the parallel kinetic energy and efficiently absorbed by the divertor plates. As observed above, this process is not as efficient for the ions because the collision frequency is lower. A possible explanation for this finding is the temperature boundary condition of 35eV that limits the temperature in the SOL. The ion-ion collision frequency scales as \( T_i^{-3/2} \) and the ion-electron collision frequency approximately as \( T_e^{-3/2} \). This implies that decreasing the ion and electron temperature boundary condition to more realistic values of 2eV would allow both the ion-ion and ion-electron collisionality to increase by a factor 73. Hence, the energy transfer between ions and electrons would become more efficient resulting in lower SOL temperatures for the ions. These findings demonstrate that collisions are essential for efficient heat transport in the SOL.

### 7.2.3. Radial electric field

The profiles of the radial electric field are shown in Fig. 7.8. The profiles are approximately flat in the confined region. The main generator of the radial electric field in the confined region is given by GAMs. They change their structure over time and average out over longer
Figure 7.8.: Figure displaying the mean radial electric field measured at the outboard mid-plane of the three different GENE-X simulations performed. In addition, the fluctuation amplitude is drawn as a shaded area around the mean. For comparison, the radial electric field of GRILLIX simulations with and without neutrals and the experimental measurements is shown. The picture in the bottom excludes the GRILLIX simulations such that a close comparison between the GENE-X results and the experiment is possible.
time periods [165]. Consequently, the profiles in the confined region are flat after averaging over 100µs. Close to the separatrix we observe a well in the radial electric field in the confined region, and a hill in the SOL. This leads to an opposite poloidal rotation of the plasma inside and outside of the separatrix. The fluctuation amplitude of the radial electric field is large and on the same order of magnitude as the value of the field itself. The simulations performed with GENE-X reproduce the shape and amplitude of the experimental measurement better than the simulations performed with GRILLIX.

7.3. Power exhaust

Next, we analyse the power balance of the simulations. As we solve for the full-\(f\) Vlasov equation, we do not prescribe any input power to the simulations. Plasma density and energy is transferred into the plasma by the core and SOL boundaries. The plasma evolves freely and establishes a quasi-steady state. As the plasma \(\beta\) of the simulation is low, the main driver of cross field heat transport in the confined region is the radial component of the \(E \times B\) heat flux. We define it as

\[
q_{E \times B} := \sum_{\sigma} \int_{-\infty}^{\infty} \int_{0}^{\infty} \left( \frac{1}{2} m_{\sigma} v_{\parallel}^2 + \mu B \right) \left( \frac{c B}{B_{\parallel}} \times \nabla \phi_1 \right) \cdot e_{\psi} f_{\sigma} dW
\]

\[
\approx \left( \frac{c B}{B} \times \nabla \phi_1 \right) \cdot e_{\psi} \sum_{\sigma} E_{\sigma},
\]

(7.14)

(7.15)

where \(E_{\sigma}\) denotes the kinetic energy of the plasma. In slight abuse of terminology, we call the radial component of the \(E \times B\) heat flux the \(E \times B\) heat flux in the following. The \(E \times B\) heat flux of the simulations with realistic mass ratio, measured at \(t = 1\)ms, is shown in Fig. 7.9. The \(E \times B\) heat flux is dominant on the low field side. We observe large radial streamers in the top and bottom of the device ranging from the core boundary to the separatrix. After crossing the separatrix the \(E \times B\) heat flux falls off rapidly. We do not observe any significant \(E \times B\) heat flux in the divertor region.

In the SOL the dominant driver of heat transport is the parallel heat flux. We define the parallel heat flux as

\[
q_{\parallel} := \sum_{\sigma} \int_{-\infty}^{\infty} \int_{0}^{\infty} v_{\parallel} \left( \frac{1}{2} m_{\sigma} v_{\parallel}^2 + \mu B \right) f_{\sigma} dW.
\]

(7.16)

Compared to the temperature, presented in Eq. (7.11), that is calculated in a frame where \(u_{\sigma} = 0\), Eqs. (7.15) and (7.16) account for the total energy flux via \(E \times B\) and parallel transport and can therefore be used to calculate the transport of power through the device. We measure the power transported over the separatrix via the \(E \times B\) drift by performing a surface integral of the \(E \times B\) heat flux over the flux surface at \(\psi = 0.999\). Like the electrostatic potential, the \(E \times B\) heat flux fluctuates over time. Therefore, we average the result over the 24 poloidal planes and the last 100µs of the simulation and calculate the fluctuation amplitude as the standard deviation. The results are shown in Tab. 7.1. We observe that
Figure 7.9.: Figure displaying the radial component of the $E \times B$ heat flux of the final state of the simulation on the poloidal plane at $\varphi = \pi$. Similar to Fig. 7.2, the separatrix, wall and divertor is shown. The $E \times B$ heat flux is dominant on the low field side and falls off rapidly after crossing the separatrix. We do not observe any significant $E \times B$ heat flux in the divertor region.
Table 7.1.: Table displaying the total power flowing over the separatrix and onto the divertor for the three simulations performed.

<table>
<thead>
<tr>
<th>μ = 1/400</th>
<th>Separatrix power</th>
<th>Divertor power</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.8MW ± 0.4MW</td>
<td>2.3MW</td>
<td></td>
</tr>
<tr>
<td>μ = 1/3600</td>
<td>1.8MW ± 0.3MW</td>
<td>0.7MW</td>
</tr>
<tr>
<td>μ = 1/3600 with collisions</td>
<td>1.4MW ± 0.3MW</td>
<td>1.1MW</td>
</tr>
</tbody>
</table>

Figure 7.10.: Figure displaying the line of measurement of the parallel heat flux onto the divertor. The separatrix is drawn in green, the device wall in blue, the divertor legs in grey and the boundary line of the divertor in red. The line of measurement, drawn in purple, is calculated by following the points on the boundary line of the divertor in opposite direction of the magnetic field by a toroidal angle of π/8. This moves the line of measurement away from the divertor such that the heat flux is not given by the chosen boundary condition.

The power transported over the separatrix is the largest for the simulation with reduced mass ratio. This finding is consistent with the increased level of turbulence that we discussed in the analysis of the plasma profiles. For the simulations with realistic mass ratio the power fluctuates around 1.4MW–1.8MW. The heating power in the experiment, determined by the power balance, is equal to 475 kW. After crossing the separatrix and leaving the confined region, the energy is transported along the field lines onto the divertor. We measure the heat flux and total power hitting the divertor. This process is not straightforward as the plasma at the divertor is set by the boundary conditions, the divertor has a complicated shape and the magnetic field lines hit the divertor at a shallow incidence angle. We solve the first problem by measuring the heat flux on a line in front of the divertor. The line is generated by following the points on the divertor along the magnetic field lines by a toroidal angle of π/8. The line of measurement is shown in Fig. 7.10. On the line of measurement we interpolate the parallel
heat flux with cubic spline interpolation. To solve the last problem and account for the reduction of heat flux onto the divertor due to the shallow incident angle, we calculate the normalised normal vector \( \mathbf{n} \) to the line of measurement and reduce the interpolated parallel heat flux by the factor \( \sin \theta = \mathbf{n} \cdot \mathbf{b} \), where \( \theta \) denotes the incidence angle of the magnetic field line. The measured \( \sin \theta \) varies between 0.03 on the lower part of the divertor and 0.01 on the upper part of the divertor.

To measure the SOL fall-off length \( \lambda_q \), we fit the function

\[
q(s) = \frac{q_0}{2} \exp \left( -\frac{s^2}{2\lambda_q} - \frac{\pi}{\lambda_q} \right) \text{erfc} \left( \frac{s}{2\lambda_q} - \frac{\pi}{s} \right) + q_{BG},
\]

(7.17)
described in [26], to the electron heat flux.

\[
\text{erfc}(x) = 1 - \text{erf}(x) = \frac{2}{\sqrt{\pi}} \int_{x}^{\infty} e^{-t^2} \, dt
\]

(7.18)
denotes the complementary error function [166]. \( q_0 \) denotes the peak heat flux, \( s \) the power spreading factor and \( q_{BG} \) the background heat flux. To remove the influence of the flux expansion from the OMP to the line of measurement, we perform the fit with the OMP mapped distance \( s \) instead of the length of the line of measurement. The OMP mapped distance is calculated by following the points on the line of measurements along the magnetic field lines until they hit the OMP. The OMP mapped distance is measured as the distance of the intersection points to the separatrix. The measured heat flux profiles and the corresponding fit for the divertor at the low field side are shown for the simulation with reduced mass ratio in Fig. 7.11, for the simulation with realistic mass ratio in Fig. 7.12 and for the simulation with realistic mass ratio and collisions in Fig. 7.13. We observe that the electron heat flux is much larger than the ion heat flux and more peaked close to the separatrix. The measured shape of the electron and total heat flux profile is well described by the Eich model presented in Eq. (7.17). The measured \( \lambda_q \) reads 12mm for the run with reduced mass ratio, 1.9mm for the run with realistic mass ratio and 6.7mm for the run with realistic mass ratio and collisions. The increase in \( \lambda_q \) for the simulations with reduced mass ratio agrees well with the observation of increased turbulence in the profile analysis above. The stronger the turbulence, the further the particles are transported into the SOL. In addition, the reduced mass ratio decreases the electron heat conductivity which slows the parallel heat transport in comparison to the radial \( E \times B \) transport and leads to a further broadening of \( \lambda_q \). The reason for the increase in \( \lambda_q \) for the simulation with collisions is similar. Due to collisions, fast electrons are decelerated which reduces the parallel heat conductivity and leads to a broadening of \( \lambda_q \) by the same mechanism [80]. The measured \( \lambda_q \) from the experiment is around 3mm–4mm and lies in between the results of our simulations with and without collisions. It seems that collisions have an important influence on the broadening of the \( \lambda_q \), and further investigations with a more sophisticated collision operator are needed. The measured peak heat flux from the experiment is around 0.6MW/m\(^2\) – 0.65MW/m\(^2\) and approximately by a factor of 2–3 smaller than the results of our simulations. This finding is consistent with the input power of the experiment, mentioned above, that is approximately by a factor of 2.9–3.8 smaller than
Figure 7.11.: Figure displaying the ion, electron and total heat flux onto the divertor for the simulation with reduced mass ratio. Furthermore, the Eich fit of the total heat flux is drawn. The heat flux is measured as a function of the OMP mapped distance to account for the flux expansion between the OMP and the divertor. The measured SOL fall-off length is $\lambda_q = 12\text{mm}$.

Figure 7.12.: Figure displaying the ion, electron and total heat flux onto the divertor, analogous to Fig. 7.11, for the simulation with realistic mass ratio. The measured SOL fall-off length is $\lambda_q = 1.9\text{mm}$.
Figure 7.13.: Figure displaying the ion, electron and total heat flux onto the divertor, analogous to Fig. 7.11, for the simulation with realistic mass ratio and collisions. The measured SOL fall-off length is $\lambda_q = 6.7\text{mm}$.

The total power absorbed by the divertor on the low and high field side is displayed in Tab. 7.1. The measurements are consistent with the $E \times B$ heat flux. Between 40%–80% of the power crossing the separatrix is absorbed by the divertor. The rest part is transported to the divertor via different mechanisms or is absorbed by the main wall.

### 7.4. Conclusion and outlook

In this chapter we presented the first GENE-X simulations in diverted geometry including a magnetic X-point. We performed the simulations with three different configurations. The first configuration used a reduced electron-to-ion mass ratio of $\mu = m_e/m_i = 1/400$, the second a realistic electron-to-deuterium mass ratio of $m_e/m_i = 1/3600$ and the third a realistic mass ratio with a BGK collision operator. All simulations were performed over a timescale of 1ms in the magnetic equilibrium and at the parameters of the ASDEX Upgrade discharge #36190.

We compared the profiles of density, temperature and the radial electric field to experimental measurements and simulations with the Braginskii fluid code GRILLIX. We found that collisions have an impact on the temperature but not on the density profile. The simulations with collisions were able to reproduce aspects of the experimental temperature profiles. The radial electric field, obtained with GENE-X, agreed well with the experimental measurement. We analysed the power balance of the simulations by measuring the $E \times B$ heat flux flowing over the separatrix and the parallel heat flux onto the divertor. In addition, we calculated $\lambda_q$ by performing an Eich fit to the parallel heat flux profile in front of the divertor. The shape of the measured heat flux profile agreed well with the theoretical prediction by the Eich model. We observed that collisions broaden $\lambda_q$. In the simulations with reduced mass ratio
the fluctuation amplitude was increased and $\lambda_q$ broadened. This indicated that a realistic mass ratio is important for realistic edge and SOL simulations.

As collisions have shown to have a significant effect on the plasma temperature and $\lambda_q$, it will be important to use a more realistic collision model in the future. As mentioned in the outlook of Chapter 2, implementing a Fokker-Planck type Lennard-Bernstein/Dogherty collision operator [114, 115] is current work in progress [116]. The SOL profiles for the ions seemed to be significantly affected by the temperature boundary condition of 35eV. For future studies it is vital to reduce the SOL temperature to more realistic values around 2eV and find solutions to efficiently represent temperature differences of two orders of magnitude on the velocity space grid. One possible candidate to achieve this could be the implementation of block structured grids in GENE-X [167, 168]. Finally, it would be interesting to study the effect of neutral physics, similar to [80], on the plasma profiles and the heat exhaust.
8. Summary and outlook

Understanding and predicting the effects of turbulence in the edge and SOL belongs to one of the most challenging problems on the path towards a magnetic confinement fusion power plant. Turbulence is characterised by a cascade like, transport of energy between different scales and is believed to play a crucial role in the L-H transition and the particle and heat exhaust of fusion devices.

In this work, we presented a new gyrokinetic turbulence code, GENE-X, that is tailored towards edge and SOL simulations. Developing such a code is a difficult task and multiple challenges had to be overcome. We presented detailed conclusions and outlooks at the end of each chapter. In the following, we summarise the highlights of this work and provide a unified outlook.

8.1. Summary

We started this work, in Chapter 2, with a precise discussion of gyrokinetic theory and the derivation of the self-consistent model implemented in the GENE-X code. The model included the full distribution function of ions and electrons and is, therefore, capable of handling large fluctuation amplitudes in SOL plasmas. The model was consistently derived from the second order gyrokinetic Lagrangian and conserves energy as well as the number of particles on the continuous level.

We continued, in Chapter 3, by discussing the coordinate system used for the simulations. The magnetic geometry of the edge and SOL is complex. Magnetic field lines are not confined to flux surfaces and can intersect with the device wall. In addition, there is a transition from closed to open magnetic flux surfaces which renders flux-coordinate systems, that have been used in fusion theory for decades, ineffective. We solved this problem by implementing a locally field-aligned coordinate system based on the FCI approach. We discussed the coordinate transformation, derived expressions for the differential operators and expressed the gyrokinetic Vlasov-Maxwell system in the new coordinate system. Using this technology we are able to perform simulations without restrictions by the magnetic geometry. This includes magnetic geometries with single and even multiple magnetic X-points.

In Chapter 4 we described the discretisation scheme used. The choice of the discretisation scheme was motivated by the locally field-aligned coordinate system and constructed with the method of lines. The spatial discretisation featured centred finite differences and the solution of the remaining, temporal initial value problem was done with a Runge-Kutta method of order four.
As the implementation of scientific codes is complex and their execution resource intensive,
we put, in Chapter 5, a special emphasis on the software design and the computational
performance of the implementation. The design principles behind the code were explained
in detail and visualised with the help of UML diagrams. The analysis of the performance
comprised two steps. First, we analysed the intra-node performance of the implementation
and verified it against the roofline model. Second, we demonstrated that the inter-node
performance is excellent by measuring the strong and weak scaling at the example of a
real application. Finally, we described the continuous integration workflow with automatic
testing that is used extensively in the GENE-X code and ensures that the implementation of
the gyrokinetic Vlasov-Maxwell system is correct.

In Chapter 6, we ensured that the numerical scheme, using the FCI approach, solves the gy-
rokinetic Vlasov-Maxwell system correctly. We performed a comprehensive list of benchmarks
testing different aspects of the code. First, we used the method of manufactured solution to
demonstrate that the numerical scheme converges to second order accuracy. The tests were
performed in slab, circular and screw pinch geometry. Second, we measured the growth rates
of ITG modes in a screw pinch geometry. This test allowed for a precise comparison between
standard flux coordinates and the locally field-aligned coordinate system. Third, we verified
the electromagnetic part of the model by measuring the frequency and Landau damping of
Alfvén in slab geometry. Finally, we probed the complete gyrokinetic Vlasov-Maxwell system
in toroidal geometry by measuring the frequency and growth rates of ITG modes in a cyclone
base case simulation.

To close this work, we presented simulations of the edge and SOL of AUG in Chapter 7
that included the separatrix and the X-point. The simulations were performed over 1ms of
plasma evolution until profile saturation. We compared the profiles of density, temperature
and the radial electric field to experimental measurements and previous GRILLIX simulations.
We found that simulation performed with a BGK collision operator were able to reproduce
aspects of the experimental profiles quite well. Further, we calculated the power exhaust
of the simulation and measured $\lambda_q$. The overall shape of the heat flux profile, measured
close to the divertor, agreed well with the theoretical prediction. We observed that collisions
broaden $\lambda_q$. The simulations performed with a reduced mass ratio of $m_e/m_i = 1/400$ showed
an increased fluctuation level and a significantly broader $\lambda_q$, which makes the application of
reduced mass ratio simulations for edge and SOL plasmas questionable.

8.2. Outlook

With the development of the GENE-X code, we have taken a first step towards a predictive tool
to study turbulence in the edge and SOL of fusion devices. There are numerous interesting
ways to improve and apply the code in the future. We split the future work into three
categories: (i) Improvements of the plasma model, (ii) improvements of the numerical scheme
and (iii) future applications of the GENE-X code.
8.2.1. Improvements of the gyrokinetic model

To improve the predictability of the code, it is important to model effects specific to the SOL more precisely. This includes:

- The implementation of a more sophisticated Fokker-Planck type collision operator. Currently, the implementation of an energy conservative, full-$f$ Lennard-Bernstein/Dogherty collision operator [114, 115] is work in progress [116].

- The implementation of a neutral physics model. A solid starting point could be the simple neutral model implemented in the GRILLIX code recently [80].

- The implementation of sheath boundary conditions [169].

- Removing the linearisation of the quasi-neutrality equation, made in Eq. (2.76), to further improve the full-$f$ capabilities of the code.

8.2.2. Improvements of the discretisation scheme

While the discretisation scheme chosen has been shown to work well, major improvements to the discretisation scheme can still be done on two fronts:

- The implementation of a new grid in velocity space that allows to represent the velocity structure of plasmas with temperatures that vary over multiple orders of magnitude. A possible candidate are block-structured grids, introduced in [167, 168].

- The development and implementation of a numerical scheme based on mimetic finite differences that conserves the particle number and if possible even the momentum and energy of the discrete system.

8.2.3. Future applications of the GENE-X code

Finally, there are many interesting ways to use the GENE-X code in the future. This includes:

- Performing simulations of H-mode plasmas to investigate the L-H transition.

- Performing a validation of the GENE-X code in X-point geometry. This could for instance be achieved by following the recent validation of Braginskii fluid codes presented in [170].

- Generalising the GENE-X code to 3D geometries to investigate turbulence in the edge and SOL of stellarator fusion devices – including magnetic islands and stochastic regions.

- Performing whole device turbulence simulations of a medium size tokamak, like the Tokamak à Configuration Variable (TCV) [171], that range all the way from the magnetic axis, over the separatrix into the scrape-off layer.
A. Supplemental derivations

A.1. Explicit form of the discretised parallel derivatives

In this section, we provide the explicit form of the derivatives of the Lagrange polynomials used in Eqs. (4.35) and (4.49). Consider five grid points in y direction $(y_{-2}, y_{-1}, y_0, y_1, y_2)$. The first derivatives of the Lagrange polynomials read

\[
\frac{d\ell_{-2}}{dy}(0) = \frac{-(y_{-1} - y_0)(y_0 - y_1)(y_0 - y_2)}{(y_{-2} - y_{-1})(y_{-2} - y_0)(y_{-2} - y_1)(y_{-2} - y_2)} \tag{A.1}
\]

\[
\frac{d\ell_{-1}}{dy}(0) = \frac{(y_{-2} - y_0)(y_0 - y_1)(y_0 - y_2)}{(y_{-2} - y_{-1})(y_{-1} - y_0)(y_{-1} - y_1)(y_{-1} - y_2)} \tag{A.2}
\]

\[
\frac{d\ell_0}{dy}(0) = \frac{y_{-1}(3y_0^2 + y_1y_2 - 2y_0(y_1 + y_2)) + y_0(-4y_0^2 - 2y_1y_2 + 3y_0(y_1 + y_2))}{(y_{-2} - y_0)(y_{-1} + y_0)(y_{-1} - y_0)(y_{-1} - y_2)} + \frac{y_{-2}(3y_0^2 + y_1y_2 - 2y_0(y_1 + y_2) + y_{-1}(-2y_0 + y_1 + y_2))}{(y_{-2} - y_0)(y_{-1} + y_0)(y_{-1} - y_0)(y_{-1} - y_2)} \tag{A.3}
\]

\[
\frac{d\ell_1}{dy}(0) = \frac{(y_{-2} - y_{-1})(y_{-1} - y_0)(y_{-1} + y_0)(y_{-1} - y_1)(y_{-1} - y_2)}{(y_{-2} - y_{-1})(y_{-1} + y_0)(y_{-1} - y_0)(y_{-1} - y_2)} \tag{A.4}
\]

\[
\frac{d\ell_2}{dy}(0) = \frac{(y_{-2} - y_0)(y_{-1} + y_0)(y_{-1} - y_1)(y_{-1} - y_2)}{(y_{-2} - y_2)(y_{-1} + y_2)(y_{-1} - y_0)(y_{-1} + y_2)} \tag{A.5}
\]

and the fourth derivatives of the Lagrange polynomials read

\[
\frac{d^4\ell_{-2}}{dy^4}(0) = \frac{24}{(y_{-2} - y_{-1})(y_{-2} - y_0)(y_{-2} - y_1)(y_{-2} - y_2)} \tag{A.6}
\]

\[
\frac{d^4\ell_{-1}}{dy^4}(0) = \frac{-24}{(y_{-2} - y_{-1})(y_{-1} - y_0)(y_{-1} - y_1)(y_{-1} - y_2)} \tag{A.7}
\]

\[
\frac{d^4\ell_0}{dy^4}(0) = \frac{-24}{(y_{-2} - y_0)(y_{-1} + y_0)(y_0 - y_1)(y_0 - y_2)} \tag{A.8}
\]

\[
\frac{d^4\ell_1}{dy^4}(0) = \frac{-24}{(y_{-2} - y_1)(y_{-1} + y_1)(y_0 + y_1)(y_1 - y_2)} \tag{A.9}
\]

\[
\frac{d^4\ell_2}{dy^4}(0) = \frac{24}{(y_{-2} - y_2)(y_{-1} - y_2)(y_0 - y_2)(y_1 - y_2)}. \tag{A.10}
\]

These relations can be used together with Eqs. (4.35) and (4.49) to calculate the first and fourth derivative on the unstructured y grid via a stencil operation.
A.2. Explicit form of the gyrokinetic Vlasov-Maxwell system

In this section, we present the explicit form of the normalised gyrokinetic Vlasov-Maxwell system, expressed in the locally field-aligned coordinate system, that is implemented in the GENE-X code. The gyrokinetic Vlasov equation reads

$$
\frac{\partial \hat{f}_\sigma}{\partial t} + \sqrt{\frac{2\hat{T}_0(\psi_0)}{\hat{m}_\sigma}} \hat{B}_\parallel \cdot \nabla \hat{f}_\sigma + \frac{\rho_{ref}}{L_{\text{ref}}T_{0\parallel}B_\parallel} \times \left( \hat{T}_0(\psi_0)\hat{\mu} \hat{\nabla} \hat{B} + q_\sigma \hat{\nabla} \hat{\phi}_1 \right) \nabla \hat{f}_\sigma
$$

$$
- \frac{\hat{B}_\parallel}{\sqrt{2\hat{m}_\sigma T_{0\parallel\parallel}} B_\parallel} \cdot \left( \hat{T}_0(\psi_0)\hat{\mu} \hat{\nabla} \hat{B} + q_\sigma \hat{\nabla} \hat{\phi}_1 \right) \frac{\partial \hat{f}_\sigma}{\partial \hat{v}_\parallel} - \frac{q_\sigma}{\sqrt{2\hat{m}_\sigma T_{0\parallel\parallel}}} \frac{\partial \hat{A}_\parallel}{\partial \hat{t}} \frac{\partial \hat{f}_\sigma}{\partial \hat{v}_\parallel} = 0,
$$

(A.11)

with the components

$$(a) = \sqrt{\frac{2\hat{T}_0(\psi_0)}{\hat{m}_\sigma}} \hat{B}_\parallel \frac{\partial \hat{f}_\sigma}{\partial \hat{y}} + \frac{2\hat{c}^2\hat{T}_0(\psi_0)}{\hat{B}_\parallel^2} \rho_{ref} \frac{1}{L_{\text{ref}} b_{yc}} \left( b_{xc} \left( \frac{\partial \hat{A}_\parallel}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{x}} - \frac{\partial \hat{B}}{\partial \hat{z}} \frac{\partial \hat{f}_\sigma}{\partial \hat{x}} \right) + b_{zc} \left( \frac{\partial \hat{A}_\parallel}{\partial \hat{z}} \frac{\partial \hat{f}_\sigma}{\partial \hat{x}} - \frac{\partial \hat{B}}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{x}} \right) \right)$$

$$
+ \frac{1}{b_{yc}} \left( b_{xc} \left( \frac{\partial \hat{A}_\parallel}{\partial \hat{z}} \frac{\partial \hat{f}_\sigma}{\partial \hat{z}} - \frac{\partial \hat{B}}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{z}} \right) \right) ;
$$

$$(b) = \frac{1}{\hat{B}_\parallel^2} \rho_{ref} \left[ \hat{T}_0(\psi_0)\hat{\mu} \frac{b_{xc}}{b_{yc}} \left( \frac{\partial \hat{B}}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{y}} - \frac{\partial \hat{B}}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{y}} \right) + b_{zc} \left( \frac{\partial \hat{B}}{\partial \hat{z}} \frac{\partial \hat{f}_\sigma}{\partial \hat{y}} - \frac{\partial \hat{B}}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{y}} \right) \right]
$$

$$
+ q_\sigma \left( b_{xc} \left( \frac{\partial \hat{\phi}_1}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{z}} - \frac{\partial \hat{\phi}_1}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{z}} \right) + b_{zc} \left( \frac{\partial \hat{\phi}_1}{\partial \hat{z}} \frac{\partial \hat{f}_\sigma}{\partial \hat{z}} - \frac{\partial \hat{\phi}_1}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{z}} \right) \right)$$

$$
+ \frac{1}{b_{yc}} \left( b_{xc} \left( \frac{\partial \hat{\phi}_1}{\partial \hat{z}} \frac{\partial \hat{f}_\sigma}{\partial \hat{z}} - \frac{\partial \hat{\phi}_1}{\partial \hat{y}} \frac{\partial \hat{f}_\sigma}{\partial \hat{z}} \right) \right) \right] \quad \text{and} \quad \text{(A.13)}
$$
Ampère's law reads

\[
T_{0\sigma}(\psi_0) \mu \left( B \frac{\partial \hat{B}}{\partial y} + \sqrt{2m_\sigma T_{0\sigma}(\psi_0)} \hat{v}_{\parallel} \left[ \frac{\rho_{\text{ref}}}{q_\sigma L_{\text{ref}}} b_{y_c} \right] \right) + \frac{1}{\sqrt{2m_\sigma T_{0\sigma}(\psi_0) B_{\parallel}^*}} \left[ \hat{\Theta}_{0\sigma} - \sum \frac{\hat{n}_{0\sigma}}{B^2} \frac{\partial}{\partial \hat{z}} \left( \sum \frac{\hat{n}_{0\sigma}}{B^2} \frac{\partial}{\partial \hat{z}} \right) \hat{\phi}_1 \right]
\]

The quasi-neutrality equation reads

\[
- \left( \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right)^2 \left( \frac{1}{K_{\text{cyl}}} \frac{\partial}{\partial \hat{x}} \left( J_{\text{cyl}} \sum_\sigma \frac{\hat{n}_{0\sigma}}{B^2} \frac{\partial}{\partial \hat{x}} \phi_1 \right) + \frac{\partial}{\partial \hat{z}} \left( \sum_\sigma \frac{\hat{n}_{0\sigma}}{B^2} \frac{\partial}{\partial \hat{z}} \phi_1 \right) \right) = \sum_\sigma \hat{n}_{0\sigma}(\psi_0) \hat{q}_\sigma \int_0^\infty \int_0^\infty \pi B^* \hat{f}_\sigma \hat{v}_{\parallel} d\hat{v} d\hat{\mu} , \quad (A.15)
\]

Ampère’s law reads

\[
- \left( \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right)^2 \left( \frac{1}{K_{\text{cyl}}} \frac{\partial}{\partial \hat{x}} \left( J_{\text{cyl}} \frac{\partial}{\partial \hat{x}} A_1 \right) + \frac{\partial^2}{\partial \hat{z}^2} A_1 \right) = \frac{\beta_{\text{ref}}}{2} \sum_\sigma \hat{n}_{0\sigma}(\psi_0) \hat{q}_\sigma \sqrt{\frac{2T_{0\sigma}(\psi_0)}{m_\sigma}} \int_0^\infty \int_{-\infty}^\infty \pi B^* \hat{f}_\sigma \hat{v}_{\parallel} d\hat{v} d\hat{\mu} \quad (A.16)
\]
and the generalised Ohm’s law reads

\[
- \left[ \left( \frac{\rho_{\text{ref}}}{L_{\text{ref}}} \right)^2 \left( \frac{1}{J_{\text{cyl}}} \frac{\partial}{\partial \hat{x}} \left( \hat{J}_{\text{cyl}} \frac{\partial}{\partial \hat{x}} \right) + \frac{\partial^2}{\partial \hat{z}^2} \right) + \beta_{\text{ref}} \sum_{\sigma} \hat{n}_{0\sigma}(\psi_0) \frac{q_\sigma^2}{m_\sigma} \int_0^\infty \int_{-\infty}^\infty \pi B_{\parallel}^* \frac{\partial f_\sigma}{\partial \hat{v}_\parallel} \hat{v}_\parallel d\hat{v}_\parallel d\hat{\mu} \right] \frac{\partial A_1}{\partial t} \\
+ \beta_{\text{ref}} \sum_{\sigma} \hat{n}_{0\sigma}(\psi_0) \frac{q_\sigma^2}{m_\sigma} \int_0^\infty \int_{-\infty}^\infty \pi B_{\parallel}^* \left( \frac{\partial f_\sigma}{\partial \hat{v}_\parallel} \right)^* \hat{v}_\parallel d\hat{v}_\parallel d\hat{\mu}.
\]

(A.17)

In the implementation we set \( \hat{T}_{0\sigma}(\psi_0) = 1 \) and \( \hat{n}_{0\sigma}(\psi_0) = 1 \).
A.3. Identities of the plasma dispersion function

The plasma dispersion function is defined as \[172\]

\[ Z : \omega \mapsto Z(\omega) = \frac{1}{\sqrt{\pi}} \int_{-\infty}^{\infty} e^{-t^2} \frac{t}{t - \omega} \, dt. \]  

(A.18)

It fulfills the identities

\[ \frac{1}{\sqrt{\pi}} \int_{-\infty}^{\infty} \frac{te^{-t^2}}{t - \omega} \, dt = \frac{1}{\sqrt{\pi}} \int_{-\infty}^{\infty} e^{-t^2} + \frac{\omega}{t - \omega} e^{-t^2} \, dt \]

\[ = 1 + \omega Z(\omega) \]  

(A.19)

and

\[ \frac{1}{\sqrt{\pi}} \int_{-\infty}^{\infty} \frac{t^2 e^{-t^2}}{t - \omega} \, dt = \frac{1}{\sqrt{\pi}} \int_{-\infty}^{\infty} (t + \omega)e^{-t^2} + \frac{\omega^2}{t - \omega} e^{-t^2} \, dt = \omega + \omega^2 Z(\omega) \]

\[ = \omega(1 + \omega Z(\omega)). \]  

(A.20)
A.4. Proof of an integral identity of the Poisson bracket

In this section, we prove
\[ \int H^\sigma \{ f^\sigma, H^\sigma \} \star dV dW = 0 \tag{A.21} \]
under the assumption that boundary integrals vanish. The gyrokinetic Poisson bracket \{ , \}⋆, defined in Subsection 2.3.6, evaluated on the two functions \( f \) and \( g \), can be written as
\[ \{ f, g \}^\star = \sum_{a=1}^{6} \sum_{b=1}^{6} J_{ab} \frac{\partial f}{\partial z_a} \frac{\partial g}{\partial z_b}, \tag{A.22} \]
where \( J_{ab} \) is a skew-symmetric matrix. Inserting Eq. (A.22) into Eq. (A.21) yields
\[
\int g \{ f, g \}^\star dV dW = \sum_{a=1}^{6} \sum_{b=1}^{6} \int J_{ab} \frac{\partial f}{\partial z_a} \frac{\partial g}{\partial z_b} dV dW \\
= \sum_{a=1}^{6} \sum_{b=1}^{6} \int \frac{1}{2} J_{ab} B^\star_B \left( g \frac{\partial f}{\partial z_a} + g \frac{\partial f}{\partial z_b} \right) \frac{1}{m_{\sigma}} dV \, d\theta \, d\mu \, dv \\
= \sum_{a=1}^{6} \sum_{b=1}^{6} \int \frac{1}{2} \left( \frac{\partial}{\partial z_b} \left( J_{ab} B^\star_B g^2 \frac{\partial f}{\partial z_a} \right) - J_{ab} B^\star_B g^2 \frac{\partial^2 f}{\partial z_a \partial z_b} \right) \frac{1}{m_{\sigma}} dV \, d\theta \, d\mu \, dv \\
= 0, \tag{A.23} 
\]
where we performed a partial integration over \( z_b \), assumed that boundary integrals vanish, used the skew-symmetry of \( J_{ab} \) and Liouville’s theorem \( \sum_{b=1}^{6} \frac{\partial}{\partial z_b} (J_{ab} B^\star_B) = 0 \) [94]. Setting \( g = H^\sigma \) and \( f = f^\sigma \) proves Eq. (A.21).
B. Supplemental results

B.1. Results of the scaling analysis

In this section, we give the measured times for the evaluation of one timestep in the analysis of the strong and weak scaling, presented in Chapter 5. The measured times for the strong scaling are contained in the first and for the weak scaling in the second table.

<table>
<thead>
<tr>
<th>Number of nodes</th>
<th>Time per timestep</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>884s</td>
</tr>
<tr>
<td>2</td>
<td>454s</td>
</tr>
<tr>
<td>4</td>
<td>229s</td>
</tr>
<tr>
<td>8</td>
<td>115s</td>
</tr>
<tr>
<td>16</td>
<td>63.2s</td>
</tr>
<tr>
<td>32</td>
<td>34.8s</td>
</tr>
<tr>
<td>64</td>
<td>17.5s</td>
</tr>
<tr>
<td>128</td>
<td>9.00s</td>
</tr>
<tr>
<td>256</td>
<td>5.05s</td>
</tr>
<tr>
<td>512</td>
<td>2.98s</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Number of nodes</th>
<th>Time per timestep</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>8.17s</td>
</tr>
<tr>
<td>2</td>
<td>7.38s</td>
</tr>
<tr>
<td>4</td>
<td>7.57s</td>
</tr>
<tr>
<td>8</td>
<td>7.50s</td>
</tr>
<tr>
<td>16</td>
<td>6.77s</td>
</tr>
<tr>
<td>32</td>
<td>7.30s</td>
</tr>
<tr>
<td>64</td>
<td>6.91s</td>
</tr>
<tr>
<td>128</td>
<td>7.23s</td>
</tr>
<tr>
<td>256</td>
<td>7.43s</td>
</tr>
<tr>
<td>512</td>
<td>7.62s</td>
</tr>
</tbody>
</table>
B.2. Solution of the electrostatic dispersion relation in screw pinch geometry

In this section, we give the results for the complex frequency \( \hat{\omega} \) obtained from the dispersion relation in Eq. (6.19) up to four significant digits. They are contained in the following table.

\[
\begin{array}{c|c}
  m & \hat{\omega} \\
  \hline
  5 & -1.725 + 1.689i \\
  6 & -1.740 + 1.760i \\
  7 & -1.744 + 1.800i \\
  8 & -1.742 + 1.819i \\
  9 & -1.735 + 1.823i \\
 10 & -1.725 + 1.817i \\
 11 & -1.713 + 1.804i \\
 12 & -1.700 + 1.786i \\
 13 & -1.686 + 1.764i \\
 14 & -1.671 + 1.740i \\
 15 & -1.657 + 1.714i \\
 16 & -1.643 + 1.688i \\
 17 & -1.629 + 1.661i \\
 18 & -1.615 + 1.634i \\
 19 & -1.601 + 1.607i \\
 20 & -1.588 + 1.580i \\
\end{array}
\]

B.3. Numerical results for the growth rate \( \text{Im} \, \hat{\omega} \) in screw pinch geometry

In this section, we give the numerical results for the growth rate \( \text{Im} \, \hat{\omega} \) obtained from simulations with \textit{GENE-X} described in Section 6.3. They are contained in the following table.

\[
\begin{array}{c|c|c}
  m & \text{Im} \, \hat{\omega} \, \text{for} \, q = \infty & \text{Im} \, \hat{\omega} \, \text{for} \, q \neq \infty \\
  \hline
  5 & 1.74 & 1.73 \\
 10 & 1.85 & 1.84 \\
 15 & 1.73 & 1.72 \\
 20 & 1.58 & 1.58 \\
\end{array}
\]
B.4. Solution of the electromagnetic dispersion relation in slab geometry

In this section, we give the results for the complex frequency $\hat{\omega}$ obtained from the dispersion relation Eq. (6.35) for the parallel mode number $\hat{k}_x = 1$, different perpendicular mode numbers $\hat{k}_y$ and different plasma beta $\bar{\beta} = \beta_{\text{ref}}/m_e$ up to four significant digits. They are contained in the following table.

<table>
<thead>
<tr>
<th>$\hat{k}<em>x\rho</em>{\text{ref}}/L_{\text{ref}}$</th>
<th>$\hat{\omega}$ for $\bar{\beta} = 0.2$</th>
<th>$\hat{\omega}$ for $\bar{\beta} = 2$</th>
<th>$\hat{\omega}$ for $\bar{\beta} = 20$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1/128</td>
<td>$189.7 - 6.095 \cdot 10^{-3}i$</td>
<td>$60.00 - 2.130 \cdot 10^{-3}i$</td>
<td>$18.97 - 2.279 \cdot 10^{-4}i$</td>
</tr>
<tr>
<td>1/64</td>
<td>$189.6 - 2.440 \cdot 10^{-2}i$</td>
<td>$60.00 - 8.517 \cdot 10^{-3}i$</td>
<td>$18.98 - 9.114 \cdot 10^{-4}i$</td>
</tr>
<tr>
<td>1/32</td>
<td>$189.7 - 9.789 \cdot 10^{-2}i$</td>
<td>$60.01 - 3.405 \cdot 10^{-2}i$</td>
<td>$18.98 - 3.646 \cdot 10^{-3}i$</td>
</tr>
<tr>
<td>1/16</td>
<td>$187.5 - 3.960 \cdot 10^{-1}i$</td>
<td>$60.05 - 1.360 \cdot 10^{-1}i$</td>
<td>$19.01 - 1.458 \cdot 10^{-2}i$</td>
</tr>
<tr>
<td>1/8</td>
<td>$181.6 - 1.635 \cdot 10^{0}i$</td>
<td>$60.19 - 5.399 \cdot 10^{-1}i$</td>
<td>$19.11 - 5.827 \cdot 10^{-2}i$</td>
</tr>
<tr>
<td>1/4</td>
<td>$164.8 - 6.687 \cdot 10^{0}i$</td>
<td>$60.73 - 2.010 \cdot 10^{0}i$</td>
<td>$19.52 - 2.324 \cdot 10^{-1}i$</td>
</tr>
<tr>
<td>1/2</td>
<td>$136.1 - 2.192 \cdot 10^{1}i$</td>
<td>$62.40 - 7.584 \cdot 10^{0}i$</td>
<td>$21.04 - 9.187 \cdot 10^{-1}i$</td>
</tr>
<tr>
<td>1</td>
<td>$107.6 - 4.826 \cdot 10^{1}i$</td>
<td>$65.37 - 2.239 \cdot 10^{1}i$</td>
<td>$25.87 - 3.512 \cdot 10^{0}i$</td>
</tr>
</tbody>
</table>

B.5. Numerical results for the complex frequency $\hat{\omega}$ in slab geometry

In this section, we give the numerical results for the complex frequency $\hat{\omega}$ obtained from simulations with GENE-X described in Section 6.4. They are contained in the following table.

<table>
<thead>
<tr>
<th>$\hat{k}<em>x\rho</em>{\text{ref}}/L_{\text{ref}}$</th>
<th>$\hat{\omega}$ for $\bar{\beta} = 0.2$</th>
<th>$\hat{\omega}$ for $\bar{\beta} = 2$</th>
<th>$\hat{\omega}$ for $\bar{\beta} = 20$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1/64</td>
<td>$189.6 - 2.343 \cdot 10^{-2}i$</td>
<td>$60.00 - 8.318 \cdot 10^{-3}i$</td>
<td>$18.98 - 8.450 \cdot 10^{-4}i$</td>
</tr>
<tr>
<td>1/16</td>
<td>$187.5 - 3.852 \cdot 10^{-1}i$</td>
<td>$60.00 - 1.362 \cdot 10^{-1}i$</td>
<td>$19.04 - 1.418 \cdot 10^{-2}i$</td>
</tr>
<tr>
<td>1/4</td>
<td>$164.2 - 6.680 \cdot 10^{0}i$</td>
<td>$60.81 - 2.110 \cdot 10^{0}i$</td>
<td>$19.51 - 2.271 \cdot 10^{-1}i$</td>
</tr>
<tr>
<td>1</td>
<td>$108.3 - 4.773 \cdot 10^{1}i$</td>
<td>$65.00 - 2.223 \cdot 10^{1}i$</td>
<td>$25.89 - 3.475 \cdot 10^{0}i$</td>
</tr>
</tbody>
</table>
B.6. Numerical results for the complex frequency $\hat{\omega}$ in toroidal geometry

In this section, we give the numerical results for the complex frequency $\hat{\omega}$ obtained from simulations with GENE-X described in Section 6.5. They are contained in the following table.

\begin{center}
\begin{tabular}{cc}
$n$ & $\hat{\omega}$ \\
5 & $0.180 + 6.79 \cdot 10^{-2}i$ \\
10 & $0.419 + 2.05 \cdot 10^{-1}i$ \\
15 & $0.661 + 3.56 \cdot 10^{-1}i$
\end{tabular}
\end{center}
Bibliography


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