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Gyrokinetic neoclassical study of the bootstrap current in the tokamak edge pedestal with fully nonlinear Coulomb collisions

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As a followup on the drift-kinetic study of the non-local bootstrap current in steep edge pedestal of tokamak plasma by Koh et al. [Phys. Plasmas 19, 072505 (2012)], a gyrokinetic neoclassical study is performed with gyrokinetic ions and drift-kinetic electrons. Besides the gyrokinetic improvement of ion physics from the drift-kinetic treatment, a fully nonlinear Fokker-Planck collision operator – that conserves mass, momentum and energy – is used instead of Koh et al.'s linearized collision operator in consideration of the possibility that the ion distribution function is non-Maxwellian in the steep pedestal. An inaccuracy in Koh *et al.*'s result is found in the steep edge pedestal that originated from a small error in the collisional momentum conservation. The present study concludes that 1) the gyrokinetic ions do not make a noticeable difference in the bootstrap current from the drift-kinetic ion results, 2) the bootstrap current in the steep edge pedestal is generally smaller than what has been predicted from the small banana-width (local) approximation [e.g., O. Sauter et al., Phys. Plasmas 6, 2834 (1999) and E. Belly et al., Plasma Phys. Controlled Fusion [50], 095010 (2008)], 3) the plasma flow evaluated from the local approximation can significantly deviate from the non-local results, and 4) the bootstrap current in the edge pedestal, where the passing particle region is small, can be dominantly carried by the trapped particles in a broad trapped boundary layer. A new analytic formula based on numerous gyrokinetic simulations using various magnetic equilibria and plasma profiles with self-consistent Grad-Shafranov solutions is constructed.

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I. INTRODUCTION

The important role of the bootstrap current in magnetic fusion plasma has been well recognized 1^{-15} . Since the bootstrap current profile is a difficult quantity to measure in experiments, analytic formulas or numerical codes have been used for its prediction and analysis. Among the analytic formulas found in the literature, the so-called "Sauter formula"¹⁴ has been popular in the modeling and experimental communities. The Sauter formula was developed based on numerous drift-kinetic simulations under various Grad-Shafranov tokamak plasma conditions using an approximate linearized collision operator in the limit of small banana orbit width compared to the radial plasma gradient length (called the "local regime"). Recently, the importance of the bootstrap current has been re-emphasized by its critical role in the stability, equilibrium, and turbulence in the steep edge pedestal plasma^{16–18}, where the ion banana width is similar to the pressure gradient scale length (called "non-local regime"). Due to the largeness of the bootstrap current density in the steep pressure gradient region, a small inaccuracy in the prediction for its magnitude and profile can sensitively affect the physics of the edge pedestal. Most of the previous theoretical and numerical studies, and analytic formulas for the bootstrap current, including the Sauter formula¹⁴ and Belli et al.'s numerical study¹⁵, have been focused on the core plasma with small trapped-particle fraction and/or using the local limit, and do not guarantee accuracy in the steep edge pedestal. A more accurate evaluation of the bootstrap current profile in the steep edge pedestal region of tokamak plasma has been in need.

Besides the local approximation, Sauter *et al*'s simulations¹⁴ and their fitting formula have been known to be less accurate at higher electron collisionality¹⁹, $\nu_{e,*} > 1$, mainly due to the approximate electron-ion collision operator used in the simulation and also due to the simplified fitting formula used at high collision frequency that does not produce the correct asymptotic behavior. Reference [20] studied an ion contribution to the bootstrap current that arises from the interaction of finite ion banana width with the strong radial electric field occurring in a pedestal plasma, however, still in the large aspect ratio limit. Koh *et al.*²¹ used a previous version of the XGC0 global drift-kinetic code²², which was equipped with an approximate linear collision operator, to study the finite banana orbit effect on the bootstrap current in the steep edge pedestal together with the magnetic separatrix effect, and they suggested a new analytic formula by adding corrections to the Sauter formula. Belli *et al.*¹⁵ compared Sauter *et al.*'s and Koh *et al.*'s analytic bootstrap current models with results of the local neoclassical code NEO^{23,24}, and found some disagreement of the NEO results with both formulas. Landreman *et al.*²⁵ investigated the bootstrap current using their new global neoclassical code PERFECT allowing for the density gradient scale length to be on the order of the ion banana width ρ_b , but the ion temperature gradient scale length was still assumed to be much larger than ρ_b . Inaccuracy of the Sauter formula at high collisionality has also been a part of the reports of the PERFECT and NEO studies in the local regime.

Due to the importance of the problem and some disagreement with the NEO code, we have re-examined Koh et al.'s results using the more advanced, non-local, gyrokinetic neoclassical code XGCa, which uses a fully nonlinear Fokker-Planck collision operator with highly accurate conservation of energy, momentum, and mass. The electrons in XGCa are still drift-kinetic since the electron gyroradius is negligibly small compared to the pressure gradient scale length even in the pedestal. The use of a fully nonlinear and conserving collision operator removes potential uncertainties due to non-Maxwellian components of the ion distribution function in the steep pedestal and due to a possible inaccuracy from the linearized Coulomb collision operator used in XGC0. A gyrokinetic neoclassical code also removes the uncertainty of the analytic gyroviscosity model in steep edge pedestal plasma, where the ion gyroradius ρ_i is a non-negligible fraction of the pressure gradient scale length $L_P \ (\rho_i \sim 10^{-1} L_P)$. The XGCa results are compared with the local Sauter formula and with the results from the local drift-kinetic solver NEO in their respective validity regimes, showing negligible differences. In the validity regime of the Sauter formula (local and weakly collisional $\nu_{e,*} < 1$), all three results agree well; while in the extended validity regime of NEO (local and collisional), NEO and XGCa agree well while the Sauter formula gives higher bootstrap current. In the process of cross-verification between XGCa and the previous version XGC0, it was discovered that XGC0 had a small inaccuracy in the total momentum conservation from the approximate linearized Monte-Carlo collision operator. Even though the inaccuracy in the momentum conservation was small when compared to the ion momentum - hence the accuracy of the ion neoclassical physics was not affected – it was enough to raise the electron bootstrap current somewhat, which accounted for the discrepancy of the XGC0 results with XGCa, and with NEO in NEO's validity regime¹⁵. After fixing the inaccuracy in the XGC0 collision operator, the improved version XGC0 agrees with XGCa in all regimes,

and with NEO in NEO's validity regime. Thus, one takeaway from the present study, as far as the bootstrap current is concerned, is that the gyrokinetic nature of the ion dynamics in XGCa does not make a noticeable difference from the drift-kinetic ions used in XGC0.

Another physics finding from this study that has not been discussed in the literature is the dominant contribution of trapped electrons to the bootstrap current in the edge pedestal in contrast to the central core plasma, where most of the previous theoretical studies have been performed. In the edge pedestal of conventional aspect-ratio or tight aspect-ratio tokamaks, the passing particle population is much smaller than the trapped particle population. In this case, it is found that the bootstrap current penetrates into the trapped particle regime through a broadened boundary layer. In the large aspect ratio limit, the usual "passing particle dominance" of the bootstrap current is recovered.

Numerous XGCa simulation results under various Grad-Shafranov plasma and magnetic equilibrium conditions led us to construct a new analytic formula – again based on the Sauter formula – by improving the accuracy of collisional physics and adding non-local effects. The new formula represents the XGCa results (and the NEO results in the local regime) much more accurately than the Sauter formula does for all the cases studied, and can be implemented easily into a modeling or analysis code as a simple modification to the Sauter formula. By using a different type of fitting, the discontinuity in Koh *et al.*'s formula at the inverse aspect ratio $\epsilon = 0.44$ has also been removed, which was intentionally located in Koh *et al.*'s formula at an inverse aspect ratio where there is no operational tokamak at the present time or unlikely to be in the future.

In this work, we assume that the ion charge number dependence of the bootstrap current given in Sauter *et al.*¹⁴ is acceptable, and we consider the improvement of the analytic formula in the limit of singly charged ions. When there are impurity particles in the plasma, the charge number Z in the ion contribution to the bootstrap current can be identified by using the description given in Koh *et al.*²¹,

$$Z = \left(Z_{\alpha}^2 \bar{Z} Z_{eff}\right)^{1/4},\tag{1}$$

where $Z_{\alpha} = 1$ for the main ion species, $\overline{Z} = n_e/n_i$, and $Z_{eff} = \sum_{\beta} n_{\beta} Z_{\beta}^2/n_e$ with β being the ion species index. For the electron bootstrap current term, $Z = Z_{eff}$ as usual. The Z-dependence will be investigated using the gyrokinetic XGCa code in the future and our new formula will be updated if there is any significant difference. The remainder of the paper is organized as follows: Section II is a brief description of the XGCa code and the computational setup. We also discuss how a small inaccuracy in the momentum conservation in the previous version of XGC0's collision operator affected Koh *et al.*'s results, and how XGC0 results changed with corrected momentum conservation. Results from a cross-verification between XGCa, XGC0, and NEO are discussed in Sec. III A. Section III B contains a description of how collisions in the pedestal increase the trapped electron fraction of the bootstrap current. The relation of this increased trapped particle contribution to conventional neoclassical theory at high aspect ratio is explained in Sec. III C. Section III D is dedicated to the discussion of finite orbit-width effects. Our improvement to the Sauter formula is discussed in Sec. IV. Section V contains summary and discussion.

II. THE XGCa CODE AND THE BASIC SIMULATION CONDITIONS

The main code used for the present study is the new non-local total-f gyrokinetic neoclassical code XGCa. We will present a brief description of XGCa in this section. A more detailed description of XGCa will be presented in a separate report. The global total-f driftkinetic neoclassical code XGC0 is also used in the present work for comparison with the previous work by Koh *et al.*²¹, who used an older version $XGC0^{21,22,26,27}$. We also use the perturbative drift-kinetic code NEO, which was developed for local physics in the plasma away from the magnetic separatrix, for cross-verification in NEO's validity regime. Detailed description of NEO can be found in Refs. [23 and 24].

XGCa is an axisymmetric version of the global total-f gyrokinetic particle-in-cell code XGC1²⁸, which includes gyrokinetic ions, drift-kinetic electrons and neutral particles. The axisymmetric Poisson solver in XGCa removes the turbulence solution but keeps the poloidal variation of the electrostatic potential that was missing in XGC0. XGCa uses a non-linear multi-species Fokker-Planck-Landau collision operator with the real electron-to-ion mass ratio²⁹, which has been generalized from the single species operator developed by Yoon and Chang³⁰. Since the gyro-averaging is performed explicitly in XGCa using the usual 4-point averaging technique³¹, there is no need for an ad-hoc gyro-averaging model (such as gyro-viscosity) like the one used in XGC0 for the steep edge pedestal physics.

XGCa results converge well at approximately 10,000 particles per configuration space

grid vertex. The simulation time needed to reach a quasi-steady state is approximately 2 pedestal ion collision times corresponding to 5000 ion time steps or 10 ion toroidal transit times. A quasi-steady state is defined in the present work as a state in which all the physical quantities including the radial and poloidal electric field, toroidal and poloidal flows, and plasma profiles are out of the initial transient state and evolve on the neoclassical transport time scale. The implicit, non-linear collision operation in XGCa exhibits good convergence behavior if the collision operation is performed approximately at every electron collision time (or more frequently) corresponding to 1-5 time steps depending on the minimum temperature of ions and electrons. A typical wall-clock time to reach a quasi-steady neoclassical state is half a day on the IBM BlueGeneQ System Mira using 16384 compute nodes and the real electron mass.

Because of the importance of the accuracy of the collision operation (especially the accuracy of momentum conservation) in calculating the bootstrap current, we discuss some details of the conservation properties of the nonlinear collision operator here. For a more complete description, we refer the readers to Refs. [29] and [30]. The nonlinear collision operation in XGCa and XGC1 is performed on a two-dimensional velocity grid with the marker particle velocity distribution from each configuration space vertex being mapped to the velocity grid. After each collision operation, the collision result is mapped back to the marker particles' weights. The velocity grids for ions and electrons are normalized to the respective thermal velocity, and have identical size and resolution in normalized velocity units. In this work, we use 40×40 cells with a high cut-off at 4 times the thermal velocity.

We examine the conservation properties of the nonlinear collision operator in a simple, but well designed, comprehensive test in order to demonstrate the accuracy needed for the bootstrap current calculation. The test presented here is the relaxation of temperature and flow to the thermal equilibrium (see Ref. [29] for more comprehensive verification). The solution of each implicit collision operation is iterated until the relative errors of mass, momentum, and energy per time step are less than 10^{-6} . The marker particles in this collision accuracy test are not time-advanced in the equation of motion; only the right hand side of the Boltzmann equation is solved. Thus, the Monte-Carlo and the particle-mesh interpolation errors are not considered in this test of the collision operator, but they are examined through convergence studies in particle number and grid size in complete XGCa simulations. The initial electron and ion temperatures have both, isotropic difference and individual anisotropy: $T_{e,\perp,0} = 390$ eV, $T_{e,\parallel,0} = 300$ eV, $T_{i,\perp,0} = 260$ eV, and $T_{i,\parallel,0} = 200$ eV. The initial parallel ion and electron flows are $u_{e,0} = 59.9$ km/s and $u_{i,0} = 1.3$ km/s, representing a typical level of electron and ion flows in the edge pedestal with the relations $u_{i,0} \sim (m_e/m_i)^{1/2}u_{e,0}$ and $u_{e,0} \sim v_{t,i}/2$, where $v_{t,i}$ is the ion thermal velocity. The errors in energy and momentum conservation are normalized relative to the initial energy and momentum. The energy and momentum conservation accuracy observed in the relaxation test is illustrated in Fig. 1. It can be seen that the time-accumulated relative momentum conservation error is at the 10^{-6} level and the time-accumulated relative momentum conservation times, which is a much longer simulation time than the time needed (~ 2 ion collision times) for measuring the bootstrap current or any other neoclassical quantities. Moreover, the energy and momentum errors set to be 10^{-6} per collision operation is sufficient to keep the accumulated conservation error to be around 10^{-5} relative to the initial condition. It is found, in general, that the energy conservation error is smaller than the momentum conservation error.

Since the momentum in the electrons is smaller than that in the ions by $(m_e/m_i)^{1/2}$, an error of less than 10^{-2} in the electron bootstrap current requires a relative error of less than $10^{-2}(m_e/m_i)^{1/2} = 1.6 \cdot 10^{-4}$ in the total momentum. In the real XGCa simulations, though, it is found that the relative error in the electron bootstrap current is less than this simple estimate since the small deviation $\delta j_{b,e}$ away from the equilibrium is damped by the collisional friction with ions. Its influence on the ions is smaller by $(m_e/m_i)^{1/2}$. When the conservation accuracy in complete XGCa simulations is reduced from 10^{-6} to 10^{-8} per time step as part of the convergence studies, the same result for the bootstrap current is found.

The low noise ratio in XGCa enables us to use a single time slice of the distribution function to evaluate the bootstrap current. No time averaging is necessary as in case of XGC0. Therefore, the choice of a time slice in quasi-steady state has little influence on the result as demonstrated in Fig. 2. This figure shows the time evolution of the bootstrap current at the location of its maximum in the pedestal from an XGCa simulation using the JET geometry along with the currents predicted by the Sauter formula, and an improved formula to be presented in Sec. IV. The time evolution of the bootstrap current as given by the formulas is due to the evolution of the plasma profiles.

The plasma and magnetic configurations we use in this report include circular limiter ge-



FIG. 1. Conservation properties of the non-linear collision operator in XGCa during a comprehensive relaxation test starting from $T_{e,\perp,0} = 390 \text{ eV}$, $T_{e,\parallel,0} = 300 \text{ eV}$, $T_{i,\perp,0} = 260 \text{ eV}$, $T_{i,\parallel,0} = 200 \text{ eV}$, $u_{e,0} = 59.9 \text{ km/s}$, and $u_{i,0} = 1.3 \text{ km/s}$): a) Relative error of total energy and b) relative error of total momentum. Time is normalized to the ion collision time. The errors are normalized relative to the initial energy and momentum.



FIG. 2. Time evolution of the bootstrap current at the radial location of the maximum of the bootstrap current profile ($\psi_N = 0.93$) shown in Fig. 8 a). Time is normalized to the local ion collision time. After an initial transient phase of approximately one ion collision time, the bootstrap current evolves only slowly in time. Comparison with the Sauter formula and the modified formula to be presented in Sec. IV demonstrates that the slow time variation is due to the slow variation of the plasma profiles caused by neoclassical transport. This indicates that the system has reached a quasi-steady state.

ometries that were generated by the equilibrium solvers ISOLVER³² and FLOW³³, as well as realistic tokamak magnetic configurations for C-Mod (discharge 1120803014)³⁴, DIII-D geometry (similar to discharge 96333)²⁸, JET (similar to discharge 85407³⁵), ASDEX-Upgrade (discharge 28093)³⁶, and NSTX (discharges 128013^{37,38} and 132543³⁹) (all devices are described in Ref. [40]). In case of the limiter configurations, the simulation domain includes only the closed field line region. In case of the divertor configurations, it includes the whole volume. Note that in this article, we do not study the separatrix effect discussed by Koh *et al.*²¹ in detail since the effect is relatively small. As a matter of fact, the improved formula we propose in Sec. IV achieves good accuracy without Koh *et al.*'s separatrix correction. The circular limiter configurations are used to study the fundamental physics effects without the complication introduced by the separatrix and scrape-off layer. The circular limiter configurations also facilitate comparison of the XGC results with other neoclassical codes that are unable to treat the open field line region. Due to the interplay of background gradients and collisionality, the peak of the bootstrap current profile usually lies between the pedestal top and the pedestal center, i.e. not at the point of maximal plasma gradient.

When using the local neoclassical drift-kinetic code NEO away from the magnetic separatrix for cross-verification with XGCa in the local limit, the following simulation conditions have been set in NEO. We used 25 grid points per flux-surface in the poloidal direction (33 for NSTX), 17 basis function in pitch-angle, and 9 basis functions in energy space. As in all XGCa simulations, we used the real electron mass in NEO. The density and temperature profiles from a selected time slice (in quasi-steady state) of the corresponding XGCa simulation were used as input to NEO. Since NEO cannot solve for the radial electric field, we used the gradient of the flux-surface averaged potential, $d\langle \phi \rangle/d\psi$ from XGCa as input for the zeroth order radial electric field. While this is merely a shift of the reference frame in the local code NEO, it enables us to compare the ion and electron contributions between NEO and XGCa in the laboratory frame.



FIG. 3. Density and temperature profiles for case CIRC1.

III. NUMERICAL RESULTS

A. Cross-verification of XGCa, XGC0, and NEO in the local regime

In order to test if the global neoclassical simulations performed with XGCa and XGC0 agree with the local NEO simulations and the local analytic formula by Sauter *et al.* in a mild gradient case (local regime), we performed a cross-verification between XGCa, XGC0, NEO and the Sauter formula. Here, we define the inverse aspect ratio ϵ as the ratio of the mean minor radius $R_{min} \equiv (R_+ - R_-)/2$ to the geometrical center $R_c \equiv (R_+ + R_-)/2$ of a flux-surface, where R_+ and R_- are the major radii at the outer- and inner-most points of a flux-surface. Thus, $\epsilon \equiv (R_+ - R_-)/(R_+ + R_-)$. The three simulation cases we discuss in this subsection cover all the representative aspect ratio regimes: a conventional aspect ratio (with a realistic tokamak equilibrium), a tight aspect ratio (circular, $\epsilon_m = 0.63$ the location of the maximal bootstrap current), and an in-between aspect ratio (circular, $\epsilon_m = 0.51$) case.

The tight aspect ratio case CIRC1 is a Grad-Shafranov equilibrium with circular plasma boundary. The inverse aspect ratio at the outer boundary $\psi_N = 1$ is $\epsilon_a = 0.84$, where ψ_N is the poloidal flux normalized to its value at the separatrix. The density and temperature profiles (Fig. 3) are taken to be simple tanh-type pedestals. At the maximum location of the bootstrap current profile, $\epsilon_m = 0.63$ and the electron collisionality ν_{e*} is 0.9.

The case CIRC2 is a Grad-Shafranov equilibrium with circular boundary as well, but the inverse aspect ratio is $\epsilon_m = 0.51$ and $\nu_{e*} = 3.0$ at the maximum location of the bootstrap current. At the outer boundary $\psi_N = 1$, the inverse aspect ratio is $\epsilon_a = 0.88$. The density



FIG. 4. Density and Temperature profiles for case CIRC2.



FIG. 5. Density and Temperature profiles for JET. The pedestal width is widened to satisfy the local approximation

and temperature profiles (Fig. 4) for CIRC2 consist of a tanh-type pedestal and increase linearly from the pedestal top towards the magnetic axis. Around $\psi_N = 0$ and $\psi_N = 1$, the profiles are flattened.

Finally, for the conventional aspect ratio case we use the EFIT⁴¹ magnetic equilibrium of the JET H-mode discharge 85407, however, with an artificially broadened pedestal width (as shown in Fig. 5) in order to satisfy the local approximation. Thus, the plasma profile is not completely consistent with the Grad-Shafranov magnetic equilibrium. At the location of the maximum of the bootstrap current, $\epsilon_m = 0.29$, $\nu_{e*} = 1.7$. Unlike the two circular cases, the JET simulation includes the scrape-off layer.

We quantify the non-locality of the density and ion temperature profiles using the ratio of the ion orbit-width scale length ρ_b (half the maximal thermal ion orbit width) to the scale lengths of the corresponding gradients measured at the outer midplane. The orbit width was determined numerically using the conservation of the canonical toroidal angular momentum

$$R^2 \nabla \varphi \cdot (m \boldsymbol{v} + e \boldsymbol{A}) = m R v_T - e \psi = \text{const.}$$
⁽²⁾

where R is the major radius, φ the geometric toroidal angle, m the particle mass, v the particle's velocity, A the vector potential, v_T the particle's toroidal velocity and ψ the poloidal magnetic flux. Usually, the velocity v_T consists of the toroidal components of parallel velocity, magnetic inhomogeneity drifts, and $E \times B$ drift. However, to get a simpler and more practical estimate of ρ_b , we retain only the parallel velocity and neglect the orbit-squeezing effect from the $E \times B$ shear. Then, the maximal half-width $\delta \psi$ of the orbit of a thermal ion with $v_{\parallel} = 0$ at the minimum of the magnetic field at $\psi = \psi_0$ becomes

$$\delta \psi = \psi - \psi_0 \approx \frac{m_i R B_{\varphi}}{e B_{min}(\psi_0)} \sqrt{1 - \frac{B_{min}(\psi_0)}{B_{max}(\psi_0)}},$$

$$\Rightarrow \rho_b(\psi_0) = \frac{r(\psi_0 + \delta \psi) - r(\psi_0 - \delta \psi)}{2},$$
(3)

where $r = R_{+} - R_{c}$ is the outer midplane radius.

The bootstrap current obtained for case CIRC1 is shown in Fig. 6 a). Due to the effect of inaccurate momentum conservation discussed earlier in Sec. II, the previous version of XGC0 with an approximate linear collision operator yields higher current than XGCa and NEO. With the improved collision operator, the new version of XGC0 agrees very well with XGCa. The local simulation result from NEO agrees reasonably with the results of the global simulations with XGCa. At the peak of the current profile at $\psi_N \approx 0.89$, the difference between XGCa and NEO is 3.7% with the non-locality parameter $\rho_b/L_{T_i} \approx 0.14$. Since $\nu_{e*} < 1$, the Sauter formula also reproduces the XGCa result reasonably well. The non-locality parameters of density and ion temperature are shown in Fig. 6 b). They are well below 1 so that the small orbit width expansion used in NEO is approximately valid.

Reasonable agreement between XGCa and NEO is found also for CIRC2 and JET. The corresponding bootstrap current profiles and non-locality parameters are shown in Figs. 7 and 8. The Sauter formula over-predicts the bootstrap current in these cases. However, given that $\nu_{e*} > 1$ is outside of the accuracy range of the Sauter formula, this is not surprising and in line with similar findings in Refs. [21] and [25].



FIG. 6. a) Bootstrap current profile for the CIRC1 case, a circular limiter configuration with tight aspect ratio and lower plasma collisionality. XGC0 with inaccurate momentum conservation gives higher bootstrap current than the other codes and the Sauter formula. With accurate momentum conservation, XGC0 agrees well with XGCa and NEO. At the maximum location of the bootstrap current, NEO deviates from XGCa by 3.7%. The Sauter formula also reproduces the XGCa results well. b) The ratios of ion orbit width ρ_b to density and ion temperature gradient scale lengths are small enough to consider this case to be in the local regime.

B. Collisional suppression of the passing electron current in the edge pedestal

In order to elucidate the role of trapped and passing particles, and the collision effect in carrying the bootstrap current, we analyze the guiding center distribution functions obtained from our XGCa simulations. Some of the simulations discussed in this section are in the non-local regime with pedestal widths comparable to the ion orbit width. However, we will show that part of the difference we find between XGCa and the Sauter formula in the collisional edge can be explained without invoking finite orbit-width effects.

The reduction of the bootstrap current compared to the Sauter formula observed in the CIRC2 case and the modified JET case [see Figs. 7 a) and 8 a)] is a robust feature in the plateau-collisional regime. The bootstrap current profiles obtained from XGCa, NEO, and the Sauter formula for C-Mod (discharge 1120803014) and NSTX (discharge 132543) shown in Figs. 9 a) and c) are further proof. The simulations for C-Mod and NSTX were initialized with the experimentally measured density and temperature profiles shown in Figs. 10 a) and b). At the maximum location of the respective bootstrap current profiles as obtained from XGCa, the inverse aspect ratio and collisionality are $\epsilon_m = 0.31$ and 0.69, and $\nu_{e*} = 3.7$ and



FIG. 7. a) Bootstrap current profile for the CIRC2 case, a circular limiter configuration with a moderately small aspect ratio and high collisionality. XGCa and NEO show good agreement while the Sauter formula over-estimates the bootstrap current. b) The ratios of ion orbit width ρ_b to density and ion temperature gradient scale lengths show that this case is in the local regime.



FIG. 8. a) Bootstrap current profile for the modified JET case (discharge 85407 with an artificially broadened plasma profile to avoid finite orbit width effects). XGCa and NEO agree very well. b) Ratio of the ion orbit width ρ_b to the density and ion temperature gradient scale lengths.

2.9, respectively. The pedestal widths in the C-Mod and NSTX simulations are comparable to the ion orbit width, which means that local neoclassical theory and simulations are no longer valid.

While NEO and XGCa agree well at radial locations in the local regime, the current densities obtained from NEO for the realistic profiles used in the C-Mod and NSTX cases (Fig. 9) tend to be higher than the results found from XGCa in the steep pedestal region. However, NEO results still reproduce at least a large part of the reduction of j_b compared



FIG. 9. Bootstrap current profiles for a) C-Mod (discharge 1120803014) and c) NSTX (discharge 132543). The XGCa simulations were initialized with experimental density and temperature profiles. The corresponding ratios of ion orbit width to gradient scale lengths are shown in b) and d). The C-Mod and NSTX pedestals are in the non-local regime. While XGCa and NEO agree very well in the local regime at smaller ψ_N , away from the pedestal, the two codes differ in the non-local regime in the pedestal by 5.5% in C-Mod and 9% in NSTX, respectively, when comparing the respective maxima of the bootstrap current. The Sauter formula over-estimates the bootstrap current when compared to XGCa.

to Sauter shown in Figs. 7 a), 8 a), and 9 a), in which the pedestal collisionality is high. Therefore, the local neoclassical theory may also contain physics that explain at least some of the reduction. This is the focus of this section. Finite orbit width effects will be discussed in Sec. IIID.

The following analysis of the behavior of trapped and passing particles will prove that the observed difference between the Sauter model and the results of our numerical simulations are in part due to collisional suppression of the passing electron contribution to j_b . Here,



FIG. 10. Density and temperature profiles for a) C-Mod (discharge 1120803014) and b) NSTX (discharge 132543).

we utilize a feature common to all the XGCa simulations (and the NEO simulations in the local regime), namely that the ion contribution to the bootstrap current (measured in the laboratory frame) is small to negligible on most flux-surfaces. This is shown in Fig. 11 using the example of the circular limiter case CIRC2 (local regime, discussed in Sec. III A). This case was chosen because it demonstrates the good agreement of the ion and electron contributions to the bootstrap current in the local regime obtained with XGCa and NEO. Therefore, we concentrate on the electron distribution function in the following analysis while suppressing the species indices.

The electron distribution function f is measured in XGCa on each vertex of the configuration space mesh as function of the perpendicular velocity v_{\perp} and the parallel velocity v_{\parallel} . For the calculation of the bootstrap current, we write $f = f_M + \delta f$. Since the Maxwellian distribution f_M has no mean flow, it is sufficient to evaluate j_b using δf .

However, δf itself is not yet useful for the analysis of the bootstrap current in velocity space because it contains the Pfirsch-Schlüter current, which has to be removed by evaluating the orbit-average of the parallel flow moment of δf . The electron bootstrap current is given by

$$j_b = \left\langle \frac{\boldsymbol{j} \cdot \boldsymbol{B}}{B_0} \right\rangle = -e \left\langle \frac{B}{B_0} \int \mathrm{d}^3 v \, v_{\parallel} \delta f \right\rangle,\tag{4}$$

where e is the elementary charge. The flux surface average removes the Pfirsch-Schlüter currents. To identify the contributions to j_b from trapped and passing particles, we need to invert the order of the configuration and velocity space integrals. We follow the derivation in Refs. [42 and 43] and use the velocity coordinates $\varepsilon = (m/2)(v/v_{th})^2 + q\delta\phi/(eT)$ and



FIG. 11. Ion and electron contributions to the total bootstrap current for the case CIRC2, a circular limiter configuration in the local regime. The ion contribution to the bootstrap current is noticeable only at the pedestal foot ($\psi_N > 0.6$). The maximum of the ion flow at $\psi_N \approx 0.67$ has important consequences for the passing electron flow [cf. Fig. 12 c)]. XGCa and NEO show good agreement.

 $\lambda = \mu/\varepsilon$, where $v_{th} = \sqrt{eT/m}$ is the thermal velocity with the temperature T in eV, μ is the magnetic moment, and $\delta \phi = \phi - \langle \phi \rangle$ is the poloidal variation of the electric potential.

In the following, all velocities are normalized to the thermal velocity v_{th} , $\delta\phi$ is normalized to the temperature, the particle charge to the elementary charge, and the magnetic field Bto its minimum B_{min} on each flux-surface. Since the result of Eq. (4) cannot depend on a constant ϕ_0 added to $\delta\phi$, we choose ϕ_0 such that $q(\delta\phi + \phi_0) \ge 0$ and use the gauge $\delta\phi \rightarrow \delta\phi + \phi_0$. With σ being the direction of the parallel velocity, the corresponding Jacobian is

$$\int d^3 v = 2\pi v_{th}^3 B \sum_{\sigma} \int_0^\infty d\varepsilon \,\varepsilon \int_0^{\lambda_{max}} \frac{\sigma d\lambda}{|v_{\parallel}|}.$$
(5)

The flux-surface average can be written as

$$\langle A \rangle = \frac{\oint \mathrm{d}\theta \mathcal{J}\mathcal{A}}{\oint \mathrm{d}\theta \mathcal{J}} = \frac{\oint \mathrm{d}\theta \mathcal{J}\mathcal{A}}{\mathcal{V}'} \tag{6}$$

with an appropriate Jacobian \mathcal{J} . When evaluating the flux-surface average numerically, the Jacobian is given by the volume of the Voronoi cells around the vertices of the configuration space mesh. By exchanging the order of the velocity and configuration space integrations in Eq. (4), one obtains

$$j_b = -2e\pi v_{th}^4 B_{min} \int_0^\infty \mathrm{d}\varepsilon \int_0^{\lambda_{max}} \mathrm{d}\lambda \varepsilon \left[\sum_{\sigma} \frac{1}{\mathcal{V}'} \int \mathrm{d}\theta \mathcal{J} B^2 \delta f \right].$$
(7)

The boundaries of the integrals over θ are the bounce points $\pm \theta_b$ defined by $\epsilon(1-\lambda B)-q\delta\phi = 0$ for trapped electrons and $\pm \pi$ for passing electrons. In the new velocity space coordinates, the parallel and perpendicular velocities can be written as

$$v_{\perp} = \sqrt{2\varepsilon\lambda B},$$

$$v_{\parallel} = \sqrt{2[\varepsilon(1-\lambda B) - q\delta\phi]}.$$
 (8)

The definition of the bounce points follows immediately from the expression for v_{\parallel} . Note that the position of the bounce points depends on the potential. Therefore, particle trapping may be modified significantly if $q\delta\phi \sim 1$. This has been investigated by Chang in Ref. [44]. The boundaries of the λ integration are also energy dependent. Since we chose a gauge that ensures $q\delta\phi \geq 0$, the lower boundary is always 0. The upper boundary is given by

$$\lambda_{max} \equiv \max\left(0, \frac{1}{B}\left(1 - \frac{q\delta\phi}{\varepsilon}\right)\right) \tag{9}$$

at each vertex of the configuration space mesh. The trapped-passing boundary λ_c is the minimum of λ_{max} on each flux-surface. We now define

$$\{\delta f\} \equiv -2e\pi v_{th}^4 B_{min} \varepsilon \left[\sum_{\sigma} \frac{\sigma}{\int \mathrm{d}\theta \mathcal{J}} \int \mathrm{d}\theta \mathcal{J} B^2 \delta f \right],\tag{10}$$

and use this quantity rather than δf to analyze the velocity space properties of the bootstrap current. The same analysis could be performed for the ion distribution function, but is not shown here. While the resulting ion current is correct, the trapped-passing boundary defined above is only approximate because the ion orbit width is finite.

We begin with the analysis of the contributions from trapped and passing electrons in the edge pedestal, which are obtained by evaluating the velocity space integral in Eq. (7) with the proper boundaries. It has been known from the large aspect-ratio theories that the passing particles carry most of the current. However, we find that this is not true at low aspect ratio, where the passing particle fraction is small. Figures 12 a)-c) show the contributions of trapped and passing particles to the electron current in the edge pedestals of C-Mod, NSTX, and the circular case CIRC2. C-Mod has the lowest inverse aspect ratio ($\epsilon = 0.31$ at $\psi_N \approx 0.96$) among these cases and, thus, the smallest but still significant trapped particle region ($f_t = 0.67$) in the edge pedestal. At the peak of the electron bootstrap current, already one fourth of the total current is carried by trapped particles in the C-Mod pedestal. At $\psi_N \gtrsim 0.98$ in the C-Mod case, the trapped particle contribution survives while the passing particle current plummets with increasing collisionality. In NSTX, the trapped particle current dominates in the whole pedestal region because of the high trapped particle fraction ($\epsilon = 0.68$, $f_t = 0.86$ at $\psi_N = 0.954$). The density and temperature profiles become steeper towards the separatrix [cf. Fig. 9 d)], providing a stronger bootstrap current drive. The passing particle contribution is reduced further toward the separatrix due to the increased collisional friction.

It is interesting to note that the passing particle current in case CIRC2 [Fig. 12 c)] is even inverted. Comparison with Fig. 11 shows that the minimum of the passing electron current at $\psi_N \approx 0.67$ coincides with a maximum of the ion current. Note that the opposite signs of the ion and electron current mean that the ion and electron flows have the same direction. This suggests that the drag due to the friction with the ion flow inverts only the passing electron flow whereas friction cancels on average for trapped electrons.

By evaluating the integral in Eq. (7) only over the energy ε , one obtains the bootstrap current as a function of ψ_N and λ , denoted by $j_b(\psi_N, \lambda)$, which allows for a more detailed analysis of the trapped and passing particle currents. Figure 13 a) shows $j(\psi_N, \lambda)$ at $\psi_N =$ 0.49 ($\nu_{e*} = 1.7$), $\psi_N = 0.58$ ($\nu_{e*} = 3.2$), and $\psi_N = 0.63$ ($\nu_{e*} = 6.0$) for case CIRC2. The respective trapped-passing boundaries are indicated by vertical dotted lines. The trapped and passing particle contributions to the bootstrap current are given by the area under the curves of \tilde{j} in the trapped and passing region. Two important observations can be made immediately in Fig. 13 a). First, at small aspect ratio, the bootstrap current contribution from the trapped region is not sharply concentrated around the trapped-passing boundary λ_c as in case of large aspect ratio but rather exhibits a long tail deep into the trapped region (In pitch-angle space $\xi = v_{\parallel}/v = (1 - \lambda)^{1/2}$, the tail decays faster towards the deeply trapped region). Naturally, the current must vanish towards $\lambda = 1$ because the deeply trapped particles with bounce angles close to the outer midplane cannot carry current. Due to the slower decay in the trapped region, the trapped particle current becomes significant at small aspect ratio. Second, while the transition between the trapped and passing regions is relatively smooth at $\psi_N = 0.49$ (where the electron collisionality is relatively low, $\nu_{e*} \sim 1.7$), a sudden drop centered around $\lambda = \lambda_c$ develops at $\psi_N > 0.58$ and $\psi_N = 0.63$, where the electron collisionality is higher. The passing particle current even changes its sign at $\psi_N = 0.63 \ (\nu_{e*} = 6.0)$. This phenomenon is due to the drag on the passing electrons exerted by the oppositely flowing ions as shown in Fig. 12 c). Other simulations analyzed for the



FIG. 12. Trapped and passing particle contributions to the (electron component of the) bootstrap current for a) C-Mod (discharge 1120803014), b) NSTX (discharge 132543), and c) the circular limiter case CIRC2. The trapped particle current is negligible only in the large aspect ratio limit. Even in C-Mod, approximately 25% of the total electron current are provided by trapped electrons at $\psi_N \approx 0.96$. Due to the large trapped particle fraction in spherical tokamaks, trapped particles dominate the bootstrap current in the pedestal region of NSTX. In case CIRC2, passing particles dominate the bootstrap current in the core but trapped particles dominate at the maximum of the current profile.

present work display analogous features.

In order to make certain that the decrease of the passing current fraction with the collisionality is a universal phenomenon in different magnetic geometries, we evaluate the ratio of the amplitudes $\alpha_p \equiv \overline{\tilde{j}_b(\lambda < \lambda_c)}$ and $\alpha_t = \max[\tilde{j}_b(\lambda > \lambda_c)]$, which are related to the trapped



FIG. 13. a) Pitch-angle (λ) dependence of the electron component of the bootstrap current for the circular limiter case CIRC2 on three different flux-surfaces. Vertical dotted lines mark the boundary between passing ($\lambda < \lambda_c$) and trapped regions ($\lambda > \lambda_c$). Horizontal and diagonal dotted lines indicate the mean amplitudes α_p and α_t in the passing and trapped regions that are connected to the passing and trapped currents via Eq. (11). With increasing collisionality, the ratio α_p/α_t decreases; α_p becomes negative. b) α_p/α_t for various XGCa simulations of circular limiter and realistic tokamak configurations. The plateau at small $\hat{\nu}_e = \nu_{e*}\epsilon^{3/2}$ indicates that the role of trapped and passing particles is determined mainly by the sizes of the trapped and passing regions in phase space. Decreasing α_p/α_t with increasing $\hat{\nu}$ indicates that collisions reduce the relative importance of the passing electrons for the bootstrap current.

and passing particle currents via

$$j_p = \alpha_p \lambda_c,$$

$$j_t \approx \frac{1}{2} \alpha_t \left(1 - \lambda_c \right).$$
(11)

Notice here that the trapped current is approximated as a triangle in λ -space. Dividing α_p by α_t removes the dependence on the plasma gradients. Figure 13 b) shows a plot of α_p/α_t versus $\hat{\nu}_e = \nu_{e*} \epsilon^{3/2}$, combining data from cases CIRC1, CIRC2, JET (cf. Sec. III A), C-Mod, NSTX (this section), DIII-D (Sec. III D) and ASDEX-Upgrade. For this plot, we used only data points in the vicinity of the pedestal. This shows that collisions reduce the relative contribution of the passing electrons to the bootstrap current in addition to the effect from the sizes of the trapped and passing regions.

C. Role of trapped and passing particles at large aspect ratio

In order to demonstrate the connection of our results to the conventional neoclassical theories, we analyze the ratio j_t/j_p , and the width of the collisional boundary layer between trapped and passing electrons using the XGCa simulation in JET geometry. In particular, we verify the XGCa results against two results from local neoclassical theory in the large aspect ratio limit. It is well known⁴, that, in the limit of large aspect ratio, the bootstrap current is carried predominantly by the passing particles. The small contribution from the trapped particles originates from a boundary layer between the passing and trapped region in velocity space, which is due to collisional mixing. The width of this boundary layer is approximately $\delta \xi \approx (\nu_{e*} \epsilon^{3/2})^{1/3}$ (Ref. [4]), where $\xi = (1 - \lambda)^{1/2}$ is a more commonly used pitch angle variable. The trapped particle contribution to the bootstrap current is a factor of ϵ smaller than the one from the passing particles⁴.

The pitch angle dependence of the bootstrap current, $\tilde{j}_b(\psi_N, \xi)$ is shown in Fig. 14 a) for the flux-surface $\psi_N = 0.08$, where $\epsilon = 0.08$ and $\nu_e = 0.26$. The trapped-passing boundary is located at $\xi_c = 0.38 \approx (2\epsilon)^{1/2}$, and the collisional boundary width is estimated to be $\delta \xi \approx 0.18$. The estimated width of the boundary layer agrees reasonably with the decay length of $\tilde{j}_b(\psi_N, \xi)$ in the trapped region. Figure 14 b) shows the ratio of trapped-topassing electron bootstrap current. In agreement with local neoclassical theory, $j_t/j_p \approx \epsilon$ to reasonable accuracy up to $\psi_N \approx 0.5$ or $\epsilon \approx 0.2$. At $\psi_N > 0.5$, j_t/j_p becomes greater than ϵ as the width of the collisional boundary layer becomes comparable to the width of the passing particle region, $\delta \xi_p \approx 1 - (2\epsilon)^{1/2}$.

D. Finite orbit width effects

In the edge pedestal of a tokamak, the ion orbit width ρ_b can easily be comparable to the density and temperature gradient scale lengths L_n and L_{Ti} . When this happens, the small orbit-width approximation used by local neoclassical codes and by analytic theories breaks down and their accuracy becomes questionable. In the simulations using the C-Mod and NSTX geometries that were discussed in the previous section, we found differences between local and global simulations. Therefore, we address the importance of finite orbit-width effects for the pedestal bootstrap current in this section.



FIG. 14. a) Pitch-angle ($\xi = (1 - \lambda)^{1/2}$) dependence of the electron component of the bootstrap current for the modified JET case at $\psi_N = 0.08$, where $\epsilon = 0.08$ and $\nu_{e*} = 0.26$. The red vertical dotted line marks the boundary between passing ($\xi > \xi_c$) and trapped regions ($\xi < \xi_c$), the blue dotted line indicates the approximate extent of the collisional boundary layer into the trapped region according to Ref. [4]. b) Ratio of trapped to passing electron current, j_t/j_p . At $\psi_N < 0.5$, $j_t/j_p \approx \epsilon$ can be seen, in agreement with the local neoclassical theories⁴. At $\psi_N > 0.5$, the width of the collisional boundary layer becomes comparable to the width of the passing particle region leading to deviations of j_t/j_p from ϵ .

In the previous section, we noted that the ion contribution to the bootstrap current is small compared to the electron contribution. Due to the smallness of the electron orbit width, one might be tempted to conclude from the absence of a sizable ion contribution to j_b that finite orbit-width effects cannot be important. We will show, however, that finite orbit-width effects enter the electron flow indirectly via the radial electric field E_r .

If one considers the source of the bootstrap current, it may seem peculiar at first sight that the ion current is much smaller than the electron current. Since the electron orbit width is a factor of $(m_e/m_i)^{1/2}$ smaller than the ion orbit width but their thermal velocity is a factor of $(m_i/m_e)^{1/2}$ larger, the current drive of ions and electrons for given gradient appears to be comparable when $T_i \approx T_e$. However, this argument does not hold in the presence of a radial electric field E_r , which is a direct consequence of the ion orbit motion, needed to keep the plasma quasi-neutral. Since it is a radial force, the electric field drives a parallel flow in the same way as a pressure gradient does. Because E_r is in the same direction as the pressure gradient, the radial electric field counteracts the ion parallel flow from ∇p but adds to the ∇p drive for electrons. This can be seen from the local neoclassical expression for the parallel ion flow⁴

$$nu_{\parallel i} = -\frac{I}{m\Omega_i} \left(\frac{\mathrm{d}\langle p_i \rangle}{\mathrm{d}\psi} + en \frac{\mathrm{d}\langle \phi \rangle}{\mathrm{d}\psi} \right) + K(\psi)B,\tag{12}$$

where Ω_i is the ion cyclotron frequency, p_i is the ion pressure, and ϕ is the electrostatic potential. Usually, the pressure and potential gradient terms – the ion diamagnetic and $E \times B$ flows – tend to cancel, resulting in E_r being in the same direction as the pressure gradient. A similar expression can be derived for the electron flow but with the opposite sign for the radial electric field term due to the negative electron charge. Therefore, the radial electric field terms cancel in the local regime when the ion and electron flows are added to calculate the current. This cancellation is exact only if the ion orbit width is small compared to the gradient scale lengths. If $\rho_b/L_P \sim 1$, the orbit-averaged electric fields experienced by ions and electrons differ, and the electric field term does not cancel any longer. Thus, the finite orbit-width effects influence the bootstrap current indirectly via the electron current due to the incomplete cancellation of the E_r -drive terms.

Figure 15 illustrates the consequence of the electric field effect. It shows the ion and electron contributions to the bootstrap current obtained from XGCa and NEO for the C-Mod case that was discussed in Sec. III B [cf. Fig. 9 a) and b)]. At $\psi_n < 0.94$ (where $\rho_b/L_{Ti} \approx \rho_b/L_n < 0.2$), local neoclassical physics are valid. The ion and electron currents of the local NEO and the global XGCa simulation agree very well. At $\psi_N > 0.94$, large differences between XGCa and NEO arise although the radial electric field obtained with XGCa was used as input for NEO. This is because, in contrast to a global code, the same electric field acts on ions and electrons in a local neoclassical code regardless of the ion orbit width. As a consequence, a local simulation cannot reproduce the ion and electron currents of a global simulation when $\rho_b/L_P \sim 1$ as can be seen in Fig. 15. The plasma flow calculation performed in a local code in an edge pedestal would be incorrect.

There is another finite orbit-width effect that can affect the ion flow directly. It can be understood using the finite orbit-width expression for the ion thermal conductivity derived by Chang⁴⁵. Since ions experience different collision frequencies and safety factors on the inner and outer legs of their orbits, the finite orbit width correction is proportional to

$$\Delta_{\psi} \left(\nu^{-1} \frac{\mathrm{d}\nu}{\mathrm{d}\psi} + q^{-1} \frac{\mathrm{d}q}{\mathrm{d}\psi} \right),\,$$



FIG. 15. Ion and electron contributions to the total bootstrap current for C-Mod (discharge 1120803014³⁴, non-local regime). The ion contribution to the bootstrap current found from XGCa is small throughout the pedestal. At $\psi_N > 0.94$ ion and electron currents obtained from NEO deviate significantly from XGCa. This deviation demonstrates the limit of local neoclassical theory and simulation. At $\psi_N > 0.95$, the ratio of ion orbit width to the gradient scale lengths exceeds 0.5.

where ν is the ion collision frequency and Δ_{ψ} the ion orbit width ρ_b in units of poloidal flux. This correction is applied to the analytic fitting formula introduced in Sec. IV.

1. Assessment of the finite orbit-width effect from density and ion temperature pedestals

We study the impact of the non-locality from the density and the ion temperature gradients using a circular limiter configuration. The magnetic field geometry is identical to the case CIRC2 discussed in Secs. III A and III B, but the density and temperature profiles are chosen differently (still as tanh-type pedestals). In the cases CIRC2a and CIRC2b, the bootstrap current is driven mainly by the ion temperature gradient while the density gradient is sub-dominant ($\rho_b/L_{Ti} \approx 0.23$, $\rho_b/L_n < 0.03$, $\epsilon = 0.49$, $\nu_{e*} = 2.6$ at $\psi_N = 0.55$ for CIRC2a, and $\rho_b/L_{Ti} \approx 0.28$, $\rho_b/L_n \ll 1$, $\epsilon = 0.42$, $\nu_{e*} = 1.6$ at $\psi_N = 0.43$ or CIRC2b). The CIRC2b case examines a stronger ion temperature non-locality than the CIRC2a case. In contrast, the density gradient dominates the current drive in the case CIRC2c ($\rho_b/L_{Ti} \approx 0.05$, $\rho_b/L_n \approx 1.3$, $\epsilon = 0.55$, $\nu_{e*} = 0.24$ at $\psi_N = 0.65$). The current profiles and the non-locality parameters ρ_b/L_n and ρ_b/L_{Ti} for cases CIRC2a, CIRC2b, and CIRC2c are shown in Figs. 16, 18, and 20.

The case CIRC2a exhibits a similar collisional reduction of the bootstrap current compared to the Sauter formula as discussed in Sec. III B. Around the maximum of ρ_b/L_{Ti} at $\psi_N \approx 0.5$, the current measured in XGCa is lower than the current obtained from NEO by approximately 10% of the maximum XGCa bootstrap current. Where the profiles are in the local regime, XGCa and NEO agree well. For case CIRC2b, the deviation between XGCa and NEO is up to 40% of the maximum XGCa current at the maximum of ρ_b/L_{Ti} at $\psi_N \approx 0.47$. In contrast, in the CIRC2c case, XGCa and NEO show good agreement with a deviation of only 4% of the maximum XGCa current even at the maximum of ρ_b/L_N in spite of the very narrow density pedestal. There are large differences between the ion flows from XGCa and NEO due to finite orbit-width effects (not shown). But because the ion flow is significantly smaller than the electron flow, its influence on the total current is small. Since the collisionality is low in case CIRC2c, the Sauter formula agrees with the XGCa simulations as well.

Based on the findings in the cases CIRC2a, CIRC2b, and CIRC2c, we conclude that one prerequisite for the occurrence of finite orbit-width effects in the bootstrap current is the existence of a non-local ion temperature gradient. This observation shares commonalities with Landreman *et al.*²⁵, who found a finite orbit width correction to the bootstrap current that is proportional to the product of density and ion temperature gradient in the case $\rho_b/L_n \sim 1$ and $\rho_b/L_{Ti} \ll 1$.

2. Assessment of the finite orbit-width effect at different collisionality

In order to investigate the collisionality dependence of the finite orbit-width effect on the bootstrap current, we ran three simulations using the geometry of DIII-D discharge 96333²⁸ with artificial temperature profiles (all with tanh-type edge pedestals) that are shown in Fig. 22 a). The pedestal top temperatures are 250, 500, and 1,000 eV. Ion (electron) collisionalities ν_{i*} (ν_{e*}) at the peaks of the corresponding bootstrap current profiles decrease with increasing pedestal height and are approximately 3.7 (5.2), 1.4 (2.0), 0.5 (0.9). The corresponding aspect ratios at the maximum bootstrap current are $\epsilon_m = 0.36$, 0.36 and 0.35. As shown in Fig. 22 b), the case with the highest collisionality has the lowest bootstrap current. All



FIG. 16. a) Bootstrap current profile in the mildly non-local ion temperature pedestal of case CIRC2a, which uses the same magnetic field data as the circular limiter case CIRC2 (Fig. 7), but the ion temperature pedestal is narrower. b) Ratio of the ion orbit width to the density and ion temperature gradient scale lengths. As long as $\rho_b/L_{Ti} \leq 0.2$, XGCa and NEO show good agreement. The deviation between XGCa and NEO at $\psi_N \approx 0.5$ is approximately 10% of the maximum of the XGC bootstrap current.



FIG. 17. Density and temperature profiles for the circular case CIRC2a.

three XGCa simulations exhibit much lower current than predicted by the Sauter formula. The local simulations with NEO can account for some of that difference but still yield larger current than XGCa, with the difference increasing as ν_{e*} decreases. The relative difference between XGCa and NEO at $T_{i,top} = 1$ keV is 12.3% at $\psi_N = 0.968$. In contrast, the relative difference between XGCa and NEO at $T_{i,top} = 250$ eV is only 5.3% at $\psi_N = 0.982$. Since the non-local effect should become weaker at higher collisionality as the ion orbits become narrower and are interrupted more frequently, this behavior is expected.



FIG. 18. a) Bootstrap current profile in the stronger non-local ion temperature pedestal of the case CIRC2b, which uses the same magnetic field data as the circular limiter case CIRC2 (Fig. 7). The ion temperature pedestal was made narrower than in the case CIRC2a to enhance finite orbit-width effects. b) Ratio of the ion orbit width to the density and ion temperature gradient scale lengths. The ion temperature gradient is dominant at the density pedestal top. The deviation between XGCa and NEO at $\psi_N \approx 0.47$ is approximately 40% of the maximum of the XGCa bootstrap current.



FIG. 19. Density and temperature profiles for the circular case CIRC2b, which exhibits a stronger non-locality in the ion temperature gradient than the case CIRC2a.

IV. IMPROVEMENT OF THE SAUTER FORMULA

We use the results of our XGCa simulations to improve Sauter's formula for non-local edge pedestals by introducing a few correction factors. The first correction is constructed such as to reproduce the collisional damping of the passing electron current discussed in



FIG. 20. a) Bootstrap current profile in the strongly non-local density pedestal of the case CIRC2c, which uses the same magnetic field data as the circular limiter case CIRC2 (Fig. 7), however, with stronger current drive from the density gradient than the ion temperature gradient. b) Ratio of the ion orbit width to the density and ion temperature gradient scale lengths. The ion temperature gradient is very small compared to the density gradient. Despite the narrow density pedestal, the deviation between XGCa and NEO at $\psi_N \approx 0.65$ is only approximately 4% of the maximum of the current profile. The result implies that the finite orbit width-effect is mostly caused by the ion temperature gradient.



FIG. 21. Density and temperature profiles for the circular case CIRC2c with the density gradient dominating the bootstrap current drive.

Sec. III B. The second correction accounts for the finite orbit-width effects as seen in Sec. III D. Finally, the third correction is to restore the correct asymptotic behavior in the limit $\nu_{e*} \gg 1$.

Our modifications are based on the Sauter formula¹⁴, which expresses the bootstrap



FIG. 22. Collisionality scan using DIII-D geometry. a) Assumed density and temperature profiles in DIII-D geometry with pedestal top temperatures of $T_{i,top} \approx 250$, 500, 1,000 eV. b) Corresponding bootstrap current profiles. With increasing pedestal height and decreasing collisionality, the difference between the bootstrap currents obtained with XGCa and NEO increases due to the less frequent interruptions of the ion orbits. The relative differences between the maxima of the XGCa and NEO bootstrap current profiles are 5.3% for $T_{i,top} \approx 250$ eV, 6.3% for $T_{i,top} \approx 500$ eV, and 12.3% for $T_{i,top} \approx 1,000$ eV. The reduction of j_b compared to the Sauter formula can no longer be explained by the local collisional effect alone as discussed in Sec. III B.

current j_b as

$$j_b \equiv \left\langle \frac{\boldsymbol{j} \cdot \boldsymbol{B}}{B_0} \right\rangle = -\frac{Ip_e}{B_0} \left(L_{31} \frac{P}{p_e} \frac{\mathrm{d}\ln P}{\mathrm{d}\psi} + L_{32} \frac{\mathrm{d}\ln T_e}{\mathrm{d}\psi} + L_{34} \alpha_i \frac{T_i}{ZT_e} \frac{\mathrm{d}\ln T_i}{\mathrm{d}\psi} \right). \tag{13}$$

In this formula, ψ is the poloidal magnetic flux, $I(\psi) = RB_{\varphi}$, B_0 is the magnetic field on the magnetic axis, Z is the ion charge number, $T_{i/e}$ are the ion and electron temperatures, p_e is the electron pressure, and P is the total pressure. The coefficients L_{3j} and α_i are defined in App. A.

A. Collisional correction in the local regime

For the correction of the Sauter formula at practical edge collisionalities ($\nu_{e*} < 100$), we multiply Eq. (13) by a Padé term of the form

$$\beta_{col} \equiv \frac{1 + c_1 c_2 \nu_{e*}^2}{1 + c_1 \nu_{e*}^2}.$$
(14)

In the limit of infinite collisionality, $\beta_{col} \rightarrow c_2$. For simplicity, we choose c_2 to be a constant. Therefore c_1 must depend on the aspect ratio or, alternatively, the trapped particle fraction: hence, $c_1 = c_1(\epsilon)$. By examining the collisional XGCa simulation results, $c_2 \approx 0.15$ was identified as initial guess. For convenience, we divide the XGCa data set into four aspect ratio bins covering the simulation data from C-Mod and ASDEX-Upgrade ($0.28 \leq \epsilon \leq$ 0.322), modified DIII-D ($0.352 \leq \epsilon \leq 0.362$, circular cases ($0.4 \leq \epsilon \leq 0.55$), and NSTX ($0.6 \leq \epsilon \leq 0.72$). For each of the bins, we varied c_1 using Newton's method to minimize the difference between the modified formula and the XGCa data. Reasonable agreement between the modified formula and XGCa is obtained for collisional cases by describing the aspect ratio dependence of c_1 using two hyperbolic tangent functions

$$c_{1}(\epsilon) = 0.5 \left\{ \left[(a_{1} - b_{1}) \tanh\left(\frac{2(\epsilon - \epsilon_{c,1})}{w_{1}}\right) + (a_{1} + b_{1}) \right] - \left[(a_{2} - b_{2}) \tanh\left(\frac{2(\epsilon - \epsilon_{c,2})}{w_{2}}\right) + (a_{2} - b_{2}) \right] \right\}$$
(15)

with $a_1 = a_2 = 0.044$, $b_1 = 0$, $b_2 = 0.006$, $\epsilon_{c,1} = 0.15$, $\epsilon_{c,2} = 0.51$, $w_1 = 0.1$, $w_2 = 0.15$, and $c_2 = 0.15$ as initial guess. Note that the ϵ -dependence of c_1 for $\epsilon < 0.28$ is heuristic, chosen to ensure that β_{col} becomes 1 in the limit of high aspect ratio. After all the corrections mentioned above are added to the Sauter formula, these coefficients are readjusted using the values described here as initial guess.

B. Finite orbit-width corrections

During the numerous attempts to fit an analytic expression to the XGCa bootstrap current data in the non-local regime, we found that the finite orbit-width corrections associated with steep ion temperature pedestals need to be added at two different places. One correction factor damps the overall strength of the bootstrap current in addition to the collisional damping factor β_{col} . Therefore, we define $\beta_{damp} = \beta_{col} + \beta_{\nabla T_i}$. The second correction factor modifies the transport coefficient L_{34} .

The best fit to the XGCa bootstrap current data is found for $\beta_{\nabla T_i}$ in the form

$$\beta_{\nabla T_i} \equiv -\frac{\Lambda_1 \nu_{i*}}{1 + \Lambda_2 \nu_{i*}^2 + \Lambda_3 \nu_{i*}^4} \left(1 - \epsilon\right)^{\Lambda_4} \Delta_R^{\Lambda_5} \frac{\Delta_\psi}{T_i} \left| \frac{\mathrm{d}T_i}{\mathrm{d}\psi} \right|,\tag{16}$$

where $\Delta_R \equiv dR_c/dR_{min}$ is the Shafranov shift, and Δ_{ψ} is the ion orbit width ρ_b in units of poloidal flux. The Shafranov shift reflects the dependence of the flow shear on $\nabla \psi$ at the outer midplane, and the aspect ratio term accounts for the observation that the deviations between XGCa and the Sauter formula becomes smaller with decreasing aspect ratio. A more detailed study, which is not discussed here, shows that the physical origin of Eq. (16) is the viscous damping of the ion flow due to finite ion orbit-width. Again, the coefficients Λ_i will be determined by a fitting procedure after all the corrections have been added to Eq. (13).

The finite orbit-width correction to L_{34} is found to be similar to the correction of the neoclassical heat conductivity derived by Chang⁴⁵. It is proportional to

$$\Delta_{\psi} \left(\nu^{-1} \frac{\mathrm{d}\nu}{\mathrm{d}\psi} + q^{-1} \frac{\mathrm{d}q}{\mathrm{d}\psi} \right) \frac{\mathrm{d}T_i}{\mathrm{d}\psi}.$$
 (17)

This correction reflects the different collision frequencies and safety factors, that ions experience between the inner and outer legs of the ion orbits. We modify the L_{34} term in Eq. (13) as follows:

$$L_{34}\alpha_i \frac{1}{ZT_e} \frac{\mathrm{d}T_i}{\mathrm{d}\psi} \to L_{34}^{\Delta_b} \alpha_i \frac{1}{ZT_e} \frac{\mathrm{d}T_i}{\mathrm{d}\psi} = L_{34}\alpha_i \frac{1}{ZT_e} \frac{\mathrm{d}T_i}{\mathrm{d}\psi} \left\{ 1 - 2\Delta_\psi \left(-\frac{3}{2}L_{T_i}^{-1} + L_n^{-1} + L_q^{-1} \right) \right\}.$$
(18)

Here, $L_X^{-1} \equiv X^{-1} dX/d\psi$ is the gradient scale length of X in units of poloidal flux.

C. Correction of the asymptotic behavior in the collisional limit

A closer look at the polynomial dependence of the coefficients L_{3j} defined in Eq. (A8) on the trapped particle fractions $f_{t,eff}^{3j}$ reveals a minor weakness of the Sauter formula. According to Hinton and Hazeltine⁴, and other theories, the coefficients L_{3j} must be proportional to ν_{e*}^{-2} in the collisional regime $\nu_{e*} \gg 1$. But the leading order term in Eq. (A8) decays as ν_{e*}^{-1} in the limit $\nu_{e*} \gg 1$. Therefore, the Sauter formula will always yield systematically too high predictions for the bootstrap current in the strongly collisional limit. We note here that Sauter's coefficients were fitted to the results of numerical simulations for a practical range of collisionalities and were not intended for impractically high ν_{e*} .

In order to correct the asymptotic behavior of the Sauter formula in the strongly collisional limit, we blend the Sauter formula with a bootstrap current formula by Helander and Sigmar⁴⁶ – in the following referred to as "Helander formula" – using the viscosity coefficients for the collisional regime by Hirshman and Sigmar⁴⁷. The Helander formula was derived for arbitrary aspect ratio and has the correct asymptotic behavior. Neglecting the loop voltage $(E_{\parallel} = 0)$, Eq. (12.51) in Ref. [46] can be expressed as

$$j_b \equiv \left\langle \frac{\boldsymbol{j} \cdot \boldsymbol{B}}{B_0} \right\rangle = -\frac{Ip_e}{B_0} \left(L_{31}^{(h)} \frac{P}{p_e} \frac{\mathrm{d}\ln P}{\mathrm{d}\psi} + L_{32}^{(h)} \frac{\mathrm{d}\ln T_e}{\mathrm{d}\psi} + L_{31}^{(h)} \alpha_i^{(h)} \frac{T_i}{ZT_e} \frac{\mathrm{d}\ln T_i}{\mathrm{d}\psi} \right) \tag{19}$$

with the coefficients $L_{31}^{(h)}$, $L_{32}^{(h)}$, and $\alpha_i^{(h)}$ defined in App. B. Notice that $L_{34}^{(h)} = L_{31}^{(h)}$.

In order to avoid confusion, we will denote the coefficients L_{3j} in the original Sauter formula by $L_{3j}^{(s)}$ in the remainder of this section. In the limit of infinite collisionality, the leading order terms in $L_{31}^{(h)}$ and $L_{32}^{(h)}$ are indeed $O(\nu_{e*}^{-2})$ as predicted in Ref. [4]. In order to correct the asymptotic behavior of coefficients of the Sauter formula, we multiply them by a Padé factor of the form

$$\gamma_{3j} = \frac{1 + c_3 \nu_{e*}}{1 + c_4 \nu_{e*}^2}.$$
(20)

For $\nu_{e*} \gg 1$, γ_{3j} becomes $c_3/(c_4\nu_{e*})$. If we denote the leading order terms of the coefficients L_{3j} in the collisional limit as $L_{3j}^{(1)}$ and $L_{3j}^{(2)}$ for terms $O(\nu_{e*}^{-1})$ and $O(\nu_{e*}^{-2})$, respectively, the obvious choice of c_3 and c_4 is

$$c_{3} = 1,$$

$$c_{4} = \frac{L_{3j}^{(s,1)}\nu_{e*}}{L_{3j}^{(h,2)}\nu_{e*}^{2}}.$$
(21)

However, this choice does not control at which ν_{e*} the bootstrap current coefficients switch to the asymptotic behavior of Helander's coefficients. Therefore, we multiply c_3 and c_4 by a common factor c_5 , which cancels in the collisional limit, but makes sure that $c_4c_5\nu_{e*,c}^2 = 1$ at the critical collisionality at which we wish to place the transition to Helander's coefficients. Since the correction terms β_{col} , $\beta_{\nabla T_i}$, and $L_{34}^{\Delta_b}$ improve the accuracy of the Sauter formula sufficiently, we place the transition between the Sauter and the Helander formula at collisionalities higher than those in our simulations, where the leading order term $L_{31}^{(h,2)}$ becomes dominant, i.e. $|L_{31}^{(h,4)}|/L_{31}^{(h,2)} = 10^{-2}$. Thus, the correct asymptotic behavior is ensured by the substitution $L_{3j}^{(s)} \to \gamma_{3j}L_{3j}^{(s)}$. The final interpolation coefficients are

$$\gamma_{3j} = \frac{1 + c_5 \nu_{e*}}{1 + c_5 (L_{3j}^{(s,1)} / L_{3j}^{(h,2)}) \nu_{e*}},$$

$$c_5 = \frac{L_{31}^{(h,2)} (\nu_{e*,c})}{L_{31}^{(s,1)} (\nu_{e*,c}) \nu_{e*,c}}.$$
(22)

The ion flow coefficient α_i is a constant in the strongly collisional limit. Therefore, the

interpolation coefficient for α_i is defined as

$$\gamma_{\alpha} = \frac{1 + c_5 \nu_{i*}^2}{1 + c_5 (\alpha_i^{(s,0)} / \alpha_i^{(h,0)}) \nu_{i*}^2}.$$
(23)

Explicit expressions for $L_{3j}^{(s,1)}$, $L_{3j}^{(h,2)}$, $\alpha_i^{(s/h,0)}$, and $\nu_{e*,c}$ are given in App. B.

One further complication in this interpolation is a singularity in $L_{31}^{(s,1)}$ arising from this coefficient's dependence on the collisionless trapped particle fraction [cf. Eq. (A9)]. The denominator of $f_{t,eff}^{31}$ contains the term $(1 - f_t)\nu_{e*}/Z$, which vanishes for $f_t = 1$ and makes $L_{31}^{(s)}$ of order $\nu_{e*}^{-1/2}$ instead of ν_{e*}^{-1} . This causes a singularity in $L_{31}^{(s,1)}$. To avoid this singularity, we choose to modify $f_{t,eff}^{31}$ slightly to

$$f_{t,eff}^{31} = \frac{f_t}{1 + (1 - 0.1f_t)\sqrt{\nu_{e*}} + 0.5(1 - 0.99f_t)(\nu_{e,*}/Z)}.$$
(24)

The deviation of the resulting $L_{31}^{(s)}$ from the original definitions in Eqs. (A8)-(A9) is largest at Z = 1 and high collisionality. However, for ν_{e*} as high as 50, it is less than 3%. This is perfectly acceptable taking into account the fact that the correction factor γ_{31} starts to become dominant at this collisionality.

D. Improved bootstrap current formula

Combining all of our modifications, the modified Sauter formula becomes

$$j_{b} = -\beta_{damp} \frac{Ip_{e}}{B_{0}} \left\{ \gamma_{31} L_{31} \frac{P}{p_{e}} \frac{\mathrm{d}P}{\mathrm{d}\psi} + \frac{\gamma_{32} L_{32}}{T_{e}} \frac{\mathrm{d}T_{e}}{\mathrm{d}\psi} + (\gamma_{34} L_{34})(\gamma_{\alpha} \alpha_{i}) \frac{1}{ZT_{e}} \left\{ 1 - 2\Delta_{\psi} \left(-\frac{3}{2} L_{Ti}^{-1} + L_{n}^{-1} + L_{q}^{-1} \right) \right\} \frac{\mathrm{d}T_{i}}{\mathrm{d}\psi} \right\}.$$
(25)

To optimize the agreement of this modified formula with XGCa, we adjusted the free parameters in β_{damp} [Eqs. (14) and (16)] using a genetic algorithm optimization. The goal of this process is to minimize the deviation of the bootstrap current predicted by Eq. (25) from the XGCa results. We define this deviation as

$$\delta j \equiv \frac{j_b^{(formula)} - j_b^{(XGCa)}}{j_b^{(XGCa)}}.$$
(26)

The fitness function used to quantify the quality of a given parameter set was chosen to be

$$\Gamma = \sqrt{\operatorname{Var}(\delta_j)} \left(1 + 10 \,\overline{\delta j} \right), \tag{27}$$

where $\overline{\delta j}$ is the arithmetic mean of δj . The inclusion of the mean error in the fitness function ensures that the optimization routine prefers parameter sets with lower mean error over those with higher mean error if their standard deviations are identical. In order to exclude small values of j_b from the optimization, all the data points with $j_b < \max(j_b)_{ped}/2$ are discarded. The fitting procedure converged to the following parameters:

$$c_{2} = 0.2947, a_{1} = a_{2} = 0.0488, b_{1} = 0, b_{2} = 0.0086,$$

$$\epsilon_{c,1} = 0.15, \epsilon_{2,c} = 0.5099, w_{1} = 0.1, w_{2} = 0.15,$$

$$\Lambda_{1} = 19.1702, \Lambda_{2} = 1.9056, \Lambda_{3} = 0.0106, \Lambda_{4} = 2.994, \Lambda_{5} = 0.9958.$$
 (28)

The new formula Eq. (25) reproduces the numerical results from XGCa much better than the original Sauter formula, Eq. (13), or Koh *et al.*'s formula²¹. Comparisons between XGCa results, NEO results, Eq. (13), and Eq. (25) are shown in Figs. 7-9, and Figs. 16-20. The improved quality of our modified formula is illustrated also in Figs. 23 a) and b), which show the bootstrap currents from the Sauter and the modified Sauter formula plotted versus the numerical result from XGCa. Large deviations from the diagonal line, which marks the line of exact agreement, can be observed for the Sauter formula. The average relative error of the original Sauter formula is $\overline{\delta j} = 23.8\%$ with a standard deviation of $\sigma = 26.1\%$. The modified formula deviates from the numerical results on average by only $\overline{\delta j} = 1.4 \cdot 10^{-3}$ with $\sigma = 4.5\%$.

V. SUMMARY AND CONCLUSIONS

The present numerical study elucidates the mechanisms that determine the bootstrap current in the steep edge pedestal plasma conditions. Using the numerical data obtained with the neoclassical codes XGC0, XGCa, and NEO, we analyzed the contributions from ions and electrons, and from trapped and passing particles. We investigated the non-local effects introduced by edge pedestal widths being comparable to the ion orbit width, and we studied the limits of local neoclassical theory. Based on the numerical results of XGCa, we modified the Sauter formula¹⁴ to improve its accuracy significantly for parameters relevant to present day tokamak experiments.

Koh *et al.*'s modification of the Sauter formula²¹ needed to be re-investigated because a benchmark of the old version of XGC0 that was used to develop that formula and the newly



FIG. 23. Statistical accuracy of Eqs. (13) and (25) as compared to the XGCa result. The horizontal axis represents the results from XGCa, the vertical axis is the result of the bootstrap formulas. The dotted lines indicate the σ and 3σ intervals. Our modified formula fits the numerical results obtained with XGCa significantly better than the Sauter formula.

developed gyrokinetic neoclassical code XGCa identified a small inaccuracy in the momentum conservation of the XGC0 collision operator. Even though the degree of inaccuracy was small, it had a noticeable impact on the electron current due to the smallness of the electron mass. Since this inaccuracy increased the bootstrap current, the enhancement of j_b in spherical tokamaks reported by Koh *et al.* disappeared in XGC0 simulations after the collision operator had been corrected. Cross-verification of XGCa, new XGC0, and NEO [Figs. 6-8] in the local regime ($\rho_b \ll L_{\nabla}$) demonstrated excellent agreement in the bootstrap current.

In the collisional regime, the Sauter formula has been found to overestimate the bootstrap current significantly even in the local regime, similar to reports in Refs. [15 and 21]. A detailed analysis of the velocity-space properties of the bootstrap current in Sec. III B revealed that this deviation is caused by friction of the passing electron current with the ions at collisionalities $\nu_{e*} > 1$. As a result of this friction, the passing electron current in the pedestal. At high enough collisionality this phenomenon occurs even in the edge of conventional aspect ratio tokamaks. Most of the current is usually carried by electrons rather than ions. We observe non-negligible ion contributions only in the steepest part of the edge pedestal when the diamagnetic and $E \times B$ flows do not cancel. In these cases, the flow of the passing electrons may even be opposite to the total bootstrap current.

The width of the edge pedestal in H-mode can become comparable to the ion orbit width, and the approximations upon which the local neoclassical analyses are based break down. Therefore, we investigated the non-local effects in the tokamak edge by comparing results from local (NEO) and global (XGCa) calculations for narrow edge pedestals in Sec. IIID. While agreement between the local and global simulations is excellent in the local regime, differences appear in the non-local regime. Usually, the current measured in the local simulation is larger than the one from the global simulation in the steep edge pedestal. The radial electric field is key to understanding how the non-locality affects the bootstrap current. In the local regime, the effect of E_r on the ions and electrons is the same and cancels in the bootstrap current. Moreover, ions experience quite different forces (from ∇p , E_r , friction, etc.) on the inner and outer legs of their orbits when the ion orbit width is comparable to the gradient scale lengths. In the end, the orbit average of those forces determines the kinetic equilibrium. Thus, the plasma flow calculated from a local equation is inaccurate in the edge pedestal as demonstrated in Fig. 15. Most of the finite orbit-width effects in the bootstrap current arise from steep ion temperature gradients and not from steep density gradients.

Based on the results of the XGCa simulations discussed in this work and the insights gained from the analysis of collisional and non-local effects, we developed a new modification to the Sauter formula that reproduces our numerical results for C-Mod, DIII-D, ASDEX-Upgrade, JET, and NSTX geometry with much higher accuracy than the original Sauter formula. In addition, we corrected the asymptotic behavior of the Sauter formula in the fluid limit. In the steep edge pedestal, the local NEO results overestimate the bootstrap current by up to 15% when compared to the XGCa results. The Sauter formula over-estimates the bootstrap current much more.

Possible deleterious effects of using a linearized collision operator in the steep edge pedestal has not been investigated in the present work.

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Appendix A: Coefficients of the Sauter bootstrap current formula

The factor α_i in Eq. (13) in the term proportional to the ion temperature gradient is given by

$$\alpha_i \equiv \left[\frac{\alpha_0 + 0.25(1 - f_t^2)\sqrt{\nu_{i*}}}{1 + 0.5\sqrt{\nu_{i*}}} + 0.315\nu_{i*}^2 f_t^6\right] \frac{1}{1 + 0.15\nu_{i*}^2 f_t^6} \tag{A1}$$

$$\alpha_0 \equiv -\frac{1.17 \left(1 - f_t\right)}{1 - 0.22 f_t - 0.19 f_t^2}.$$
(A2)

The collisionless trapped particle fraction f_t is defined by

$$f_t \equiv 1 - \frac{3}{4} \left\langle B^2 \right\rangle \int_0^{1/B_{max}} \frac{\lambda d\lambda}{\left\langle \sqrt{1 - \lambda B} \right\rangle},\tag{A3}$$

and the ion and electron collisionalities by

$$\nu_{i*} = \frac{e^2}{12\pi^{3/2}\epsilon_0^2} \frac{qRn_i Z^4 \ln \Lambda_{ii}}{T_i^2 \epsilon^{3/2}},\tag{A4}$$

$$\nu_{e*} = \frac{\sqrt{2}e^2}{12\pi^{3/2}\epsilon_0^2} \frac{qRn_e Z \ln \Lambda_e}{T_e^2 \epsilon^{3/2}},\tag{A5}$$

where q is the safety factor, $n_{i/e}$ are the ion and electron densities, ϵ is the inverse aspect ratio, and temperatures are in eV. The Coulomb logarithms in Eqs. (A4) and (A5) are

$$\ln \Lambda_{ii} = 30 - \ln \left(Z^3 \frac{\sqrt{n_i}}{T_i^{3/2}} \right), \tag{A6}$$

$$\ln \Lambda_e = 31.3 - \ln \left(\frac{\sqrt{n_e}}{T_e}\right). \tag{A7}$$

The coefficients L_{31} , L_{32} , and L_{34} are¹⁴

$$L_{31} = F_{31} \left(X = f_{t,eff}^{31} \right) = \left(1 + \frac{1.4}{Z+1} \right) X - \frac{1.9}{Z+1} X^2 + \frac{0.3}{Z+1} X^3 + \frac{0.2}{Z+1} X^4,$$

$$L_{32} = F_{32,ee} \left(X = f_{t,eff}^{32,ee} \right) + F_{32,ei} \left(Y = f_{t,eff}^{32,ei} \right),$$

$$F_{32,ee}(X) = \frac{0.05 + 0.62Z}{Z(1+0.44Z)} \left(X - X^4 \right)$$

$$+ \frac{1}{1+0.22Z} \left[X^2 - X^4 - 1.2 \left(X^3 - X^4 \right) \right] + \frac{1.2}{1+0.5Z} X^4,$$

$$F_{32,ei}(Y) = \frac{0.56 + 1.93Z}{Z(1+0.44Z)} \left(Y - Y^4 \right)$$

$$+ \frac{4.95}{1+2.48Z} \left[Y^2 - Y^4 - 0.55 \left(Y^3 - Y^4 \right) \right] + \frac{1.2}{1+0.5Z} Y^4,$$

$$L_{34} = F_{31} \left(X = f_{t,eff}^{34} \right)$$
(A8)

with the effective collisional trapped particle fractions $f_{t,eff}^{3j}$ (j = 1, 2, 4)

$$f_{t,eff}^{31} = \frac{f_t}{1 + (1 - 0.1f_t)\sqrt{\nu_{e*}} + 0.5(1 - f_t)(\nu_{e,*}/Z)},$$

$$f_{t,eff}^{32,ee} = \frac{f_t}{1 + 0.26(1 - f_t)\sqrt{\nu_{e*}} + 0.18(1 - 0.37f_t)(\nu_{e*}/\sqrt{Z})},$$

$$f_{t,eff}^{32,ei} = \frac{f_t}{1 + (1 - 0.6f_t)\sqrt{\nu_{e*}} + 0.85(1 - 0.37f_t)\nu_{e*}(1 + Z)},$$

$$f_{t,eff}^{34} = \frac{f_t}{1 + (1 - 0.1f_t)\sqrt{\nu_{e*}} + 0.5(1 - 0.5f_t)(\nu_{e,*}/Z)}.$$
(A9)

Appendix B: Coefficients for the improved bootstrap current formula

By comparing the definition of the Helander formula from Eq. (12.51) in Ref. [46] to our definition in Eq. (19), one can identify $L_{31}^{(h)} = d_1/D$ and $L_{32}^{(h)} = d_2/D$. The friction coefficients l_{ij} are given in the appendix of Ref. [46]. They are evaluated using the approximation $m_e/m_i \to 0$. To evaluate $L_{31}^{(h)}$, $L_{32}^{(h)}$ and $\alpha_i^{(h)}$, we use the viscosity coefficients μ_{sj} for the collisional regime calculated by Hirshman and Sigmar⁴⁷, given by Eqs. (4.20)-(4.22) and (4.31)-(4.40) therein. We express the viscosity coefficients as

$$\mu_{ij} = c_{ij} n_i T_i \tau_{ii}, \quad \mu_{ej} = c_{ej} (Z n_i) T_e \tau_{ee}, \tag{B1}$$

where j = 1, 2, 3, and the collision frequencies are defined in Eq. (4.7) in Ref. [47]:

$$\frac{1}{\tau_{ss'}} = \frac{\sqrt{2}n_{s'}Z_s^2 Z_{s'}^2 e^4 \ln(\Lambda)}{12\pi^{3/2}\epsilon_0^2 \sqrt{m_s} (eT_s)^{3/2}}.$$
(B2)

The comparison of Eqs. (A4) and (A5) with Eq. (B2) yields the ion and electron collision times

$$\tau_{ii} = \frac{qR}{\nu_{i*}\epsilon^{(3/2)}\sqrt{2eT_i/m_i}}, \quad \tau_{ee} = \frac{qRZ}{\nu_{e*}\epsilon^{(3/2)}\sqrt{eT_e/m_e}}.$$
(B3)

Also in the limit of $m_e/m_i \rightarrow 0$, the coefficients c_{sj} are

$$c_{i1} = 1.357,$$

$$c_{i2} = 2.192,$$

$$c_{i3} = 6.919,$$

$$c_{e1} = \frac{(10Z^3/12)(408 + 289.914Z^3)}{192 + 425.678Z^3 + 178Z^6},$$

$$c_{e2} = \frac{(10Z^3/12)(912 + 468.105Z^3)}{192 + 425.678Z^3 + 178Z^6},$$

$$c_{e3} = \frac{(10Z^3/12)(2592 + 1477.85Z^3)}{192 + 425.678Z^3 + 178Z^6}.$$

Following Ref. [46], we define the geometry coefficient c_B by

$$\hat{\mu}_{sj} = \frac{3\langle (\boldsymbol{b} \cdot \nabla B)^2 \rangle}{\langle B^2 \rangle} \mu_{sj} = c_B \mu_{sj}.$$
(B4)

After substituting these coefficients in Helander's formula and evaluating the limit $\nu_{s*} \gg 1$, we obtain the leading terms

$$L_{31}^{(h,2)} = \frac{c_B q^2 R_0^2 Z^2 ((4\sqrt{2} + 13Z)c_{e1} + 6Zc_{e2})}{4(\sqrt{2} + Z)\epsilon^3 \nu_{e*}^2},$$

$$L_{32}^{(h,2)} = \frac{c_B q^2 R_0^2 Z^2 ((4\sqrt{2} + 13Z)c_{e2} + 6Zc_{e3})}{4(\sqrt{2} + Z)\epsilon^3 \nu_{e*}^2},$$

$$\alpha_i^{(h,0)} = \frac{c_{i2}}{c_{i1}}.$$
(B5)

Using the effective trapped particle fraction from Eqs. (A9) and (24), the leading order terms of $L_{3j}^{(s)}$ become

$$L_{31}^{(s,1)} = \frac{2f_t Z(2.4 + Z)}{(1+Z)(1-0.99f_t)\nu_{e*}},$$

$$L_{32}^{(s,1)} = \frac{f_t \left(\frac{0.05+0.62Z}{0.18-0.0666f_t}\sqrt{Z} - \frac{0.56+1.93Z}{(0.85-0.3145f_t)(1+Z)}\right)}{(1+0.44Z)Z\nu_{e*}},$$

$$L_{34}^{(s,1)} = \frac{f_t Z(2.4 + Z)}{(1+Z)(0.5-0.25f_t)\nu_{e*}},$$

$$\alpha_i^{(s,0)} = 2.1.$$
(B6)

The critical collisionality $\nu_{e*,c}$ needed for the interpolation factors $\gamma_{3j}^{(s)}$ is obtained from the relation $|L_{31}^{(h,4)}|/L_{31}^{(h,2)} = 10^{-2}$, which yields

$$\nu_{e*,c} = \frac{5\sqrt{c_B}qR_0Z}{\epsilon^{3/2}}\sqrt{\frac{a}{b}},$$

$$a = 32c_{e1}^2 + 8\sqrt{2}\left(13c_{e1}^2 + 9c_{e1}c_{e2} + 2c_{e2}^2\right)Z$$

$$+ \left[\left(13c_{e1} + 9c_{e2}\right)^2 + 7c_{e2}^2 + 6c_{e3}\left(6c_{e1} + 4c_{e2}\right)\right]Z^2,$$

$$b = 6c_{e2}Z\left(\sqrt{2} + Z\right) + c_{e1}\left(8 + 17\sqrt{2}Z + 13Z^2\right).$$
(B7)

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