Tractable flux-driven temperature, density, and rotation profile evolution with the quasilinear gyrokinetic transport model QuaLiKiz

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Abstract

Quasilinear turbulent transport models are a successful tool for prediction of core tokamak plasma profiles in many regimes. Their success hinges on the reproduction of local nonlinear gyrokinetic fluxes. We focus on significant progress in the quasilinear gyrokinetic transport model QuaLiKiz (Bourdelle et al 2016 Plasma Phys. Control. Fusion 58 014036), which employs an approximated solution of the mode structures to significantly speed up computation time compared to full linear gyrokinetic solvers. Optimisation of the dispersion relation solution algorithm within integrated modelling applications leads to flux calculations $\times 10^6$–$7$ faster than local nonlinear simulations. This allows tractable simulation of flux-driven dynamic profile evolution including all transport channels: ion and electron heat, main particles, impurities, and momentum. Furthermore, QuaLiKiz now includes the impact of rotation and temperature anisotropy induced poloidal asymmetry on heavy impurity transport, important for W-transport applications. Application within the JETTO integrated modelling code results in $1$ s of JET plasma simulation within $10$ h using $10$ CPUs. Simultaneous predictions of core density, temperature, and toroidal rotation profiles for both JET hybrid and baseline experiments are

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presented, covering both ion and electron turbulence scales. The simulations are successfully compared to measured profiles, with agreement mostly in the 5%–25% range according to standard figures of merit. QuaLiKiz is now open source and available at www.qualikiz.com.

Keywords: tokamak, turbulence, gyrokinetics, integrated modelling

(Some figures may appear in colour only in the online journal)

1. Introduction

An accurate and predictive model for turbulent transport fluxes driven by microinstabilities is a vital component of predictive tokamak plasma simulation. This enables the interpretation and optimisation of present-day experiments, extrapolation to future machine performance, and design of control systems.

We report on significant progress made in the tractability and validation of the quasilinear gyrokinetic transport model QuaLiKiz. The basis of the QuaLiKiz model is extensively described in [1–4].

Optimisation of the dispersion relation solver has accelerated the calculation time by a factor 50 compared to the QuaLiKiz version reported in [4, 5]. The dispersion relation for a single wavenumber is now solved within ~1 s. This allows tractable simulation of flux-driven dynamic profile evolution including all transport channels: ion and electron heat, main particles, impurities, and momentum. QuaLiKiz is now of comparable speed as the TGLF gyrofluid quasilinear model [6, 7], used for similar multi-channel applications [8]. These numerical improvements are summarised in section 2 and appendix A.

Additional physics has been added, widening the applicability of the model. This includes poloidal asymmetries, important for heavy impurity transport, reproducing previous results obtained with GKW [9]. The ETG model has been recalibrated, and includes a rudimentary multi-scale model. The eigenfunction solver has been improved, resulting in further validation of the impact of E × B shear and momentum transport. These additions are summarised in section 3.

QuaLiKiz has been coupled to the JETTO-SANCO [10, 11] integrated modelling suite. The code speed-up now allows, for the first time with QuaLiKiz, tractable heat, particle, and momentum transport simulations for main ion, impurity, temperature, and toroidal rotation core profile predictions. Validation was carried out on JET hybrid and baseline pulses. Agreement with measured profiles were mostly in the 5%–25% range according to standard figures of merit. The integrated modelling simulations are described in section 4.

The motivation for development of quasilinear transport models such as QuaLiKiz is the prohibitive computational cost of direct numerical simulation with massively parallel nonlinear gyrokinetic codes over discharge timescales. Even for gradient-driven, $\delta f$ simulations using codes such as GENE [12], GKW [13], GYRO [14], and GS2 [15], the computational cost for either ion or electron scale simulations is $\sim 10^4$ CPU h for fluxes at a single radius, and is significantly larger for multi-scale simulations [16]. Assuming $O(10^5)$ flux calls per 1 s of JET-scale plasma evolution, and 10 radial points, the computational cost for 1 s of discharge profile evolution is $\sim 10^8$ CPU h. Use of implicit time-step solvers [17] or steady-state iteration schemes [18] can reduce the computation time, and provides unique opportunities for validation of nonlinear predictions. Nevertheless, routine application of nonlinear turbulence codes for core profile prediction remains elusive, especially when including both ion and electron turbulence scales.

The quasilinear approximation provides significantly increased tractability. This has proven to be a successful tool for tokamak turbulence model reduction, successfully reproducing nonlinear simulations over a large range of parameters (see [4] and references therein, and [6, 19, 20]). The quasilinear approximation has also been successfully applied for stellarator turbulence modelling [21]. The approximation is valid in the plasma confinement zone where the density fluctuations are small—$\delta n/n \sim O(\%)$. Their success hinges on the reproduction of local nonlinear gyrokinetic fluxes [22].

The specific assumptions made in the QuaLiKiz linear gyrokinetic solver lead to a ~6 order of magnitude computational speed-up compared to local $\delta f$ nonlinear gyrokinetics. This brings the computational cost of 1 s of JET-scale plasma evolution down to $\sim 100$ CPU h, which is routinely feasible. These assumptions are as follows:

- Shifted circle ($\hat{s} - \alpha$) geometry with a small inverse aspect ratio expansion. This allows the bounce averaging of the trapped species to be expressed as elliptic integrals, speeding up the dispersion relation calculation time. Our neglect of shaping parameters such as triangularity and elongation is expected to lead to systematic overpredictions of the transport level. However, in the plasma core for standard tokamak shapes, only a relatively minor flux overprediction ($<\sim 30\%$) and critical stability threshold underprediction ($<\sim 10\%$) are expected [23, 24]. While the strong dependence of H-mode pedestal, and hence global confinement, on plasma shaping is well established [25], pedestal confinement is out of the scope of QuaLiKiz model. In the integrated modelling discussed here, we apply the measured pedestal height as boundary conditions for core modelling with QuaLiKiz. For more extreme plasma shapes, for example the low aspect ratio typical of spherical tokamaks, or high ($\alpha > \sim 2$) elongation, we do not expect QuaLiKiz to be valid. Generalisation to a more complete geometry description is the subject of future work.

- Electric potential fluctuations only. To save computation time, we only solve the Poisson field equation (quasineutrality) and not Ampere’s law. This is equivalent to assuming $\beta = 0$. The relevance of QuaLiKiz for spherical
tokamak configurations is thus further reduced, since microtearing modes can dominate the electron heat transport in high-$\beta$ regions [26, 27]. Finite-$\beta$ stabilisation of ITG is also not included in QuaLiKiz, which can be important for high performance scenarios [28–30]. However, since the nonlinear stabilisation (flux reduction) of ITG due to finite-$\beta$ is greater than would be expected by the linear $\beta$-stabilisation [31, 32], future work will explore modifying the QuaLiKiz saturation rule to capture this nonlinear effect.

- Shifted Gaussian (ballooned) eigenfunction ansatz. A significant gain in computational speed ($\sim$2 orders of magnitude) is achieved by a non-self-consistent solution of the eigenvalue (real frequency and growth rate) and eigenfunction (radial and poloidal structure of the electrostatic potential) in the dispersion relation. The eigenfunction is first calculated through expanding the dispersion relation under a strongly driven assumption. See appendix B for further details. This eigenfunction is then inserted into the kinetic dispersion relation which is solved using a weak formulation of Poisson’s equation [3]. The eigenfunction ansatz is restricted to strongly ballooned Gaussian structures. This limits the dispersion relation solution accuracy in cases where broad poloidal eigenfunctions are observed in self-consistent calculations. This limits the accuracy of trapped electron mode calculations. Furthermore, slab-like ITG is not captured, as observed at low magnetic shear or significant Shafranov shift. For examples, see [33] and section 3.5.1 of this paper. However, in spite of its relative simplicity, the symmetry breaking induced by the shifted Gaussian mode structure in the presence of rotation has proven successful for QuaLiKiz calculations of momentum transport and $E \times B$ suppression of transport [5].

2. Dispersion relation solver optimisation

The QuaLiKiz calculation is comprised of three steps. The first step is the eigenvalue solver that calculates the instability frequencies and growth rates. The second step calculates the quasilinear flux integrals. These two steps are typically carried out for a spectrum of unstable modes. The third step is the saturation rule. This builds the final transport fluxes by combining the spectrum of quasilinear flux integrals with a modelled spectral form factor and amplitude level based on nonlinear simulations.

The bulk of time spent for the QuaLiKiz calculation is in the first step, the eigenvalue solver. This is a root finding algorithm in the complex plane, of the dispersion relation $D(\omega) = 0$ found by setting the quasineutrality constraint on the perturbed plasma densities from the linearised and gyro-averaged Vlasov equation. See [3, 4] for further details of the form of $D(\omega)$ solved in QuaLiKiz. $\omega$ is the eigenvalue $\omega_r + i\omega_i$, where $\omega_r$ is the real frequency and $\gamma$ the growth rate. The QuaLiKiz algorithm only searches for solutions in the upper half of the complex plane, i.e. for unstable modes (positive $\gamma$), and not for damped modes (negative $\gamma$).

The QuaLiKiz eigenvalue solver algorithm has been optimised. This led to a calculation speed-up by an approximate total factor of 50–100. This was key to transform QuaLiKiz into a pragmatic tool for integrated modelling applications. Per wavenumber, the QuaLiKiz calculation time is now $\sim$0.5–1 s when not including rotation, and $\sim$2–4 s when including rotation (longer, due to a loss of symmetry in the integrations). We briefly summarise the various steps taken for this optimisation. Full details are found in appendix A.

- The root finding algorithm in the QuaLiKiz eigenvalue solver follows the Davies method [34]. The contour paths have been extensively optimised, minimising the number of individual $D(\omega)$ calculations. This included modifying the contour shapes, contour overlap, and contour ranges. A significant speed-up of factor $\sim$8 is achieved compared to the previous QuaLiKiz version. A similar level of robustness in the root finding is achieved as in the previous version, since we maintain full coverage of the physically relevant regions in the complex plane.

- Improved numerical techniques for integral functions within $D(\omega)$. For the plasma dispersion function, the standard ‘WofZ’ method [35] was replaced by the faster Weideman algorithm [36]. Furthermore, the Carlson method [37] for calculating elliptic integrals was replaced by the Fukushima method [38]. Together, a further factor $\sim$3 speed-up was achieved compared to the previous QuaLiKiz version.

- Within integrated modelling simulations, at a given timestep, the eigenvalue solution from the previous timestep is used to accelerate the calculation. The previous solution is used as a first guess for a Newton solver to converge to the new eigenvalues. The validity of this approach is through the relatively small modification in background plasma profiles between the $\Delta t \sim$ 1 ms timesteps typical of integrated modelling simulations. At every $\approx$10 ms, the full contour solution is again sought out, since new eigenvalues may appear. This procedure accelerates the integrated modelling simulation by a factor $\sim$5.

3. Extensions to the QuaLiKiz physics model

The physics content in QuaLiKiz has been extended and further validated. This includes an improved eigenfunction solver in the presence of rotation, further validation of momentum transport, allowing an arbitrary number of active or trace ion species, poloidal asymmetry (important for heavy impurity transport), and a new saturation rule for ETG turbulence. These points will be discussed in this section, together with a discussion of caveats and limitations of the $\alpha$-stabilisation and $E \times B$ shear turbulence suppression model in QuaLiKiz.
3.1. New eigenfunction solver

A key feature of QuaLiKiz is the use of an eigenfunction calculated from an expansion of the linear gyrokinetic equation arising from a strongly driven (mode frequency ordered larger than drift and transit frequencies) assumption. This allows for significantly faster eigenvalue calculations. This approximated eigenfunction has been compared to self-consistent gyrokinetic eigenfunctions calculated by GENE, GKW, GYRO, and agrees in regimes where the eigenfunction is ballooned [5, 33].

Including the eigenfunction shift due to rotation effects breaks the eigenfunction symmetry leading to angular momentum flux. The success of applying approximated shifted eigenfunctions for momentum transport predictions is not trivial. This approach in QuaLiKiz has been validated through comparisons against angular momentum flux predictions with non-approximated eigenfunctions [5].

The fluid eigenfunction solution in the present version of QuaLiKiz has been rederived and improved compared with [5]. Particular care has been taken in setting up a consistent and less restrictive ordering of the various terms. The model is also refined by consistently carrying out pitch angle integrations. An extensive overview of the revised derivation is presented in appendix B.

3.2. Verification of momentum transport model

Verification of the QuaLiKiz momentum transport model using the new eigenfunction solver is reported here. First, we show that we recover previous predictions of the perpendicular $E \times B$ shear impact on turbulence suppression and momentum transport. This is seen in figure 1, and carried out for GA-standard-case (GASTD) parameters, with 8 toroidal modes at ITG length-scales ($k \rho_{ik} = [0.1, 0.175, 0.25, 0.325, 0.4, 0.5, 0.7, 1.0]$), where $\rho_i \equiv \sqrt{T_i/m_i/Z_i} B_i$. Similarly to both [5] (figure 9 therein) and [19] (figure 4), the impact of $\gamma_E$ on the fluxes leads to full suppression at approximately $2 \times$ the maximum linear growth rate, and a similar degree of $\gamma_E$ driven momentum flux.

QuaLiKiz also predicts increasing momentum pinch with trapped electron drive, as predicted by analytic theory and gyrokinetic simulations [39]. In figure 2, we show the Prandtl number $\equiv \chi_i/\chi_p$ (left panel) and momentum Pinch number $\equiv RV_i/\chi_p$ (right panel) for a GASTD case $R/L_m$ scan. $\chi_i$ is the ion heat diffusivity. $V_i$ is the momentum diffusivity, $V_i$ is the momentum pinch, as defined by the momentum flux decomposition:

$$\Pi_i = R \sum m_i n_i \left( -\chi_i \frac{\partial u_i}{\partial r} + V_i u_i \right) + \Pi_{\text{RS}},$$

where $\Pi_{\text{RS}}$ is the residual stress. Note that the GASTD case has a single ion species, so the summation is not relevant here. The pinch is calculated from carrying out the $R/L_m$ scan with $M \equiv u_i/\nu_{hi} = 0.2$ and $M' = 0$. The momentum diffusivity is calculated from a $R/L_m$ scan with $M = 0$ and $M' = 1$, where $M' \equiv -R \frac{d\Pi_i}{dr}$. As expected, while the Prandtl number is nearly constant, the Pinch number increases (in absolute terms) with increasing $R/L_m$, i.e. increasing trapped electron drive. This scan is identical to that shown in [5] (see figure 10 therein). However, with the new eigenfunction model, the Prandtl number is now closer to unity, and the Pinch number scan is upshifted by approximately 1.5 units, both in disagreement with full gyrokinetic modelling (see figures 5 and 7 in [39]). Nevertheless, the new eigenfunction model assumptions and ordering are more consistent compared to that presented in [5]. Therefore, we maintain the new
model in spite of the reduced accuracy in comparison to full gyrokinetic modelling in this particular scan.

Finally, beyond the scope of the previous studies [5], we also explored the sensitivity of the Pinch number to local inverse aspect ratio \( r/R \), in an \( \epsilon \) scan based on GASTD case parameters. This is effectively a scan of the trapped electron drive, due to the proportionality of the trapped electron fraction to \( \epsilon^{0.5} \). The results are shown in figure 3. As expected, a clear increase in momentum pinch is observed with increasing \( (r/R)^{0.5} \). This further corroborates the QuaLiKiz rotation model.

3.3. Poloidal asymmetries for heavy impurities

The constraint of parallel force balance in the presence of centrifugal forces, and/or temperature anisotropy (e.g. due to NBI and ICRH heating), leads to asymmetry of particle density \( n_j \) on a flux surface [40–42]. Following the Bi-Maxwellian temperature anisotropy model of [43], we obtain, at a given radius, for species \( j \), the poloidal density profile \( n_j(\theta) \):

\[
n_j(\theta) = n_{j,LFS} \frac{T_j(\theta)}{T_{j,LFS}} \times \exp \left[ \frac{Z_j \Phi(\theta) - \frac{1}{2} m_j \Omega^2 (R(\theta)^2 - R_{LFS}^2) T_j}{T_{j,LFS}} \right]
\]

(2)

with:

\[
\frac{T_j(\theta)}{T_{j,LFS}} = \left( \frac{1 - \frac{T_{j,LFS}}{T_{j,LFS}}} {B_{LFS} B(\theta)} \right)^{-1},
\]

(3)

where LFS denotes the low field side values, \( \Phi(\theta) \) is the equilibrium electrostatic potential, \( \Omega \) the angular toroidal velocity, \( T_{\perp} \) the perpendicular temperature, and \( T_{\parallel} \) the parallel temperature. \( \Phi(\theta) \) is solved numerically at each poloidal location by invoking the quasineutrality constraint. This has now been implemented in QuaLiKiz and successfully verified against the similar quasineutrality solver in GKW. The inputs into the system are the toroidal velocity, temperature anisotropy at any single poloidal location (here defined as LFS for convenience, but this can be generalised), and density ratios at any single poloidal location (similarly defined here as the LFS for convenience).

While the density asymmetry is small for light ions, it can be non-negligible for heavy impurities. The high mass leads to significant centrifugal forces, while the high charge leads to a significant response to the equilibrium potential. The density asymmetry can significantly modify the resultant diffusion and

Figure 2. Scan of impact of trapped electron drive on QuaLiKiz momentum transport predictions via a \( R/L_n \) scan based on GASTD case parameters. The Prandtl number is shown in the left panel, and the Pinch number in the right panel.

Figure 3. Scan of impact of trapped electron drive on QuaLiKiz momentum pinch predictions via a \( r/R \) scan based on GASTD case parameters.

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pinch for both turbulent and neoclassical transport, and is a critical effect to include for heavy ion impurity transport predictions such as for W [44-46].

For turbulent transport in QuaLiKiz, we closely follow the model of [9]. The impact of the density asymmetry is not included self-consistently in the dispersion relation, which maintains the particle densities as flux functions. However, when evaluating the quasilinear particle flux integrations, we include the density asymmetry. The asymmetries lead to a poloidally dependent density gradient, leading to new terms in the quasilinear integral which effectively emerge as new pinch terms, which can become significant for heavy impurities. These terms are then evaluated by flux surface averaging, and are additionally weighted by the eigenfunction shape. Full details are found in appendix C. The flux function inputs into the QuaLiKiz dispersion relation are taken from the LFS values. At low impurity densities (trace values) this choice is of negligible importance. However, for non-trace heavy impurities this choice can have an impact on the dispersion relation predictions. Due to the $n_s \frac{V_q^2}{T}$ prefactors in the dispersion relation, even experimentally relevant low densities of high-Z impurities can have a non-negligible impact on the instability calculation. Future work will focus on how best to approximate the O(2D flux function density and density gradient inputs based on the 1D poloidal profile, for such non-trace cases.

The QuaLiKiz poloidal asymmetry model for heavy impurity transport was validated against quasi-linear GKW predictions, using parameter sets defined in [9]. A comparison was made of the predicted impurity normalised logarithmic density gradient $R/L_n$. Assuming no core sources, this is provided by the zero-flux constraint: $R/L_n = -\frac{\phi_i}{D}$, where $V$ is the convective velocity (pinch) and $D$ the diffusivity. Collisionless runs were carried out, scanning over the impurity mass $A$ (in amu). The impurity charge throughout the scan was set at $Z = A/2$ for $A < 92$, and $Z = 46$ for $A > 92$. Two basic datasets were employed: an ITG dataset with $R/L_{\ell_i} = 9$, $R/L_{Te} = 6$, $R/L_n = 2$, $q = 1.4$, $\hat{s} = 0.8$, $T_e/T_i = 1$, and a TEM dataset with identical parameters apart from $R/L_{Te} = 9$, $R/L_{Ti} = 3$, $T_e/T_i = 1$. $\hat{s}$ is magnetic shear, and $q$ the safety factor. For both cases, we simulate a single wavenumber $k_\parallel \rho_i = 0.3$. The poloidal asymmetries are from centrifugal effects, with $M = 0.2$, $M' = 1$ with respect to the main ion. For the ITG case, we also carried out runs at varying $\hat{s} = 0.3, 1.3$. The comparison is shown in figure 4. In panels (a)–(c), the ITG comparison is shown. QuaLiKiz captures the GKW predicted $R/L_n$ with an rms error of $(\Delta \hat{L}/L_n)_{\text{rms}} \approx 0.5$ throughout the dataset. The TEM data has a larger rms error of $(\Delta \hat{L}/L_n)_{\text{rms}} \approx 0.8$, with a significant deviation between the QuaLiKiz and GKW predictions at high $A$ when including poloidal asymmetry effects. This is likely due to the known deficiencies of the QuaLiKiz eigenfunction ansatz in fully reproducing gyrokinetic TEM eigenfunctions [5]. Nevertheless, the GKW behaviour can be well captured by QuaLiKiz following a moderate increase of $M'$ from 1 to 1.5, as shown in figure 4(d). This illustrates the sensitivity of the effective 2D roto-diffusion pinch.

3.4. Quasilinear ETG saturation rule

The QuaLiKiz ETG nonlinear saturation rule has been recalibrated. Complete isotropisation is assumed, with $k_\parallel^2 = 2k_\perp^2$. The constant prefactor which sets the QuaLiKiz ETG-scale saturation level was tuned by comparison to single-scale ETG nonlinear GENE simulations of JET discharge L-mode discharge 78834 [47]. Separate ITG/TEM+ETG single scale nonlinear simulations of this discharge led to heat flux matching in both the ion and electron channels when compared to experimental power balance. Approximately 50% of

Figure 4. Validation between QuaLiKiz and GKW predictions of impurity $R/L_n$ (assuming no sources) for an ITG dataset at various magnetic shear (panels (a)–(c)), and for a TEM dataset (panel (d)). GKW data is depicted by the square symbols, and QuaLiKiz data by the diamond symbols. In panel (d), the curve with circular symbols signifies a QuaLiKiz run with increased $M'$. The curves including poloidal asymmetry due to centrifugal effects are shown with solid symbols, and when neglecting poloidal asymmetries, with open symbols.
the electron heat flux was from ETG turbulence. In the GENE simulations, ETG saturation was achieved by increasing the prescribed level of perpendicular \( E \times B \) shear until heat flux convergence was reached, due to ETG streamer shearing leading to a reduction of box-scale effects. This technique was assumed to be a proxy for ion-scale eddies shearing the ETG streamers. Since the GENE and QuaLiKiz ETG thresholds differ in this case by approximately 10%, the calibration was carried out on the stiffness level (gradient of flux with respect to driving gradient). This is seen in figure 5, where the GENE and QuaLiKiz predictions are parallel. The dimensionless parameters for the simulation input is summarised in table 1. The dimensional reference parameters for defining the SI fluxes and collisionality were \( B_{\text{ref}} = 3.5 \text{T} \), \( T_{\text{ref}} = 1.45 \text{keV} \), \( n_{\text{ref}} = 1.95 \times 10^{19} \text{ m}^{-3} \), and \( L_{\text{ref}} = 2.94 \text{ m} \). The QuaLiKiz ETG saturation rule contains a rudimentary multi-scale effect. The emergence of significant ETG fluxes depends on the relative weight of ion-scale and electron-scale instability. For strongly driven ion-scale modes (significantly above threshold), ETG turbulence is quenched according to observations from multi-scale simulations [16, 48]. A ‘rule of thumb’ based on a dataset of multi-scale simulations, suggests a threshold for non-quenched ETG flux at \( \lambda \equiv \max(\gamma_{\text{ETG}}/\gamma_{\text{TEM}}) > \sqrt{m_e/m_i} \) [49]. \( \gamma \) is the growth rate ratio determined by the maxima of the ion-scale and electron-scale instability spectra. Such a rule is set in QuaLiKiz, where the single-scale ETG saturation factor is multiplied by the sigmoid function:

\[
\frac{1}{1 + e^{-(\lambda - \hat{\lambda})/\delta}}.
\]

The sigmoid is employed to ensure smoothness around the transition, to avoid numerical instabilities in integrated modelling frameworks.

This QuaLiKiz ETG model is not valid in regimes with fully absent or suppressed ion-scale turbulence, where significant ETG zonal flows may arise, leading to an extensive ETG threshold shift and strong collisionality dependence [50]. Future work will aim towards a more complete QuaLiKiz multi-scale model for ETG saturation, similar to [20].

3.5. Caveats regarding \( \alpha \) and \( E \times B \) stabilisation

3.5.1. \( \alpha \)-stabilisation. The validity of the \( \alpha \)-stabilisation model in QuaLiKiz was investigated, by comparison with linear GENE gyrokinetic simulations. This stabilisation is due to the modification (and even sign reversal) of the magnetic drift frequency [51], typically for \( \hat{s} = \alpha < 0.5 \), where \( \alpha = q^2 \sum \beta_i \left( \frac{R}{L_{\theta}} + \frac{\hat{s}}{L_T} \right) \), and \( \beta_i \) is the species dependent
shear, at more inner radii (left panel), then the low $\hat{s} - \alpha$ means that at finite $\alpha$, QuaLiKiz significantly overpredicts the ITG instability threshold compared to GENE. Simply setting $\alpha = 0$ in QuaLiKiz then leads to significantly improved agreement for this case.

At low $\hat{s} - \alpha$, the GENE threshold predictions show a relatively weak dependence (see figure 6). Therefore, as a workaround to the limitations of the QuaLiKiz model at low $\hat{s} - \alpha$, we modify the inputs. If $\hat{s} - \alpha < 0.2$ for $\hat{s} > 0.2$, then $\alpha$ is modified such that $\hat{s} = \alpha = 0.2$. For $\hat{s} < 0.2$, then $\alpha$ is simply modified to 0. This is motivated by the slab-like characteristics of the instability at low $\hat{s} - \alpha$ observed in the GENE calculations, which seem to diminish the impact of the strong toroidal ITG branch stabilisation observed in QuaLiKiz. Future work will focus on a more self-consistent solution to this QuaLiKiz limitation.

3.5.2. $E \times B$-stabilisation. Parallel velocity gradient (PVG) destabilization can be a significant factor in compensating the $E \times B$ flow shear stabilisation in tokamaks, particularly at large $q/e$ [55–59]. However, when using the present QuaLiKiz eigenfunction model, the pure PVG destabilization is not captured, when compared to linear GENE simulations. This is shown in figure 9, which is a $M' = -\frac{R}{v_A a} \frac{d \Omega}{d r}$ scan of linear growth rate predictions at GASTD case parameters with $k_0 \rho_s = 0.3$ and $\alpha = 0$. The growth rate spectrum is directly related to the transport fluxes through the quasilinear response and mixing length rules. While the GENE results show a ~30% increase in linear growth rate when scanning from $M' = 0 \rightarrow 7$, the QuaLiKiz results do not show this increase, and the growth rates even slightly decrease with higher $M'$. This is due to the stabilising impact of the eigenfunction shift having a stronger impact than the additional driving $M'$ term in the diamagnetic frequency. Note that experimental $M'$ typically does not exceed $M' \sim 4$.

This disagreement may have implications for $E \times B$ turbulence suppression predictions at inner radii, where $q/e$ (and hence $M'$ for toroidal rotation) is high. Anticipating a QuaLiKiz overprediction of $E \times B$ suppression of turbulence in this regime, we have implemented an optional setting to neglect the impact of rotation at inner radii in integrated modelling applications. When this assumption is applied, then in the dispersion relation solver, rotation effects are only included in a reduced $\rho_{\text{norm}} > 0.4$ zone, with a linear reduction in impact from $\rho_{\text{norm}} = 0.6 \rightarrow 0.4$. For the quasilinear flux integrals, the full rotation eigenfunction calculation is always included at all radii. Therefore, for $\rho_{\text{norm}} < 0.4$, momentum transport is still predicted (due to symmetry breaking) even if rotation was not included in the growth rate spectrum calculation. We note that further motivation for the $\rho_{\text{norm}} > 0.4$ assumption is provided by nonlinear electromagnetic simulations, showing that turbulence suppression by $E \times B$ shear is significantly weakened at high $\beta$ [28], which is more likely at inner radii.

Further caveats are as follows. The QuaLiKiz rotation model assumes a small Mach number ordering. This ordering is broken for heavy impurities, due to the higher Mach number. Thus, we set $M = 0$ for all impurities with $A > 20.5$.
heavier than Ne). $M'$ is not modified, since the ordering assumption for $M'$ is not necessarily broken at high $M$. Finally, due to observed spurious localised particle transport peaking when applying the QuaLiKiz rotodiffusion model, we suppress rotodiffusion from particle flux outputs. The provenance of this behaviour will be investigated in future revisions of the QuaLiKiz model. For main ion transport, rotodiffusion should be an insignificant feature.

4. Validation with integrated modelling of JET discharges

We present QuaLiKiz validation within the JETTO-SANCO [10, 11] suite of integrated modelling codes, for C-wall JET hybrid scenario 75225 [60], and ILW baseline scenario 87412 [61]. These simulations include the first ever QuaLiKiz integrated modelling simulations with combined heat, particle and momentum transport, together with the impact of rotation. All source calculations are from PENCIL (NBI) and PION (ICRH). Neoclassical transport, resistivity and bootstrap current are calculated by NCLASS [62]. The current profile is prescribed from predictive poloidal flux evolution modelling, using experimentally measured kinetic profiles. In JETTO-SANCO 1 s of JET plasma simulation with QuaLiKiz costs $\sim$10 h walltime using 10 CPUs. All simulations covered both ion and electron scales with a wavenumber spectrum of $k_{r,q} = [0.1, 0.175, 0.25, 0.325, 0.4, 0.5, 0.7, 1, 1.8, 3, 9, 15, 21, 27, 36, 45]$.

QuaLiKiz has shifted-circular geometry, while JETTO maintains shaped flux surface geometry for all source calculations, and for setting the metrics in the transport PDEs. This necessitates certain choices and transformations when passing information between the codes. The radial coordinate chosen to calculate the input gradients into QuaLiKiz is $r = (R_{\text{max}} - R_{\text{min}})/2$, where $R_{\text{max}}$ is the maximum radius of a flux surface, and $R_{\text{min}}$ the minimum radius. The QuaLiKiz output fluxes (per m²) are (circular) flux-surface-averaged, and the full flux surface area is used to calculate the total transport rates within JETTO. The effective diffusivities and heat conductivities calculated from these fluxes within QuaLiKiz are transformed into JETTO coordinates, using general geometry information from JETTO. For example for particle

Figure 8. Comparison between nonlinear GENE and QuaLiKiz ion heat flux predictions for DEMO1 parameters, at mid-radius (right panel) and inner-radius (left panel). For both parameter sets, the QuaLiKiz calculations are compared to the GENE results when including (red curves) and neglecting (blue curves) $\alpha$ effects.

Figure 9. Comparison between linear growth rates calculated by QuaLiKiz (blue curve) and linear-GENE for a $M'$ scan based around the GASTD case with $k_{r,\rho_s} = 0.3$. 

(heavier than Ne). $M'$ is not modified, since the ordering assumption for $M'$ is not necessarily broken at high $M$. Finally, due to observed spurious localised particle transport peaking when applying the QuaLiKiz rotodiffusion model, we suppress rotodiffusion from particle flux outputs. The provenance of this behaviour will be investigated in future revisions of the QuaLiKiz model. For main ion transport, rotodiffusion should be an insignificant feature.
diffusivities, $D_{\text{JETTO}} = c_D D_{\text{QuaLiKiz}}$, where:

$$c_D = \frac{\partial n}{\partial \rho} \langle |\nabla \rho| \rangle.$$  \hspace{1cm} (5)

In practice, $c_D < \sim 1.2$.

For evaluating the accuracy of the QuaLiKiz predictions, we apply a simple standard deviation figure of merit [63, 64], where, for a quantity $f$:

$$\sigma = \sqrt{\int_0^{\rho_{\text{norm}}} \left(f_{\text{sim}} - f_{\text{exp}}\right)^2 \, dx} / \sqrt{\int_0^{\rho_{\text{norm}}} f_{\text{exp}}^2 \, dx}.$$  \hspace{1cm} (6)

Where $f_{\text{sim}}$ is the simulated quantity, and $f_{\text{exp}}$ is the measured quantity. We do not take into account the multiple sources of potential systematic error in the modelling, such as the heating and current drive models, $Z_{\text{eff}}$ measurements, neoclassical model, and systematic kinetic profile measurement errors. This is out of the scope of the present paper.

4.1. JET hybrid scenario 75225

In figure 10 we display a JETTO+QuaLiKiz simulation of JET C-wall hybrid discharge 75225. The measured kinetic profiles, calculated source deposition profiles, and equilibrium data from EFIT were averaged between 6 and 6.5 s. These averaged kinetic profiles were set as the initial conditions for JETTO+QuaLiKiz modelling, carried out for 1 s, which corresponds to approximately 5 energy confinement times and sufficient for profile convergence. The QuaLiKiz predicted profiles are then compared to the averaged measured profiles. The boundary condition is at normalised toroidal flux coordinate $\rho_{\text{norm}} = 0.8$. The modelling includes
heat, particle, impurity, and momentum transport simulta-
neously. Rotation effects are always included in the $\rho_{\text{norm}} > 0.4$ zone, unless otherwise specified. The C-impurity transport and radiation is evolved separately within SANCO, with transport coefficients passed from QuaLiKiz. The initial $C$ condition is a flat profile, with $Z_{\text{eff}} = 1.7$ as measured by Bremsstrahlung. The fits of the experimental profiles are from a combination of edge and core charge exchange for $T_e$ and $V_{\text{tor}}$, high resolution thompson scattering (HRTS) and LIDAR for $T_e$, and from HRTS for $n_s$.

The simulation figures of merit per transport channel are summarised in table 2. When restricting the comparison to $\rho_{\text{norm}} > 0.4$, multi-channel agreement is achieved at a $\sim 10\%$ level for each channel. However, at $\rho_{\text{norm}} < 0.4$, QuaLiKiz underpredicts the value of $T_e$ for this discharge, as expected; we did not include fast ions in the simulation, and furthermore QuaLiKiz is an electrostatic code and does not include nonlinearly enhanced electromagnetic stabilisation of ITG in the saturation rule. Fast-ion-enhanced electromagnetic stabilisation of ITG turbulence is an important effect for high performance hybrid scenarios at inner radii [29–31, 65, 66]. Pertinently, nonlinear simulations have shown electromagnetic stabilisation to be a critical effect in inner radii for this specific discharge [28].

The impact of the various rotation assumptions are shown in figure 11, with the figures of merit listed in table 3. These simulations were carried out with heat and particle transport only. The inclusion of rotation clearly improves agreement with experiment for $\rho > 0.4$. However, when including rotation throughout the full radius, then the $T_e$ and $n_s$ profiles peak at $\rho < 0.4$ with gradients significantly (particularly for $T_e$) in disagreement with the measured profiles.

The impact of including ETG scales is shown in figure 12. The figures of merit are listed in table 4. These simulations include heat, particle and momentum transport. Ignoring the ETG scales leads to an overprediction of the $T_e$ gradients at $\rho_{\text{norm}} > 0.5$. This is consistent with previous observations of an electron heat flux shortfall at outer radii in DIII-D modelling, using a previous version of TGLF [6, 7] with only $\sim 10\%$ electron heat flux from ETG scales [67]. Interestingly, excluding ETG scales leads to cross-channel transport phenomena, where $V_{\text{tor}}$ is also significantly modified, likely due to $\nabla T_e$ or $T_e/T_e$ dependencies in the momentum transport coefficients.

We summarise the 75225 simulation results. Encouraging agreement in heat, particle, and momentum transport has been achieved. The most outstanding disagreement remains $T_e$ underprediction in inner radii, consistent with missing electromagnetic physics in QuaLiKiz. Both the inclusion of rotation at $\rho_{\text{norm}} > 0.4$ (where the rotation model is valid), and ETG scales, improves agreement with experiment.

### Table 2. Standard deviation figures of merit for the JETTO + QuaLiKiz JET hybrid scenario 75225 simulations shown in figure 10.

| Zone | $\sigma_{T_e}$ | $\sigma_{V_{\text{tor}}}$ | $\sigma_{\text{fl}}$ | $\sigma_{\text{LIDAR}}$ |
|------|----------------|--------------------------|---------------------|---------------------|
| $\rho_{\text{norm}} = [0.4–0.8]$ | 7.9% | 8.3% | 7.7% | 7.3% |
| $\rho_{\text{norm}} = [0.1–0.8]$ | 24.1% | 5.0% | 13.2% | 18.8% |

### 4.2. JET baseline scenario 87412

#### 4.2.1. Heat, particle and momentum transport.

A similar validation exercise was carried out for JET ILW baseline discharge 87412. Measured kinetic profiles, calculated source deposition profiles, and equilibrium data from EFIT were averaged between 10 and 10.5 s. These averaged kinetic profiles were set as the initial conditions for JETTO + QuaLiKiz modelling, carried out for 1 s, approximately 4 energy confinement times. The QuaLiKiz predicted profiles are then compared to the averaged measured profiles. The source profiles were prescribed constant during the simulation. The boundary condition is at normalised toroidal flux coordinate $\rho_{\text{nom}} = 0.85$. Heat, particle, and momentum transport were included. Rotation effects are always included in the $\rho > 0.4$ zone, unless otherwise specified. Due to significant uncertainties in the core CX measurements for this discharge, we assume $T_i = T_e$ in the core, an assumption strengthened by the high density typical of baseline scenarios [68]. For the edge parameters, this is validated by comparing the $T_e$ HRTS and $T_i$ measurements from edge CX, as seen in the top left panel in figure 13. The Be-impurity transport and radiation is evolved separately within SANCO, with transport coefficients passed from QuaLiKiz. The initial Be condition is a flat profile, with $Z_{\text{eff}} = 1.2$ as measured by Bremsstrahlung. No W was included in these simulations, which concentrate on a phase without W-accumulation.

The full comparison is shown in figure 13. We summarise the prediction figures of merit in table 5. We did not include the (infrequent) sawteeth in the simulations, which may lead to spurious inner core peaking in the predictions. We thus limit the figure of merit radial range to $\rho_{\text{norm}} = [0.2–0.85]$. Note the excellent agreement in the heat and particle channels. As opposed to the hybrid scenario 75225, no $T_i$ underprediction is seen here. This is consistent with electromagnetic effects being of less importance in the lower $\beta$ baseline scenarios. The overprediction of $V_{\text{tor}}$ may also be affected by NTV torque from magnetic islands. 3/2 and 4/3 modes are present during the studied time window. Their impact was not taken into account. Nevertheless, all profiles are in a stationary state by the end of the simulation. This is not trivial considering that a potential feedback mechanism could occur when including momentum transport. Overpredicted $E \times B$ shear may stabilise the turbulence, leading to reduced momentum transport and a further buildup of ExB shear, forming an internal transport barrier. This did not occur for the two discharges simulated here.

The impact of neglecting the ETG scales is shown in figure 14, and the figures of merit are summarised in table 6. Neglecting ETG scales leads to a deterioration of $T_e$ agreement in the outer-half radius, again underlying the importance of ETG scales.

#### 4.2.2. Heat and particle transport only.

We examine the sensitivity of the QuaLiKiz transport predictions for JET 87412 to rotation. Since momentum transport is identically zero with the QuaLiKiz rotation model turned off (no
symmetry breaking), this sensitivity scan was carried out with heat and particle transport only.

In figure 15, we present the rotation sensitivity test. The figures of merit are listed in table 7. The inclusion of rotation has a striking impact on all profiles, significantly improving agreement when rotation is included at $\rho_{norm} > 0.4$. However, when including rotation throughout the full radius, then the $T_e$ and $n_e$ profiles peak at $\rho < 0.4$ with gradients significantly exceeding those of the measured profiles. These observations support the default choice of only including rotation effects in the dispersion relation for $\rho_{norm} > 0.4$.

### 4.2.3. Time dependent simulations

The validations discussed so far concentrated on stationary state. However, an important application for integrated modelling is profile dynamics. For example, the onset of W-accumulation is often set by a threshold out for discharge 87412, starting from an initial condition at $t_{initial} = 9$ s, approximately 300 ms following the LH transition. The $T_e$, $T_i$, and $n_e$ boundary conditions at $\rho_{norm} = 0.85$ evolve in time following the experimental profile fits.

The results are shown in figure 16. The particle confinement time is significantly longer than the heat confinement time, as evidenced from the relatively small variation in $T_e$ and $T_i$ profiles over the 1.5 s of simulation time. However, the initial $n_e$ profile is hollow, and slowly evolves towards a peaked profile. This essential behaviour is reproduced by QuaLiKiz, similar to previous work within CRONOS [69, 70]. However, the QuaLiKiz $n_e$ evolution is slower than the measured $n_e$ evolution for this case, with the predicted degree of peaking less than observed.

Further analysis shows that the $n_e$ discrepancy is limited to the time between 9.1 and 9.6 s. This is shown in figure 17, which displays $n_e$ time-traces at mid-radius, from measurements and the QuaLiKiz prediction starting at $t_{initial} = 9$ s. The significant observed increase in $n_e$ between $t = 9.1–9.6$ s is not captured by QuaLiKiz. Outside of this range, the experimental trend is well followed. We did not include the (inrequent) sawteeth in the simulations, which are observed for this discharge. This may play a role in accelerating the filling in of the hollow density profile. Future work will include such MHD effects. Nevertheless, the essential behaviour of the inversion from hollow to peaked profiles is captured.

### 5. Conclusions and outlook

The first-principle-based quasilinear gyrokinetic transport code QuaLiKiz is numerically optimised. Eigenvalue calculations take ~1 CPUs, and flux calculations take ~10 CPUs. This demands approximately 100 CPUh computation time for 1 s of JET plasma evolution within the JETTO integrated modelling suite, comparable to the TGLF quasilinear transport model. The physics content of the code has been extended and validated, including poloidal asymmetry effects important for heavy impurity transport, and a recalibrated ETG model. Successful validation was carried out within the JETTO modelling suite, for JET hybrid scenario 75225 and baseline scenario 87412, including the first QuaLiKiz integrated modelling simulations with combined heat, particle and momentum transport. Both the impact of rotation and ETG scales was shown to be important for improving agreement with experiments.
QuaLiKiz is now ready for extensive integrated modelling applications, including for W-transport, which is ongoing work. Such multi-channel integrated modelling is a powerful tool for self-consistent kinetic profile prediction, including the dynamics of interacting turbulent transport channels (heat, particle and momentum) as well as setting the background profiles for neoclassical impurity transport. Future work on the QuaLiKiz physics model will concentrate on alleviating the observed inconsistencies in $\alpha$-stabilisation and PVG destabilization, as seen in comparisons with linear gyrokinetic modelling using more complete models such as GENE and GKW. Extension to shaped geometries will be explored. Furthermore, a model for nonlinear electromagnetic stabilisation must be incorporated in the ion-scale turbulence saturation rule. Further computation acceleration aimed at extensive scenario optimisation and real-time applications is being carried out by emulation of the QuaLiKiz model using neural network regression [71].

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Appendix A. Numerical optimisation of QuaLiKiz dispersion relation

The optimisation has 3 components: tailoring the contour paths in the $D(\omega)$ root finding algorithm, improved numerical techniques for integral functions within $D(\omega)$, and taking advantage of the existence of previous solutions from previous timesteps within integrated modelling applications.

A.1. Contour path optimisation

The root finding algorithm in the QuaLiKiz eigenvalue solver follows the Davies method [34]. This involves calculating

![Figure 13. Comparison between JETTO+QuaLiKiz predictions for JET baseline scenario 87412 including heat, particle and momentum transport. The QuaLiKiz predicted and measured profiles are compared, for: ion temperature (top left panel), electron temperature (top right panel), electron density (bottom left panel) and toroidal velocity (bottom right panel). The experimental data points are averaged between 10 and 10.5 s. The experimental error bars are statistical (and are reduced following the averaging), and do not include potential systematic errors. The QuaLiKiz boundary condition is at $\rho_{nrm} = 0.85$.](image)

![Table 5. Standard deviation figures of merit for the JETTO +QuaLiKiz baseline 87412 simulation shown in figure 13. The figures of merit are calculated in the ranges $\rho_{nrm} = [0.2–0.85]$.](table)

| $\sigma_T$ | $\sigma_{T_e}$ | $\sigma_n$ | $\sigma_V$ |
|------------|----------------|------------|------------|
| 7.9%       | 6.6%           | 4.2%       | 33.1%      |
contours of $D(\omega)$ in the complex plane and invoking the argument principle to identify and set the initial guess for any roots found within each given contour. A Newton method is then applied to converge on the root.

Optimising the precise contour paths is key to reducing the computation time, by minimising the number of individual $D(\omega)$ evaluations carried out. This optimisation involves: setting the boundaries of the eigenvalue search region in the complex plane, splitting the search region into a number of separate contours, setting the degree of contour overlap, and parametrising the contour shape. Previously, the contour paths were optimised for robustness and not for speed in integrated modelling applications, meaning extensive limits and significant contour overlap was in place. We now summarise the newly applied contour path choices, which significantly speed up the solution time while maintaining a high level of robustness.

Search region limits

Only solutions in the upper (unstable) half of the complex plane are sought out. The ceiling of the contours in the imaginary plane is set by calculating simple analytical estimates of both slab and interchange modes, calculated by a simplified high $\omega$ expansion of the dispersion relation [72]. The actual ceiling then corresponds to $\max(\gamma_{\text{interchange}}, \gamma_{\text{slab}})/1.5$. The analytic growth

Table 6. Standard deviation figures of merit for the JETTO +QuaLiKiz baseline 87412 simulations shown in figure 14. The figures of merit are calculated in the ranges $\rho_{\text{norm}} = [0.2\ldots0.85]$. Results are shown following 1 s of simulation time, including and neglecting ETG scales.

| ETG assumption | $\sigma_T$ | $\sigma_T$ | $\sigma_n$ | $\sigma_{\text{tot}}$ |
|----------------|------------|------------|------------|----------------------|
| With ETG       | 7.9%       | 6.6%       | 4.2%       | 33.1%                |
| No ETG         | 6.1%       | 7.3%       | 3.2%       | 24.1%                |

Figure 14. Comparison between JETTO+QuaLiKiz predictions for JET baseline scenario 87412 including heat, particle and momentum transport. The QuaLiKiz predicted and measured profiles are compared, for: ion temperature (top left panel), electron temperature (top right panel), electron density (bottom left panel) and toroidal velocity (bottom right panel). The QuaLiKiz boundary condition is at $\rho_{\text{norm}} = 0.85$. Two QuaLiKiz cases are compared, neglecting (blue curve) and including (red curve) ETG scales. The results are shown following 1 s simulation time.
we apply a squircular mapping from the normalised circle radius to the zone most likely to contain the instabilities. For ETG scales \( k \rho_i > 2 \), only instabilities in the electron diamagnetic direction are considered, saving CPU time due to avoiding the calculation of unnecessary contours.

Contour splitting and overlap
In total, 2 squircle contours are employed on both the positive and negative sides of the real axis. In the central region spanning the real axis, a more narrow contour is applied, with a 10% overlap with each of the neighbouring contours. This central contour increases the algorithm robustness, since numerical evaluation of \( D(\omega) \) is more difficult near the real axis, while most instabilities do not have frequencies with \( \omega_r \sim 0 \).

An example of the contour paths are shown in figure 15, for the GA standard case parameter set [55] at \( k_0 \rho_i = 0.5 \). An ITG mode is present, indicated in the figure. Each separate contour is signified by a different colour. The frequencies and growth rates are in units of \( n \omega_i \equiv \frac{2 \pi f}{\Omega} \), where \( \Omega \equiv 1 \text{ keV} \).

The optimised contour choices led to a significant speed-up of factor \( \sim 8 \) compared to the previous QuaLiKiz version.

A.2. Improved numerical methods for integral functions
The plasma dispersion function is extensively called during the evaluation of \( D(\omega) \), taking up a significant fraction of the computation time. This calculation time was reduced by switching from the standard ‘WofZ’ method [35] to the faster Weideman algorithm [36]. The Weideman method is based on a rational expansion whose coefficients are pre-calculated and tabulated. This speed-up does not lead to a reduction in accuracy. A comparison between the WofZ and Weideman methods for plasma dispersion function calculations resulted in relative agreements within 0.005%, with these maximum differences only reached in the vicinity of the function zeros [74]. In the same reference, local toroidal ITG calculations using either Weideman and WofZ resulted in negligible relative growth rate and frequencies differences of \( \sim 10^{-7} \).
Considering that the plasma dispersion relations are essentially identical between WofZ and Weideman, it is reasonable to assume that similar results are achieved for TEM and ETG calculations. In QuaLiKiz, a factor 2 speed-up in the eigenvalue solver was achieved when switching from WofZ to Weideman. Following a profiling of the code performance, a bottleneck due to elliptic integral evaluations was found. Hence, further speed-up was achieved by switching to a faster algorithm for elliptic integral calculations, from the Carlson method \cite{37} to the Fukushima method \cite{38}. Elliptic integrals emerge in QuaLiKiz due to bounce averaging of trapped particles. The algorithm improvement was of particular importance for trapped electrons in the presence of collisions. This is because the trapped electron energy integral then no longer takes a form that can be reduced to a plasma dispersion function. The velocity space integral must then be calculated numerically, involving multiple calls of the elliptic integrals. Switching to the Fukushima algorithm led to a further factor 1.5 speed-up in the QuaLiKiz eigenvalue solver calculation.

A.3. Code acceleration within integrated modelling applications

A significant further speed-up in computation time can be gained specifically in integrated modelling applications. This is based on the timescale separation between the $\Delta t$ timestep in the transport PDE solver, and the energy confinement and current diffusion times. While the transport coefficients are calculated (i.e. a QuaLiKiz call) every $\Delta t$ (~1 ms in a JET simulation), only relatively small modifications of the input parameters (kinetic profiles and magnetic equilibrium) occur within this time. Thus, instead of calculating the full contour solution at every QuaLiKiz call, the full calculation is only carried out every $n$th call. In the intermediate calls, QuaLiKiz skips the contour evaluation step, and proceeds straight to the Newton solver, using the previous eigenvalue solution from the $n-1$ step (saved within memory), as the initial guess. In this manner, the evolution of the eigenvalues are tracked on the complex plane. At each timestep, the displacement of the solution due to the evolving input parameters is small enough such that the Newton solver easily converges, based on the previous solution as the initial guess. The full contour solution is still necessary at periodic intervals, to allow the capture of new eigenvalues which may have appeared in the interim period, due to the evolving input profiles. It is not feasible in QuaLiKiz to calculate damped modes and follow them into the instability region, due to singularities in the dispersion relation integrals on the real axis which would excessively increase the computation time.

A subtle point for the intermediate steps (straight-to-Newton) must be taken into consideration if no new solution is found, e.g. if the mode is stabilised due to reduced driving gradients since the last call. In that case, while that
solution is passed to the saturation rule, resulting in zero or lowered fluxes, the last unstable solution for that wavenumber must still be kept as the initial guess for the next step. If this does not occur, then once a mode is predicted stable, it will remain stable until the next call when the full contour solution is once again carried out.

We have determined that an n corresponding to ~10 ms of plasma evolution provides a reasonable balance between speed-up and robustness. For small integrated modelling timesteps, a maximum n = 20 is set. The procedure is summarised in the flow chart in figure A2. This method leads to a further factor ~5 speed-up of QuaLiKiz application within an integrated modelling environment.

Appendix B. Improved eigenfunction solution method

B.1. Ion scales

The starting point is the local dispersion relation:

\[
D(\omega) = \sum_s \left( \int \frac{f_0 q_s^2}{T_s} \left( 1 - \frac{j_0^2}{\omega} - \frac{m \omega^*}{k ||V|| - n \omega_d} \right) \phi(x) \right) = 0.
\]

(B.1)

\( \int \) signifies integration over velocity space, s is a species identifier, \( n = \frac{k_\theta q}{q} \) is the toroidal mode number corresponding to the poloidal wavenumber \( k_\theta \), \( f_0 \) is the background Maxwellian, \( q_s \) the species charge, \( T_s \) the species temperature, \( j_0 \) the Bessel function from orbit averaging (whose arguments will be discussed below), \( \omega \) the Doppler shifted mode frequency in the rotating frame \( \omega - k_x E_x \) (where \( x \) is the distance from the mode surface, and \( \gamma_x \) the perpendicular mean flow shear), \( k || \) the parallel wave number \( k_g \) (with \( s \) the magnetic shear, \( q \) the q-profile, \( x \) the distance from the resonant surface, \( R \) the major radius), and \( V || \) the parallel velocity. \( \omega^*_S \) is the diamagnetic frequency:

\[
\omega^*_S = -\omega_d \left[ \frac{R}{L_{ns}} + \frac{R}{L_{T_s}} \right] \left( \xi - \frac{3}{2} - M(2V || - M) \right)
\]

(B.2)

\( \omega_d \equiv \frac{\gamma_x}{q R / a} \) is the Mach number \( U||/V_Ts \), \( U || \) being the mean flow in the background Maxwellian, and \( V_{Ts} \) the thermal velocity. \( \xi = \frac{V || + V_s}{V_s} \), the normalised kinetic energy. \( R \) is the major radius, and \( L_{ns}, L_{T_s} \) are the normalised logarithmic gradients of the temperature,
density, and velocities respectively. \( \dot{V}_i \) is the normalised velocity (phase space coordinate) \( V_i / V_T \), \( \omega_d \) is the magnetic drift frequency, whose expression differs for the trapped and passing species due to the trapped particle bounce averaging, as will be discussed below. \( \phi(\kappa) \) is the electrostatic potential, whose functional form we seek as the eigenfunction solution of \( D(\omega) = 0 \), together with the eigenvalue \( \omega \).

The key point in the approximated eigenfunction solution is a large \( \omega \) expansion, where \( \omega / \omega_d, \omega / k_i V_i \ll 1 \). The resonant denominators are expanded to second order, leading to:

\[
D(\omega) = \left[ \frac{n_e}{T_i} \left( 1 - \left( \left( 1 - \frac{n_i^0}{\omega} \right) \left( 1 + \frac{n_{\omega_d}}{\omega} + \frac{n^2 \omega^2_d}{\omega^2} \right) \right)_i \right) \right.
\]

\[
+ \sum_i \frac{n_i^0 T_i}{T_i} \left( 1 - \left( \left( 1 - \frac{n_i^0}{\omega} \right) \left( 1 + \frac{n_{\omega_d}}{\omega} + \frac{n^2 \omega^2_d}{\omega^2} \right) \right)_i \right)
\]

\[
\times \left( 1 - \frac{k_i^2 \omega^2}{2} \right) \left( 1 - \frac{k_i^2 \rho^2}{2} \right) \right) \right) - \left( 1 - \frac{n_i^0}{\omega} \right)
\]

\[
\times \left( 1 + \frac{n_{\omega_d}}{\omega} + \frac{k_i V_i}{\omega} + \frac{k_i^2 V_i^2}{\omega^2} + \frac{2 n_{\omega_d} k_i V_i}{\omega^2} + \frac{n^2 \omega^2_d}{\omega^2} \right)
\]

\[
\left( 1 - \frac{k_i^2 \rho^2}{2} - \frac{k_i^2 \rho^2}{2} \right) (\rho) \phi(\kappa).
\]

We have assumed a small wavenumber expansion in the Bessel functions. The Bessel functions are \( J_0 (k_i \rho) \) for the passing particles, and \( J_0 (k_i \rho) J_1 (k_i \rho) \) for the trapped particles following both bounce and gyro-orbit averaging [75]. Since \( \delta_i \ll \delta_i \) and \( \rho_i \ll \rho_i \), we assume \( J_0 (k_i \rho_i) \approx 1 \) and \( J_0 (k_i \rho_i) \approx 1 \) for the electron terms.

For passing particles:

\[
\langle A(\xi, \kappa) \rangle_p \equiv \frac{2 \eta}{\sqrt{\pi}} \int_0^\infty A(\xi \eta) e^{-\xi^2} \psi \int_0^\infty K(\kappa) \kappa
\]

\[
\times (1 + 2 M e / |\sqrt{\xi} (1 - \lambda \beta) d \xi). \quad \text{(B.4)}
\]

\( \kappa \) is related to the pitch angle \( \lambda \) by \( \kappa \equiv 1 - 2 \xi \kappa^2 \). \( K \) is the complete elliptic integral of the first kind, originating from bounce averaging in the small inverse aspect ratio expansion. We note that the poloidal dependence in \( V_i \) above, together with any poloidal dependence in \( \Lambda \), must first be bounce averaged in the trapped particle integral. This is discussed below.

For passing particles:

\[
\langle A(\xi, \kappa) \rangle_p \equiv \frac{2 \eta}{\sqrt{\pi}} \int_0^\infty A(\xi \eta) e^{-\xi^2} \psi \int_0^\infty K(\kappa) \kappa
\]

\[
\times (1 + 2 M e / |\sqrt{\xi} (1 - \lambda \beta) d \xi). \quad \text{(B.5)}
\]

where, in the small inverse aspect ratio expansion, the critical angle \( \lambda_c = 1 - 2 \xi \), and in the Jacobian:

\[
T(\lambda) = 2 \int_0^{\pi} \frac{1}{\sqrt{1 - \lambda (1 + 2 \xi \sin^2(\theta/2)))} d \theta. \quad \text{(B.6)}
\]

Originating from the magnetic trapping perturbation to the transit frequency in the \( dJ_2 = dE / \omega_2 \) term in the action-angle variable Jacobian, where \( \omega_2 \) is the passing particle angular transit frequency [72].

The vertical drift frequency \( n_{\omega_d} \) in the QuaLiKiz shifted circle geometry is:

\[
n_{\omega_d} = -n_{\omega_d} \xi (2 - \lambda \beta) (\cos \theta + \sin \theta (\delta \theta - \cos \theta)). \quad \text{(B.7)}
\]

For the trapped particle integration, this expression is bounce averaged over the poloidal angle, becoming \( n_{\omega_d} = -n_{\omega_d} \xi \psi(\kappa) \), with:

\[
f(\kappa) = \frac{2 E(\kappa)}{K(\kappa)} (1 - \frac{4}{3} \kappa^2) - 1
\]

\[
+ \left( 4 \delta + \frac{4}{3} \alpha \right) \left( \kappa^2 - 1 + E(\kappa) / K(\kappa) \right). \quad \text{(B.8)}
\]

where \( \alpha = 9 \sum_i \beta_i \gamma \left( \frac{R}{l_m} + \frac{R}{l_{\perp}} \right) \) and \( K(\kappa) \) and \( E(\kappa) \) are complete elliptic integrals of the first and second kinds, respectively.

For the passing ions, we assume that the eigenfunction is strongly ballooned, and expand \( \theta \) to second order, as well as invoking the small inverse aspect ratio expansion \( \xi^2 \to 0 \), leading to:

\[
n_{\omega_d} = -n_{\omega_d} \xi (2 - \lambda + \theta^2) (\delta - \alpha - \frac{1}{2} (2 - \lambda) - \frac{\lambda \xi}{2}). \quad \text{(B.9)}
\]

The form of the eigenfunction \( \phi(\kappa) \) is assumed to be a shifted Gaussian:

\[
\phi(\kappa) = \frac{\phi_0}{(\pi R(\kappa^2))^{1/4}} e^{-\left( \frac{k - x_0}{w^2} \right)^2}. \quad \text{(B.10)}
\]

From the definition of the \( k_\perp \) operator, \( k_\perp^2 = -\frac{\partial^2}{\partial \theta^2} \). Furthermore, with the strong ballooning approximation, the \( \theta \) coordinate is related to the Fourier transform of the radial coordinate, leading to \( \theta^2 = k^2 \rho^2 \), where \( d \) is the distance between rational surfaces. Assuming small shifts, \( x_0 \ll w^2 \), the above operators act on \( \phi(\kappa) \) as follows:

\[
\frac{\partial^2}{\partial \theta^2} \phi(\kappa) = \left( \frac{x^2}{w^2} - \frac{2 x_0 x}{w^2} - \frac{1}{w} \right) \phi(\kappa). \quad \text{(B.11)}
\]

Finally, we also assume low flow shear, \( \gamma_{\perp} \ll \omega \), leading to \( \omega^2 \approx \omega^2 - 2 \omega k_\theta \gamma_{\perp} \) and \( \omega^3 \approx \omega^3 - 3 \omega^2 k_\theta \gamma_{\perp} \).

With all the above definitions and assumptions, the approximated eigenfunction solution consists of inserting all
the expressions into equation (B.3), carrying out the energy
and pitch-angle integrations, and solving for \( w, x_0, \) and \( \omega. \)
The energy integrals are carried out analytically, util-
ing: \( \xi = \frac{3}{7} \mathcal{E}_{\text{c}}, \quad \xi^2 = \frac{15}{8}, \mathcal{E}_{\text{c}}, \quad \xi^4 = \frac{945}{16} \). The pitch-
angle integrations are carried out numerically.

The dispersion relation can be written in the form:

\[
D(\omega) = \left( -\omega^3 + 3 \omega^2 k_0 \gamma_e \varepsilon \nu_{\nu_T} \right) n_{\nu_T} \frac{\gamma_p z_i^2}{T_{\nu_T}} + \sum_i \left( -\omega^3 + 3 \omega^2 k_0 \gamma_e \varepsilon \nu_{\nu_T} \right) n_{\nu_T} z_i^2 \tau_i + \sum_i F_{\nu_T} n_{\nu_T} z_i^2 \tau_i = 0, \tag{B.12}
\]

where \( F_{\nu_T}, F_{\nu_T}, F_{\nu_T} \) contain all terms associated with the
velocity space integrations for trapped electrons, trapped ions,
and passing ions respectively.

We summarise the ordering. Taking \( \varepsilon \) as the small parameter, we assume:

\[
\mathcal{O}(k_0 x) \sim \mathcal{O}(d/w) \sim \mathcal{O}(\omega^2 / \omega) \sim \mathcal{O}(1),
\]

\[
\mathcal{O}(x_0 / w) \sim \mathcal{O}(\delta_i / \omega) \sim \mathcal{O}(\varepsilon^{0.25}),
\]

\[
\mathcal{O}(k_0 \gamma_i) \sim \mathcal{O}(\rho_i / w) \sim \mathcal{O}\left( \max \frac{L_T}{\nu_T}, \frac{L_n}{\nu_n} \right) \sim \mathcal{O}(\varepsilon^{0.5}),
\]

\[
\mathcal{O}(\omega_d / \omega) \sim \mathcal{O}(k_0^2), \quad \mathcal{O}(x_0 / \omega) \sim \mathcal{O}(\varepsilon^{0.75}),
\]

\[
\mathcal{O}(M_i / M_i) \sim \mathcal{O}(\delta_i / \omega) \sim \mathcal{O}(\varepsilon^{2}),
\]

We discard all terms \( \mathcal{O}(\varepsilon^2) \) and higher. This leads to:

\[
F_{\nu_T} = f_1 \left( \omega^3 - 3 k_0 \gamma_e \omega^2 x + \omega^2 2 k_0 \gamma_e \omega x \right) W_{010} + \frac{3}{2 f_T} (\omega^2 - 2 k_0 \gamma_e \omega x) W_{01e}^T
\]

\[
- \frac{3}{2 f_T} (\omega - k_0 \gamma_e \omega x) W_{01e}^T \left( W_{d1} + W_{d2} \right)
\]

\[
+ \frac{3}{2 f_T} W_{V10}^T \left( \omega + W_{d1} + W_{d2} \right) = F_{\nu_T,0} + F_{\nu_T,1x}, \tag{B.13}
\]

\[
F_{\nu_T} = f_1 \left( \delta_i \omega^3 \hat{\rho}_i - 3 k_0 \gamma_e \omega^2 x + \omega^2 \hat{\rho}_i \left( W_{d10} + W_{d11} \right) \right)
\]

\[
- 2 k_0 \gamma_e \omega x W_{d10} \delta_i - 3 f_T \omega^2 \delta_i - 3 \hat{k}_i \omega x W_{V10}^T
\]

\[
- \frac{3}{2 f_T} \omega^2 W_{V10}^T W_{d10} + \frac{3}{2 f_T} W_{V10}^T \left( \omega + W_{d1} + W_{d2} \right)
\]

\[
+ \frac{3}{2 f_T} \omega W_{V10}^T W_{d10} - 3 \frac{f_T}{2} \omega W_{d10} x + \frac{3}{2 f_T} \omega W_{d10} x^2 - 3 \frac{f_T}{2} \omega W_{d11} x^2
\]

\[
= F_{\nu_T,0} + F_{\nu_T,1x} + F_{\nu_T,2x^2}, \tag{B.14}
\]

The definitions above are as follows. For the trapped species:

\[
W_{V10}^T = n \omega d \langle f(k) \rangle \lambda, \quad W_{V10}^T = n^2 \omega d (f(k)) \lambda, \quad V_T \equiv \left\langle \frac{2 \bar{E}}{K(k)} \right\rangle_{\lambda},
\]

where \( E(\sin^{-1} \kappa, 1/\kappa) \) is an incomplete elliptic integral of the second kind, arising from the bounce averaged \( V_T^2 \):

\[
V_T^2(\lambda) = \frac{\bar{E}}{\beta \sqrt{1 - \lambda^2 b}}
\]

\[
= \frac{2 \bar{E}}{\beta \sqrt{1 - \lambda^2}} E(\sin^{-1} \kappa, 1/\kappa) K(k),
\]

\[
= 2 \pi \left( \frac{E(\kappa)}{K(\kappa)} - (1 - k^2) \right). \tag{B.15}
\]
where we have utilised a convenient identity linking the incomplete elliptic integral of the second kind with the specific arguments we have obtained, to complete elliptic integrals of the first kind [76].

For the passing species:

\[ W_{V1\alpha} = n\omega_{f\alpha} \left( 2 - \frac{\langle \lambda \rangle_{\alpha}}{f_p} \right), \]

\[ W_{V2\alpha,0} = n\omega_{f\alpha} \left( \hat{s} - \alpha - \frac{1}{2} \right) \left( 2 - \frac{\langle \lambda \rangle_{\alpha}}{f_p} \right), \]

\[ W_{V2\alpha,1} = -n\omega_{f\alpha} \frac{\langle \lambda \rangle_{\alpha}}{2f_p}, \]

\[ W_{V3\alpha} = n\omega_{f\alpha} (1 - \lambda)(2 - \lambda), \]

\[ W_{V4\alpha,0} = n\omega_{f\alpha} \left( \left( \hat{s} - \alpha - \frac{1}{2} \right) (2 - \lambda)(1 - \lambda) \right), \]

\[ W_{V5\alpha} = n^2\omega_{f\alpha}^2 (2 - \lambda)^2, \]

\[ W_{V6\alpha} = n^2\omega_{f\alpha}^2 \left( 2(2 - \lambda) \left( \hat{s} - \alpha - \frac{1}{2} \right) (2 - \lambda) \right). \]

\[ V_1 = 1 - \langle \lambda \rangle_{\alpha}, \]

\[ V_2 = -\frac{\hat{s}}{2}, \]

\[ k_s = k_0 V_{T\alpha}, \]

\[ \hat{\rho} = 1 - \frac{k_{0\alpha}^2 \rho_{\alpha}^2}{2}. \]

Following insertion of equations (B.13)-(B.15) into equation (B.12), the final step is to separate the terms proportional to \( x^0 \) (i.e., 1, \( x \), and \( x^2 \)), such that:

\[ D(\omega) = D_0(\omega) + D_1(\omega)x + D_2(\omega)x^2 = 0. \] (B.17)

With:

\[ D_0(\omega) = -\omega^3 \left( 1 + Z_{ei} \frac{T_e}{T_i} \right) + \sum_i \left( F_{T_i,0}(\omega) + F_{P_i,0}(\omega) \right) \frac{n_i Z_i^2}{T_i}. \] (B.18)

\[ D_1(\omega) = 3\omega^2 k_0 \gamma_{Te} \frac{n_i}{T_e} \left( 1 + Z_{ei} \frac{T_e}{T_i} \right) + \sum_i \left( F_{T_i,1}(\omega) + F_{P_i,1}(\omega) \right) \frac{n_i Z_i^2}{T_i}. \] (B.19)

\[ D_2(\omega) = \sum_i \left( F_{T_i,2}(\omega) + F_{P_i,2}(\omega) \right) \frac{n_i Z_i^2}{T_i}. \] (B.20)

We have assumed all ion temperatures to be equal, allowing the emergence of \( Z_{ei} \) in the ion summation.

Following [77], we solve equation (B.17) in a perturbative manner. The eigenvalue \( \omega_{0i} \) is first solved from a modified form of \( D_0(\omega) = 0 \), related to the local toroidal ITG dispersion relation at \( \theta = 0 \), neglecting rotation, and setting \( k_v = 0 \). Following multiplication by \( T_i/n_i \) for convenience, this results in the following 3rd degree polynomial:

\[ C_3 \omega^3 + C_2 \omega^2 + C_1 \omega + C_0 = 0. \] (B.21)

Defining \( \alpha_i \equiv \frac{Z_i^2 n_i}{n_i T}, \) we obtain:

\[ C_3 = -1 + f_i + \sum_i \alpha_i (-1 + \hat{\rho}_i), \] (B.22)

\[ C_2 = f_i W_{d_1,0} - \frac{3}{2} W_{V_{1,0}} + \sum_i \alpha_i \frac{\hat{\rho}_i}{\hat{\rho}}, \] (B.23)

\[ C_1 = -\frac{3}{2} W_{V_{1,0}} + W_{d_2,0} + \frac{3}{2} W_{V_{1,0}} + \frac{3}{2} W_{V_{1,0}} \]

\[ + \sum_i \alpha_i \left( -\frac{3}{2} W_{V_{1,0}} \hat{\rho}_i (W_{d_1,0} + W_{d_2,0}) + \frac{3}{2} W_{V_{1,0}} \right) - \frac{3}{2} f_i \hat{\rho}_i (W_{d_1,0} + W_{d_2,0}) W_{V_{1,0}} + \frac{15}{4} W_{V_{1,0}}. \] (B.24)

\[ C_0 = \frac{3}{2} W_{V_{1,0}} (W_{d_1,0} + W_{d_2,0}) + \sum_i \alpha_i \left( \frac{3}{2} W_{V_{1,0}} (W_{d_1,0} + W_{d_2,0}) \right) + \frac{15}{4} (W_{d_1,0} + 2W_{d_2,0}) W_{V_{1,0}}. \] (B.25)

The \( V \) expressions are geometric terms related to integrations over even powers of \( V_{\alpha} \). The \( W_{V} \) term arise from integrations including the vertical drift frequency. The \( W_i \) terms arise from integrations over the diamagnetic frequencies. The subscript system is convenient due to the ordering of the various terms.
Equations (B.20) and (B.19) then provide the solutions for the eigenfunction width and shift respectively, following substitution of the eigenvalue solution $\omega_0$ from equation (B.21). Here we maintain the rotation terms and finite $\theta$. This leads to the following expression for the eigenfunction width $w$.

$$w = \left( \sum_i \alpha_i u_0^2 + \omega^2 A_{1i} + \omega^2 A_{2i} \right)^{0.25} \left( \sum_i \kappa_i^2 \left( \omega B_{1i} + B_{0i} \right) \right)$$  

(B.26)

with

$$A_{1i} = f_i \delta^2 + f_p \rho_i^2 \frac{1}{2},$$  

$$A_{2i} = f_i \delta^2 \left( W_{i0,0} + W_{i1,1} - \frac{3}{2} W_{i1} \right)$$

$$+ f_p \rho_i^2 \left( W_{i1,0} + f_p 3/2 d^2 W_{i2,0} \right),$$

$$A_{1i} = -\delta_i \frac{3}{2} W_{i1} + \frac{3}{2} f_p d^2 (W_{i1,0} + W_{i2,0}) W_{i2,0},$$

$$B_{1i} = \frac{3}{2} V_i,$$

$$B_{0i} = \frac{3}{2} V_i (W_{i0,0} + W_{i2,0}).$$

(B.27) - (B.31)

The eigenfunction shift is given by:

$$x_0 = -\frac{\gamma_F (U_{i0} + \sum_i \alpha_i U_{i1}) + \sum_i \alpha_i U_{i2}}{\sum_i \alpha_i D_i}.$$

(B.32)

There are two tuning factors in this process. Firstly, the eigenfunction width, before insertion into equation (B.32), is multiplied by a factor $\frac{0.1}{k_i} \omega^{0.25}$. This factor was tuned by comparison to full gyrokinetic predictions at the GA-standard case. Then, following this, the width is normalised by the factor $\max(0, 0.66d/w)$, where $d$ is the distance between neighbouring rational surfaces. This is to reduce the occasional emergence of non-physical narrow width solutions, which can significantly slow down the full kinetic dispersion relation solution procedure.

**B.2. Electron scales**

For the electron scale fluid eigenfunction solution, we assume adiabatic ions, and now include passing electrons. Electron scales are defined in QuaLiKiz as $k_i \rho_i > 2$. The ordering is similar, apart from $O(k_i/r) \sim O(\hat{\epsilon}^{0.25})$, $O(k_i \rho_i) \sim O(\hat{\epsilon}^{0.5})$, $O(\omega_i/\omega) \sim O(\hat{\epsilon})$, and $O(\gamma_F/\omega) \sim O(\hat{\epsilon}^2)$. The dispersion relation is now simpler:

$$D(\omega) = \left[ \left( 1 - \frac{n_{\omega_i}}{\omega} \right) + \frac{n_{\omega_i}}{\omega} \right] \left[ \frac{\hat{\epsilon}^2 \rho_i}{\omega^2} + \left( 1 - \frac{n_{\omega_i}}{\omega} \right) \frac{k_i V_i}{\omega} + \frac{k_i^2 V_i^2}{\omega^2} \right] - 1 - \frac{T_e}{T_i} \frac{Z_{\text{eff}}}{4} = 0.$$  

(B.37)

Which can be written as:

$$D(\omega) = F_{e}^{\text{ETG}} + F_{p}^{\text{ETG}} - \omega^3 (1 + \tau_e) = 0.$$  

(B.38)

With $\tau_e \equiv T_e / T_i Z_{\text{eff}}$. The expressions for $F_{e}$ and $F_{p}$ are also relatively simpler, due to the ordering out of any rotation related terms

$$F_{e}^{\text{ETG}} = f_i \left[ \omega \hat{\epsilon} + \omega^2 \hat{\epsilon}^2 (W_{i1} - \frac{3}{2} \omega^2 W_{i1}) \right]$$

$$+ \frac{3}{2} k_i \rho_i \left( W_{i1} + \frac{3}{2} k_i \rho_i (W_{i1,0} + W_{i2,0}) \right)$$

$$k_i \rho_i (W_{i1,0} + W_{i2,0}) - \frac{3}{2} f_p \omega^2$$

$$+ \delta_i \frac{3}{2} k_i \left( W_{i1} + \frac{3}{2} k_i \rho_i (W_{i1,0} + W_{i2,0}) \right)$$

$$+ \frac{3}{2} k_i \rho (W_{i1,0} + W_{i2,0}) \left( W_{i1} + \frac{3}{2} k_i \rho_i (W_{i1,0} + W_{i2,0}) \right).$$

(B.39)

$$F_{e}^{\text{ETG}} = f_i \left[ \omega \hat{\epsilon} - \frac{3}{2} \omega^2 W_{i1} - \frac{d^2}{w^2} \left( \frac{x^2}{w^2} - 1 \right) \right]$$

$$+ \frac{3}{2} k_i \rho_i \left( W_{i1} + \frac{3}{2} \omega (W_{i1,0} + W_{i2,0}) \right)$$

$$\times \left( W_{i1} - \frac{d^2}{w^2} \left( \frac{x^2}{w^2} - 1 \right) \right)$$

$$+ \frac{3}{2} k_i \rho_i \left( W_{i1,0} + W_{i2,0} \right) \frac{x^2}{w^2}$$

$$+ \frac{\rho_i^2}{w^2} \left( \frac{x^2}{w^2} - 1 \right) \left( \omega^3 + \omega^2 W_{i1,0} \right).$$

(B.40)
The rest of the ETG solution proceeds as for the ion scales. Equation (B.38) is solved by separating the terms proportional to $x^0$ and $x^2$, and setting each resultant equation to zero. In the perturbative scheme, the $x^0$ polynomial equation is solved at $\theta = 0$ and setting $k_p = 0$, providing the toroidal ETG eigenvalue. The full equation with terms proportional to $x^2$ then provides the eigenfunction width, when substituting the eigenvalue from the polynomial solution. The coefficients in the polynomial equation are as follows:

$$C_2 \omega^2 + C_1 \omega + C_0 = 0$$  \hspace{1cm} (B.41)

with:

$$C_2 = -1 - \tau_v + \bar{\rho}_e, \hspace{1cm} (B.42)$$

$$C_i = W_{\text{d}1} f_i \bar{\rho}_e - \frac{3}{2} W_{\text{d}11} \bar{\rho}_e - \frac{3}{2} f_p W_{\text{v}11} + W_{\text{d}10} f_p \bar{\rho}_e, \hspace{1cm} (B.43)$$

$$C_0 = -\frac{3}{2} (W_{\text{d}0} + W_{\text{d}22}) W_{\text{v}11}^R - \frac{3}{2} f_p (W_{\text{d}10} + W_{\text{d}22}) W_{\text{v}11}^R. \hspace{1cm} (B.44)$$

For the width equation, we obtain:

$$w = \left( -\frac{\omega^3 A_3 + \omega^2 A_2 + \omega A_1}{k_f^2(\omega B_1 + B_0)} \right)^{0.25} \hspace{1cm} (B.45)$$

with:

$$A_1 = f_l \frac{\rho_e^2}{2} + f_p \frac{\rho_p^2}{2}, \hspace{1cm} (B.46)$$

$$A_2 = f_l \frac{\rho_e^2}{2} W_{\text{d}10} + f_p \frac{\rho_p^2}{2} W_{\text{d}10} + \frac{3}{2} f_p d^2 W_{\text{v}20}, \hspace{1cm} (B.47)$$

$$A_1 = \frac{3}{2} f_p d^2 (W_{\text{d}10} + W_{\text{d}22}) W_{\text{v}20}, \hspace{1cm} (B.48)$$

$$B_1 = \frac{3}{2} V_l, \hspace{1cm} (B.49)$$

$$B_0 = \frac{3}{2} V_l (W_{\text{d}10} + W_{\text{d}22}). \hspace{1cm} (B.50)$$

### Appendix C. Trace heavy impurity transport coefficients with asymmetries

The impact of asymmetries on the transport coefficients is through the weighted flux surface average of the modified $n$ and $\frac{dn}{dr}$ in the quasilinear transport integrals. The averages are weighted by the electrostatic potential eigenfunction.

We closely follow the procedure outlined in [9], with two generalisations. We include temperature anisotropies, and also build the transport coefficients from a spectrum of quasi-linear weights.

The modified density is:

$$n_j(r, \theta) = n_{R,\text{LFS}}(r) P_j(r, \theta) \exp \left( \frac{-\epsilon_j(r, \theta)}{T_j(r)} \right) \hspace{1cm} (C.1)$$

with

$$\epsilon_j = Z_j e \phi(r, \theta) - \frac{m_j \Omega_j^2(r)}{2} (R^2(\theta, r) - R_{\text{LFS}}^2(r)) \hspace{1cm} (C.2)$$

and

$$P(r, \theta) \equiv \frac{T_{\text{L}}(r, \theta)}{T_J(r)} = \frac{T_{\text{L,\text{LFS}}}(r)}{T_{J,\text{LFS}}(r)} + \left( 1 - \frac{T_{\text{L,\text{LFS}}}(r)}{T_{J,\text{LFS}}(r)} \right) B_{\text{LFS}}(r, \theta) \right]^{-1}, \hspace{1cm} (C.3)$$

where $\Phi(r, \theta)$ is solved by imposing the quasineutrality constraint at each poloidal location.

We define the following weighted flux surface averaging:

$$\langle A(\theta) \rangle \equiv \frac{\oint A(\theta)(1 + \tau \cos \theta)e^{-w^2 d^2 d\theta}}{\oint (1 + \tau \cos \theta)e^{-w^2 d^2 d\theta}} \hspace{1cm} (C.4)$$

$d$ is the distance between neighbouring rational flux surfaces, and $w$ is the eigenfunction width as calculated by QuaLiKiz.

The following expressions are then convenient. They are similar to those in [44], and are related to various contributions to $n$ and $\frac{dn}{dr}$.

$$e_0 \equiv \langle P e^{-\epsilon/T} \rangle, \hspace{1cm} e_1 \equiv \langle \frac{Z e \Phi P e^{-\epsilon/T}}{T} \rangle, \hspace{1cm} e_2 \equiv \langle R_0 \frac{Z e \Phi P e^{-\epsilon/T}}{T} \rangle, \hspace{1cm} e_3 \equiv \frac{1}{R_0^2} \langle (R^2(\theta) - R_{\text{LFS}}^2) P e^{-\epsilon/T} \rangle, \hspace{1cm} e_4 \equiv \frac{1}{R_0} \left( R \frac{R \left( R \frac{dR_{\text{LFS}}}{dr} - R_{\text{LFS}} \frac{dR_{\text{LFS}}}{dr} ight) \right) P e^{-\epsilon/T} \rangle, \hspace{1cm} e_6 \equiv \frac{1}{R_0} \left( \frac{dP}{dr} e^{-\epsilon/T} \right), \hspace{1cm} e_7 \equiv \frac{Z e}{T} \langle P e^{-\epsilon/T} \phi \frac{d\phi}{d\tau} \rangle, \hspace{1cm} e_8 \equiv \langle (P e^{-\epsilon/T} (1 + \hat{\tau} \cos \theta)(\cos \theta + \hat{\tau} \sin \theta)) \rangle, \hspace{1cm} e_9 \equiv \frac{2}{R_0} \left( -R_{\text{LFS}} \frac{dR_{\text{LFS}}}{dr} P e^{-\epsilon/T} \right), \hspace{1cm}$$

Where $R_0$ is the normalising major radius.

The particle transport coefficients (for a given species) in the QuaLiKiz output, are then defined as follows, with respect to the input low-field-side densities and density gradient:

$$\Gamma = -D \frac{dn_{\text{LFS}}}{dr} + (V_T + V_C + V_{\text{TCF}} + V_{\text{UCF}} + V_{\text{PCF}}) n_{\text{LFS}}, \hspace{1cm} (C.5)$$

where $D$, $V_T$, and $V_C$ are the poloidal asymmetry modified diffusivity, thermodiffusive pinch, and compressional pinch.
respectively.

\[ D = \sum_k D_k e_{ik}, \]

\[ V_T = \sum_k V_k e_{ik}, \]

\[ V_C = \sum_k V_{ik} e_{ik}. \]

\( D_k, V_k, \) and \( V_C \) are the original wavenumber dependent diffusivity and pinch terms, as calculated from the quasilinear integrals without poloidal asymmetries, and already include all nonlinear saturation rule spectral weights and normalisations. The summation over \( k \) is necessary due to the wavenumber dependent eigenfunction weighting in \( e_{ik} \).

Borrowing from the terminology in [9], \( V_{TCF}, V_{UCF}, \) and \( V_{PCF} \) can be interpreted as additional thermodiffusive pinch, rotational diffusive pinch, and pure pinch terms, arising due to the poloidal asymmetries. Physically, they arise from a modifica-
tion of the effective diffusion coefficient due to poloidal asymmetry in the density gradient. The terms are the follows:

\[ V_{TCF} = \sum_k \frac{D_k}{K_0} \left( \frac{R}{L_T} \right) \left( e_{ik} - \frac{m \Omega^2 R_i^2}{2T} e_{ik} \right), \]

\[ V_{UCF} = \sum_k \frac{2D_k M}{K_0} \frac{\partial e_{ik}}{\partial r}, \]

\[ V_{PCF} = \sum_k \frac{D_k}{K_0} \left( e_{ik} - e_{ik} - 2M^2 \left( e_{ik} + e_{ik} + e_{ik} \right) \right). \]

The main differences between this formulation and [9] are: the summation over \( k \) for the total quasilinear flux; the inclusion of the \( P \) and \( e_{ik} \) temperature anisotropy terms.

The QuaLiKiz outputted transport coefficients are all with respect to \( \rho_{LFS} \) and \( \frac{d\rho_{LFS}}{dr} \). In a transport code, such as JETTO, where the evolved densities are rather the flux-surface-average FSA quantities, then the transport coefficients must be transformed to be respective to the FSA quantities.

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