An accurate treatment of scattering and diffusion in piecewise power-law models for cosmic ray and radiation/neutrino transport

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ABSTRACT

A popular numerical method to model the dynamics of a ‘full spectrum’ of cosmic rays (CRs), also applicable to radiation/neutrino hydrodynamics, is to discretize the spectrum at each location/cell as a piecewise power law in ‘bins’ of momentum (or frequency) space. This gives rise to a pair of conserved quantities (e.g. CR number and energy) that are exchanged between cells or bins, which in turn give the update to the normalization and slope of the spectrum in each bin. While these methods can be evolved exactly in momentum-space (e.g. considering injection, absorption, continuous losses/gains), numerical challenges arise dealing with spatial fluxes, if the scattering rates depend on momentum. This has often been treated either by neglecting variation of those rates ‘within the bin,’ or sacrificing conservation – introducing significant errors. Here, we derive a rigorous treatment of these terms, and show that the variation within the bin can be accounted for accurately with a simple set of scalar correction coefficients that can be written entirely in terms of other, explicitly evolved ‘bin-integrated’ quantities. This eliminates the relevant errors without added computational cost, has no effect on the numerical stability of the method, and retains manifest conservation. We derive correction terms both for methods that explicitly integrate flux variables (e.g. two-moment or M1-like) methods, as well as single-moment (advection-diffusion, FLD-like) methods, and approximate corrections valid in various limits.

Key words: MHD – plasmas – methods: numerical – cosmic rays – ISM: structure – galaxies: evolution.

1 INTRODUCTION

Understanding cosmic ray (CR) propagation and dynamics in the interstellar medium (ISM) and circum/intergalactic medium remains an unsolved problem of central importance in space plasma physics (Zweibel 2013, 2017; Amato & Blasi 2018; Kachelr¨uss & Semikoz 2019), with major implications for fields ranging from astrophysics, planet, star, and galaxy formation (e.g. Chen, Bryan & Salem 2016; Girichidis et al. 2016; Pakmor et al. 2016; Salem, Bryan & Corlies 2016; Simpson et al. 2016; Ruszkowski, Yang & Zweibel 2017; Wiener, Pfrommer & Oh 2017; Butsky & Quinn 2018; Farber et al. 2018; Jacob et al. 2018; Chan et al. 2019; Su et al. 2020; Hopkins et al. 2020; Ji et al. 2020, 2021; Bustard & Zweibel 2021).

In models which seek to dynamically evolve the CR population on large scales [as opposed to either historical semi-analytical models, which solve for the equilibrium CR distribution function (DF) in a static analytical Galaxy model, e.g. Korsmeier & Cuoco 2016; Evoli et al. 2017; Amato & Blasi 2018; Liu, Yao & Guo 2018, or particle-in-cell type simulations which model the dynamics of individual CRs], a central challenge is the high dimensionality of the DF $f(x, p, t)$ as a function of position $x$, CR momentum $p$, and time $t$. Recently, a number of studies (Girichidis et al. 2022; Hanasz, Strong & Girichidis 2021; Hopkins et al. 2022b; Ogrodnik, Hanasz & W˚olta´nski 2021) have addressed this by implementing variations of the method proposed in Girichidis et al. (2020) (with broadly similar methods used earlier in e.g. Jun & Jones 1999; Miniati 2001, 2007; Miniati et al. 2001; Jones & Kang 2005; Mimica et al. 2009; Yang & Ruszkowski 2017; Winner et al. 2019 as well), wherein the isotropic part of the DF $f_i$ is represented as a piecewise power-law function of momentum, in ‘bins’ of $p$ spanning some dynamic range; one can then integrate (to arbitrary precision) bin-to-bin fluxes of conserved CR number and energy (representing e.g. continuous loss or gain processes) or source/sink terms (injection or catastrophic losses or secondary production) in momentum space.

The method has many advantages. (1) Because real CR spectra are smooth and power-law-like over a wide dynamic range, these studies have shown that the spectrum over some very wide dynamic range can be represented accurately with a relatively small number of bins per species, imposing modest computational and memory cost. (2) The momentum-space and coordinate-space (advection/streaming/diffusion) operations can be operator-split, allowing the spatial part of the equations to be integrated with standard, well-studied and high-order numerical methods (exactly identical to previous treatments that considered just a single CR ‘fluid’ or bin or total energy density scalar field, e.g. Salem et al. 2016; Ruszkowski et al. 2017; Butsky & Quinn 2018; Chan et al. 2019; Hopkins et al. 2020; Ji et al. 2020, 2021; Su et al. 2020; Bustard & Zweibel 2021). (3) Conservation of number and energy is manifest, which ensures robustness of many results even in highly noisy conditions or in extreme injection/loss events. (4) It is accurate and converges efficiently in momentum-space. (5) It
trivially generalizes for methods that evolve either the ‘two-moment’ equations for the CR DF (where one evolves both the isotropic part of the DF and its flux, or equivalently the mean CR pitch angle), or ‘one-moment’ equations (where one assumes the flux is in local steady-state, so evolve just the isotropic part of the DF subject to a diffusion+streaming equation), as well as to even-further-simplified models (e.g. replacing the correct anisotropic diffusion with isotropic diffusion). These and other advantages have led, for example, to the first simulations simultaneously evolving multispecies CR spectra alongside ‘live’ fully coupled MHD dynamics on Galactic scales (Hopkins et al. 2022b).

However, while the momentum-space properties of this class of piecewise-power-law methods are very well defined (and easy to demonstrate), there is a known conceptual challenge in coordinate-space. Specifically, given some piecewise-power-law representation of $f$ in a ‘bin,’ the spatial flux of $f$ should depend on momentum, varying ‘across the bin.’ But since the flux depends itself on gradients of various moments of the DF itself, a naive attempt to integrate or average the flux over the bin leads to expressions of the form $\int d^3p \, \mathcal{F}[p, f[p], \ldots] \cdot \mathcal{G}[p, f[p], \ldots]$, where $\mathcal{F}$ and $\mathcal{G}$ are some arbitrary tensor functions. These are not just complicated, but appear at first to require ‘sub-binning’ of $f$ into infinitesimally small bins, each of which has a separately computed gradient, in order to evaluate accurately (Girichidis et al. 2020). As a result, most studies above have adopted the ‘bin-centred’ approach, wherein one assumes that all quantities of relevance for computing spatial fluxes are assumed to be constant over the momentum-width of a bin. This retains advantages (1), (2), (3), and (5) above, but leads to well-known artefacts in the spectrum when spatial transport (e.g. diffusion) dominates the escape time, sacrificing some of (4). Alternative approaches have been discussed (e.g. Girichidis et al. 2022), but (as noted by these authors) these generally sacrifice all of (2), (3), and (5); in particular the proposed non-bin-centred methods sacrifice conservation and consistency (they cannot be derived from the underlying DF equations) and can potentially lead to numerical instability or unphysical behaviours when momentum-space terms (e.g. losses) dominate.

In this paper, we derive a consistent treatment of these terms that resolves all of the challenges above and retains all of advantages (1–5) above. By considering a two-moment pitch-angle expansion of the Vlasov equation on scales large compared to CR gyro-radii, we show that the key conceptual ingredient required to resolve these issues is a consistent treatment of how the mean CR pitch angle varies across a ‘bin’. But we also show that the structure of the equations imposes consistency conditions that determine this at the level of approximation needed for the piecewise-power-law reconstruction. With this properly treated, we show the corrected numerical method is structurally identical to the ‘bin-centred’ approximation with appropriate scalar correction coefficients that are determined entirely in terms of already-evolved numerical quantities. We further show that the correction coefficients can be (self-consistently) even-further simplified if either (1) only the one-moment equation for the CRs is dynamically evolved, or (2) one only needs to capture the exact behaviour in all relevant limits of the local-steady-state flux equation (e.g. one is interested primarily in time-scales long compared to CR scattering times).

While our primary motivation in this paper is focused on applications to CRs, this qualitative method, and the challenges above, also apply in principle to analogous methods which evolve spectra of other collisionless species (e.g. radiation or neutrinos) as piecewise-power laws in similar fashion (e.g. Baschek et al. 1997). In this context, most ‘moment-based’ multigroup methods for radiation-hydrodynamics have focused on evolving just the radiation/neutrino energy in each ‘bin’ (e.g. Castor 2007), effectively equivalent to representing the spectrum as piecewise-constant, rather than a piecewise power law. Although conceptually simpler, the piecewise-constant approach requires an order of magnitude larger number of ‘bins’ across some frequency or energy range in order to represent spectra with steep or dynamically evolving power-law slopes, and sacrifices the ability to simultaneously conserve number and energy. A method like the piecewise-power-law scheme above for neutrinos has been discussed in e.g. Rampf & Janka (2002) and Müller, Janka & Dimmelmeier (2010) (their ‘simultaneously number-and-energy-conserving scheme,’ although it is described in different language than we use here), but similar conceptual difficulties (see Mezzacappa et al. 2020) have limited its application.

2 A METHOD FOR HANDLING FLUXES OF PIECEWISE-POWER-LAW SPECTRA

2.1 Set-up and definitions

Consider a population of CRs with some phase-space DF $f = dN_{\text{ei}}/d^3x \, d^3p$, with polar momentum coordinates $p = |p|$, pitch angle $\mu \equiv \cos \theta \equiv \hat{p} \cdot \hat{b}$ (where $\hat{b} \equiv B/|B|$ is the magnetic field direction), and phase angle $\phi_p$. The comoving evolution equations for the spatial or coordinate-space part of the first two $\mu$-moments of $f$ can be written (Hopkins, Squire & Batusky 2022a):

$$D_t \tilde{f}_0 = -\nabla \cdot (v \, \tilde{f}_1) + \ldots,$$

$$D_t \tilde{f}_1 + v \cdot \nabla \cdot (v \, \tilde{f}_0) = -\tilde{D}_{\mu \nu} \tilde{f}_1 - \tilde{D}_{\mu \rho} \partial_\rho \tilde{f}_0 + \ldots,$$

where $\tilde{f}_0 \equiv (4\pi)^{-1} \int d^3m \, \phi_p \, \mu^2 \, f$, so $\tilde{f}_0$ is the isotropic part of the DF and $\tilde{f}_1 \equiv (\mu) \, \tilde{f}_0$; $\tilde{D}_i X = \tilde{D}_i \left( \frac{\partial X}{\partial \rho} \right) \equiv \tilde{D}_i \left( \frac{\partial X}{\partial \rho} \right)$; $\chi \equiv (1 - \eta/2)$ and $\tilde{D} = \chi \hat{\nabla} + (1 - 3 \chi) \hat{b} \otimes \hat{b}$; $\tilde{D}_{\mu \rho} \equiv \tilde{v} \varepsilon_\mu$ is the pitch-angle averaged scattering rate (at the given $p$ and $x$); and $\tilde{D}_{\mu \nu} \equiv \varepsilon_\mu \varepsilon_\nu \, \varepsilon_\kappa / e$ in terms of the CR velocity $v = \beta c$ and $\varepsilon_\kappa \equiv v_\kappa / (v_\kappa - v_\parallel) / (v_\parallel + v_\perp)$ in terms of the ‘forward’ and ‘backward’ scattering coefficients $v_\kappa$ and phase speed $v_\parallel$ of gyro-resonant Alfvén waves (those with wavelength $\sim r_{\text{gyro}}$). We stress that equations (1)–(2) are valid for any arbitrary gyrotrropic DF: different ‘closure’ assumptions relate to how $\mu^2$ is specified (see Hopkins et al. 2022a), which is not important for our purposes.

In equations (1)–(2), the ‘...’ refers to terms which do not propagate CRs in coordinate space (e.g. injection & catastrophic losses $D_{\text{inj}} = j$, and continuous energy loss/gain processes $D_{\text{loss}} = p^{-2} \partial_\beta (p^2 \ldots)$). These can be operator-split and solved accurately with methods like those in Section 1 (Girichidis et al. 2020, 2022; Hanasz et al. 2021; Hopkins et al. 2022b; Ogrodnik et al. 2021), which model the spectrum as a piecewise-power law. In these methods, within infinitesimally small volume domain $j$, for each CR species $s$, within some ‘bin’ $m$ defined over a momentum interval $p^- < p < p^+$, we

\footnote{For our purposes here, different species of CR are linearly independent so it is sufficient to consider the DF for a single species (the total DF can then be reconstructed by simply summing over species).}

\footnote{Equation (1) formally follows from the Vlasov equation, with the standard quasi-linear scattering terms from Schlickeiser (1989), assuming the DF is approximately gyrotrropic, expanding to leading order in $\mathcal{O}(r_{\text{gyro}}/L_{\text{max}})$ (the ratio of gyro radius to resolved macroscopic scales) and $\mathcal{O}(\eta / e)$ (ratio of background MHD bulk velocities to $e$).}
assume that \( \bar{f}_0 \) can be represented as a power law with slope \( \alpha_{\bar{f}_0} \), i.e.:

\[
\bar{f}_{0,j,m,s} \approx (\bar{f}_0)_{b0,j,m,s} \left( \frac{p}{p_{0,j,m,s}} \right)^{\alpha_{\bar{f}_0}},
\]

where for analytical convenience we define \( p_{0} \equiv (p^+ p^-)^{1/2} \) as the geometric mean momentum of the ‘bin’. It is immediately obvious that the spatial part of equations (1)–(2) is independent for each ‘bin’ \( m \) and species \( s \) (i.e. there is no cross-term in equations 1–2 coupling different species or momenta), so we only need to consider one such bin to completely specify the numerical method. We therefore drop the \( j, m, s \) notation for brevity, with the understanding that all quantities considered here can (and should) depend on \( s, m \), and spatial location.

For reference below we also define \( \xi \equiv p^+/p^- \) as a dimensionless ‘bin width’.

### 2.2 Conserved quantities and the spatial flux

Given our power-law representation of \( \bar{f}_0 \) in equation (3) with two parameters \((\bar{f}_0)_{b0} \) and \( \alpha_{\bar{f}_0} \), we can clearly represent or evolve exactly two independent conserved scalar quantities of the DF (and their associated fluxes as we show below) associated with each bin. These are typically chosen to be the CR number and (kinetic) energy, with volumetric densities \( n, e \). We can define the density of any such scalar quantity in the bin by

\[
q = \int d^3 p \phi_q f = \int_{p^-}^{p^+} 4\pi p^2 dp \phi_q \bar{f}_0,
\]

for \( q = (n, e) \) we have \( \phi_q = 1, \ T(p) \) (with \( T \equiv (p^2 c^2 + m_e^2 c^4)^{1/2} - m_e c^2 \) for rest mass \( m_e \)). So evolving \((\bar{f}_0), \alpha_{\bar{f}_0} \) is equivalent to evolving \((n, e)\). Returning to equation (1), multiplying by \( 4\pi p^2 dp \phi_q \) and integrating we immediately have:

\[
D_t q = -\nabla \cdot \mathbf{F}_q + \ldots
\]

which is a standard hyperbolic conservation equation that can be integrated to desired accuracy, provided an expression for \( \mathbf{F}_q \).

Conversely, since the DF in equation (3) has two parameters which vary in space and time: \((\bar{f}_0)_{b0} \) and \( \alpha_{\bar{f}_0} \), in order to update both in a time-step self-consistently in a manifestly conservative manner, we must update both \( (q, q') = (n, e) \), which requires computing both fluxes \((\mathbf{F}_q, \mathbf{F}_{q'})\). The updated \((n, e) \) in some next-time step then immediately give the new \((\bar{f}_0), \alpha_{\bar{f}_0} \). For details, see Girichidis et al. (2020).

In principle, any ‘basis function’ representation of \( f(p) \) in the bin with two free parameters (of which a power law is simply most convenient, given the real shape of the CR DF) should allow us to conserve two scalar quantities (CR number, energy) from evolving equation (5). If we also explicitly evolve the corresponding flux equations \( D_t \mathbf{F}_q \) (derived below), then we should also conserve both of their fluxes (i.e. the CR number and energy flux, which correspond to the CR current and momentum density fields).

### 2.3 The flux evolution equation

So, taking equation (2), multiplying by \( 4\pi p^2 dp \phi_q \) and integrating, we have for the flux equation:

\[
D_t \mathbf{F}_q + \mathbf{b} \cdot \nabla \cdot I_{\mathbf{v},q} = -I_{0,q} - I_{1,q}
\]

where we made use of various definitions above. Now define, for any quantity \( X \) which might vary as a function of \( p, X_0 \equiv X[p = p_0] \) (i.e. \( X_0 \) is the value of \( X \) at the bin centre). We can then immediately define the integral \( I \) terms in the following convenient form:

\[
I_{\mathbf{v},q} \equiv \bar{\omega} \bar{v} \mathbf{D}_0 q
\]

\[
I_{0,q} \equiv \bar{\omega}_0 \bar{v}_0 \chi \alpha_b \bar{v} \mathcal{A}_0 \bar{q}
\]

\[
I_{1,q} \equiv \bar{\omega}_1 \bar{v}_1 \mathcal{A}_1 \bar{q}
\]

which places the complicated integrals into the dimensionless functions \( \bar{\omega} \) (define by the above relations to \( I \)). This allows us to write the flux equation in familiar form:

\[
D_t \mathbf{F}_q + \mathbf{b} \cdot \nabla \cdot (\bar{\omega} \bar{v} \mathcal{F}_q) = -v_{\text{eff}} \bar{\omega} \bar{F}_q - v_{\text{eff}} \bar{v} \mathcal{F}_{\text{eff}} q
\]

with the modified ‘effective’ coefficients:

\[
\mathcal{D}_{\text{eff}} \equiv \bar{\omega} \bar{v} \mathcal{D}_0
\]

\[
v_{\text{eff}} \bar{\omega} \equiv \bar{v}_0 \chi \alpha_b \bar{v} \mathcal{A}_0 \bar{q}
\]

\[
v_{\text{eff}} \bar{v} \equiv \mathcal{A}_1 \bar{q}
\]

### 2.4 The bin-centred approximation

As discussed in Section 1, equation (14) has largely been evolved according to the ‘bin-centred’ approximation, which evaluates \( F_q \) as if we had an infinitesimally narrow bin centred at \( p = p_0 \), i.e. taking \( \omega_{\text{eff}} = \omega_{\text{eff}} = \omega_{\text{eff}} = 1 \). This has obvious advantages: (1) it is numerically straightforward: in fact the spatial (advection+flux) equations for a single CR ‘bin’ become numerically exactly identical to the ‘single-bin’ CR equations (wherein one integrates over the entire CR spectrum and simply evolves a ‘total CR energy’); (2) it is fairly trivially stable and robust (any integration method which can handle the two-moment equations for single-bin CRs, or radiation, or the one-moment diffusion + streaming equation, is trivially numerically stable and robust here); (3) it is simple; (4) it still retains manifest conservation: one still evolves both \( F_q \) and \( F_{\text{eff}} \) (so e.g. can manifestly conserve CR number and energy as desired),
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(18)

This can in principle be integrated numerically to arbitrary precision. But recalling that we have already parametrized the spectrum as a piecewise power law, it is useful to parametrize other quantities such as $v$ and $\phi_q$ as approximate power-law functions of $p$ over the domain of the bin, e.g. take

$$\alpha_q \equiv \frac{\partial \ln X}{\partial \ln p} \bigg|_{p=p_0} \approx \frac{\Delta \ln X}{\Delta \ln p} = \frac{\ln (X^+ / X^-)}{\ln (p^+ / p^-)},$$

where $X^\pm \equiv X[p = p^\pm]$. So e.g. $\alpha_q = 0$ exactly for $q = n$. For CRs with $p > m_1$, $c$, $\alpha_q \approx 0$ and $\alpha_q \approx 1$ for $q = \epsilon$ (for $p < m_1$, $c$, $\alpha_q \approx 1$ and $\alpha_q \approx 2$ for $q = \epsilon$), so these are close to exact power laws regardless, and so long as the spectral bins are small enough that there is no substantial spectral curvature within a bin (a necessary assumption for a piecewise-power-law treatment to be valid in the first place), approximating non-power-law behaviour with equation (19) introduces no significant errors beyond our original piecewise power-law approximation. We can then immediately write:

$$\omega_{\alpha,q} \approx \frac{(\alpha + \alpha_x) (\xi^{\alpha} + 2 \alpha_x \xi^{-\alpha_x})}{(\alpha + \alpha_x) (\xi^{\alpha} + 2 \alpha_x \xi^{-\alpha_x}) - (\xi^{\alpha} + \xi^{-\alpha}) - 1} \xi^{-
u_e}$$

$$\approx 1 + \frac{a_x}{2} (\alpha + \alpha_x + \alpha_q + \alpha_q) (\ln |\xi| + O(|\ln |\xi||^4))$$

(20)

with $\xi \equiv p^\pm / p^\epsilon = \exp (|\ln (p^+ / p^-)|)$ and the second expression above is a series expansion in $|\ln \xi|$. Note that with the definition in equation (19), $\xi^{\nu_e} = X^+ / X^- \propto \nu_e$ for any $X$, so we could equivalently write:

$$\xi^{\nu_{0} \alpha_{f_{0}}} \xi^{2 \alpha_{s} + 2 \alpha_{e}} = \left( \frac{p^{\pm}}{p^{\epsilon}} \right)^{3} \left( \frac{f_{0}}{p^{\epsilon}} \right) \left( \frac{\nu_{0} \psi_{q}}{\psi_{q}} \right) \left( \nu_{e} \right)^{2}$$

(21)

if the latter is more convenient.

With these assumptions, we can note,

$$\mathcal{I}_{\text{cr},q} \approx \chi_0 \alpha_{f_{0}} \nu_{0} \bar{v}_{A,0} \int_{p_{0}^{\epsilon}}^{p_{\epsilon}} 4\pi p^2 dp \bar{v} \phi_{q} f_{0},$$

and immediately follow a

$4$More specifically, for the various $\alpha$ terms that appear in this paper, $\bar{f}_{0}$ and $\langle \mu \rangle$ or $f_{1}$ within a bin are assumed to be exact power laws by construction, so $\alpha_{f_{0}}$ and $\alpha_{q}$ have exact values but these can (and will) vary across cells and in time. The scattering rate $\nu_{e}$ is often assumed to be an exact power-law constant in time, but does not have to be (it could have curvature and/or vary with local plasma properties). Of course $\alpha_{e} = 0$ identically for $q = n$, but for $q = \epsilon$, $\alpha_{e}$ is approximate (but it is fixed across all time and cells for a given bin). Likewise for $\alpha_{f_{0}}$. One could numerically evaluate all integrals presented here for the relevant $\omega$ terms exactly, without approximating terms such as $\epsilon$ as piecewise power laws; but in our numerical tests this provides no appreciable improvement in accuracy compared to using the simpler, analytical power-law approximations we provide.

$5$In this expression, we also take $\bar{v}_{A}$ outside the integral. This depends implicitly on $p$ through $\nu_{e}$, as $\bar{v}_{A} = \nu_{A} (\bar{v}_{A} - \bar{v}_{A}) / (\bar{v}_{A} - \bar{v}_{A})$. But like with $\chi$, this is almost always in one of two limits, either of which is $p$-independent. As shown in Hopkins et al. (2022c), if extrinsically turbulence strongly dominates CR scattering and is forward/backward symmetric in the Alfvén frame, then $\bar{v}_{A} \rightarrow 0$ is small and constant (and the term will be unimportant regardless). If otherwise (e.g. self-confinement dominates or the scattering is asymmetric) then $\bar{v}_{A} = \nu_{A}$ independent of $p$. So we can generally safely neglect the $p$-dependence of this within the bin here, especially as we later show that the steady-state behaviour of this ‘streaming’ term reduces to the bin-centred approximation.

2.5 Towards a better approximation

To do better, we must evaluate the correction terms $\omega$ for finite $\ln \xi$. By definition, most of the necessary inputs ($\phi_q$, $f_0$, $v$) and their dependence on $p$ are specified. However, the challenge is that all three $\omega$ terms depend on powers of $\mu$ (through $f_1$ or $\chi$, $\nu_0$). This introduces new variables whose dependence on $p$ (via $\mu$) is not a priori specified.

2.5.1 Terms which depend weakly on pitch angle

Let us begin with $\alpha_{\nu_e}$. This depends only on specified inputs as above and $\nu_0$, which depends on $\langle \mu^2 \rangle$ through $\chi$. But here we can make use of the limiting behaviours of $D$: for DFs which are near isotropic (hence $\langle \mu \rangle \rightarrow 0$ is small), $\chi \rightarrow 1/3 + O(\mu^2)$ so $D \rightarrow 1/3 + O(\mu^2)$, while for DFs which are near maximally anisotropic/coherently free-streaming from a source ($\langle \mu \rangle \rightarrow -1$), $\chi \rightarrow O[\langle \mu \rangle^2]$, so $D \rightarrow \tilde{D} \otimes \delta + O[\langle \mu \rangle^2]$. In either regime, the dependence on $\langle \mu \rangle$ is quite weak, so even if $\langle \mu \rangle$ varies across the bin, it will produce very little variation in $D$. So long as we do not see a very rapid transition from confinement to free-streaming across a single bin (which we do not expect), then it is almost always safe to neglect the term in $\chi$ and $D$ across any reasonable spectral bin size, i.e. take $\mathcal{I}_{\nu_e,q} \approx \mu \int_{p_{0}^{\epsilon}}^{p_{\epsilon}} 4\pi p^2 dp \nu^2 \phi_q f_0$. If we do this,
similar procedure to obtain $\omega_0$:

$$
\omega_{0,q} \approx \frac{\int_{p^2}^{p^*-p} dp \ p^2 \ \tilde{v} \ f_0 \ \Phi_q}{\int_{p^2}^{p^*-p} dp \ p^2 \ \tilde{v}_0 \ f_0 \ \Phi_q} \approx \frac{(3 + \alpha_{f_0} + \alpha_{q} + \alpha_{v})}{(3 + \alpha_{f_0} + \alpha_{q} + \alpha_{v})} \left( \frac{\xi^{3+\alpha_{f_0}+\alpha_{q}+\alpha_{v}-1}}{\xi^{3+\alpha_{f_0}+\alpha_{q}+\alpha_{v}}-1} \right) \approx 1 + \frac{\alpha_{v}}{12} \left( 3 + \alpha_{f_0} + \alpha_{q} + \alpha_{v}/2 \right) \ln|\xi|^2 + \mathcal{O}(\ln|\xi|^4). \quad (22)
$$

Note that $\alpha_{v}$ corresponds to $\tilde{v} \propto p^{\alpha_{v}}$; for the commonly adopted phenomenological assumption in modelling Galactic and Solar system CR observables that the diffusivity scales as $\propto R^0$ for CR rigidity $R$ (at energies where $\beta \approx 1$), we have $\alpha_{v} \approx -\delta$, so those observations imply $-0.7 \approx \alpha_{v} \approx -0.4$ (Blasi & Amato 2012; Vladimirov et al. 2012; Gaggero et al. 2015; Cummings et al. 2016; Guo, Tian & Jin 2016; Johansson et al. 2016; Korsmeier & Cuoco 2016; Evoli et al. 2017; Amato & Blasi 2018; De La Torre Luque et al. 2021; Hopkins et al. 2022b).

2.5.2 Terms which depend strongly on pitch angle

Now consider $\omega_{1,q}$. Here, we cannot neglect the implicit $\mu$-dependence, because the fluxes $F_q$ are directly proportional to $\langle \mu \rangle$. So make the ansatz, like above, that we can approximate $\langle \mu \rangle \propto p^{\alpha_{\mu}}$ over the (relatively narrow) width of the bin, giving:

$$
\omega_{1,q} = \frac{\int_{p^2}^{p^*-p} dp \ p^2 \ \tilde{v} \langle \mu \rangle \ v \ f_0 \ \Phi_q}{\int_{p^2}^{p^*-p} dp \ p^2 \ \tilde{v}_0 \langle \mu \rangle \ v \ f_0 \ \Phi_q} \approx \frac{(3 + \alpha_{f_0} + \alpha_{q} + \alpha_{v} + \alpha_{\mu})}{(3 + \alpha_{f_0} + \alpha_{q} + \alpha_{v} + \alpha_{\mu} + \alpha_{v})} \left( \frac{\xi^{3+\alpha_{f_0}+\alpha_{q}+\alpha_{v}}}{\xi^{3+\alpha_{f_0}+\alpha_{q}+\alpha_{v}+\alpha_{\mu}}-1} \right) \approx 1 + \frac{\alpha_{\mu}}{12} \left( 3 + \alpha_{f_0} + \alpha_{q} + \alpha_{v} + \alpha_{\mu} + \alpha_{v}/2 \right) \ln|\xi|^2 + \mathcal{O}(\ln|\xi|^4). \quad (23)
$$

Here, as in the expressions above and in various expressions below, the first (explicit integral) expression is exact. The second makes the power-law substitution, and is exact to the extent that the power-law approximation for the quantities inside the integrand is exact over the width of the bin.\(^7\) The third is a series approximation in $|\ln \xi|$, which is generally not necessary for our numerical evaluations in a code implementation of these methods, but is convenient here for intuition-building and understanding different limits discussed below.

Equation (23) would allow us to evolve equation (14), except now we have introduced a new parameter $\alpha_{v}$, which is not a priori specified. However, it is not actually the case that $\alpha_{v}$ is unconstrained. Since our update to the DF (equation 3) requires evolving both of a pair ($q, q'$) $\Rightarrow (n, e)$ with associated fluxes ($F_q, F_{q'}$), then by combining the definitions of ($q, q', F_q, F_{q'}$), one can show there is one independent consistency relation that must be satisfied:

$$
\Psi_{qq'} \equiv \left( \frac{F_q}{q \ v_0} \right) / \left( \frac{F_{q'}}{q' \ v_0} \right) = \frac{q}{q'} F_q / F_{q'} = \frac{\int_{p^2}^{p^*-p} dp \ p^2 \ \tilde{v} \ f_0 \ \Phi_q}{\int_{p^2}^{p^*-p} dp \ p^2 \ \tilde{v}_0 \ f_0 \ \Phi_q} \approx \frac{(3 + \alpha_{f_0} + \alpha_{q})}{(3 + \alpha_{f_0} + \alpha_{q} + \alpha_{v})} \left( \frac{\xi^{3+\alpha_{f_0}+\alpha_{q}}}{\xi^{3+\alpha_{f_0}+\alpha_{q}+\alpha_{v}}-1} \right) \approx \frac{1 + \alpha_{\mu}}{12} \left( 3 + \alpha_{f_0} + \alpha_{q} + \alpha_{v} + \alpha_{\mu} \right) \ln|\xi|^2 + \mathcal{O}(\ln|\xi|^4). \quad (24)
$$

Once again we give the exact integrals, solution making the power-law replacement, and series approximation in turn.

This is sufficient to specify $\alpha_{\mu}$ and therefore $\omega_{1,q}$, according to the different integration methods described below.

2.5.3 Solution methods

With expressions for $\omega$, equation (14) can be numerically integrated with exactly the same numerical methods as used for the ‘bin-centred’ method above — the $\omega$ terms only amount to a scalar renormalization of $\xi_0$, $v_0$, and $\nu_0$ which are arbitrary anyway from the point of view of the numerical method. The added complication comes almost entirely from determining $\alpha_{\mu}$ consistently to evaluate these terms. Consider three methods to do so:

(i) Exact: One option is to exactly update ($q, q', F_q, F_{q'}$) subject to the constraint $\Psi_{qq'}$ (equation 24). One can think of this as ‘replacing’ the value of $\alpha_{\mu}$ with that determined by $\Psi_{qq'}$ in the original equations for ($q, q', F_q, F_{q'}$). While do-able in principle, this (a) is extremely non-linear and involves inverting several complicated and numerically stiff functions of four variables; (b) couples the ($q, q', F_q, F_{q'}$) variables explicitly so we are forced to update all simultaneously with a single implicit step, i.e. we cannot operator-split as is usually desired; and (c) can sometimes lead to non-invertible expressions if great care is not taken with numerical errors.

(ii) Approximate, Integrated: Alternatively, if the numerical method explicitly integrates the variables $F_q$ and $F_{q'}$ (e.g. two-moment methods), then we can insert the values of ($q, q', F_q, F_{q'}$) at some point in the time-step (at the beginning of the step or ‘drifted’ to a half-step for a standard explicit method, or their exact values at step-end for implicit integration) into equation (24) and solve for $\alpha_{\mu}$ from that expression, then use this value of $\alpha_{\mu}$ in equation (14) to calculate the update to $F_q$ and $F_{q'}$. This is similar to how the other variables in equation (14) appear and is numerically straightforward [the single numerical inversion of equation 24 for a given $\Psi_{qq'}$ value is straightforward as well]. We find this works quite well.\(^8\)

\(^7\)Technically, we have to be careful about the case where the integrand with $dp$ scales exactly as $p^{-1}$, in which case the power-law expressions should evaluate to $\ln$ instead of those shown. But for any case where the index is not exactly negative one this is can be solved without issue and if constructing a numerical interpolation one can interpolette across this boundary without divergences.

\(^8\)Some numerical caution is still always needed. For example, if one adopts the power-law approximations given above, then one needs to treat the regime around certain values where some expressions would seemingly produce divergences carefully. Specifically, this arises when the integrand in the original exact expression takes values $\sim f \ p^{-1} dp$, so the power-law solutions should be replaced with logarithmic solutions: for example if $3 + \alpha_{f_0} + \alpha_v = 0$ in equation (24), which could numerically give a 0/0 error. For power-law indices close to these critical values we recommend either using the exact integral solutions (ideally), or a lookup table designed.
And in one-moment methods, we must determine the appropriate $F_{\mu}$, $F_{\nu}$ self-consistently and simultaneously, which we discuss below.

(iii) ‘Local-Steady-State’ Values: A still simpler, but even more numerically robust method is to not solve for $\alpha_{\mu}$ from the constraint equation (24) exactly, but to instead adopt the value it would have for the corresponding terms in equation (14) if the flux equation were in local steady-state. We derive this and further define below. This has the advantage that it is extremely robust and trivially numerically stable (provided whatever integration method used for the ‘bin centred’ approximation is also stable). It sacrifices manifest consistency between equations (24) and (14) for $q$, $q'$, but we are guaranteed that when the flux equations $D_i F_{\mu}$ are close to local steady-state (which is usually the case), the consistency relations are satisfied.

We also note that while it is generally advisable to use the full numerical expressions for $\alpha_{\mu}$, the series expressions we show (expansions in $|\ln \xi|$) work surprisingly well for even large $\xi$, valid to better than $\sim 10$ per cent for all $\omega$ terms for any $\xi \lesssim 3$, and for some of the terms (especially in the ultra-relativistic limit) the series expression works well up to $\xi \lesssim 100$ (assuming the underlying terms could, in fact, be approximated as power laws reliably over that dynamic range).

2.6 Local flux steady-state behaviours

Consider the case where the flux equations (equation 2) reach approximate local-steady-state, i.e. $|D_i F_{\mu}| \rightarrow 0$ (or $|D_i F_{\mu}| \ll |v F_{\mu}|$). This occurs on approximately the scattering time $\sim v^{-1}$, which is very short in the Galactic ISM (from observations, $v^{-1} \sim 30$ yr for $\sim 1$ GeV CRs; see Hopkins et al. 2022b). Thus even if we explicitly evolve $F_{\mu}$, we expect it to be close to this ‘local flux steady-state’ value in many regimes. Moreover, the ‘one-moment’ numerical methods assume this is exactly true, to directly solve for $F_{\mu}$ and insert it into equation (1) to directly obtain a diffusion-streaming equation for the CRs (see e.g. Zweibel 2013, and references therein). Noting that this implies the scattering-limiter, so the CRs are nearly isotropic ($\chi \rightarrow 1/3$, $D \rightarrow 1/3$), we immediately obtain from equation (14):

$$F_{\mu} \rightarrow v_{\text{eff}}(\xi, q) = \frac{v_{\text{eff}}}{3 \xi^2} \nabla \cdot (\omega_{\nu, q} \mathbf{q}),$$

(25)

where $V_{\parallel} \equiv \mathbf{b} \cdot \nabla$. So up to the ‘effective’ coefficients being slightly modified by the $\omega$ terms, this is just the usual streaming/diffusion expression, with streaming speed $v_{\text{eff}}$, and effective anisotropic diffusivity $\kappa = \kappa_{1} \sim \frac{\xi^2}{3} v_{\text{eff}}$ (if we assume isotropically tangled magnetic fields on small scales, this can be further approximated as an isotropic diffusivity $D_0 \sim \kappa_{1}(3)$).

2.6.1 The ‘Alfvénic streaming-dominated’ limit

Consider the case where the Alfvénic streaming term dominates in equation (25), $F_{\nu} \rightarrow v_{\text{eff}}(\xi, q)$ (this can occur in e.g. self-confinement models when $\mathbf{b} \rightarrow \infty$). Then equation (24) becomes

$$\Psi_{q'} = (\omega_{\nu, q}, \omega_{\mu, q})/(\omega_{\nu, q}, \omega_{1}, q).$$

This is solved exactly if and only if $\alpha_{\mu} \rightarrow -\alpha_{\nu}$, i.e. the CR drift velocity $v_{\text{drift}} = (\mu) v \propto p^0$ is independent of momentum (as it must be, since they are drifting, to be interpolated over the relevant range, rather than taking the power-law expressions directly at face value).

by definition in this limit, at the momentum-independent streaming speed across the bin). Inserting this into the expressions for $v_{\text{eff}}$, we immediately have:

$$\frac{\alpha_{\text{eff}}}{\alpha_{\nu, q}} \rightarrow 1, \quad v_{\text{eff}}(\xi, q) = -\frac{3}{3} \nabla \cdot \mathbf{q}_{\nu, q},$$

(26)

In this limit, because the drift velocity is constant (across the bin), and the gradient/term and diffusive terms are irrelevant, we see that we have recovered exactly the same $F_{\mu}$ that we would have in the bin-centred approximation.

2.6.2 The diffusive or super-Alfvénic limit

Now consider the case where the ‘diffusive’ term dominates in equation (25), so $F_{\nu} \rightarrow (v_{\text{eff}}/3 v_{\text{eff}}^2) \nabla (\omega_{\nu, q} q)$. Note that when some literature refers to ‘super-Alfvénic streaming,’ this still comes from this particular term (and there is no distinction, for our purposes here). The constraint equation then becomes $\Psi_{q'} = (\omega_{1, q}, \omega_{q})/(\omega_{1, q}, \omega_{q})$, where

$$\xi_{\Omega} \equiv \frac{q}{v_{\nu, \Omega, q}} q,$$

(27)

Solving for $\alpha_{\mu}$ from this constraint gives a highly non-linear equation to be solved for $\alpha_{\mu} \rightarrow \alpha_{\mu}/(\xi_{\Omega}, \omega_{\nu, q}, \alpha_{\mu}, \omega_{\nu, q}, \alpha_{\mu}, \alpha_{\nu}, \omega_{\nu, q})$. It is more instructive to parameterize $\xi_{\Omega}$ in a similar piecewise-power-law manner: let us define $\xi_{\Omega} \equiv \xi_{\Omega}(\xi_{\nu, q})$, where $\xi_{\nu, q} = \xi_{\nu} p^0$ is defined over an infinitesimally small range of $p$, and let us assume this scale similarly as $\xi_{\nu} \propto p^{\mu}$.10 If we combine this with the steadystate expressions for $F_{\nu}$ in terms of the relevant gradients, and use equations (28)–(29), we see that the constraint relations are satisfied exactly for $\alpha_{\mu} \rightarrow -\nu_{\nu} + \alpha_{\nu} - \alpha_{\nu}$, which we can immediately insert in equation (23).

With these definitions and some similar algebra, it is also conven-

tient to note that we can write:

$$F_{\mu} \rightarrow \nabla \cdot \mathbf{q} = -\left(\frac{\omega_{\nu, q}}{3 \xi^2} \right) \nabla \cdot \mathbf{q}$$

(30)

$$\omega_{\nu, q} \approx \frac{3 + \alpha_{\nu} + \alpha_{\nu}}{3 + \alpha_{\nu} + \alpha_{\nu} - \alpha_{\nu} + 2 \alpha_{\nu} - \alpha_{\nu}}$$

(31)

which is the effective diffusivity is simply modified by a correction factor $\omega_{\nu, q}^\mu$. For $\approx \text{GeV}$ CRs, where we have empirically typical $\alpha_{\nu} \approx -4.7$ (from direct observation; e.g. Cummings et al. 2016), $\alpha_{\nu}^\mu$.

10If we take $\alpha_{\mu} \rightarrow -\nu_{\nu} + \alpha_{\nu} - \alpha_{\nu}$ and, for compactness, write $\alpha_{\nu, q} \equiv 3 + \alpha_{\nu} + \alpha_{\nu}$, we have:

$$\xi_{\nu}^\mu \approx \frac{\alpha_{\nu, q} q}{\alpha_{\nu, q} q + \Delta \alpha_{\nu}} \left(\frac{\alpha_{\nu} q + \Delta \alpha_{\nu} q}{\alpha_{\nu} q + \Delta \alpha_{\nu} q - 1} \right) \left(\frac{\xi_{\nu} q + \Delta \alpha_{\nu} q}{\xi_{\nu} q + \Delta \alpha_{\nu} q - 1} \right)^{-1},$$

(28)

with which can then be solved for $\Delta \alpha_{\nu}$.
$\sim -0.6$ (from modelling of primary-to-secondary ratios and similar constraints; De La Torre Luque et al. 2021; Hopkins et al. 2022b; Korsmeier & Cuoco 2021), $\alpha_{e} \sim 0$ (from the fact that these are ultra-relativistic), $\alpha_{e} \lesssim 0.1$ (from modelling spatially resolved Galactic $\gamma$-ray profiles at different energies; e.g. Tibaldo et al. 2015; Accro et al. 2016; Yang, Aharonian & Evoli 2016; Hopkins et al. 2022b), we obtain $\alpha_{\nu,q} = -1 - 0.05 \left(1.4 - \alpha_{q} + \alpha_{l}\right)|\ln \xi|^{2} + O\left(\ln \xi|^{4}\right) \sim -0.08, 0.03\right)|\ln \xi|^{2}$ for $q = (n, e)$. The 'mean' correction (both are $<0$ because for these energies, most of the CR number and energy is biased towards the lower-$p$ end of the 'bin,' where the effective $\kappa$ is smaller) is modest and not so important, given the CR and energy is biased towards the lower-$p$ end of the 'bin,' where the effective $\kappa$ is smaller. We can therefore use the right correction: the CR number flux is somewhat modified (because CR number is more strongly dominated by the low-$p$ end of the bin), and the (small) difference here causes the spectral slope to steepen within the bin as CRs diffuse. 

Note that if we must still evaluate $\xi_{q}/\xi_{q}$ to determine $\alpha_{l}$ for equation (30) above, then it is not necessarily more computationally useful than just using $F_{q} \to \left(v_{q}^{3}/v_{q}^{3}\right) \bar{\nabla} \left(\alpha_{\nu,q} q\right)$ as we would have previously, but it is still useful to guide our intuition. Moreover, we can note that in the limit where the diffusive term dominates the flux, with negligible losses, and the CR (n, e) equations are themselves close to steady-state (assuming also $\nu$ and the source injection spectrum do not vary strongly with spatial location), then $\alpha_{l} \to 0$. Since that is precisely the regime where it matters most to get this correction 'right,' we can assume this without much loss of accuracy given our other significant simplifications above.

2.6.3 The 'local-steady-state' approximation for flux corrections

With all this in mind, if one adopts a two-moment method (evolving $F_{q}$ explicitly) with the primary goal of capturing the exact behaviour in the three possible limits of equation (2) (free-streaming/weak-scattering, or near-isotropic/strong-scattering/diffusive, or trapped/advecive/Alfvénic-streaming), then it is sufficient to adopt the 'local-steady-state' approximation for $\alpha_{\mu}$ in equation (14) using the appropriate value of $\alpha_{\mu}$ each would have if it were dominant. This gives

$$D_{q}F_{q} + v_{q}^{3} \bar{b} \cdot \bar{\nabla} \left(\alpha_{\nu,q} q\right) = -\nu^{*} \left(F_{q} - v_{\nu,0} q\right),$$

$$v_{\nu,0} \equiv -\nu \alpha_{\nu} \bar{v}_{A,0},$$

$$v^{*} \equiv v_{0} \alpha_{\nu,q} \left(\alpha_{e} - \alpha_{v} - \alpha_{l}\right)$$

with $\alpha_{\nu,q}$ from equation (20) and $\alpha_{\nu,q}$ from equation (23). One can immediately verify this reduces correctly to any of the relevant local-steady-state limits above.

If one evolves a 'one-moment' method – e.g. evolving the CRs according to a single streaming + diffusion or Fokker–Planck type approximation (valid only in the strong-scattering limits), then we can approximate the limits of interest via

$$F_{q} \to -\frac{\nu_{q}}{3} \bar{v}_{A,0} q - \left(\alpha_{\nu,q} \frac{v_{0}^{2}}{3 \bar{v}_{0}}\right) \nabla_{q}$$

(with $\alpha_{\nu,q}$ from equation (31)) where $\alpha_{l}$ in $\alpha_{\nu,q}$ can be computed or (for even greater simplicity), approximated as $\approx 0$ without severe loss of accuracy.

3 SIMPLE NUMERICAL TESTS

In Fig. 1, we consider a simple illustrative numerical test of the proposed methods. To isolate the interesting behaviour and construct a simple, analytically tractable test problem, we consider transport of a power-law injection spectrum in a plane-parallel atmosphere, analogous to classic thin disc or leaky-box type models for CRs. Specifically, consider an infinitely thin source plane in the $xy$ axis, in a homogeneous, stationary background (e.g. $\nu_{\ast} = 0$, $\bar{b} = \bar{z} = \text{constant}$) with space-time-independent $\bar{\varepsilon}_{A} = \text{constant}$ and $\nu \propto p^{-0.5}$. To $\propto p$ and $\beta \approx 1$ (e.g. the ultra-relativistic limit, though this choice has no effect on our conclusions), ignoring all non-spatial transport terms (e.g. catastrophic or radiative losses) except for injection in the source plane at a constant rate per unit area $J_{\nu} = dN_{\nu}/dt \, dA \, d^{2}p$. Numerically, we integrate this on a domain with 10 spatial cells in the vertical direction from $z = 0$ (with an inflow/injection boundary) to $z = 1$ (with an outflow boundary) in arbitrary code units, and injection slope $j_{\nu,0} \propto p^{-2}$ similar to expected physical values, using the finite-volume two-moment method (evolving $n, \nu, F_{\nu}$, $F_{\nu}$, $F_{\nu}$) in the code GIZMO (Hopkins 2015; Hopkins & Raives 2016; Hopkins 2017; Hopkins et al. 2022b), with the $\omega$ values determined according to the different proposed methods described in the text. We discretize the momentum domain with 10 bins over 2 dex (though again, given the simplifications of our problem, the dynamic range of $p$ is not important to our conclusions). We set the normalization of $\nu_{A}$ and $\nu$ to two different values to compare two limits.

First, we consider a 'streaming-dominated' limit, obtained by setting $\nu$ to a very large value ($\sim 10^{8} \, p^{-0.5}$ in code units) with $\nu_{A} = 1$ (and effective diffusion coefficient $v^{2}/3 \nu$ set to an arbitrarily small value), so analytically $\bar{f}_{1} \to \left(D_{\nu} / D_{\nu}\right) \partial_{p} \bar{f}_{0}$. This has a simple constant-flux steady-state solution with $f_{1} = -\nu \alpha_{\nu} \bar{\varepsilon}_{A}$, so $f_{0} \to J_{\nu} / \left|\alpha_{\nu} \bar{\varepsilon}_{A}\right|$ is spatially uniform and proportional to the injection spectrum (i.e. $f_{0} = \bar{\varepsilon}_{A} = -4.2$). As predicted in Section 2.6.1, the injection spectrum is simply advected here, so all methods (including the simple bin-centred approximation) reproduce the exact solution in Fig. 1 in this limit.

Second, we consider a 'diffusion-dominated' case, setting $\nu_{A} = 0$ with finite $\nu$ (evolved to several times the effective diffusion time). In steady-state now $\nu \bar{f}_{1} \to (v^{2}/3 \nu) \bar{\partial}_{p} \bar{f}_{0} = J_{\nu}$, so $\bar{f}_{0} \to \left(3 \bar{\nu} / v^{2}\right) J_{\nu} = \text{constant in space and } \nu \propto p^{-0.5}$. Because higher energy CRs have a lower $\bar{\nu}$, and correspondingly larger effective diffusivity $v^{2}/3 \nu$, they escape faster and their steady-state abundance (relative to injection) is reduced, steepening the spectrum by one power of $\nu$. All the numerical methods in Fig. 1 capture this effect 'on average' across bins. But for the 'bin-centred' method, as anticipated in Section 2.4, we effectively ignore the variation of $\nu$ within each bin (taking the bin-centred $\nu_{0}$ as constant across each bin). This means we very slightly overestimate the total value of $\nu$ (leading to a small underestimate of the mean $\nu_{0}$, averaged over the bin), but more importantly the method conserves the spectral slope within

\[\text{Even relatively sophisticated closure schemes for evolving } (\mu^{2}) \text{ proposed in the literature focus primarily on the behaviour in these three limits, as opposed to intermediate cases; see Hopkins et al. (2022a) for a review.}

\[\text{We have also tested these problems implementing the 10-element discretization in 1D, solved via a Crank–Nicholson scheme in PYTHON using either the two-moment equations or (since we consider the steady-state solutions) directly integrating the single-moment streaming + diffusion equation in Section 2.6, which gives indistinguishable results to those shown in Fig. 1.}\]
each bin, producing the ‘step’ structures seen. On the other hand, introducing the scalar $\omega$ correction terms as proposed in this paper, with either method in Fig. 1, leads to excellent agreement with the exact solutions (with the slope in each bin numerically agreeing with the exact solution to better than ~1 percent).

4 APPLICATIONS TO RADIATION/NEUTRINO DYNAMICS

It is natural to ask whether the methodology above can be cross-applied to radiation or neutrino transport, where one can easily imagine situations in which a similar piecewise-power-law reconstruction of the radiation spectrum would be useful.

For the sake of consistency with the large radiation/neutrino transport literature, in this section we will consider a different set of variable definitions matching the convention in those fields. Let $\nu$ refer to the radiation frequency (so $h\nu$ is energy, analogous to $p$ for CRs), so the specific intensity $I_\nu(n, \nu, \mathbf{x}, t)$ is equivalent to the DF $f$ in terms of the radiation direction unit vector $\mathbf{n}$, the mean/iso-intensity $J_\nu = (4\pi)^{-1} \int d\Omega$ is analogous to $j_0$, $\Delta \nu$ is analogous to $f_0$, $\Omega = (4\pi J_\nu)^{-1} \int d\Omega \mathbf{n} \otimes (\mathbf{n} I_\nu)$ is the Eddington tensor, $c, \kappa, \rho$ in terms of the opacity $\kappa_\nu$ and gas density $\rho$ is akin to the CR scattering rate $\nu$, $q_\nu = dq/d\nu \approx 4\pi \phi_\nu J_\nu$ is defined such that for photon number and energy $q = (n_e, e_\nu)$ we have corresponding $\phi_\nu = (1/h\nu, 1)$, and $F_\nu \equiv \phi_\nu c \int d\Omega \mathbf{n} I_\nu = c (n_e) q_\nu$. $(\nabla \nu)_i$ with $\nu \equiv (4\pi J_\nu)^{-1} \int d\Omega \mathbf{n} I_\nu$ is the flux term. With these definitions, the spatial part of the first two moments of the non-relativistic radiation-MHD moments equations, as usually written in the lab frame, are (Mihalas & Mihalas 1984)

$$\frac{\partial \nu_\nu}{\partial t} = -\nabla \cdot F_\nu + \ldots$$

$$\frac{\partial F_{\nu}}{\partial t} + c^2 \nabla \cdot (\nabla \nu)_i q_\nu = -c \kappa_\nu \rho \left[ F_{\nu} - q_\nu v_{\nu}\cdot(\mathbf{I} + \nabla \nu) \right] + \ldots$$

Note that the equations in the co-moving frame (to leading order in $\mathcal{O}(v_{\nu}\rho/c)$) are equivalent to taking $\partial_t \rightarrow D_t$ and dropping the $v_{\nu}\rho$ term above, so our discussion here applies equally to both cases. Equation (34) is again just advection, and integrating over a frequency interval from $\nu^{-}$ to $\nu^{+}$, we immediately have $\frac{\partial q_\nu}{\partial t} = -\nabla \cdot F_\nu (with \sim \int_{\nu^{-}}^{\nu^{+}} d\nu q_\nu$, $F_\nu \equiv \int_{\nu^{-}}^{\nu^{+}} d\nu F_\nu)$, so we only need to consider equation (35).

4.1 The strong-scattering and flux-limited diffusion-like limit

In the strong-scattering ‘local-steady-state’ limit for the flux, we have the usual diffusive approximation with $\nabla \nu \rightarrow 1/3$, $F_\nu \rightarrow (4/3) v_{\nu}\rho q_\nu - (c/3 \kappa_\nu \rho) \nabla q_\nu$. Integrating this, we immediately obtain:

$$F_{\nu} \rightarrow \frac{4}{3} v_{\nu}\rho q_\nu - \omega_{\nu, q} \frac{c}{3 \kappa_\nu \rho} (\nabla q)_i$$

$$\omega_{\nu, q} \equiv \frac{\int_{\nu^{-}}^{\nu^{+}} d\nu \kappa_\nu^{-1} \phi_\nu (\nabla J_\nu)_i}{\int_{\nu^{-}}^{\nu^{+}} d\nu \kappa_\nu^{-1} \phi_\nu (\nabla J_\nu)_i} \approx \frac{(1 + \alpha_j + \alpha_q - \alpha_{\ell_i})}{(1 + \alpha_j + \alpha_q - \alpha_{\ell_i} - \alpha_{\ell_i})} \left( \frac{\xi^{1+\alpha_j+\alpha_q-\alpha_{\ell_i}} - 1}{\xi^{1+\alpha_j+\alpha_q-\alpha_{\ell_i} - 1} - 1} \right) \xi^{\alpha_{\ell_i}/2} \approx 1 - \frac{\alpha_q}{12} \left( 1 + \alpha_j + \alpha_q - \alpha_{\ell_i} - \alpha_{\ell_i} - \alpha_{\ell_i}/2 \right) ||\ln \xi||^2 + \mathcal{O} (||\ln \xi||^3),$$

where $J_\nu \propto v_{\nu}\phi_\nu \propto v_{\nu}\kappa_\nu \propto v^{\alpha_q}$, and $(\nabla J_\nu)_i = J_{\nu, i}/\ell_i$ with $\ell_i \propto v^{\alpha_{\ell_i}}$ for each gradient component.

---

**Figure 1.** Numerical tests (Section 3) of our proposed correction terms for spatial transport of piecewise-power-law spectra. We consider a homogeneous, 1D stratified atmosphere with a continuous injection spectrum $f_0 \propto p^{-4.2}$ at the lower boundary and outflow from the upper boundary, constant streaming speed $v_A$ and scattering rate $\gamma \propto p^{-0.5}$, discretized into 10 momentum intervals, and evolved until steady-state using the numerical methods described in the text. We compare exact analytical steady-state solutions to numerical solutions using either (1) the ‘bin-centred’ approximation ($\omega_{\nu, q} = \omega_{\ell, q} = \omega_{\ell, q} = 0$; Section 2.4), (2) the ‘approximate-integrated’ method (Section 2.5.3) to solve for the $\omega$ terms (given the $c$-in indices evolved values of $n, \epsilon, F_n, F_\ell$ to calculate $\omega_{\ell, q}$, from equation (24)), and (3) the ‘local-steady-state’ approximation for $\omega$ terms (Section 2.6.3), using the local-steady-state values of $\omega_{\ell, q}$. Top: Parameters chosen so the transport is streaming-dominated ($\gamma$ very large). We plot the steady-state spectrum compensated by $p^{3.2}$ and in units such that the exact solution equals unity. In the streaming-dominated limit, the transport speed is momentum-independent so the spectrum is simply advected without change in spectral slope, and the different approximations behave identically. Bottom: Parameters chosen so the transport is diffusion-dominated ($\gamma = 0$). The ‘bin-centred’ approximation introduces well-known step artefacts, as a result of assuming the scattering rate and $\mu$ are constant within each bin, which conserves the injection slope within each momentum bin. Both our proposed methods for including the $\omega$ terms produce the correct spectral slopes within bins.
We can in principle solve for each value of $\alpha_{ij}$ as in Section 2.6.2 above\textsuperscript{13}, but if we assume that either the dependence of gradient scale length on wavelength in the bin is small ($\alpha_{i,j} \sim 0$) or just that the gradient direction does not strongly depend on wavelength across the bin ($\alpha_{i,j} \approx \alpha_{i,j} \approx \alpha_{i,j}$), we can write this in terms of a scalar ‘effective’ $\kappa$.

\[
F_q = \frac{4}{3} \frac{\rho}{\rho \kappa} \nabla q = \frac{3}{3} \gamma_{\text{eq}} q - \frac{c}{3 \kappa_{\text{eq}} \rho} \nabla q
\]

(39)

(i.e. just equation 37 with $\alpha_{i,j} \rightarrow \alpha_{ij}$).

Now if we assume $J_q$ is blackbody-like, so $|\nabla J_q| \rightarrow |dJ_q/dT|/|\nabla T|$, and consider the equation for the radiation energy density $q = p_\text{c}$ (so $\rho_\text{eq} = 1$), equation (40) becomes immediately recognizable as the usual Rosseland mean opacity (the $|\nabla T|$ term factors out, being independent of $\nu$). So essentially, we have just generalized this convention for (1) an arbitrary non-blackbody intensity, and (2) other conserved radiation quantities such as $n_p$ ($\phi_0 = 1/\hbar \nu$, $\alpha_{eq} = 1$), needed if we wish to correctly evolve the radiation spectrum as a piecewise power law with two degrees of freedom.

4.2 The weak-scattering and M1-like limits

Now consider cases where one wishes to evolve the flux equation (35) explicitly, in e.g. first-moment (M1) or variable Eddington tensor or other related moments-based methods. Integrating, in component form, we can write:

\[
\frac{1}{c^2} \frac{\partial F_{q_i}^l}{\partial t} = \left[ \nabla \cdot \left( \int_{\nu_\text{c}}^{\nu_\text{v}} d\nu \kappa_{\nu_\text{c}} q / |\nabla J_q| \right) \right] I
\]

\[- \left[ \int_{\nu_\text{c}}^{\nu_\text{v}} d\nu \kappa_{\nu_\text{c}} \rho \phi_{\nu_\text{c}} J_q \left\langle (\mathbf{n})_\nu - \frac{\mathbf{v}_{\text{eq}}}{c} \cdot \hat{\mathbf{b}} \right. \right] I
\]

\[= -\alpha_{i,j} \rho_\text{eq} \left[ |\nabla q_0| \left( \int_{\nu_\text{c}}^{\nu_\text{v}} d\nu \kappa_{\nu_\text{c}} F_{q_i}^l - \alpha_{i,j} \rho_\text{eq} q_0 \mathbf{v}_{\text{eq}} \cdot \hat{\mathbf{b}} \right) \right].
\]

(41)

If we assume $(\mathbf{n})_\nu \propto \nu^{b_{\nu,\text{eq}}}$ (analogous to $\alpha_\mu$ for CRs), then we can write:

\[
\alpha_{i,j} \rho_\text{eq} \frac{\int_{\nu_\text{c}}^{\nu_\text{v}} d\nu \kappa_{\nu_\text{c}} J_q \mathbf{F}_{q_i}^l}{\int_{\nu_\text{c}}^{\nu_\text{v}} d\nu \kappa_{\nu_\text{c}} F_{q_i}^l}
\]

\[\approx \left( 1 + \alpha_j + \alpha_j + \alpha_{n,i} \right) \left( \xi_{1+\alpha_j+\alpha_j+\alpha_{n,i}+\alpha_{n,i}} - 1 \right) \xi^{-\alpha_{n,i}/2}
\]

\[\approx 1 + \frac{\alpha_{n,i}}{12} \left( 1 + \alpha_j + \alpha_j + \alpha_{n,i} + \alpha_{n,i}/2 \right) \ln |\xi|^2 + O(\ln |\xi|^3)
\]

(42)

and we have an analogous consistency relation which determines $\alpha_{n,j}$ for each component of $\mathbf{F}_{q_i}$:

\[
\Psi_{q_i}^{l} \equiv \left( \frac{F_{q_i}^l}{q \nu_0} \right) / \left( \frac{F_{q_i}^l}{q v_0} \right) = \frac{q^{l} F_{q_i}^l}{q F_{q_i}^l}
\]

\[\approx \left( \int_{\nu_\text{c}}^{\nu_\text{v}} d\nu (\mathbf{n})_\nu \phi_\nu J_q \right) \left( \int_{\nu_\text{c}}^{\nu_\text{v}} d\nu \phi_\nu J_q \right)
\]

\[\int_{\nu_\text{c}}^{\nu_\text{v}} d\nu \phi_\nu J_q
\]

\[\approx \left( 1 + \alpha_j + \alpha_j \right) \left( 3 + \alpha_j + \alpha_j + \alpha_{n,i} \right) \left( 3 + \alpha_j + \alpha_j + \alpha_{n,i} \right) \left( 3 + \alpha_j + \alpha_j + \alpha_{n,i} \right)
\]

\[\times \left( 3 \xi_{1+\alpha_j+\alpha_j+\alpha_{n,i}} - 1 \right) \left( 3 \xi_{1+\alpha_j+\alpha_j+\alpha_{n,i}} - 1 \right)
\]

\[\approx 1 + \frac{\alpha_{n,i}}{12} \left( \xi - \alpha_{n,i} \right) \ln |\xi|^2 + O(\ln |\xi|^3).
\]

(43)

If the fluxes $F_q$ are explicitly evolved, we can then use $\Psi_{q_i}^{l}$ to determine $\alpha_{n,i}$ and thus $\alpha_{i,j}^{l}$, just as in our ‘exact’ and ‘integrated, approximate’ methods from Section 2.5.3 above. If instead we wish to replace $\alpha_{i,j}^{l}$ with its ‘local flux steady-state’ value we see from Section 4.1 we would have $\alpha_{n,i} \rightarrow -\alpha_{i,j} + \alpha_{k}$ in $\alpha_{i,j}^{l}$.

The real challenge arises with the treatment of the Eddington tensor ($\mathbf{D}_\nu$) terms in $\alpha_{i,j}^{l}$ and $\alpha_{i,j}^{l} q_0$. For CRs, it is worth emphasizing that the relation we wrote in Section 2.1, $\mathbf{D} = \chi (1 + (1 - \chi) \mathbf{b} \otimes \mathbf{b}$ is not some approximate closure: it is the most general possible form of $\mathbf{D}$ for a gyrotropic DF, and depends on a single scalar degree of freedom $\langle \mu^2 \rangle$ (and likewise, its parallel gradient $\mathbf{b} \cdot \nabla \mathbf{D}$ introduces only a single scalar degree of freedom). Moreover, gyrotropy means that even for an arbitrarily anisotropic CR DF, $\mathbf{D} \propto \mathbf{b} \otimes \mathbf{b}$ depends on the magnetic field direction $\mathbf{b}$, which is of course CR-momentum-independent. On the other hand, for radiation $\mathbf{D}$ has, in general, five independent degrees of freedom, and the $\mathbf{D}_\nu$ term introduces ~10 more.\textsuperscript{14} So the problem is rather severely underconstrained. Moreover, even in the simplest possible highly anisotropic case, where the radiation at a given $\nu$ is perfectly coherent (free-streaming in a single direction), we have $\mathbf{D}_\nu \sim \langle \mathbf{n}_\nu \rangle \otimes \langle \mathbf{n}_\nu \rangle$. But this (unlike $\mathbf{b} \otimes \mathbf{b}$) depends on an evolved property of the radiation flux itself ($\langle \mathbf{n}_\nu \rangle$), so it can depend on $\nu$, which means that we have no formal justification to neglect the variation in $\mathbf{D}_\nu$ across the bin.

This is not a new problem: defining a robust ‘closure’ for $\mathbf{D}_\nu$ is arguably the central challenge for moments-based radiation or neutrino-hydrodynamics schemes (see e.g. Wilson et al. 1975; Levermore 1984; Gnedin & Abel 2001; Rosdahl & Teyssier 2015; Murchikova, Abdikamalov & Urbatsch 2017; Foucart 2018). And many of the most popular numerical methods use highly approximate closures that only approach the exact solutions in very specific regimes (e.g. when $\mathbf{b}$ is nearly isotropic, or the radiation is a perfectly coherent 1D beam, etc.). So it is not clear if, in practice, we could solve for the correct $\alpha_{i,j}^{l}$ and $\alpha_{i,j}^{l} q_0$ even if we specified exactly some simple functional closure relation for $\mathbf{D}_\nu$. Thus, lacking another way to make progress, we will briefly consider – without justification, we stress – what we would obtain if we neglect the variations in $\mathbf{D}_\nu$ across each bin.

\textsuperscript{13} Specifically computing $\ell_q^l$, $\ell_q^l$ and using

\[
\ell_q^l \approx \alpha_{i,j} \left( \xi_{1+\alpha_j+\alpha_j} - 1 \right) \xi^{-\alpha_j/2}
\]

\[
\alpha_{i,j} \left( \xi_{1+\alpha_j+\alpha_j} - 1 \right) \xi^{-\alpha_j/2}
\]

\[\approx 1 + \frac{\alpha_{n,i}}{12} \left( \xi - \alpha_{n,i} \right) \ln |\xi|^2 + O(\ln |\xi|^3)
\]

(38)

\textsuperscript{14} These come from the dependence of $I_\nu$ on $\hat{\mathbf{n}}$ (ray direction), and its (arbitrary) gradient. Since $\mathbf{D}$ is a symmetric 3x3 matrix (in 3D) normalized to have trace unity (as it is defined by the moments of $I_\nu$) it has 5 degrees of freedom, and we have a similar number of degrees of freedom for each component of the vector gradient of $I_\nu$ which appears in $\mathbf{D}_\nu \cdot \langle \mathbf{n}_\nu \rangle$.
For the $\alpha_{\ell,0,q}$ term, if we neglect variation in $D_\ell$ across the bin we can calculate it directly as $\alpha_{\ell,0,q} \approx (1 + \alpha_0 - \alpha_\ell)/(1 + \alpha_0 - \alpha_q)$, for the simple reason that for a non-relativistic $v_{\text{gas}}$ (the valid limit of our expressions), the ‘advection’ term in $v_{\text{gas}}$ is only ever important in the strong-scattering, tightly coupled regime, where we can (quite accurately) assume the ‘local-steady-state’ approximation from above and, just like with CRs, this term reduces exactly to its ‘bin-centred’ version, with $\alpha_{\ell,0,q}/\alpha_{\ell,1,q} \to 1$.

The $V \cdot (\nabla, q_\ell)$ term becomes trivial with $\alpha_{\ell,0,q,v,q} \to 1$ if we neglect variations in $D_\ell$ across the bin. But we caution that while simple, this is much less ‘safe’ an assumption than neglecting the variations for $\alpha_{\ell,0,q}$. That is because this term is the dominant term controlling $\partial_t F_\ell$ in the weak-scattering regime, which is precisely where we said earlier it is not always safe to neglect variations in $D_\ell$ with $v$. For many moments-based methods, this regime is also where $D_\ell$ is estimated rather poorly. So this may not be a significant source of error relative to those pre-existing errors for methods like M1, but that remains to be tested.

5 CONCLUSIONS

We derive and test a simple improvement to numerical methods that dynamically evolve the CR spectrum, representing it as a piecewise power-law across momentum-space with standard advection/diffusion behaviour in coordinate-space. Previous attempts to do so generally allow for smooth and exact evolution of the piecewise-power-law slopes under momentum-space operations (e.g. continuous and catastrophic losses, injection, etc.), but for the spatial terms adopted the ‘bin-centred’ approximation that leads to errors in the local spectral shape when CR diffusion is important (or these methods sacrificed conservation or consistency with the underlying flux equations). We show that these errors are formally second order in momentum-space, but they can be eliminated, allowing for smooth evolution of the CR spectra under diffusion, maintaining consistency with the underlying Vlasov equations and manifest conservation of CR number and energy (and current and momentum, in two-moment methods).

The modification amounts to a set of three simple, scalar correction factors which, once computed, can be immediately applied (as e.g. a correction to the ‘effective’ bin-centred diffusion coefficient, or to the scalar quantities whose gradients are calculated), which can be computed exactly entirely as a function of actual evolved quantities in-code (i.e. there is no need to invoke new assumptions, or to implicitly evolve or take gradients of a ‘finer grained’ DF). They require no fundamental modification to the numerical method adopted (and have no effect on its stability properties).

The important conceptual addition is that the definitions of the conserved quantities and structure of the underlying equations for the DF impose a consistency requirement for how the mean pitch-angle ($\mu_0$) must vary across the bin, which allows us to derive these correction factors. We consider both exact formulations of this constraint, and even simpler, approximate versions which still maintain manifest conservation and ensure consistency in all relevant limits when the CR flux equations are in local steady-state (e.g. on time/spatial scales larger than the CR scattering time/mean-free-path). We test these in a simple idealized problem and show they recover the desired behaviours, with negligible difference in computational expense. All of the above applies both to one-moment methods which evolve a single scalar diffusion + streaming/advection equation (or Fokker–Planck type equation) or two-moment methods which explicitly evolve the CR flux.

We also extend this idea to similar methods that evolve radiation or neutrino hydrodynamics (again treating the spectrum as a piecewise power law, attempting to simultaneously conserve both photon number and energy). We show that in the ‘local flux steady-state’ or ‘single-moment’ limit (aka the advective-diffusive limit for radiation transport), in which the intensity is close-to-isotropic, the appropriate correction terms can be derived and represent a generalization of the usual Rosseland mean opacity to arbitrary non-thermal spectra and other conserved quantities (e.g. photon number). However, in the weak-scattering limit, the usual ambiguity in the form of the Eddington tensor makes the problem underdetermined. The key difference is that we can safely assume CRs have a close-to-gyrotropic DF with respect to the magnetic field direction (which is, of course, CR momentum-independent) – but there is no analogous constraint for radiation.

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DATA AVAILABILITY

The data supporting this article are available on reasonable request to the corresponding author.

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