Predictions for Gravity-mode Periods and Surface Abundances in Intermediate-mass Dwarfs from Shear Mixing and Radiative Levitation

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Abstract

The treatment of chemical mixing in the radiative envelopes of intermediate-mass stars has hardly been calibrated so far. Recent asteroseismic studies demonstrated that a constant diffusion coefficient in the radiative envelope is not able to explain the periods of trapped gravity modes in the oscillation spectra of $\gamma$ Doradus pulsators. We present a new generation of MESA stellar models with two major improvements. First, we present a new implementation for computing radiative accelerations and Rosseland mean opacities that requires significantly less CPU time. Second, the inclusion of shear mixing based on rotation profiles computed with the 2D stellar structure code ESTER is considered. We show predictions for the mode periods of these models covering stellar masses from 1.4 to 3.0 $M_\odot$, across the main sequence, computed for different metallicities. The morphology of the chemical mixing profile resulting from shear mixing in combination with atomic diffusion and radiative levitation does allow for mode trapping, while the diffusion coefficient in the outer envelope is large (>10$^5$ cm$^2$ s$^{-1}$). Furthermore, we make predictions for the evolution of surface abundances for which radiative accelerations can be computed. We find that the N/C and C/O abundance ratios correlate with stellar age. We predict that these correlations are observable with precisions $\lesssim$0.1 dex on these ratios, given that a precise age estimate can be made.

Unified Astronomy Thesaurus concepts: Asteroseismology (73); Stellar abundances (1577); Stellar evolution (1599); Stellar oscillations (1617)

1. Introduction

A complete and calibrated theory of chemical mixing inside stars remains an outstanding problem in stellar structure and evolution theory (Salaris & Cassisi 2017). The chemical mixing on a microscopic level is ascribed to atomic diffusion (including radiative levitation) and can be derived from first principles, whereas macroscopic mixing, caused by turbulent flows, is often parameterized as the combined effect of convection, (convective) core boundary mixing (CBM), and radiative envelope mixing (REM), each with their own free parameters. Convective mixing can be described by mixing-length theory (MLT; Böhm-Vitense 1958), where the free parameter $\alpha_{\text{MLT}}$, setting the typical length scale, is often fixed to a value calibrated to the Sun (Choi et al. 2018). A commonly used parameterization for the CBM is based on convective penetration (Zahn 1991; Augustson & Mathis 2019), where the fully mixed core is extended over a distance described by a dimensionless parameter $\alpha_{\text{ov}}$, times the local pressure scale height. In this extended region, the temperature gradient is the adiabatic one. This form of CBM was found to give a more accurate description than diffusive exponential overshooting with a radiative temperature gradient from a sample of 26 slowly pulsating B-type (SPB) stars (Pedersen et al. 2021). Beyond the CBM zone, REM takes over, which can be induced by several mechanisms, such as internal gravity waves (IGWs); Rogers & McElwaine (2017), meridional flows (Meynet & Maeder 2002), or shear instabilities due to differential rotation (Maeder 2009).

The advent of asteroseismology has made it possible to probe the internal physics of pulsating stars through their oscillation spectrum (Aerts et al. 2010). For nonrotating, nonmagnetic stars the period difference between gravity ($g$) modes with equal spherical degree, $\ell$, azimuthal order, $m$, and consecutive radial order, $n$, is constant in the asymptotic regime where $n \gg \ell$ (Tassoul 1980). Yet departures from a constant period spacing are induced by the changes in the local composition (Miglio et al. 2008), by rotation (Bouabid et al. 2013), and by the presence of a magnetic field (Prat et al. 2019; Van Beeck et al. 2020). The study of a star’s period spacing pattern (i.e., mode period $P_n$ vs. the period spacing $\Delta P_n = P_{n+1} - P_n$), or equivalently the precise mode periods themselves, allows one to probe the deep internal stellar structure (see Aerts 2021, for a review).

Different prescriptions for REM have been tested on a sample of SPB stars in the work by Pedersen et al. (2021), comparing the observed periods of $g$-mode oscillations with those predicted by models. Their work shows that an REM profile based on vertical shear is preferred for the majority of the sample containing 26 stars. Similarly, Mombarg et al. (2021) measured the value of $f_\text{ov}$, the parameter defining the extent of the CBM region (Fretyag et al. 1996), for a sample of 37 $\gamma$ Doradus ($\gamma$ Dor) stars. In their work, a constant value $D_{\text{macro}} = 1$ cm$^2$ s$^{-1}$ for the macroscopic mixing in the radiative envelope was chosen. Although for a large part of the sample this simple prescription seems sufficient, there are also several
stars that show signatures of mode trapping, such that modes are confined in a narrow cavity owing to wave reflection caused by a gradient in the local mean molecular weight. The clearest example of a γ Dor star with trapped modes is KIC 11294808, which shows pronounced dips in the period spacing pattern (Van Reeth et al. 2015) that could not be explained with the physics used in the models of Mombarg et al. (2021). These authors find that similar amplitudes of the dips, compared to those observed in KIC 11294808, are only reproduced at a level of \( D_{\text{macro}} = 0.05 \text{ cm}^2 \text{ s}^{-1} \) for the macroscopic mixing. However, such low diffusion coefficients throughout the radiative envelope are four to six orders of magnitude lower than typical values assumed for macroscopic mixing induced by turbulent diffusion in stellar models (Miglio et al. 2008; Christophe et al. 2018; Ouazzani et al. 2019). Furthermore, due to gravitational settling, KIC 11294808 should have a low surface metallicity (if radiative levitation is relatively weak), which is not supported by spectroscopic observations (Gebruers et al. 2021).

Moreover, the work by Mombarg et al. (2020) demonstrated that the inclusion of atomic diffusion and radiative levitation has large effects on the resulting predicted periods of the γ-modes. Similar results were obtained for p-modes by Deal et al. (2020). Yet, due to large amounts of computation time required for calculations related to radiative levitation and the Rosseland mean opacity, the inclusion of radiative levitation in the modeling of γ-modes in dwarfs has so far been limited to two slowly rotating γ Dor stars (Mombarg et al. 2020).

The gravitational settling of helium near the core boundary reduces the local chemical gradient and therefore inhibits mode trapping (Théado et al. 2009). Since it is well known that atomic diffusion occurs in stars (Michaud et al. 2015), this gives us extra constraints on the macroscopic mixing, as the effect of atomic diffusion needs to be counteracted in some parts of the stars when mode trapping is observed. Similarly, a form of turbulent diffusion that is scaled with the local atomic diffusion coefficient of helium was introduced by Richer et al. (2000) to counteract the gravitational settling in AmFm stars, in order to reconcile with observed surface abundances.

In this work, we demonstrate that (macroscopic) shear mixing in combination with microscopic mixing due to atomic diffusion (including radiative levitation) can explain the observed γ-mode periods of the γ Dor stars, and we investigate the implications of including these two mixing phenomena for the evolution of the surface abundances. In Section 2, we present an improved method to compute radiative accelerations and the Rosseland mean opacity from monochromatic opacity tables, reducing the computation time drastically by more than a factor of 4. In Section 3, we discuss a new implementation of macroscopic mixing induced by vertical shear and combine this with microscopic mixing to study when mode trapping can occur across the γ Dor mass regime for different ages and metallicities (Section 4). Moreover, we show the predicted evolution of surface abundances in Section 5, discuss the numerical sensitivity in Section 6, and finally conclude in Section 7.

2. Improved Routines for Radiative Levitation

We start by discussing the numerical implementation to compute radiative accelerations (i.e., the acceleration of elements induced by the process of radiative levitation) and the improvements we have made to reduce the computation time. The change in the local mass fraction of a chemical species \( i \) over time is described by

\[
\frac{\partial X_i}{\partial t} = \left( \frac{dX_i}{dt} \right)_{\text{nuc}} + \frac{1}{\rho r^2} \frac{\partial}{\partial r} \left( \rho r^2 \left[ D_{\text{macro}}(t, r) \frac{\partial X_i}{\partial r} - \nu_{\text{diff},i}(t, r) \right] \right),
\]

where the first term on the right-hand side describes the change in composition due to nuclear reactions, and the second term describes the change induced by both macroscopic and microscopic mixing. Here \( \rho \) is the local density, \( r \) is the distance from the stellar center, \( D_{\text{macro}} \) is the diffusion coefficient for macroscopic turbulent mixing, and \( \nu_{\text{diff},i} \) is the diffusion velocity of species \( i \).

We make use of the open-source stellar structure and evolution code\( \text{MESA} \), r11701 ( Paxton et al. 2011, 2013, 2015, 2018, 2019), where the default routines for computing radiative accelerations are based on the work by Hu et al. (2011). The diffusion velocity of a species \( i \) is determined in\( \text{MESA} \) by solving the Burgers equations (Burgers 1969). We refer to Mombarg et al. (2020) and references therein for details on the numerical implementation. The effectiveness of radiative acceleration enters the Burgers equations via the local acceleration induced by momentum absorption of photons generated in the stellar core (Hu et al. 2011),

\[
g_{\text{rad},i}(r) = \frac{\mu \varepsilon c}{\mu_i c} \mathcal{F}(r) \gamma_i(r),
\]

where \( \mu_i \) and \( \mu \) are the molecular weights of species \( i \) and the mean molecular weight, respectively, \( \kappa_R \) is the Rosseland mean opacity, \( \mathcal{F}(r) \) is the local radiative flux, \( c \) is the speed of light, and

\[
\gamma_i(r) = \int_0^\infty \frac{\sigma_i(u) \left[ 1 - e^{-u} \right] - a_i(u)}{\sum u f_k(r) \sigma_i(u)} du,
\]

where \( u = h \nu / k_B T \) (with \( h \) the Planck constant and \( k_B \) the Boltzmann constant). This integral depends on the local mixture that is given by the fractional abundances \( f_k \), with \( \sum f_k = 1 \), and its computation requires the monochromatic cross sections \( \sigma_i \) and correction terms \( a_i \).\( \text{MESA} \) relies on the tables provided by the OP project (Seaton 2005) for these quantities. These OP monochromatic tables provide data for H, He,6 C, N, O, Ne, Na, Mg, Al, Si, S, Ar, Ca, Cr, Mn, Fe, and Ni, which are equally spaced in

\[
\nu(u') = \frac{15}{4 \pi^4} \int_0^{u'} \frac{u' e^{-u' d} du}{(1 - e^{-u'})^3},
\]

where \( u \) ranges from \( 10^{-3} \) to 20.

As a result of atomic diffusion, the Rosseland mean opacity can no longer be evaluated for the same mixture in each cell of the model since

\[
\kappa_R(r) = \frac{1}{\mu} \left( \int_0^\infty \frac{1}{\sum u f_k(r) \sigma_i(u)} du \right)^{-1},
\]

For H and He, the correction terms \( a_i \) are not provided.
where the factors $f_k(r)$ for the metals are no longer constant throughout the star. Consistently computing $\kappa R$ requires a lot of additional computation time compared to when the fractional metal abundances are constant throughout the star.

As a first step to reduce the number of redundant computations, we pre-process the OP mono tables where we compute the cross section for momentum transfer to an atom (i.e., the numerator in Equation (3)). The data from the OP project are tabulated in temperature $T$ and electron number density $N_e$. The latter has to be converted to a mass density as per

$$\log \rho = \log N_e + \log \mu - \log \Xi - \log N_{A},$$  \hspace{1cm} (6)

where $\Xi = \sum df_k \Xi_k$ is the average number of electrons per atom and $N_{A}$ is Avogadro’s number. For the temperature $T(r)$ and density $\rho(r)$ in each cell, $\kappa R$ and $\gamma$ are computed by means of a bicubic interpolation, where we use the following scheme to select the 16 points on which the interpolation is done. First, select the closest point in the table by minimizing

$$\Delta = \sqrt{\left(\frac{1}{0.0025} \frac{\left(\log T(0) - \log T(r)\right)^2}{0.25} + \left(\frac{\left(\log \rho(0) - \log \rho(r)\right)^2}{0.25}\right)\right)},$$ \hspace{1cm} (7)

where the two constants are used to account for the difference in spacing of the tabulated $T(0)$ ($\log T(0)$ given in steps of 0.05 dex) and $\rho(0)$ ($\log \rho(0)$ given in steps of 0.5 dex). Subsequently, a grid of $4 \times 4$ points is constructed, centered around $T(r)$ and $\rho(r)$. It occasionally happens that a cell has temperature and/or density close to the edge of the OP mono tables. In that case, the $4 \times 4$ grid is moved one step at the time in temperature/density until a $4 \times 4$ grid is found that encloses $T(r)$ and $\rho(r)$. For one location in the OP mono tables (relevant for intermediate-mass main-sequence (MS) stars), the grid selection is hard-coded.

To further speed up the computation of $\kappa R$, the quantity $\kappa R(0)$ is only recomputed if

$$\max_k \left(\frac{|f_k^{(0)} - f_k^{(\rho)}|}{f_k^{(\rho)}}\right) > 10^{-4},$$ \hspace{1cm} (8)

where the superscript “$(\rho)$” indicates the mixture for which the grids have been precomputed, and “$(c)$” indicates the current mixture. The threshold value of $10^{-4}$ was found to be an optimal value by Hui-Bon-Hoa (2021), using a similar technique to speed up computations of the Rosseland mean opacity in the Toulouse-Geneva evolution code (Hui-Bon-Hoa 2008; Théado et al. 2012). We choose to evaluate the mixture in the convective core, as this is the largest zone in the star where the mixture is homogeneous. After the first time step for which $\kappa R(0)$ is computed, the $4 \times 4$ grids selected for each cell and the corresponding $T(r)$, $\rho(r)$, and $f_k$ are saved for the next time step. For the following time steps, if Equation (8) is true, the following conditions are evaluated:

$$\max_k \left(\frac{|f_k^{(c)} - f_k^{(\rho)}|}{f_k^{(\rho)}}\right) > 10^{-4},$$ \hspace{1cm} (9)

$$|\log T^{(c)}(r) - \log T^{(\rho)}(r)| > 10^{-2},$$ \hspace{1cm} (10)

$$|\log \rho^{(c)}(r) - \log \rho^{(\rho)}(r)| > 10^{-1},$$ \hspace{1cm} (11)

where the superscript “(c)” indicates the values of the previous time step. If any of these three conditions is fulfilled for a specific cell, the grid points for interpolation are recomputed for the new local mixture and stored for the next time step. Otherwise, the grid points of the previous time step are used for the interpolation of the opacity and its derivatives with respect to temperature and density.

Similarly, for the radiative accelerations we precompute a grid containing $\gamma^{(c)}$ for each temperature-density point in the OP mono data, using the initial mixture. Yet, as we only need to compute the radiative accelerations in the radiative envelope, a slightly different methodology is used, compared to the one used to compute the opacity. The radiative envelope is divided into two zones with an equal number of cells based on the number of cells for which $g_{\text{rad},i}$ needs to be computed. In each of these two zones, we define a mixture, where $f_k = \sum df_k$ is the average value of the fractional abundance of element $k$ across all cells (with cell index $i$) in the zone, and renormalize afterward such that $\sum df_k = 1$.

The value of $\gamma$ is then interpolated in temperature and density from one of the precomputed grids. A precomputed grid is recomputed for the current average mixture $f_k^{(c)}$ if the condition in Equation (8) is true. To ensure a smooth transition between these two zones, within 15 cells on each side of the boundary we blend the two values of $\log g_{\text{rad},i}$ for the two average mixtures according to

$$\log g_{\text{rad},i} = \beta_i \log g_{\text{rad},i}^{(1)} + (1 - \beta_i) \log g_{\text{rad},i}^{(2)},$$ \hspace{1cm} (12)

where we vary $\beta_i$ linearly between 0 and 1 in this transition zone and the superscripts denote the two zones that each have an average mixture.

In summary, the quantities that are saved for the next time step are listed in Table 1, and a flowchart is shown in Figure 1. Hence, by reusing previously made computations and allowing a small error on the precision of the opacity and radiative accelerations, much CPU time is saved. To evolve a 1.7 $M_\odot$ model starting at the pre-MS to a core hydrogen mass fraction $X_C = 0.005$, including radiative levitation for all elements for which OP mono data are available, takes 30 minutes on 36 CPUs with the routine presented here. In comparison, this takes 130 minutes with the current routines in MESA. Additionally, models computed in this work experience almost no failures to converge on the pre-MS, which was until now, besides the long wall times, also a bottleneck for us to compute grids of models with radiative levitation included. Figure A1 in Appendix A shows the difference in the predicted radiative accelerations and Rosseland mean opacity between the method presented in this paper and those of Hu et al. (2011). Overall, the differences are small and the differences in the surface temperature and luminosity are well within typical precisions that can be achieved observationally.
3. Implementation of Shear Mixing in MESA

In this work, we focus on macroscopic mixing from rotation-induced turbulence, henceforth referred to as shear mixing. It should be noted that this does not refer to the various forms of shear instabilities implemented in MESA (see Heger et al. 2000), which cause very spiky mixing profiles and, as such, are not suitable for g-mode asteroseismology (Truytart 2016). We follow the formalism by Zahn (1992, Equation 2.14), where we scale the local diffusive mixing coefficient as per

$$D_{\text{macro}}(r, t) = \eta K \left( \frac{r}{N} \frac{d\Omega}{dr} \right)^2,$$

where $\eta$ is a free parameter, $N$ is the (weighted) Brunt–Väisälä frequency, and $K$ is the thermal diffusivity,

$$K = \frac{16 \sigma_{\text{SB}} T^3}{3 \kappa_R \rho c_p},$$

with $\sigma_{\text{SB}}$ the Stefan–Boltzmann constant and $C_p$ the heat capacity. The radial derivative of the rotation frequency, $\Omega$, is computed using the 2D ESTER code (Espinosa Lara & Rieutord 2013; Rieutord et al. 2016) that computes a steady model at a given mass, $M_*; \ell$ the fraction of the initial core hydrogen mass fraction left, $X_c/X_{\text{ini}}; \ell$ and the rotation rate expressed as a fraction of the Keplerian critical rotation rate, $\Omega_{\text{bc}}$. For stars with masses below $\sim 3 M_\odot$, the convergence of ESTER becomes delicate. As such, we compute an ESTER model for $3 M_\odot$, $X_c/X_{\text{ini}} = 0.95$, and $\Omega_{\text{bc}} = 0.2$ and apply simple scaling relations for $M_*$ and $X_c/X_{\text{ini}}$ to scale the rotation profile from this model to the appropriate mass and age of the model we want to compute. We take a rotation profile averaged over the colatitude, starting from the boundary of the convective core, as shown in the top panel of Figure 2. As can be seen in the bottom panel of Figure 2, the global shear profile of $\frac{d\Omega}{dr}$ remains roughly the same along the evolution.

Therefore, we choose to scale the normalized $\left( \frac{d\Omega}{dr} \right)_{\text{norm}}$ profile of the aforementioned ESTER model as per

$$\xi_{M_*} = a_1 M_*^2 + a_2 M_* + a_3,$$  \quad (15)

$$\xi_{X_c/X_{\text{ini}}} = a_4 (X_c/X_{\text{ini}})^2 + a_5 (X_c/X_{\text{ini}}) + a_6.$$  \quad (16)

That is,

$$\frac{d\Omega}{dr}(M_*, X_c/X_{\text{ini}}) = \xi_{M_*} \cdot \xi_{X_c/X_{\text{ini}}} \cdot \left( \frac{d\Omega}{dr} \right)_{\text{norm}}.$$  \quad (17)

The values of the constants $a_i$ are listed in Table 2. At each cell of the MESA model, the local value of $D_{\text{macro}}$ is interpolated from the normalized $\frac{d\Omega}{dr}$ profile of the 3 $M_\odot$ ESTER model that is defined on an interval $[r_{cc}, R_*]$, that is, from the convective core boundary to the surface. In Figure 3, we show rotation profiles for three different masses at the zero-age MS (ZAMS). In addition, the rotation profile for $0.5 \Omega_{\text{bc}}$ is also shown for the most massive model, demonstrating that the morphology of the normalized rotation profile barely changes for different rotation rates.

The Brunt–Väisälä frequency can be decomposed into a part related to structure and a part related to composition,

$$N_{\text{structure}}^2 \approx \frac{g \delta}{H_P} (\nabla_{\text{ad}} - \nabla),$$  \quad (18)

$$N_{\text{composition}}^2 \approx \frac{g \bar{\rho}}{H_P} \nabla,$$  \quad (19)
Figure 2. Top panel: rotation profiles for a 3 M_☉ ESTER model with a rotation rate of Ω = 0.2Ω_k. The profiles are normalized by the rotation frequency at the core boundary (indicated by the dotted vertical lines). Bottom panel: the evolution of the shear profile (absolute value) for a 3 M_☉ ESTER model (solid lines) and the scaled approximations used for the mixing in the MESA models (dotted lines). The inset shows a zoom-in close to the core boundary (i.e., where r = 0). The radial derivative of the local rotation frequency shown here is dimensionless, as Ω is normalized by \( \sqrt{\rho/(\Omega R)} \), and r is normalized by the size of the radiative envelope, R_env, where r = 0 at the convective core boundary.

Table 2

| Coefficient | Value |
|-------------|-------|
| a₁         | 0.05159189 |
| a₂         | -0.30399420 |
| a₃         | 0.63434263 |
| a₄         | 0.14382828 |
| a₅         | -0.39459215 |
| a₆         | 0.43750162 |

where \( \nabla_\text{ad}, \nabla, \) and \( \nabla_\mu \) are the adiabatic, temperature, and chemical gradient, respectively. Moreover, g is gravitational acceleration, \( H_P \) is the local pressure scale height, \( \delta = -(\partial \ln \rho/\partial \ln T)|_{\rho,\mu}, \) and \( \phi = -(\partial \ln \rho/\partial \ln \mu)|_{\rho,\mu}. \) In the formalism of Mathis et al. (2004), the mixing coefficient scales with \( K + \Delta h, \) where \( \Delta h \) is the horizontal diffusion coefficient. Here we do not compute \( \Delta h, \) but instead approximate

\[ K + \Delta h = \eta K \]

and construct a weighted Brunt–Väisälä frequency,

\[ N^2 = N^2_{\text{str}} + \frac{\eta}{\eta - 1} N^2_{\text{comp}}. \]  

Figure 3. The variation of the normalized rotation profile with stellar mass for ZAMS models. For the 3.0 M_☉ model, the rotation profile at 0.5Ω_k is shown by the dashed–dotted line (almost the same as the profile for 0.2Ω_k). Note that in this plot r ∈ [0, R].

\[ K + \Delta h = \eta K \]

and construct a weighted Brunt–Väisälä frequency,

\[ N^2 = N^2_{\text{str}} + \frac{\eta}{\eta - 1} N^2_{\text{comp}}. \]  

Hence, our implementation has one free parameter, \( \eta, \) to control the global scaling of the mixing profile. This weighting of the different components follows from the fact that while both components of the buoyancy force are weakened by horizontal diffusion, only the structure (thermal) component is additionally weakened through thermal diffusion (Talon & Zahn 1997). The point where the transition is made from CBM to REM is chosen at the smallest radius where \( D_{\text{REM}}(r) > D_{\text{ov}}(r). \) An example of a profile for \( D_{\text{macro}}(r) \) is shown in Figure 4. The mixing profile based on vertical shear used by Pedersen et al. (2021) comes from a profile computed by Georgy et al. (2013), relying on 1D stellar models. Pedersen et al. (2021) then scale this profile by a free parameter that sets the diffusion coefficient at the edge of the overshoot zone. In this work, we rely on 2D ESTER models to compute the rotation profile, which are in addition scaled with the stellar mass and the hydrogen mass fraction in the core. We find similar mixing profiles to those derived by Mathis et al. (2004), although the drop in the diffusion coefficient close to the overshoot zone (around \( r/R = 0.1 \) in Figure 4) is much more pronounced in our models.

One particularly interesting star in the sample of Van Reeth et al. (2015) is KIC 11294808 (\( f_{\text{rot}} = 0.77 \) days⁻¹, \( M_* = 1.655 M_☉, X_\alpha = 0.19, \)) for which period spacing patterns of dipole \( (ℓ = 1, m = 1) \) modes and quadrupole \( (ℓ = 2, m = 2) \) modes were observed. The periods of the quadrupole modes exhibit clear recurring dips, indicative of trapped modes. Therefore, we use this star to find a reasonable value for \( \eta, \) which produces dips in the period spacing pattern for a model with the parameters found by Mombarg et al. (2021). We use a scaling of \( \alpha_{\text{ov}} \approx 10f_{\text{ov}} \) (Claret & Torres 2017), as in this work we model the CBM with convective penetration. On one hand, when \( \eta \) is set too low (≤20; see Figure 5), helium settling is not counteracted efficiently, thereby stabilizing the chemical gradient near the core boundary. On the other hand, if \( \eta \) is
set too high (around 5000, not shown), chemically homogeneous evolution occurs. As shown in Figure 5, a value of \( \eta = 60 \) provides a good estimate for KIC 11294808. We stress that the predicted locations of the dips in the period spacing pattern are extremely sensitive to the stellar mass, age, and input physics, and that the aim of this paper is not to precisely match the periods but to investigate whether models with shear mixing and radiative levitation can reproduce the global morphology of the observed period spacing pattern.

4. Oscillation Spectra

We have computed MESA models for masses of 1.4, 1.7, 2.0, and 3.0 \( M_\odot \) covering the entire \( \gamma \) Dor mass regime and the lowest part of the SPB mass regime. For each model, we have computed the predicted periods at \( X_c/X_{\text{ini}} = 0.95, 0.45, \) and 0.10. Moreover, we have tested three different values for the initial metallicity, \( Z_{\text{ini}} \), namely, 0.010, 0.015, and 0.020. We refer to Mombarg et al. (2021) and Mombarg et al. (2020) for more details on our MESA setup. One difference between the setup used in this work and the aforementioned ones is that here we describe the CBM by convective penetration, following the results from Pedersen et al. (2021). It should be noted, however, that no clear distinction between an exponential and step overshoot profile\(^7\) has yet been made for \( \gamma \) Dor stars (Mombarg et al. 2019, 2021). Our focus is on the prograde dipole modes and radial orders \( n \in [-100, -10] \), as these are the most commonly observed mode geometry in \( \gamma \) Dor and SPB pulsators (e.g., Van Reeth et al. 2016; Li et al. 2020; Pedersen et al. 2021). The mode periods were computed using the stellar pulsation code \texttt{GYRE} (v5.2; Townsend 2003; Townsend & Teitler 2013). In Figure 6, we show the predicted period spacings, \( \Delta P_n = P_{n+1} - P_n \), as a function of period, \( P_n \), for the three different values of \( X_c/X_{\text{ini}} \). To produce mode trapping when atomic diffusion (including radiative levitation) is taken into account, a low mixing efficiency close to the core boundary is required, and a high efficiency further outward. Our models show that this can be realized by the prescription given in Equation (13), as we observe mode trapping for all stellar masses studied here when we choose \( \eta = 60 \). The prescription for REM used in this work also requires much higher diffusion coefficients outside the shear zone, solving the stark contrast between theoretical predictions and the very low diffusion coefficients that Van Reeth et al. (2016) and Mombarg et al. (2021) needed to explain the mode trapping in some stars in their sample, and the low values found by Moravveji et al. (2016), when the value of the diffusion constant was fixed throughout the star.

In Figure 7, the effect of the initial metallicity is demonstrated for models with \( X_c/X_{\text{ini}} = 0.10 \) (see Figure B1 in Appendix B for models with \( X_c/X_{\text{ini}} = 0.45 \)). Since the initial helium abundance in our models is scaled with the metallicity, according to the chemical enrichment rate derived by Verma et al. (2019), a higher \( Z_{\text{ini}} \) means that the helium settling near the core diminishes the chemical gradient to a larger extent, and...

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as such, the dips in the period spacing pattern caused by mode trapping are less pronounced. Interestingly, we observe three sinusoidal modulations in the period spacings of the high radial order modes of the evolved $1.7 \, M_\odot$ model in the lowest-metallicity case, whereas for higher metallicities a nearly constant slope is seen. In the model with $Z_{\text{ini}} = 0.010$, additional dips in the REM profile are formed, which create several mode cavities in which modes can get trapped (see Figure C1 in Appendix C).

The signatures of mode trapping for models with combined atomic diffusion and shear mixing are similar to those predicted by models with no (or very low) micro- and macroscopic REM, as either set of physics introduces an oscillatory behavior of the dips in the period spacings, as shown by Miglio et al. (2008) and Bouabid et al. (2013). Besides mode trapping, dips in the period spacing pattern can also be caused by coupling between gravito-inertial modes in the envelope and pure inertial modes in the convective core at specific mode frequencies (Ouazzani et al. 2020; Saio et al. 2021).

5. Surface Abundances

Additionally, the inclusion of REM allows us to study the evolution of the surface abundances, which provide extra constraints on the global efficiency of the chemical mixing. Dotter et al. (2017) already showed that atomic diffusion and radiative levitation affect the surface abundances throughout stellar evolution significantly for stars with masses between 0.5 and $1.5 \, M_\odot$. In Figure 8, we show for our models the predicted surface abundances as a function of age for elements He, C, N, O, Na, Mg, Al, Si, S, Ca, Fe, and Ni. The predicted evolution of the $[\text{Fe}/H]$ abundance of the $1.4 \, M_\odot$ model from Deal et al. (2020), who also include radiative levitation and rotational mixing in their models, shows a similar trend compared to our findings. That is, the $[\text{Fe}/H]$ abundance decreases during the first part of the MS and increases again during the second part of the MS. The same behavior is seen for elements heavier than oxygen, which do not partake in the CNO cycle and thus have a constant bulk abundance. Around halfway through the MS, the radiative acceleration of iron-peak elements exceeds the local gravitational acceleration, preventing these elements from sinking toward the core, resulting in an increase of the metal surface abundances. Yet, for the models with $1.7 \, M_\odot$ and up, the radiative levitation of calcium is strong relatively near the surface, and a constant increase in the surface abundance along the MS is observed. The fact that we see changes in surface abundances throughout the evolution indicates that even the high mixing efficiencies in the envelope used in this work are not enough to completely dominate over the effects of radiative

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**Figure 6.** Period spacing patterns for models with shear mixing and atomic diffusion (including radiative levitation) at $X_c/X_{\text{ini}} = 0.95$ (top panel), $X_c/X_{\text{ini}} = 0.45$ (middle panel), and $X_c/X_{\text{ini}} = 0.10$ (bottom panel). For all models $Z_{\text{ini}} = 0.015$, $\alpha_{ov} = 0.15$, and $\Omega = 0.2\Omega_{bk}$.

**Figure 7.** Same as Figure 6, but for varying metallicity. For all models $X_c/X_{\text{ini}} = 0.10$.

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8 Presented as $[X/H] = \log(X/X_\odot) - \log(A_X/A_\odot) - \log \epsilon_{X,\odot} + 12$, with $X_\odot$ the mass fraction and $A_X$ the atomic mass. For the values of $\log \epsilon_{X,\odot}$, the solar composition from Asplund et al. (2001, $Z_{\odot} = 0.0139$) is used.
Figure 8. Predicted evolution of the surface abundances (scaled to the Sun) for models with $Z_{\text{ini}} = 0.015$ and $\eta = 60$. The dotted lines correspond to a model with $1.4 \, M_\odot$ and $\eta = 2$. The dashed–dotted lines are for a $1.4 \, M_\odot$ model without any macroscopic mixing ($\eta = 0$).
levitation. Yet the typical excursions in \([X/H]\) are about a factor of 10 less than what was predicted in the absence of strong REM by Mombarg et al. (2020). For the other two metallicity values \((Z_{\text{ini}} = 0.010 \text{ and } 0.020)\), we find similar behavior of the surface abundances. As the typical precision for \([X/H]\) derived in F-type stars is at best on the order of 0.1 dex (Gebran et al. 2010; Niemczura et al. 2015; Takada-Hidai et al. 2017; Gebruers et al. 2021), we expect that most \(\gamma\) Dor stars will not show clear signs of heavy-element depletion at the surface, as most have moderate to fast rotation frequencies (Van Reeth et al. 2016, 2018). Indeed, spectroscopic studies of \(\gamma\) Dor and SPB stars show that most of them have surface compositions similar to the Sun (Kahraman Aliçavuş et al. 2016; Gebruers et al. 2021).

Aside from microscopic mixing, efficient REM due to rotational mixing can transport the CNO elements from the core toward the surface. Hydrogen burning via the CNO cycle occurs through a chain of several reactions, where the conversion of nitrogen to oxygen is slow and causes a buildup of nitrogen and a depletion of carbon, while the amount of oxygen stays roughly the same. We predict that this change can be observed, as the gravitational settling of these elements is not efficient enough to cause depletion of the CNO elements at the surface. In Figure 9, the ratios of N/C and C/O are shown, as well as Mg/Fe and O/Fe. Observational measurements of \(\log(N/C)\) or \(\log(C/O)\) could serve as additional constraints on the stellar age, if determined with precisions of \(\sim 0.2\) and 0.1 dex, respectively. The ratio Mg/Fe is typically the abundance ratio that can be determined most precisely in F-type stars. Our models predict that this ratio remains almost constant throughout the MS, which is consistent with ratios inferred by Gebruers et al. (2021) for a sample of 91 \(\gamma\) Dor stars.

Additionally, the predicted evolution of the surface abundances for a 1.4 \(M_\odot\) model with \(\eta = 2\) (i.e., the horizontal diffusion coefficient is of the same order as the thermal diffusivity) is shown in Figure 8 (dotted lines). We find that the excursions in \([X/H]\) of O, Na, Mg, Al, Si, S, Fe, and Ni are not significantly increased, compared to typical precisions that can be achieved with spectroscopy. The lower mixing efficiency at the outer part of the envelope can no longer drive Ca to the surface around mid-MS, resulting in Ca depletion when the star evolves. The diffusion coefficient just below the surface convection zone of the model with \(\eta = 2\) is about two orders lower than that of the model with \(\eta = 60\) (i.e., the thermal diffusivity is smaller than the horizontal diffusion). For C and N, we find that the mixing becomes too low to induce any changes in the abundances at the surface. In more evolved stars, the C/H ratio could be used to put a lower limit on the mixing just outside of the CBM zone, when the bulk metallicity is well known and the initial composition is assumed to be close to the solar composition. For models without shear mixing, the maximum depletion of the heavier elements is stronger, and underabundances in Mg and Fe (Mg/Fe ratio remains stable) could be observed in mid-MS stars when precisions of \(\sigma_{[X/H]} \leq 0.1\) dex can be achieved. The 1.4 \(M_\odot\) model without macroscopic mixing (dashed–dotted lines in Figure 8) does not show complete helium depletion, even though the radiative levitation for this element is extremely weak. The settling of helium at the surface of this model is impeded by the formation of a surface convection zone (see

![Figure 9](image_url)

**Figure 9.** Predicted evolution of several surface abundance ratios for models with \(Z_{\text{ini}} = 0.010\) (solid lines), 0.015 (dashed–dotted lines), and 0.020 (dotted lines). The most precise measurements achieved by Gebruers et al. (2021) for SPB and \(\gamma\) Dor stars are indicated with a gray square and circle, respectively. The most precise measurements of the Mg/Fe ratio are around 0.6 dex and are thus much larger than the predicted change of this ratio along the evolution.
Verma & Silva Aguirre 2019), as is shown in Figure D1 in Appendix D.

6. Numerical Uncertainties

As the profile of the Brunt–Väisälä frequency is used in the computation of the mixing profile, the evolution becomes sensitive to the mesh resolution of the stellar model. For simple models without radiative levitation or shear mixing, the difference in the predicted mode periods at some point becomes smaller than the typical observational uncertainty on the period when more cells are added to the model. However, when radiative levitation is taken into account, the mesh resolution is limited, since each cell has its own composition, which makes it significantly more difficult for the solver to converge to a model within the tolerances. We computed the best model of KIC 11294808 with three different mesh grids with 2845 (M1), 3010 (M2), and 3109 (M3) cells,9 where the extra cells are placed around the large change in chemical composition just outside the core. The differences in the predicted mode periods by model M2 and M3 are roughly the same as the period differences between model M1 and M2, as shown in Figure E1 in Appendix E, indicating that our predictions of the mode periods are limited to a numerical uncertainty of ~60 s on the low radial order (shorter-period) modes and ~30 s for the high radial order (longer-period) modes. In Figure E2, we show rotational kernels overplotted on the g-mode cavity to illustrate that these the dips in the period spacing patterns are indeed caused by trapped modes and are not the result of theoretical uncertainties. We emphasize that these uncertainties are caused by the choice of the meshing, and that it is not the noise on the individual periods computed from a single equilibrium model. Even though these numerical uncertainties are, compared to the mode periods, smaller than 0.2%, they are still several times larger than the extremely small uncertainties typically achieved with the Kepler nominal mission data.

7. Discussion and Conclusion

This study demonstrates the added value of integrating a 1D stellar evolution code and a 2D stellar structure code to provide a more accurate description of the evolution of chemical mixing in the envelope of rotating stars with a convective core. The predictions presented in this work serve as a valuable guide for more detailed calibration of the envelope mixing in intermediate-mass gravity-mode pulsators, as Pedersen et al. (2021) already showed the need of varying diffusion coefficients throughout the envelope in order to explain the observed pulsations of a sample of 26 SPB stars. The treatment of chemical mixing presented in this work allows for mode trapping, even when microscopic diffusion in the form of atomic diffusion (including radiative levitation) is taken into account. Moreover, it results in realistic and much larger diffusion coefficients in the outer envelope, compared to when microscopic diffusion and shear mixing are ignored. The inclusion of radiative levitation (and consistent Rosseland opacity) makes the morphology of the period spacing pattern more dependent on the bulk metallicity. When shear mixing is the dominant source of macroscopic mixing, the iron abundance is expected to be relatively stable over the MS evolution and thus can be used as a tracer for the bulk metallicity of the star. We predict that in some cases stars with subsolar metallicity ([M/H] < −0.15) show a very characteristic modulation of the period spacings in the asymptotic regime. The estimated mass and age of the star KIC 7380501 by Mombarg et al. (2021) are close to those of the 1.7 M⊙ model in Figure 7, and its metallicity is subsolar ([M/H] = −0.21 ± 0.06; Gebruers et al. 2021). The morphology of the period spacing pattern of KIC 7380501 is particularly complex to model (see Figure F1 in Appendix F), where some parts of the period spacing pattern, consisting of three modes, show a much larger slope compared to the general slope of the pattern. This could be explained by the pattern predicted from the aforementioned equilibrium models, if only a few consecutive modes are observed, and so radial orders are suppressed.

Besides mixing induced by rotation, it has been shown that IGWs generated by turbulent core convection can also efficiently transport material throughout the radiative envelope (Lecoanet & Quataert 2013; Rogers & McElwaine 2017). The work of Rogers & McElwaine (2017) predicts the efficiency of the mixing induced by IGWs to scale as \( D_{\text{IGW}}(r) \propto \rho(r)^{-5/3} \), where \( \rho \) is somewhere between \( \zeta \) and 1. Hence, we expect that for stars with trapped modes IGW mixing is not dominant, as the current simulations do not predict a large variation of the mixing coefficient near the core. Yet, for stars that show no signs of mode trapping in their oscillation spectra, mixing induced by IGWs might be the dominant source. A density-dependent scaling of the diffusion coefficient (that is constant in time) is hard to distinguish from a constant diffusion coefficient, based on the periods of gravity modes (Pedersen et al. 2018). In that case, the surface abundances may hold more discriminating power when microscopic mixing is treated consistently, as done in this work. The effect of IGW mixing on the mode periods and surface abundances will be addressed in a future paper.

The NASA TESS mission (Ricker et al. 2015) covers a much larger part of the sky compared to Kepler (Borucki et al. 2010) and therefore provides a unique opportunity to study stars in different metallicity regimes. Period spacing patterns in 128 γ Dor stars in the TESS Southern Continuous Viewing Zone have been extracted by S. Garcia et al. (submitted). These stars are overall brighter than those observed by Kepler, making them suitable for high-resolution spectroscopic follow-up studies.

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9 MESA adapts the total number of cells at each time step. The numbers listed are specifically for the mentioned models.
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Software: MESA (r11701; Paxton et al. 2011, 2013, 2015, 2018, 2019), MESASDK (Townsend 2019a, 2019b), GYRE (v5.2; Townsend & Teitler 2013; Townsend et al. 2018), matplotlib (Hunter 2007), numpy (Harris et al. 2020).

Appendix A
Benchmark Results

In Figure A1, we show a benchmark test of our novel implementation for computing radiative accelerations and consistent Rosseland mean opacities, compared to the implementation in MESA r11701.

Figure A1. A comparison between the radiative accelerations (top panel) and Rosseland mean opacity (bottom panel) computed with the method presented in this paper (solid lines) and the routines from Hu et al. (2011) implemented in MESA r11701 (dashed–dotted lines). The computations are based on a 1.7 $M_\odot$ model at $X_c = 0.325$ ($Z_{ini} = 0.015$). The extent of the convective core is highlighted by the shaded region. The local gravitational acceleration is indicated by the gray dotted line.

Appendix B
Predicted Period Spacings Mid-MS

In Figure B1, we show the effect of the initial metallicity on the predicted period spacing patterns for models of mass 1.4, 1.7, 2.0, and 3.0 $M_\odot$, roughly halfway through the MS.

Figure B1. Predicted period spacing patterns for models with shear mixing and radiative levitation, for different metallicities. For all models $X_c/X_{ini} = 0.45$.

Appendix C
Rotational Kernels

Figure C1 shows the predicted mixing profile, overplotted with the rotational kernel of a $g$-mode with $(\ell, m, n_{gg}) = (1, 1, -75)$. For the model with $Z_{ini} = 0.010$ (top panel) this mode is trapped, whereas for the model with $Z_{ini} = 0.015$ mode trapping is not observed.

Appendix D
Chemical Profiles of Helium

In Figure D1, we show the helium mass fraction throughout the star at different times during the MS for a 1.4 $M_\odot$ model without macroscopic mixing (i.e., only microscopic mixing; dashed–dotted lines in Figure 8).
Appendix E

Theoretical Uncertainty

Figure E1 demonstrates how the predicted mode periods (top and middle panels) and the Brunt–Väisälä frequency (bottom panel) are influenced by the number of cells of the MESA equilibrium model. In Figure E2, we show the corresponding rotational kernels of the shortest-period modes of the model with 2845 cells that is shown in Figure E1.

Appendix F

KIC 7380501

In Figure F1, we show the $(\ell, m) = (1, 1)$ period pattern of KIC 7380501 that was extracted by Van Reeth et al. (2015). The modulations observed in this period spacing pattern are particularly ill reproduced by the best model found by Mombarg et al. (2021).

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Figure E2. Rotational kernels normalized to the largest maximum amplitude of the kernels for the shortest-period modes of the model with 2845 cells. The corresponding mode period (in days) and period spacing (in seconds) are also shown. The modes that correspond to the dips in the period spacing pattern shown in the top panel of Figure E1 are highlighted in red. The gray dashed lines indicate the Brunt–Väisälä frequency, normalized to 1.

Figure F1. The observed period spacing pattern of KIC 7380501 found by Van Reeth et al. (2015). The uncertainties of the observed values are typically smaller than the symbol size.

References

Aerts, C. 2021, RvMP, 93, 015001
Aerts, C., Christensen-Dalsgaard, J., & Kurtz, D. W. 2010, A Steroseismology (Heidelberg: Springer)
Ashplund, M., Amarsi, A. M., & Grevesse, N. 2021, A&A, 653, A141
Augustsson, K. C., & Mathis, S. 2019, ApJ, 874, 83
Böhm-Vitense, E. 1958, ZaP, 46, 108
Borucki, W. J., Koch, D., Basri, G., et al. 2010, Sci, 327, 977
Bouabid, M.-P., Dupret, M.-A., Salmon, S., et al. 2013, MNRAS, 429, 2500
Burgers, J. M. 1969, Flow Equations for Composite Gases (New York: Academic Press)
Choi, J., Dotter, A., Conroy, C., & Ting, Y.-S. 2018, ApJ, 860, 131
Christophe, S., Ballot, J., Ouazzani, R. M., Antoci, V., & Salmon, S. J. A. J. 2018, A&A, 618, A47
Claret, A., & Torres, G. 2017, ApJ, 849, 18
Deal, M., Goupil, M. J., Marques, J. P., Reese, D. R., & Lebreton, Y. 2020, A&A, 633, A23
Dotter, A., Conroy, C., Cargile, P., & Asplund, M. 2017, ApJ, 840, 99
Espinosa Lara, F., & Rieutord, M. 2013, A&A, 552, A35
Freytag, B., Ludwig, H. G., & Steffen, M. 1996, A&A, 313, 497
Gebran, M., Vick, M., Monier, R., & Fossati, L. 2010, A&A, 523, A71
Gebruers, S., Straumit, I., Tkachenko, A., et al. 2021, A&A, 650, A151
Heger, A., Langer, N., & Woosley, S. E. 2000, ApJ, 528, 368
Hu, H., Tout, C. A., Glebbeek, E., & Dupret, M.-A. 2011, MNRAS, 418, 195
Hui-Bon-Hoa, A. 2008, Ap&SS, 316, 55
Hui-Bon-Hoa, A. 2021, A&A, 646, L6
Hunter, J. D. 2007, CSE, 9, 90
Kahraman Aliçavuş, F., Niemczura, E., De Cat, P., et al. 2016, MNRAS, 458, 2307
Lecoanet, D., & Quataert, E. 2013, MNRAS, 430, 2363
Li, G., Van Reeth, T., Bedding, T. R., et al. 2020, MNRAS, 491, 3586
Maeder, A. 2009, Physics, Formation and Evolution of Rotating Stars (Berlin: Springer)
Mathis, S., Palacios, A., & Zahn, J. P. 2004, A&A, 425, 243
Meynet, G., & Maeder, A. 2002, A&A, 390, 561
Michaud, G., Alecian, G., & Richer, J. 2015, Atomic Diffusion in Stars (Switzerland: Springer International Publishing)
Miglio, A., Montalbán, J., Noels, A., & Eggenberger, P. 2008, MNRAS, 386, 1487
Mombarg, J. S. G., Dotter, A., Van Reeth, T., et al. 2020, ApJ, 895, 51
Mombarg, J. S. G., Van Reeth, T., & Aerts, C. 2021, A&A, 650, A58
Mombarg, J. S. G., Van Reeth, T., Pedersen, M. G., et al. 2019, MNRAS, 485, 3248
Moravveji, E., Townsend, R. H. D., Aerts, C., & Mathis, S. 2016, ApJ, 823, 130
Niemczura, E., Murphy, S. J., Smalley, B., et al. 2015, MNRAS, 450, 2764
Ouazzani, R. M., Marques, J. P., Goupil, M. J., et al. 2019, A&A, 626, A121
Paxton, B., Bildsten, L., Dotter, A., et al. 2011, ApJS, 192, 3
Paxton, B., Cantiello, M., Arras, P., et al. 2013, ApJS, 208, 4
Paxton, B., Marchant, P., Schwab, J., et al. 2015, ApJS, 220, 15
Paxton, B., Schwab, J., Bauer, E. B., et al. 2018, ApJS, 234, 34
Paxton, B., Smolec, R., Schwab, J., et al. 2019, ApJS, 243, 10
Pedersen, M. G., Aerts, C., Pápics, P. I., et al. 2021, NatAs, 5, 715
Pedersen, M. G., Aerts, C., Pápics, P. I., & Rogers, T. M. 2018, A&A, 614, A128
Prat, V., Mathis, S., Buysschaert, B., et al. 2019, A&A, 627, A64
Richer, J., Michaud, G., & Turcotte, S. 2000, ApJ, 529, 338
Ricker, G. R., Winn, J. N., Vanderburg, A., et al. 2015, JATIS, 1, 014003
Rieutord, M., Espinosa Lara, F., & Putigny, B. 2016, JCoPh, 318, 277
Rogers, T. M., & McElwaine, J. N. 2017, ApJL, 848, L1
Saio, H., Takata, M., Lee, U., Li, G., & Van Reeth, T. 2021, MNRAS, 502, 5856
Salaris, M., & Cassisi, S. 2017, RSOS, 4, 170192
Seaton, M. J. 2005, MNRAS, 362, L1
Takada-Hidai, M., Kurtz, D. W., Shibahashi, H., et al. 2017, MNRAS, 470, 4908
Talon, S., & Zahn, J. P. 1997, A&A, 317, 749
Tassoul, M. 1980, ApJS, 43, 469
Théado, S., Alecian, G., LeBlanc, F., & Vauclair, S. 2012, A&A, 546, A100
Théado, S., Vauclair, S., Alecian, G., & LeBlanc, F. 2009, ApJ, 704, 1262

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Townsend, R. 2019a, MESA SDK for Linux, v20190503, Zenodo, doi:10.5281/zenodo.2669541
Townsend, R. 2019b, MESA SDK for Mac OS, 20190503, Zenodo, doi:10.5281/zenodo.2669543
Townsend, R. H. D. 2003, MNRAS, 340, 1020
Townsend, R. H. D., Goldstein, J., & Zweibel, E. G. 2018, MNRAS, 475, 879
Townsend, R. H. D., & Teitler, S. A. 2013, MNRAS, 435, 3406
Truyeart, K. 2016, Master’s thesis, KU Leuven, Belgium

Van Beeck, J., Prat, V., Van Reeth, T., et al. 2020, A&A, 638, A149
Van Reeth, T., Mombarg, J. S. G., Mathis, S., et al. 2018, A&A, 618, A24
Van Reeth, T., Tkachenko, A., & Aerts, C. 2016, A&A, 593, A120
Van Reeth, T., Tkachenko, A., Aerts, C., et al. 2015, ApJS, 218, 27
Verma, K., Raodeo, K., Basu, S., et al. 2019, MNRAS, 483, 4678
Verma, K., & Silva Aguirre, V. 2019, MNRAS, 489, 1850
Zahn, J. P. 1991, A&A, 252, 179
Zahn, J. P. 1992, A&A, 265, 115