The role of mass transfer and common envelope evolution in the formation of merging binary black holes

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\section{Introduction}

In the last five years, the discoveries from ground-based gravitational-wave detectors grows, increasingly accurate theoretical models are required to compare them to the observed sample and disentangle contributions from multiple channels. In formation models involving isolated binary stars, important uncertainties remain regarding the stability of mass transfer and common-envelope evolution. To study some of these uncertainties, we have computed binary simulations using the \textsc{mesta} code consisting of a 30\,$M_\odot$ star in a low metallicity ($Z/10$) environment with a black-hole companion. We have developed an updated prescription to compute mass transfer rates including the possibility of outflows from outer Lagrangian points, as well as a method to self-consistently determine the core-envelope boundary in cases where there is common-envelope evolution. We find that binaries survive common-envelope evolution only if unstable mass transfer happens after the formation of a deep convective envelope, resulting in a narrow range (0.2 dex) in period for successful envelope ejection. All cases where binary interaction is initiated with a radiative envelope have large binding energies ($\sim 10^{53}$ erg), and they result in mergers during the common-envelope phase even under the assumption that all the internal and recombination energy of the envelope, as well as the energy from an inspiral, is used to eject the envelope. This is independent of whether or not helium is ignited in the core of the donor, conditions under which various rapid-population synthesis calculations assume a successful envelope ejection is possible. Moreover, we find that the critical mass ratio for instability is such that across a large range in initial orbital periods ($\sim 1$–1000 days), merging binary black holes can be formed via stable mass transfer. A large fraction of these systems undergo overflow of their L$^c$ equipotential, in which case we find that stable mass transfer produces merging binary black holes even under extreme assumptions of mass and angular momentum outflows. Our conclusions are limited to the study of one donor mass at a single metallicity, but they suggest that population synthesis calculations overestimate the formation rate of merging binary black holes produced by common-envelope evolution and that stable mass transfer could dominate the formation rate from isolated binaries. This is in agreement with a few other recent studies. Further work is required to extend these results to different masses and metallicities as well as to understand how they can be incorporated into rapid population synthesis calculations.

\textbf{Key words.} binaries – close – stars: massive – stars: black holes – gravitational waves – X-rays: binaries

\section{ABSTRACT}

As the number of merging binary black holes observed with ground-based gravitational-wave detectors grows, increasingly accurate theoretical models are required to compare them to the observed sample and disentangle contributions from multiple channels. In formation models involving isolated binary stars, important uncertainties remain regarding the stability of mass transfer and common-envelope evolution. To study some of these uncertainties, we have computed binary simulations using the \textsc{mesta} code consisting of a 30\,$M_\odot$ star in a low metallicity ($Z/10$) environment with a black-hole companion. We have developed an updated prescription to compute mass transfer rates including the possibility of outflows from outer Lagrangian points, as well as a method to self-consistently determine the core-envelope boundary in cases where there is common-envelope evolution. We find that binaries survive common-envelope evolution only if unstable mass transfer happens after the formation of a deep convective envelope, resulting in a narrow range (0.2 dex) in period for successful envelope ejection. All cases where binary interaction is initiated with a radiative envelope have large binding energies ($\sim 10^{53}$ erg), and they result in mergers during the common-envelope phase even under the assumption that all the internal and recombination energy of the envelope, as well as the energy from an inspiral, is used to eject the envelope. This is independent of whether or not helium is ignited in the core of the donor, conditions under which various rapid-population synthesis calculations assume a successful envelope ejection is possible. Moreover, we find that the critical mass ratio for instability is such that across a large range in initial orbital periods ($\sim 1$–1000 days), merging binary black holes can be formed via stable mass transfer. A large fraction of these systems undergo overflow of their L$^c$ equipotential, in which case we find that stable mass transfer produces merging binary black holes even under extreme assumptions of mass and angular momentum outflows. Our conclusions are limited to the study of one donor mass at a single metallicity, but they suggest that population synthesis calculations overestimate the formation rate of merging binary black holes produced by common-envelope evolution and that stable mass transfer could dominate the formation rate from isolated binaries. This is in agreement with a few other recent studies. Further work is required to extend these results to different masses and metallicities as well as to understand how they can be incorporated into rapid population synthesis calculations.

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\section{1. Introduction}

In the last five years, the discoveries from ground-based gravitational wave (GW) detectors have driven a large effort to understand the origin of compact binary black holes (BHs). With 50 detections with a high significance (Abbott et al. 2019, 2020a), the majority of the observed GW sources correspond to binary BH mergers covering a large range of masses \texttilde 5–100\,$M_\odot$. Important discoveries contained in the first Gravitational Wave Transient Catalog (Abbott et al. 2019) include the discovery of BHs more massive than those detected through electromagnetic observations in the Galaxy (Abbott et al. 2016) and the direct association of short gamma-ray bursts with binary neutron star mergers (Abbott et al. 2017). The increased size of the sample on the second Gravitational Wave Transient Catalog (Abbott et al. 2020a) points to additional features in the properties of merging binary BHs, indicative of the contribution of multiple formation channels (Abbott et al. 2021; Zevin et al. 2021). Accurate predictions from different formation channels are then necessary to understand their relative contributions.

A large number of formation scenarios have been proposed to form the merging binary BHs observed by ground-based detectors. Scenarios that involve binary systems include evolution through a common-envelope (CE) phase (e.g., Paczynski 1976; van den Heuvel 1976; Tutukov & Yungelson 1993; Belczynski et al. 2002; Dominik et al. 2012; Stevenson et al. 2017; Giacobbo & Mapelli 2018), chemically homogeneous evolution (Mandel & de Mink 2016; Marchant et al. 2016; de Mink & Mandel 2016; du Buisson et al. 2020; Riley et al. 2021), stable mass transfer (MT; van den Heuvel et al. 2017; Neijssel et al. 2019; Bavera et al. 2021) and Population III stars (Belczynski et al. 2004; Kinugawa et al. 2014;
Inayoshi et al. (2017). Dynamical processes are also predicted to contribute to the observed sample, including isolated triple systems (Thompson 2011; Antonini et al. 2017; Vigna-Gómez et al. 2021) and interactions in globular (e.g., Kulkarni et al. 1993; Sigurdsson & Hernquist 1993; Portegies Zwart & McMillan 2000; Rodriguez et al. 2015; Di Carlo et al. 2019) and nuclear clusters (Antonini & Perets 2012). Additional formation scenarios include the pairing and growth of stellar mass BHs in the disks of active galactic nuclei (McKernan et al. 2014; Stone et al. 2017; Bartos et al. 2017), and primordial BH formation (Bird et al. 2016; Sasaki et al. 2018).

Our focus is on the formation of merging binary BHs through both CE evolution and stable MT. The majority of predicted binary BHs that formed through CE evolution or stable MT in an isolated binary involve an initial phase of stable MT. This removes the hydrogen envelope of the more massive star and leads to the formation of a wide single degenerate binary (Dominik et al. 2012; Langer et al. 2020). When the secondary evolves and thus fills its Roche lobe, depending on its response to mass loss and the mass ratio of the system, it can undergo a CE phase which ejects the envelope of the secondary at the cost of hardening the binary (cf. Belczynski et al. 2016). If the second phase of MT is stable, depending on the mass ratio of the system, a phase of nonconservative mass transfer can harden the binary without the need for CE evolution, as van den Heuvel et al. (2017) argue, thus potentially allowing for the formation of compact object binaries that can merge within a Hubble time. Recent studies have claimed that the contribution from stable mass transfer to the observed sample of merging binary BHs can be comparable or even larger than those that formed through CE evolution (Neijssel et al. 2019; Bavera et al. 2021).

However, binary evolution models have large uncertainties associated with the CE phase (see Ivanova et al. 2013 for a recent review). Important open issues include the unknown efficiency of CE evolution (Zorotovic et al. 2010; Davis et al. 2012; Toonen & Nelemans 2013), the mass boundary at which CE evolution terminates (the core-envelope boundary; Han et al. 1994; Dewi & Tauris 2000; Ivanova 2011), and the actual conditions for the onset of CE evolution (Hjellming & Webbink 1987; Soberman et al. 1997; Pavlovskii et al. 2017). Observational and theoretical constraints for the outcome of CE evolution in massive stars are also limited as the vast majority of observed post-CE systems as well as multidimensional hydrodynamical simulations of CE evolution correspond to low and intermediate-mass stars (see Iaconi & De Marco 2019 for a recent compilation of CE simulations and observed post-CE systems).

One uncertainty of CE evolution that is studied in population synthesis calculations is the impact of the evolutionary stage of the star at the onset of CE evolution. Belczynski et al. (2010) proposed that in low metallicity environments, a larger number of massive stars are expected to survive CE evolution, thus leading to an enhanced formation rate of merging binary BHs. This is argued as a consequence of halted expansion after core-hydrogen burning, with stellar models undergoing core-helium burning as blue supergiants and covering a larger range of radii after core-helium ignition. Belczynski et al. (2010) argued that interaction after core-helium ignition favors the ejection of the envelope as, at that point, a well defined core-envelope structure is formed. In various population synthesis codes, this argument is encoded in “pessimistic” and “optimistic” options for CE survival. In the pessimistic approach, all systems undergoing CE evolution before core-helium ignition are assumed to merge during CE evolution, while the optimistic approach allows for their survival. Recently, Klencki et al. (2020, 2021) have argued that this approach likely overestimates the number of merging binary BHs that formed as most of the systems predicted to survive CE in the optimistic scenario would initiate CE evolution with a radiative envelope that has a binding energy too large to eject through an inspiral.

The aim of this work is to study, in detail, how the evolutionary stage of the stellar progenitor of a BH affects MT stability and the outcomes of CE evolution. We do this by computing detailed models with the MESA code (Paxton et al. 2011, 2013, 2015, 2018, 2019) of a low metallicity massive star with a BH companion, meant to reproduce the evolution of a binary system after the formation of the first BH. Recent work on the formation of binary BHs using detailed binary evolution models (in contrast to rapid population synthesis calculations) has been presented by Eldridge & Stanway (2016) and Pavlovskii et al. (2017), covering a broad range of donor masses and metallicities. Our work, in contrast, focuses on a single donor mass of 30 $M_{\odot}$ at a metallicity of Z=10, but with a large resolution in the initial mass ratio and period. This allows us to study, in detail, how the stability of MT and the outcomes of CE evolution are affected by the evolutionary stage of the donor. For these calculations, we have made improvements to the commonly used MT prescription of Kolb & Ritter (1990, hereafter KR90) and implemented a method to model CE evolution which self-consistently determines the core-envelope boundary. Our methods are described in Sect. 2 and we present our results in Sect. 3. We conclude by discussing our results in Sect. 4. All input files necessary to reproduce our simulations as well as associated data products are available for download online.

2. Methods

We performed our simulations using version 15140 of the MESA stellar evolution code. Our models consist of binary systems with a 30 $M_{\odot}$ donor star and a BH companion. The objective of these simulations is to model a MT event happening after the formation of the first BH in the system, which is argued to be the most important MT phase where a CE can lead to the formation of a merging binary BH (cf. Dominik et al. 2012). Since we did not model the previous evolution of the system, we approximated the starting state of the donor star in the system as a zero-age main-sequence (ZAMS) star of its mass. Evolution was computed until either core carbon was depleted or until a merger between the donor star and the BH occurred.

Mass transfer rates were computed using the method described in Sect. 2.1, while the actual accretion rate into the BH was limited to its Eddington rate as described in Marchant et al. (2017). Our model for CE evolution is described in Sect. 2.2. Mass loss, either due to stellar winds from the donor or from mass accreted from the vicinity of the BH by MT above the Eddington limit, was assumed to take an amount of angular momentum corresponding to the specific orbital angular momentum of each component. We also accounted for the possibility of a star growing to the point that not only did it overfill its Roche lobe, but also overflowed its outer Lagrangian point. Whenever this happened, we accounted for the loss of mass and angular momentum as described in Sect. 2.1.5. Angular momentum loss from GW radiation was also included (Peters 1964), but it does not play a role in the timescales of evolution of our models which are of the order of millions of years. Our simulations do not take stellar rotation or spin-orbit coupling into account.

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Our stellar models were computed at a low metallicity of $Z = Z_\odot/10$, where we took the relative metal mass fractions from Grevesse & Sauval (1998) and the solar abundance to be $Z_\odot = 0.0142$ (Asplund et al. 2009). Opacities were computed using opacity tables from the OPAL project (Iglesias & Rogers 1996), together with the low temperature opacity tables of Ferguson et al. (2005). The equation of state used by MESA consists of a combination of OPAL (Rogers & Nayfonov 2002), HELM (Timmes & Swesty 2000), PC (Potekhin & Chabrier 2010), and SCVH (Saumon et al. 1995). Nuclear reaction rates were taken in order of preference from Cyburt et al. (2010) and Angulo et al. (1999).

Convection was modeled using the mixing length theory of Böhm-Vitense (1958) as described by Cox & Giuli (1968), using a mixing length parameter $\alpha = 2$. Convective regions were determined using the Ledoux criterion (Ledoux 1947). We included overshooting of the hydrogen burning convective core using a step-overshooting scheme, extending the size of the convective core by $\alpha_{\text{osc}} = 0.335$ pressure scale heights following the calibration of Brott et al. (2011). Overshooting from other convective regions is not well understood, so for convective cores after the main-sequence, we included a small amount of exponential overshooting (Herwig 2000) given by a length scale for exponential decay of the mixing coefficient of $f = 0.01^5$. We also included semi-convective mixing (Langer et al. 1983) and thermohaline mixing (Kippenhahn et al. 1980). Following Schootemeijer et al. (2019), we adopted a large value for the efficiency parameter for semiconvection $\alpha_{\text{osc}} = 100$, while for thermohaline mixing we adopted an efficiency parameter of unity.

Stellar winds were accounted for using a combination of different mass loss rate prescriptions as described by Brott et al. (2011). This includes the line-driven mass loss rates derived by Vink et al. (2001) for stars with a hydrogen surface mass fraction of $X > 0.7$ and the Wolf–Rayet mass loss rate of Hamann et al. (1995) for $X < 0.4$, scaled by a factor of 10 to account for wind clumping (Yoon et al. 2010). For surface hydrogen mass fractions between $X = 0.7$ and $X = 0.4$, we interpolated between the rate of Vink et al. (2001) and Hamann et al. (1995) to provide a continuous transition. At temperatures below that of the bi-stability jump (as derived by Vink et al. 2001), we took the maximum between the rate of Nieuwenhuijzen & de Jager (1990) and the one resulting from the combination of the Vink et al. (2001) and Hamann et al. (1995) rates we just described. Additionally, we scaled the Nieuwenhuijzen & de Jager (1990) rate by the same factor $(Z/Z_\odot)^{0.85}$ predicted by Vink et al. (2001) for line-driven winds. This significantly lowers the mass loss rates of red-supergiants in low metallicity environments, although observations suggest there is only a weak dependence on metallicity (van Loon et al. 2005; Goldman et al. 2017). Nevertheless, we kept this scaling for the sake of comparison, as it is commonly used in population synthesis calculations.

### 2.1. Mass transfer

We modeled MT using an extension of the method developed by KR90, which accounts for MT from an extended atmosphere as described in Ritter (1988), hereafter R88 as well as the case

3 In practice, since mixing length theory predicts no mixing at the edge of a convective region, overshooting in MESA is specified by a pair of variables $f$ and $f_0$, where $f_0$ determines a distance in units of pressure scale heights into the convective region, from which the mixing coefficient from overshooting is computed. For step overshooting, we took $f = 0.345$ and $f_0 = 0.01$, while for exponential overshooting we used $f = 0.01$ and $f_0 = 0.005$. where optically thick regions overflow the Roche lobe of the donor. In particular, this does not assume that the photosphere of the star operates as a hard rim, potentially leading to significant amounts of overflow during MT. Here we describe the method of KR90 in detail and our modifications to it, which include the possibility of outflows from the outer Lagrangian point of the donor. For the case of overflow from optically thick regions, we also discuss similarities between our method and that of Pavlovskii & Ivanova (2015).

#### 2.1.1. The Roche potential

The computation of the MT rate depends on the Roche potential $\Phi$, which is taken to be that of two point masses in a circular orbit:

$$
\Phi = \frac{G M_d}{a} \left( \frac{1}{r_d} - \frac{q}{r_a} - \frac{q + 1}{2} \left( \hat{x} - \hat{x}_{\text{cm}} \right)^2 + \hat{y}^2 \right),
$$

(1)

where $M_d$ is the mass of the donor and the mass ratio is defined as $q \equiv M_d/M_a$ with $M_a$ being the mass of the accretor. We used a coordinate system centered on the donor star (see Fig. 1), denoting distances normalized by the orbital separation $a$ as $\hat{x} = x/a$; $\hat{r}_d$ and $\hat{r}_a$ are the normalized distances of a point to the center of each star, and $\hat{x}_{\text{cm}} = q/(q + 1)$ indicates the $x$-coordinate of the center of mass of the system. The $x$-coordinate of the first Lagrangian point is given by $X_{L_1}$. The $x$-coordinate of the outer Lagrangian point of the donor (which can be either $L_2$ or $L_1$ depending on the mass ratio) is given by $X_{L_{out}}$, which is defined as being negative. The Roche lobe radius of the system $R_{\text{RL}}$ is defined as the volume equivalent radius of the region below the equipotential at the first Lagrangian point $\Phi_{L_1}$, and we used the fitting formula from Eggleton (1983) to approximate it:

$$
R_{\text{RL}} \equiv \frac{0.49 q^{2/3}}{0.6 q^{2/3} + \ln (q^{-1/3} + 1)},
$$

(2)
Fig. 2. Ratio between the volume equivalent radius associated with the outer Lagrangian point of the donor and its Roche lobe radius. Bottom panel: result from numerical integration of the Roche potential together with the fit given in Eq. (3). Top panel: relative error between the fit and the data, with the small variations being due to the finite precision of the volume calculation.

Since we considered stars with significant overflow, potentially reaching the outer Lagrangian point of the donor, we also defined volume-equivalent radii beyond the equipotential of $L_1$ by dividing space with a plane crossing $L_1$, which is perpendicular to the line joining both stars, as depicted in Fig. 1. We refer to this plane as the $L_1$ plane. The volume equivalent radius for an arbitrary value of $\Phi$ is then defined as $V(\Phi) = 4 \pi R(\Phi)^3/3$, where $V(\Phi)$ is the volume contained within the equipotential $\Phi$ and on the side of the $L_1$ plane where the donor is located. In particular, we care about the radius associated with the equipotential of the outer Lagrangian point of the donor, for which we find that the following equation provides a fit with an error <0.15% in the range of $-10 < \log_{10} q < 10$:

$$\frac{R_{\text{out}}}{R_{\text{RL}}} = 1 + \frac{2.74}{1 + [(\ln q + 1.02)/\sigma]^2} \times \frac{1}{7.13 + q^{0.386}}.$$  \hspace{1cm} (3)

Figure 2 shows the value of $R_{\text{out}}/R_{\text{RL}}$ for different values of the mass ratio. In the limit of $\log_{10} q \rightarrow \pm \infty$, we have that $R_{\text{out}}/R_{\text{RL}} \rightarrow 1$, while in the range of $-1 < \log_{10} q < 1$, which is typical of interacting binary stars, we find that $R_{\text{out}}/R_{\text{RL}}$ is between ~1.2 and 1.3. Thus, a donor star growing beyond ~20%–30% of its Roche lobe radius is expected to also fill its outer Lagrangian point.

2.1.2. Mass transfer through the $L_1$ point

Restricting the discussion first to flows through the $L_1$ point, the MT rate can be computed as an integral over the $L_1$ plane,

$$M_{\text{mt},L_1} = \int \rho v \, dA,$$  \hspace{1cm} (4)

where $\rho$ and $v$ are the density and velocity of the fluid in the $L_1$ plane. The flow is assumed to be steady in which case the Bernoulli equation is satisfied,

$$\frac{1}{2} v_i^2 + \int_i^f dP = \frac{1}{2} v_f^2 + \Phi_i,$$  \hspace{1cm} (5)

where the integral is done along a streamline of the fluid with $i$ and $f$ denoting an initial and final point along a streamline. Following the work of Lubow & Shu (1975), the velocity near the Lagrangian point is expected to be equal to the sound speed. Depending on whether the photosphere of the star is inside the Roche lobe or has expanded beyond it, the flow is assumed to be isothermal (R88) or adiabatic (KR90), respectively. Combining either the adiabatic or isothermal approximation with Eq. (5) allows for the computation of $\rho$ and $v$ in the vicinity of the Lagrangian point as a function of the Roche potential $\Phi$.

Either in the isothermal or adiabatic approximation, the surface integral in Eq. (6) can be expressed as an integral over the potential,

$$M_{\text{mt},L_1} = \int \rho v \frac{dA}{d\Phi},$$  \hspace{1cm} (6)

where $A(\Phi)$ is the area in the $L_1$ plane below the equipotential $\Phi$. Figure 1 illustrates the area $A(\Phi_{\text{out}})$. The $dA/d\Phi$ term describes how the area enclosed by an equipotential in the $L_1$ plane changes with $\Phi$ and encodes the properties of the Roche geometry. Along the $L_1$ plane, given the symmetry of $\Phi$ and the $L_1$ point being a local minimum, the Roche potential can be approximated as

$$\Delta \Phi = C_1 y^2 + C_2 z^2 + C_3 y^4 + C_4 y^2 z^2 + C_5 z^4,$$  \hspace{1cm} (7)

where $\Delta \Phi \equiv \Phi - \Phi_{L_1}$. As shown in Appendix A, $C_3 = C_5 = C_4/2$ and the value of $dA/d\Phi$ can be computed up to first order in $\Delta \Phi$ as

$$\frac{dA}{d\Phi} = \frac{\pi}{\sqrt{C_1 C_2}} \left(1 - \frac{C_3(C_2^2 + 2 C_1 C_2 + 3 C_2^2)}{4 C_1^2 C_2^2} \Delta \Phi + O[(\Delta \Phi)^2] \right).$$  \hspace{1cm} (8)

Both R88 and KR90 approximate $dA/d\Phi$ as a constant, but as we want to consider the case where the star can significantly overflow its Roche lobe, we included the term of order $\Delta \Phi$ when computing MT in the optically thick case. R88 and KR90 also rely on fits to the coefficients $C_1$ and $C_2$ as a function of the mass ratio. To avoid relying on fitting formulae with a limited range of validity, we directly computed the location of the $L_1$ point, $L_{1\text{L}}$, and computed all $C_1$ coefficients explicitly from the Roche potential\(^4\). The variation of $dA/d\Phi$ with increasing overflow has also been taken into account for optically thick Roche lobe overflow (RLOF) by Pavlovskii & Ivanova (2015), but using tables with numerical integrations of the area in the $L_1$ plane instead of a higher-order analytical approximation.

2.1.3. Optically thin mass transfer ($R_{\text{ph}} < R_{\text{RL}}$)

For the case of MT from a star with a photosphere radius $R_{\text{ph}}$ below its Roche lobe radius $R_{\text{RL}}$, R88 considers the flow to have

\(^4\) R88 defines a function $F(q)$ in their Eq. (A8), which contains the product $C_1 C_3$, and provides a fit for this function. The expression provided by R88 has a typo which is corrected in Eq. (A3) of KR90. Despite this typographical error in R88, we have verified that the fit to $F(q)$ provided in Eq. (A9) of R88 matches the correct expression.
a constant temperature equal to $T_{\text{eff}}$, in which case the relevant sound speed is the isothermal one,

$$v_{\text{th}}^2 = \frac{\partial P}{\partial \rho} \bigg|_T = \frac{kT_{\text{eff}}}{\mu m_\text{h}},$$  \hspace{1cm} (9)

where $k$ is the Boltzmann constant, $\mu$ is the mean molecular weight, and $m_\text{h}$ is one atomic mass unit. This sound speed corresponds to the equation of an ideal gas plus radiation,

$$P = \frac{\rho kT_{\text{eff}}}{\mu m_\text{h}} + \frac{\rho a_{\text{rad}} T_{\text{eff}}^4}{3},$$  \hspace{1cm} (10)

where $a_{\text{rad}}$ is the radiation constant. We considered streamlines that start with negligible velocity from the photosphere of the star by taking $v_1 = 0$ and $\Phi_1 = \Phi_{\text{ph}}$ in Eq. (5), where $\Phi_{\text{ph}}$ is the value of the Roche potential at the photosphere. Combining Eqs. (5), (9), and (10), one can solve for the density of the flow at the L$_1$ point,

$$\rho_{\text{L1}} = \rho_{\text{ph}} \frac{\left(\Phi_{\text{ph}} - \Phi_{\text{L1}}\right)}{v_{\text{th}}^2},$$  \hspace{1cm} (11)

where the ph and L1 subscripts indicate properties at the photosphere and the Lagrangian point.

Equation (11) can be evaluated in the vicinity of the L$_1$ point to express the variation in density across the L$_1$ plane as a function of the variation in $\Phi$ and $v$,

$$d\rho = -\rho_{\text{L1}} \frac{v_{\text{th}}}{v_1} \left(\frac{d\Phi}{v_1} + \frac{\left(\rho_{\text{ph}} - \rho_{\text{L1}}\right)}{v_{\text{th}}} d\sqrt{v_1^2} \right).$$  \hspace{1cm} (12)

As for a steady flow, the L$_1$ point operates as a nozzle, and acceleration perpendicular to the flow is expected to vanish on the L$_1$ plane,

$$dP = -\rho d\Phi \rightarrow d\rho = \frac{\rho}{v_1} \frac{d\Phi}{d\sqrt{v_1^2}}.$$  \hspace{1cm} (13)

which combined with Eq. (12) results in $dv = 0$ (and thus $v = v_{\text{th}}$) in the vicinity of L$_1$.

These results can be used to evaluate the MT rate using Eq. (6). Transforming the integral into one over the density with Eq. (13) results in

$$M_{\text{thin}} = -\int_{\rho_{\text{L1}}}^{\rho_{\text{ph}}} \frac{\sqrt{v_1^2}}{d\Phi} d\rho.$$

As the density decays exponentially with the potential, see Eq. (11), only the regions in the very vicinity of L$_1$ are expected to contribute to the MT rate, so we approximated $dA/d\Phi$ by its value at L$_1$ to obtain

$$M_{\text{thin}} = M_{0\text{thin}} \exp \left(\frac{\Phi_{\text{ph}} - \Phi_{\text{L1}}}{v_{\text{th}}^2}\right),$$

where $M_{0\text{thin}}$ is the MT rate for a star just filling its Roche lobe,

$$M_{0\text{thin}} = \frac{\rho_{\text{ph}} v_{\text{th}}^3}{\sqrt{V/L_{\text{L1}}}}.$$  \hspace{1cm} (16)

The factor that is still required to compute the MT rate in an evolutionary code is the difference $\Phi_{\text{ph}} - \Phi_{\text{L1}}$, R88 opted to use the derivative of the potential with respect to volume equivalent radii $r_v$ covered by different equipotentials and approximated

$$\Phi_{\text{ph}} - \Phi_{\text{L1}} = \left(\frac{d\Phi}{dr_v}\right)_{L_{\text{L1}}} \left(P_{\text{LL}} - P_{\text{ph}}\right).$$  \hspace{1cm} (17)

Instead, we followed the method of Jackson et al. (2017) who developed an expansion of the Roche potential around the donor as a function of the equivalent radius $r_v$,

$$\Phi(r_v) = -\left(\frac{GM_s}{a} + \frac{GM_a}{2a(M_d + M_s)}\right) - \frac{GM_d}{r_v} \left[1 - a_1 \left(\frac{r_v}{a}\right)^3 + a_2 \left(\frac{r_v}{a}\right)^6 + O\left(\frac{r_v}{a}\right)^9\right],$$

with $a_1$ and $a_2$ given by

$$a_1 = \frac{1}{3} \left(\frac{M_d + M_s}{M_s}\right),$$

$$a_2 = \frac{4}{45} \left(\frac{(M_d + M_s)^2 + 9M_s^2 + 3M_d(M_d + M_s)}{M_s^3}\right).$$

Using this, we computed $\Phi_{\text{L1}}$ with the Roche lobe radii from Eq. (2) and computed $\Phi_{\text{ph}}$ using the photospheric radius of our stellar model.

In all our simulations, only a small fraction of the total MT comes from the contribution of this thin MT rate. However, it still operates as a physically motivated mechanism to smoothly turn on MT which helps prevent numerical instabilities in the calculations when a star initiates RLOF.

### 2.1.4. Optically thick mass transfer ($R_{\text{ph}} > R_{\text{RL}}$)

Following KR90, if layers of the star below the photosphere are overflowing the Roche lobe, then the MT rate is computed as

$$M_{\text{mt,th}} = M_{0\text{th}} + M_{\text{thick}},$$

where $M_{\text{thick}}$ is the contribution to the integral in Eq. (6) from the overflowing, optically thick regions,

$$M_{\text{thick}} = \int_{\Phi_{\text{thick}}}^{\Phi_{\text{ph}}} \frac{\rho dA}{d\Phi}.$$  \hspace{1cm} (19)

To calculate the density and velocity of the flow at L$_1$, we again made use of the Bernoulli equation, Eq. (5), by assuming the flow moves parallel to the equipotential surfaces and is adiabatic such that the pressure and sound speed are given along a streamline by

$$P = k\rho^{\Gamma_1}, \hspace{1cm} v^2 = \frac{\Gamma_1 P}{\rho}.$$  \hspace{1cm} (22)

The constant $k$ can be computed in terms of a reference pressure and density for each streamline, $k = \rho_0^{1/\Gamma_1}$. This assumes that $\Gamma_1$ is constant along the streamline; Pavlovskii & Ivanova (2015) considered the case of a variable $\Gamma_1$ and found that the resulting MT rates are only modified by a few percent.

We assumed that along each equipotential surface, far from the Lagrangian point, the fluid is near hydrostatic equilibrium ($dP_0 \approx -\rho d\Phi$ and $v \ll v_{\text{th}}$) and took the value of $P_0$ and $\rho_0$ there to compute $k$. Combining Eq. (22) with Eq. (5), and taking $\Phi_{\text{thick}} = \Phi_{\text{ph}}$, $v_1 = 0$ and $v_{\text{th}}$, one can compute the density of the flow in the L$_1$ plane for a given value of $\rho_0$,

$$\rho = \rho_0 \left(\frac{2}{\Gamma_1 + 1}\right)^{1/(\Gamma_1 - 1)}.$$  \hspace{1cm} (23)

Combining this density with Eq. (21) gives the contribution to the MT rate from the overflowing optically thick layers as

$$M_{\text{thick}} = \int_{\rho_0}^{\rho_{\text{thick}}} \rho^{1/2} \left(F_3(\Gamma_1) \frac{dA}{d\Phi} \right) dP_0.$$  \hspace{1cm} (24)
where

$$F_\Gamma (\Gamma_1) = \Gamma_1^{1/2} \left( \frac{2}{\Gamma_1 + 1} \right)^{(\Gamma_1 + 1)/(2(\Gamma_1 - 1))} \quad (25)$$

In our calculations, we considered $P_0$ and $\rho_0$ to be the density in the overflowing layers of our hydrostatic model, from which we also obtained $\Gamma_1$. In this case, $P(R_{RL})$ and $P(R_\text{orb})$ are the pressure of the stellar model in the layer were $R = R_{RL}$ and $R = R_\text{orb}$, respectively.

Equation (24) is exactly as derived by KR90, but in our calculations there are two important differences as to how we computed it. First, KR90 made the assumption of an ideal gas, replacing the $P_0/\rho_0$ ratio in Eq. (24) by $kT/\mu m_b$. In layers dominated by radiation pressure, this underestimates the mass loss rate significantly, so we did not make this assumption and computed $P_0/\rho_0$ from our stellar model. The second difference is that we did not assume $dA/d\Phi$ to be constant, but instead evaluated it using Eq. (8), with $d\Phi$ being computed in the overflowing layers of the model as

$$d\Phi = \int_{P(R_{RL})}^{P} \frac{dP_0}{\rho_0}. \quad (26)$$

### 2.1.5. Mass loss from the outer Lagrangian point of the donor

Models of interactive massive stars using the prescription of KR90 can exhibit radii much larger than their Roche lobe radii. In particular, in the simulations presented in this work involving a 30 $M_\odot$ donor, we find cases where the outer Lagrangian point would overflow ($R > R_{\text{out}}$). For these cases, we need a model for the overflow from this outer Lagrangian point, which is $L_3$ if $M_d > M_a$ and $L_2$ otherwise.

The entire method described in Sects. 2.1.3 and 2.1.4 can be applied for an outflow through the outer Lagrangian point by replacing the Roche lobe radius with $R_{\text{out}}$ using Eq. (3) and evaluating $dA/d\Phi$ factor at the outer Lagrangian point instead of $L_1$. However, an assumption needs to be made as to what happens to this outflow. We assume the material is ejected from the system carrying the specific angular momentum that corresponds to the outer Lagrangian point,

$$j_{\text{out}} = \frac{2\pi}{P_{\text{orb}}} a^2 \left( \frac{M_a}{M_a + M_d} - \dot X_{\text{out}} \right)^2. \quad (27)$$

In order for this to be satisfied, the outflow needs to have a velocity much larger than the orbital velocity, otherwise tidal forces from the binary can modify the angular momentum content of ejected material. This has been studied in the case of overcontact binary systems that overflow the $L_2$ point (Shu et al. 1979; Pejcha 2014; Pejcha et al. 2016) or have their stellar winds torqued as they accelerate (MacLeod & Loeb 2020). If the energy of the outflowing material is not sufficient to unbind it from the binary, it can also accumulate in a circumbinary disk.

Owing to these uncertainties, we also experimented with models where the specific angular momentum in Eq. (30) is increased by factors of 4 and 9, corresponding to points corotating with the orbit at 2 and 3 times the distance of $L_{\text{out}}$ from the center of mass. However, any of our simulations that undergo stable MT while overflowing $L_2$ or $L_3$ should be considered with care, since this is an extreme regime where three-dimensional hydrodynamic effects become important.

### 2.1.6. Outflows from $L_2$ when $M_d/M_a < 1$

An alternative approach to the problem of large overflow in simulations was taken by Misra et al. (2020). Considering systems with neutron star accretors, they computed the equivalent radii associated with the full $L_2$ equipotential (including both the donor and the accretor), and they assumed that whenever a stellar model exceeded this radius, then MT becomes unstable owing to the formation of an outflow with a high specific angular momentum. However, since for a compact object accretor there is not a hydrostatic structure within the Roche lobe of the accretor, the volume associated with the full $L_2$ equipotential cannot be directly compared to the radii of a hydrostatic stellar model in order to assess overflow of the outer Lagrangian points.

Nevertheless, in cases where $M_d > M_a$, some of the material streaming through $L_1$ could potentially be ejected from the vicinity of $L_2$ before the donor overflows $L_3$. In this case, our simulations would underestimate angular momentum loss from the system, since the specific angular momentum associated with $L_2$ cannot be up to 47% larger than that of $L_3$, as is shown in Fig. 3. Figure 3 also indicates the volume equivalent radii for the donor and the first three Lagrangian points as a function of the mass ratio. Bottom: volume equivalent radii for the donor corresponding to the $L_2$ and $L_3$ equipotentials. Volume equivalent radius for $L_2$ is taken from the fit of Marchant et al. (2016), which due to small errors in the fit results in a slightly larger value than the radius for the $L_3$ equipotential at $q = 1$.

$$R_{\text{L}} = 1 + 0.299 \tan^{-1}(1.84q^{0.397})q^{0.520}. \quad (28)$$

Although for a broad range of mass ratios, the donor can expand significantly before filling out the $L_2$ equipotential, much less expansion beyond the Roche lobe is required to fill out $L_2$.

In order to assess potential uncertainties during phases of large overflow, where material streaming through $L_1$ could be ejected from the vicinity of $L_2$ when $M_d/M_a < 1$, we also considered an alternate model. In this model, instead of accounting for overflows from the donor’s outer Lagrangian point, we assumed part of the mass streaming through $L_1$ is ejected from $L_2$. To
do this, we computed an additional optically thin rate from Sect. 2.1.3 by replacing $R_{\text{RL}}$ with $R_{L_2}$ and evaluating $dA/d\Phi$ at $L_2$ rather than at $L_1$. Whenever $R_{\text{th}} > R_{L_2}$, we considered the contribution to the optically thick mass transfer rate transferred through the $L_1$ plane, but above the equipotential of $L_2$, which is given by Eq. (24) with a modified lower interval for the integral:

$$M_{\text{th}, \text{above} L_2} = \int_{P_{R_{L_2}}}^{P_{R_{th}}} \left( \frac{P_0}{P_i} \right)^{1/2} F_2(\Gamma) \frac{dA}{d\Phi} dP_0. \quad (29)$$

Since this expression describes a flow through $L_1$, $dA/d\Phi$ has to be evaluated at $L_1$. We assumed both the optically thin component through $L_2$ and the optically thick component through $L_1$ above $L_2$ equipotential are ejected from the system, taking the specific angular momentum that corresponds to the $L_2$ point,

$$j_{L_2} = \frac{2\pi}{P_{\text{orb}}} a^2 \left( \frac{M_a}{M_a + M_d} - \tilde{X}_{L_2} \right)^2. \quad (30)$$

Similar to what was discussed in the previous section, material could be torqued or form a circumbinary disk before being ejected, so we also consider extreme cases where we increased the specific angular momentum removed by factors of 4 and 9.

The default setup of our simulations is the one described in Sect. 2.1.5, but we also performed simulations with this modified model with $L_2$ outflows to test the robustness of our results. For simplicity, in the remainder of the manuscript we do not use the subscript 0 to refer to properties from our hydrostatic stellar models and refer to the photospheric radius of a star as $R$.

### 2.2. Common envelope evolution

Whenever our MT prescription gives a MT rate exceeding a given threshold $M_{\text{th}}$, we assume evolution proceeds through a CE phase. We treated CE evolution following the standard energy prescription (Tutukov & Yungelson 1979; Webbink 1984) which equates the binding energy of ejected layers to the difference in orbital energy product of an inspiral,

$$E_{\text{bind}} = \alpha_{\text{CE}} \Delta E_{\text{orb}}, \quad (31)$$

where $\alpha_{\text{CE}}$ is a free parameter that represents the efficiency with which the orbital energy ejects the envelope. Using subscripts $i$ and $f$ to represent the pre- and post-CE properties of the system, the difference in orbital energy is

$$\Delta E_{\text{orb}} = -\frac{GM_{\text{af}} M_{\text{sf}}}{2a_i} + \frac{GM_{\text{df}} M_{\text{ia}}}{2a_i}, \quad (32)$$

and we considered $M_{\text{af}}$ to be the core mass of the donor $M_{\text{core}}$, while ignoring accretion into the accretor such that $M_{\text{sf}} = M_{\text{ia}}$. The binding energy depends on the value of $M_{\text{core}}$ and was computed by adding up the internal and gravitational potential energy of the removed layers at the onset of CE,

$$E_{\text{bind}} = \int_{M_{\text{core}}}^{M_{\text{af}}} \left( -\frac{Gm}{r} + \alpha_{\text{th}} u \right) dm, \quad (33)$$

where $u$ is the specific internal energy of the gas and we included an additional free parameter, $\alpha_{\text{th}}$, which was introduced by Han et al. (1995) to represent the efficiency with which thermal energy can be used to eject the envelope. In this work, we include the contribution of recombination energy from hydrogen and helium on $u$. For simplicity, we also assume $\alpha_{\text{th}} = 1$ throughout.

Computing the binding energy requires one to know $M_{\text{core}}$, which is a nontrivial problem that can lead to large variations in the predicted binding energy (see for instance the discussion in Ivanova et al. 2013). Our goal is to determine $M_{\text{core}}$ self-consistently in our simulations by modeling the stripping of the stellar envelope. For this, at the onset of CE, we computed the value of $E_{\text{bind}}$ for all choices of $M_{\text{core}}$ in the stellar model. The CE phase was then modeled by artificially removing mass from the star and computing, at each step, the binding energy one would have obtained from the pre-CE model if $M_{\text{core}}$ were assumed to be the current mass of the star. In this way, Eq. (32) can be used to determine the final orbital energy product of an inspiral, the mass losing stellar model can be used to determine the point at which the star would contract inside its Roche lobe.

We did not simply force a rapid mass loss rate on our simulations until we found $R < R_{\text{RL}}$, but instead softly turned off the mass loss rate as the star went inside its Roche lobe to determine the mass coordinate at which the star would be undergoing contraction in the absence of mass loss. In particular, we forced a mass loss rate from CE evolution of

$$\log_{10} M_{\text{CE}} = \begin{cases} \log_{10} M_{\text{high}} \\ \log_{10} M_{\text{high}} + \frac{1 - \delta}{\delta} \log_{10} \left( \frac{M_{\text{th}}}{M_{\text{th, high}}} \right) \end{cases} \quad \frac{\dot{\rho}}{\rho_{\text{ref}}} > 1, \quad 1 - \delta < \frac{\dot{\rho}}{\rho_{\text{ref}}} < 1, \quad (34)$$

where we interpolated between a high MT rate $M_{\text{high}}$ meant to reproduce a near adiabatic response, and a small MT rate, $M_{\text{low}}$, which should be well below the thermal timescale of the envelope. In particular in our simulations, we chose $M_{\text{high}} = M_{\text{th}}$, $M_{\text{low}} = 10^{-5} M_{\text{th}}$, and $\delta = 0.02$. Our choice of $M_{\text{high}}$ was based on the thermal timescale of the envelope estimated as $\tau_{\text{th}} = E_{\text{bind}}/L$ and the associated MT rate $M_{\text{th}} = M/\tau_{\text{th}}$. For the 30 $M_{\odot}$ donor considered in this work evolved as a single star, and assuming a core-envelope boundary at the innermost point where $X > 0.1$ (Dewi & Tauris 2000), we find that $M_{\text{th}}$ does not exceed 0.1 $M_{\odot}$ yr$^{-1}$. Similarly, our choice of $M_{\text{low}}$ is such that it is comparable to MT happening on a nuclear timescale. The dependence of our results on the free parameters $M_{\text{high}}, M_{\text{low}}$, and $\delta$ is discussed in Appendix B, where we show how the outcome of some of our CE models are affected by these choices.

If the star contracts below $R/R_{\text{RL}} = 0.98$, then we assumed the CE phase finished and proceeded with the evolution of the model in the standard way, applying our MT formalism described in the previous subsections. As pointed out by Ivanova (2011), a star can significantly contract during a phase of rapid mass loss, but re-expand by orders of magnitude on a few thermal timescales if mass loss is suddenly shut off. Because of this, they determined the core mass of a stellar model by performing rapid mass loss simulations and finding at which mass coordinate the resulting star would not re-expand after mass loss is shut off. In our simulations, we did find re-expansion after CE, which, according to Ivanova (2011), would indicate that the core boundary is at a deeper mass coordinate. However, we find that the ensuing MT phase is stable when using our MT prescription.

This method is similar to the one presented by Deloye & Taam (2010) and Ge et al. (2010), who computed the adiabatic response of the stellar radii to rapid mass loss and compared it to the evolution of the Roche lobe radii in a binary to determine at which point a binary would detach. The main difference is that our method determines the mass coordinate at which detachment would happen without mass loss, capturing the nonadiabatic response of the star near the end of CE. Another similar method is the one presented by
Eldridge et al. (2017), which was integrated into detailed binary evolution calculations but does not consider the contribution of thermal or recombination energy.

3. Results

In this work, we focus on the evolution of a 30 $M_{\odot}$ star at a metallicity of $Z_{\odot}/10$, separated into different evolution phases. Dots indicate equal time intervals of $10^5$ years. The Hertzsprung gap is defined as the evolutionary stage between the terminal-age main sequence and the moment where the ratio of the nuclear burning luminosity and the surface luminosity satisfies $\log_{10}(L_{\text{nuc}}/L) < 0.001$ for at least a thermal timescale estimated as $GM^2/R L$, or the star develops a convective envelope $> 1 M_{\odot}$. Top: evolution under our default set of physical inputs. Middle: evolution with core overshooting increased from 0.335 to 0.535 pressure scale heights. Bottom: evolution with wind mass loss rates tripled.

If one computes the binding energy using, for example, the fit provided by Claeyts et al. (2014) rather than the structure of the model, the resulting binding energy is always underestimated, with more that an order of magnitude difference in late stages. In particular, this fit to the binding energy is used in population synthesis calculations of GW sources, including the MOBSE (Giacobbo & Mapelli 2018) and COSMIC (Breivik et al. 2020) codes. Another parametrization of the binding energy that is used in published rapid population synthesis results is that of Xu & Li (2010), which is an option in the STARTRAK (Dominik et al. 2012) and COMPAS (Neijssel et al. 2019) codes. In this case, we also find a consistent underestimation of the binding energy.

Before discussing grids of binary simulations, it is instructive to study how our 30 $M_{\odot}$ donor evolves as a single star. The top panel of Fig. 4 shows the evolution in the Hertzsprung-Russell (HR) diagram for this star under the physical assumptions described in Sect. 2, for which we find that expansion is halted before becoming a red supergiant. Most of the helium-burning phase of this star would be spent as a blue supergiant with a convective envelope developing only near core helium depletion. This type of evolution is sensitive to physical assumptions in the model. In particular, Fig. 4 shows the effect of increasing overshooting during core hydrogen burning, as well as increasing wind strengths. With increased overshooting, the star undergoes rapid expansion after the main sequence and becomes a red supergiant before core helium ignition. The model with a higher mass loss rate slows down its expansion before becoming a red supergiant, but it expands to a larger radius during the HG than our default model. These variations are not captured in rapid population synthesis codes, in particular those reliant on the fits to single star evolution of Hurley et al. (2000) which are based on detailed stellar models without mass loss and with a fixed amount of overshooting. Stellar winds may be overestimated in our calculations (Björkklund et al. 2021), while observations of massive stars suggest that our choice for overshooting from a convective hydrogen-burning core is underestimated for stars with masses $> 15 M_{\odot}$ (Castro et al. 2014). Keeping in mind that due to uncertainties in stellar evolution, a halted expansion in the HG is uncertain, we can still study what would be the consequences in the evolution of a star if this effect is real.

Focusing only on our model that uses our default set of assumptions, we assessed if core helium ignition has a significant impact on the core-envelope structure of the star which could favor CE ejection. Figure 5 shows a Kippenhahn diagram for the 30 $M_{\odot}$ model, indicating how the stellar radius and binding energy of the envelope evolve. Assuming that the core-envelope boundary is located at the points where the hydrogen mass fraction drops below either $X = 0.1$ or $X = 0.3$ (similar to Dewi & Tauris 2000), we find that the binding energy of the star remains at $\sim 10^{50}$ erg from terminal-age main sequence (TAMS) to the point where a convective envelope is developed. This happens despite a $\sim 30$ fold increase in the stellar radius. This model experiences the formation and merger of multiple intermediate convection zones during the HG phase, so we performed convergence tests to verify that our results are not significantly affected by changes in temporal and spatial resolution during this phase (see Appendix D).

If one computes the binding energy using, for example, the fit provided by Claeyts et al. (2014) rather than the structure of the model, the resulting binding energy is always underestimated, with more that an order of magnitude difference in late stages. In particular, this fit to the binding energy is used in population synthesis calculations of GW sources, including the MOBSE (Giacobbo & Mapelli 2018) and COSMIC (Breivik et al. 2020) codes. Another parametrization of the binding energy that is used in published rapid population synthesis results is that of Xu & Li (2010), which is an option in the STARTRAK (Dominik et al. 2012) and COMPAS (Neijssel et al. 2019) codes. In this case, we also find a consistent underestimation of the binding energy.
except for late stages near core-carbon burning where the envelope becomes almost fully convective. Both Claeys et al. (2014) and Xu & Li (2010) include internal energy in their fits, but not recombination energy. A systematic underestimation of the binding energy in population synthesis calculations implies an overestimation of post-CE orbital separations, or a prediction of successful envelope ejection in systems that would merge during CE. We also compared the binding energies of our models with different overshooting and wind strengths to the results obtained from the Claeys et al. (2014) and Xu & Li (2010) fits, and find that the fits consistently underestimate the binding energy of the envelope as well (see Fig. 6). Binding energies computed using the Xu & Li (2010) fit experience a jump when the star transitions out of the HG, which for the track with extra overshooting does not happen until the star begins to rise in the Hayashi line.

The constancy of the binding energy while the star expands with a radiative envelope is also illustrated by Fig. 7, which shows the full binding energy profiles at different evolutionary phases. In particular, between the end of the HG and the point where a deep convective envelope starts to develop, the radius of the star increases by more than an order of magnitude, but this expansion only happens in the outermost mass layers of the star. It is only after the formation of the convective envelope that a large fraction of the envelope mass is pushed to large radii with correspondingly low binding energies that would make envelope ejection during CE feasible. All of this points to the ignition of helium at the stellar core having no impact on the binding energy of the envelope, and also since the ignition of helium does not modify the mode of energy transport in the outer layers, the stability of MT should not be impacted in a fundamental way either.

The near constancy and high value of the binding energy before the formation of a convective envelope has also been studied by Kruckow et al. (2016) and Klencki et al. (2021) comprehensively, covering a large range of masses and metallicities. However, the binding energies computed by Kruckow et al. (2016) and Klencki et al. (2021) as well as those shown in Fig. 5 rely on an assumed boundary for the core mass. By performing binary simulations using the method described in Sect. 2.2 to determine the core-envelope boundary self-consistently, we can verify if these computed binding energies with an assumed boundary are a good approximation.

3.1. Grid of simulations

Using our prescription for MT and our method to calculate the outcome of CE evolution, we computed a grid of models consisting of a 30 $M_\odot$ donor with a BH accretor in a circular orbit. Our method for CE evolution still leaves the efficiency $\alpha_{\text{CE}}$ as a free parameter, and we computed simulations with both $\alpha_{\text{CE}} = 1$, 0.4, 0.2, and 0.1. The grid spans a period range between $-0.3 < \log_{10}(P_i/d) < 3.5$ and initial mass ratios $q = M_{\text{BH}}/M_{i,j}$ between 0.01 and 0.59. This includes BHs with masses as low as 0.3 $M_\odot$ that are not expected to form through stellar evolution, but they were included to test the limits of MT stability. This allowed us to cover the entire parameter space where CE happens, as well as to cover systems from the boundary of RLOF at ZAMS, to effectively single star evolution at large periods. Owing to the small amount of radial expansion occurring in the phases where the star has a convective envelope, we adopted two different resolutions in the period for our grid. For $\log_{10}(P_i/d) < 3.2$, we used a spacing of $\Delta \log_{10}(P_i/d) = 0.1$, while at higher initial orbital periods we increased the resolution

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**Fig. 5.** Kippenhahn diagram for our default 30 $M_\odot$ model, separating different evolutionary stages as well as showing the evolution of the stellar radii. “He dep.” and “C dep.” stand for core helium and carbon depletion respectively. In addition, we show the result of computing $E_{\text{bind}}$ under the assumption that the core–envelope boundary $M_{\text{crit}}$ at which CE would finish is at the innermost mass coordinates where the hydrogen mass fraction is $X = 0.1$ or $X = 0.3$. We only show $E_{\text{bind}}$ after the formation of the helium core, so no information is given on the main sequence. To provide a comparison point to rapid population synthesis codes, we also show the resulting $E_{\text{bind}}$ from the fits of Claeys et al. (2014) and Xu & Li (2010).
in the period to $\Delta \log_{10}(P_i/d) = 0.01$. The resolution in mass ratio of the grid is $\Delta q = 0.02$.

The final outcomes of our grid with $\alpha_{CE} = 1$ are shown in Fig. 8. As a simple approximation to determine if a simulation can produce a merging binary BH, for stars that reach core carbon depletion we assumed the donor star collapses directly to form a BH with a mass equal to its baryonic mass. Our $30 M_\odot$ donor produces final helium core masses $\sim 14 M_\odot$, for which the formation of a BH through direct collapse is indeed expected (Fryer 1999). From our simulations, we did not find any cases of successful envelope ejection from CE evolution for initial orbital periods below $\log_{10}(P_i/d) < 3.2$. Below this period range, the boundary between systems undergoing stable MT and those merging during CE evolution varies continuously with the period, going from $q \sim 0.5$ for systems interacting close to ZAMS, down to $q \sim 0.15$ for cases where interaction happens right before the formation of a convective envelope. The boundary is not sensitive to whether an interaction happens before or after the TAMS, or before or after the end of the HG. This is to be expected as the radial response to rapid mass loss is dependent on the mode of energy transport in the stellar envelope (Hjellming & Webbink 1987), and the envelope of the star across all these phases is radiative. Pavlovskii et al. (2017) also find stable MT between a star and a BH for such extreme mass ratios for different donor masses, but owing to their coarse mass ratio sampling (no more than three different mass ratios for each donor mass), they cannot describe in detail how the boundary for stability shifts with the initial orbital period.

The lack of systems surviving CE evolution between the end of the HG and the formation of a convective envelope can be understood in terms of the binding energy of the envelope. As was discussed for the single $30 M_\odot$ model, the binding energy during this phase is $\sim 10^{50}$ erg. Considering evolution between the end of the HG and the formation of a convective envelope, we find that CE evolution only happens for BH masses $\lesssim 5 M_\odot$, and that stripping the hydrogen envelope of the $30 M_\odot$ star would result in a $\sim 14 M_\odot$ helium core. For these masses and assuming a fully efficient use of the orbital energy to eject the envelope, in order to release $10^{50}$ erg, the orbital separation needs to be reduced below $1.3 R_\odot$, an orbit for which a helium star of $14 M_\odot$ would overfill its Roche lobe. Population synthesis calculations that underestimate the binding energy during this phase and find successful CE ejection, for example, by using fits such as those of Claeys et al. (2014) or by assuming a fixed value of the $\lambda$ parameter, potentially overestimate the rate of formation of merging binary BHs through CE evolution (Gallegos-Garcia et al., in prep.).

The boundary for unstable MT is also right at the place where it allows for a narrow band of systems undergoing stable MT that produce a binary BH merging in less than a Hubble time. In this case, the shrinkage of the orbit from stable MT is sufficient to produce a compact BH binary (van den Heuvel et al. 2017), and we find this outcome for a wide range of initial orbital periods ranging from 1 to 1000 days. Multiple systems undergoing stable MT fill their outer Lagrangian point during that process, but remain stable despite the increased angular momentum loss. For wider orbital periods, where MT occurs after the development of a deep convective envelope, the critical mass ratio for stability shifts upwards (up to $q \sim 0.5$ for systems interacting after core helium depletion). The majority of our models with an assumed efficiency of $\alpha_{CE} = 1$ that manage to eject their envelopes during CE evolution result in wide BH binaries, for which GW radiation is insufficient to produce a merger in less than a Hubble time. Figure 9 shows the outcome of our simulations with CE efficiencies of $\alpha_{CE} = 0.4, 0.2,$ and 0.1, in which case we find that most systems that produce wide binary BHs at $\alpha_{CE} = 1$ would...
merge during CE evolution rather than produce merging binary BHs at lower efficiencies.

One important concern in our simulations that form merging binary BHs through stable MT is that an important fraction of these undergo overflow of $L_2$ or even $L_1$. The methods we have developed to account for this are necessarily an approximation, as they rely on hydrostatic one-dimensional stellar models. Moreover, the simulations shown in Fig. 8 do not account for potential outflows through $L_2$ when $M_2 < M_{\Delta}$, which could remove significant mass and angular momentum from the system. Using the alternate method described in Sect. 2.1.6 for $L_2$ outflows, we can test if the systems that form merging binary BHs through stable MT are sensitive to the potential presence of such outflows. Figure 10 shows simulations done using this method that cover the region where we find that stable MT leads to merging binary BHs, including cases where we assume that the specific angular momentum removed by material outflowing $L_2$ is up to nine times the specific angular momentum of the $L_2$ point. Compared to our simulations that only account for overflow of the outer Lagrangian point of the donor, we find that the boundary for instability is pushed to higher BH masses. However, we still find systems across a large range of periods that form merging binary BHs, as additional angular momentum losses lead to the formation of more compact binaries post-MT. Given this, the conclusion that stable MT can lead to the formation of merging binary BHs across a large range of periods appears to be robust.

As our models do not include the evolution prior to the formation of the first BH, we cannot assess the relative likelihood of each of our individual simulations to occur in nature. However, if we make the simplistic assumption that after BH formation the orbital period is distributed flat in $\log_{10} P$ and the mass ratio distribution is flat in $q$, our grid with $\alpha_{CE} = 1$ gives a ratio between the number of merging binary BHs that formed through CE evolution to those that formed via stable MT of 0.017. The actual outcome of our simulations for systems surviving CE evolution should be considered with care, since we approximated a complex three-dimensional phenomenon with one-dimensional hydrostatic models. However, even under the extreme case that all our systems that survive CE evolution would produce a binary BH that merges in less than a Hubble time, we find that the ratio between merging binary BHs that formed through CE evolution to those that formed via stable MT is 0.35. This resembles the results from population synthesis calculations of Neijssel et al. (2019), who find that depending on the metallicity distribution of the star formation history, most merging binary BHs that will be observed by current generation GW detectors at design sensitivity would have been formed through stable MT. Our results suggest that the yield of merging binary BHs from CE evolution is overestimated at low metallicities, which would further increase the relative contribution of stable MT relative to CE evolution in population synthesis calculations such as those of Neijssel et al. (2019).

### 3.2. Mass transfer and outer Lagrangian point overflow

We consider here two cases from our simulation grid that lead to the formation of a merging binary BH: a system interacting before the end of HG ($M_{BH} = 7.5 M_\odot$, $P_1 = 31.6$ days) and one interacting after it ($M_{BH} = 4.5 M_\odot$, $P_1 = 1000$ days). The short period system evolves without overflowing its outer Lagrangian point, while the long period one is expected to overflow it, so we can use it to study how the resulting outcome is modified when accounting for this on our MT prescription. To evaluate the impact of the changes we have made to the MT prescription of (KR90), we used four different setups for our simulations:

1. A full implementation of our MT prescription including overflows from the outer Lagrangian point as described in Sect. 2.1.5.
2. Same as 1, but without the assumption $P/\rho = kT/\mu m_H$ made in KR90 (see the discussion in Sect. 2.1.4)
3. Same as 1, but considering the area term $dA/d\Phi$ to be constant and ignoring the term linear in $\Delta \Phi$ of Eq. (8).
4. Using the prescription of KR90 without modification and without outflows from the outer Lagrangian point.

Figure 11 shows the evolution in the HR diagram for both the $P_1 = 31.6$ day and $P_1 = 1000$ day system using our updated MT prescription. In both cases, MT first proceeds rapidly, stripping the majority of the hydrogen envelope in a few thousand years, followed by a longer phase ($\sim 10^7$ yr) driven by the evolution of the core during core-helium burning. In both cases, MT
Fig. 9. Same as Fig. 8, but for CE efficiency parameters of $\alpha_{\text{CE}} = 0.4, 0.2,$ and 0.1.

Fig. 10. Same as Fig. 8, but for a subset of models computed using the method described in Sect. 2.1.6 that considers outflows from L$_2$ rather than L$_1$ when $M_s < M_d$. Each panel indicates simulations performed with different specific angular momentum assumed for the material outflowing through L$_2$, which is considered to be $j_{L2}$, $4j_{L2}$, or $9j_{L2}$. The purple dotted line indicates the boundary between stable MT and CE evolution as determined by our models that account only for outflows from the outer Lagrangian point of the donor.

Fig. 11. Evolution in the HR diagram for a $30 M_\odot$ donor star in two systems leading to the formation of a merging binary BH through stable MT. Dots in the tracks indicate intervals of $10^5$ years, while thicker lines indicate periods of RLOF.

period of 1.8 days. After it first interacts, the 1000 day system always remains near or at RLOF, reaching carbon depletion with a surface hydrogen mass fraction $X_s = 0.36$, a radius of $7.5 R_\odot$, and an orbital period of 1.5 days. The resulting binary BHs have large delay times for a merger due to GW radiation, with the 31.6 day system being expected to merge after 6.4 Gyr and the 1000 day systems after 6.0 Gyr.

Focusing on the 31.6 day system, which we do not predict to overflow its outer Lagrangian point, Fig. 12 shows the evolution of the MT rate and the Roche filling factor with the variations to the MT prescription described before. For our updated

lowers the mass of the donor to $\sim 13 M_\odot$ and carbon is depleted while still undergoing RLOF. The 31.6 day system depletes carbon while still retaining hydrogen in its surface (surface mass fraction of $X_s = 0.40$), with a radius of $8.1 R_\odot$ and an orbital
Fig. 12. Evolution of the MT rate and overflow for a system with a 30\(M_\odot\) donor star, a 7.5\(M_\odot\) BH, and an initial orbital period of 31.6 days. Top: ratio of the computed MT rate under different assumptions, compared to the result from our full MT prescription. Middle: ratio of the stellar radius to the Roche lobe radius and to the volume equivalent radius of the L\(_2\) and L\(_3\) equipotentials (for the L\(_2\) equipotential, we only include the results of our default model for ease of visualization). Bottom: MT rate.

prescription, and in the absence of outflows from the outer Lagrangian point, the computed MT rate is expected to be higher than that of KR90. This is because dropping the assumption of \(P/\rho = kT/\mu m_H\) of KR90 and accounting for the linear term in \(dA/d\Phi\) increases the value of the integrand in Eq. (24). As shown in Fig. 12, we do find that both of these assumptions result in a lower MT rate and a higher amount of RLOF compared to our full MT scheme. This is very noticeable in the slower second MT phase, where our new MT prescription predicts a maximum MT rate, while the KR90 prescription comes from dropping the assumption that \(P/\rho = kT/\mu m_H\), while the contribution of the linear term in \(dA/d\Phi\) has a much smaller impact. Without our updated MT prescription, the KR90 prescription predicts that the system overflows its outer Lagrangian point.

The case for the 1000 day system, which undergoes overflow of its outer Lagrangian point in our model, is shown in Fig. 13.

In this case, we also find that our updated prescription keeps the star significantly more compact, with the KR90 prescription reaching a Roche filling factor of \(R/R_{RL} = 1.48\), while our model reaches \(R/R_{RL} = 1.34\). With the updated MT scheme, the donor reaches an overflow of the outer Lagrangian point of \(R/R_{L\text{out}} = 1.04\), resulting in an outflow from L\(_3\) that at its maximum accounts for less than 9\% of the total mass loss rate of the star. Since most mass is still transferred through L\(_1\), the additional angular momentum loss from mass loss through L\(_3\) only produces a small change in the final orbital separation, with the model using the prescription of KR90 resulting in a final orbital period of 1.6 days compared to the 1.5 day result when using our updated MT prescription. For this binary system, we find that during overflow of the outer Lagrangian point including the linear term in \(dA/d\Phi\) has an effect comparable to that of assuming \(P/\rho = kT/\mu m_H\). As discussed in Appendix A, it is indeed expected that the linear term in \(dA/d\Phi\) has a larger impact on mass outflows through L\(_3\) than from L\(_1\).

As the specific angular momentum that would be carried by material being ejected through the outer Lagrangian point is uncertain (we assume by default that it corresponds to the location of the Lagrangian point), we also repeated the 1000 day simulation but increased the angular momentum loss rate associated with this effect by factors of 4 and 9. In both cases, we still find the system to be stable, but they result in much shorter final periods of 1.2 day and 0.83 day. This corresponds to delay times of 3.6 Gyr and 1.2 Gyr, compared to the 6.0 Gyr delay time for our default model. This additional uncertainty could then allow for stable MT to produce systems with short delay times, where tidal spin-up can result in a high spin second formed BH (cf. Bavera et al. 2020). Another potential source of mass and angular momentum loss is the occurrence of L\(_2\) outflows when \(M_\odot < M_\odot\) for which our default model does not account. For the 100 day system, we find that the donor slightly overfills the L\(_2\) equipotential, exceeding the L\(_2\) equivalent radius by 0.2\%. This small amount of overflow is not expected to play a significant role, and indeed in our experiments with L\(_2\) outflows shown in
Fig. 14. Evolution of the radius and Roche lobe radius for two models undergoing successful envelope ejection from CE evolution. Both models correspond to a simulation of a 30 \( M_\odot \) star with a 14.1 \( M_\odot \) BH with an initial orbital period of 2344 days, with the only difference between the simulations being the efficiency of CE, \( \alpha_{\text{CE}} = 1 \) or 0.1. The shaded blue region indicates pre-CE evolution which is identical for both cases. For each case, empty circles denote the moment our algorithm determines that CE evolution would end as the star contracts within its Roche lobe in the absence of mass loss from CE evolution.

In summary, we find that our modifications to the MT prescriptions of R88 and KR90 do not have a significant effect on the MT rate or the final masses after MT concludes. The changes, however, do affect the size of the donor through the MT phase, resulting in more compact stars with less overall mass loss from CE evolution. Systems where we predict overflow of the outer Lagrangian point result in smaller orbital periods than when using the prescriptions of R88 and KR90, but we do not find that the additional source of angular momentum loss has a significant impact on the stability of these systems.

3.3. Common envelope in donors with convective envelopes

To illustrate the result of our simulations that survive CE evolution, we consider a particular model that results in successful envelope ejection in both our \( \alpha_{\text{CE}} = 1 \) and \( \alpha_{\text{CE}} = 0.1 \) grids, which has a 30 \( M_\odot \) star with a 14.1 \( M_\odot \) BH in an initial orbital of 2344 days. This model undergoes RLOF after core helium depletion and the evolution of their stellar and Roche lobe radii is shown in Fig. 14 for both CE efficiencies considered. Before the onset of RLOF, wind mass loss reduces the mass of the donor to 28.2 \( M_\odot \) and before hitting the threshold value of \( M_{\text{thigh}} = 1 \ M_\odot \ yr^{-1} \), which we first modeled to be a common envelope, while the mass of the donor is reduced to 24.7 \( M_\odot \). The onset of CE evolution for both slightly overflows its outer Lagrangian point \((R/R_{\text{lou}} = 1.08)\), at which point our MT prescription predicts 53\% of the mass to be lost by the star going through L1. During the CE calculation, we artificially removed the mass and adjusted the separation according to the binding energy of the removed layers, resulting in a sharper decrease in mass of the Roche lobe radius for the lower CE efficiency. The stellar radius remained larger than 10\(^3\) \( R_\odot \) until the mass of the star was lowered below 15 \( M_\odot \), at which point the radius began to decrease sharply with the mass. The point at which we found detachment without any enforced mass loss from CE, see Eq. (34), varies with efficiency, with the \( \alpha_{\text{CE}} = 1 \) simulation detaching at a mass of 14.3 \( M_\odot \) and the \( \alpha_{\text{CE}} = 0.1 \) model detaching at 13.7 \( M_\odot \).

Figure 15 shows the binding energy profiles of the star at different phases of evolution. Between the beginning of RLOF and the onset of CE evolution, the donor loses \( \sim 3 \ M_\odot \) of mass which modifies the binding energy of the convective envelope but not of the deeper layers. The binding energy of the ejected envelope changes significantly between the \( \alpha_{\text{CE}} = 1 \) and \( \alpha_{\text{CE}} = 0.1 \) simulations with \( E_{\text{bind}} = -3.9 \times 10^{47} \) erg and \( E_{\text{bind}} = -4.3 \times 10^{46} \) erg, respectively. For the fully efficient simulation, all ejected mass formed part of the convective envelope at the onset of CE. This factor of about 10 difference on the binding energy, coupled with the factor of 10 difference in CE efficiency leads to very different outcomes post-CE. The \( \alpha_{\text{CE}} = 1 \) simulation terminates CE evolution with an orbital separation of 570 \( R_\odot \), while the \( \alpha_{\text{CE}} = 0.1 \) simulation finishes CE with an orbital separation of 8.4 \( R_\odot \). As discussed by Deloye & Taam (2010), the core-envelope boundary is not uniquely defined, resulting in different values depending on the mass ratio and efficiency of the process. One caveat of our simulations is that the amount of mass loss previous to CE evolution is sensitive to our choice of \( M_{\text{thigh}} \), which modifies the binding energy of the outer layers of the star (see Appendix B). This modifies the post-CE separation for systems we predict to remain wide, such as the \( \alpha = 1 \) system discussed here. Accurate predictions for these wide post-CE models requires a better description of how the star transitions from stable MT to CE evolution.

After detachment, for both efficiencies we find that the donor undergoes a thermal pulse and re-expands, filling its Roche lobe but undergoing stable MT rather than developing a CE again. Figure 16 shows the evolution in the HR diagram both before and after CE evolution. For the case of fully efficient CE evolution, during this additional phase of MT the star would appear as an extended blue or yellow supergiant with a surface hydrogen mass fraction of \( X_c \sim 0.4 \). The duration of this MT phase for the case of \( \alpha_{\text{CE}} = 1 \) is \( \sim 2000 \) years, during which mass is transferred at a rate above the Eddington limit of the 14.1 \( M_\odot \) BH companion.
For the $\alpha_{CE} = 0.1$ case, the donor star has a radius of just $2R_\odot$ with a surface hydrogen mass fraction of $X_H \sim 0.3$, and it transfers mass at super-Eddington rates for $\sim 800$ years. Given that the Eddington luminosity of a $14.1 M_\odot$ BH is $\sim 2 \times 10^{59}$ erg s$^{-1}$, during this final MT phase the system can potentially appear as an ultra-luminous X-ray source (see Kaaret et al. 2017 for a recent review), but owing to the short time before core carbon depletion, the number of observable sources could be negligible compared to those that formed by other evolutionary pathways.

Not all our simulations that survive CE evolution have short lifetimes as X-ray sources. For the system we have discussed so far, this is the case because CE evolution happens after core helium depletion. If we consider a system with a $7.5 M_\odot$ BH at an initial orbital period of $2000$ days and $\alpha_{CE} = 1$, we find that the onset of CE happens when the donor has a central helium mass fraction of $Y_H = 0.02$, and for the entirety of its post-CE lifetime until core carbon depletion ($260000$ years), it is a Roche-lobe filling binary. This matches the results of Quast et al. (2019) who argue that post-CE systems can undergo MT on the nuclear timescale of the donor. We expect that in cases where the star does not halt its expansion in the HG, CE evolution is initiated with a much larger value of $Y_H$, leading to a longer lifetime as an X-ray source post-CE.

The single model we considered with different values of $\alpha_{CE}$ illustrates how the binding energy is dependent on the efficiency of CE evolution, but using the models from our grid we can see how the resulting binding energy of systems surviving CE evolution depends on the mass ratio as well. Figure 17 illustrates the binding energy as a function of the radius at RLOF resulting from models at three different initial mass ratios, $\alpha_{CE} = 1$, and varying initial orbital periods. For most of these simulations, we find that CE evolution terminates near or above the mass coordinate of the bottom of the convective envelope at RLOF, resulting in binding energies $<10^{48}$ erg. These low binding energies are below those predicted by the fits of Xu & Li (2010) and Claeyts et al. (2014), although for a mass ratio of $q = 0.17$ we find binary models with binding energies that are smaller than the value of these two fits. The $q = 0.17$ simulations with binding energies $>10^{48}$ erg behave similarly to the simulation with $\alpha_{CE} = 0.1$ shown in Fig. 15, where the binary only detaches after removing layers with a high binding energy that were radiative at the onset of CE evolution. This results in an order of magnitude variation of the binding energy with a varying mass ratio.

### 3.4. Common envelope in donors with radiative envelopes

As all of our simulations where CE evolution is triggered for a star with a radiative envelope are expected to merge during CE evolution, we cannot properly define a core-envelope boundary to compute the binding energy. However, we can artificially produce systems that eject their outer envelope by using artificially high values of $\alpha_{CE}$. Here, we consider systems consisting of a $30 M_\odot$ donor and a $3 M_\odot$ BH with initial orbital periods between $10^4$ days and $10^5$ days, and values of $\alpha_{CE}$ of 10 and 20. The low mass of the BH, which would fall in the putative lower BH mass gap (Ozel et al. 2010; Farr et al. 2011; Kreidberg et al. 2012; Abbott et al. 2020b), was chosen such that MT would be unstable across the range of chosen orbital periods.

All of these simulations managed to eject their envelopes during CE evolution, and Fig. 18 shows the resulting masses post-CE as well as the binding energy of the ejected layers. We find that CE terminates when the surface hydrogen mass fraction is relatively high ($X \sim 0.4–0.5$) and that the binding energy of the envelope varies between $1.3 \times 10^{49}$ erg and $6.6 \times 10^{49}$ erg, a factor of 2 variation despite a change by a factor of 20 in radius at RLOF. Figure 18 also illustrates that whether RLOF occurs before or after the end of the HG has no impact on the binding energy. Estimating the binding energy using either the Claeyts et al. (2014) or Xu & Li (2010) fits would underestimate the binding energy of the envelope for all these simulations. Since the binding energy just changes by a factor of 2 despite over an order of magnitude change in the radius of the star, population synthesis calculations that approximate $E_{\text{bind}}$ using a fixed value of the dimensionless $\lambda$ parameter cannot reproduce the variation of $E_{\text{bind}}$ with radius, independent of the choice of $\lambda$. 

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Fig. 16. HR diagram showing the evolution of a system undergoing CE evolution composed of a $30 M_\odot$ donor with a $14.1 M_\odot$ BH companion at an initial period of 2344 days. Results are shown for two values of the efficiency of CE evolution, $\alpha_{CE} = 1$ and $\alpha_{CE} = 0.1$, with the evolution prior to CE being identical.

Fig. 17. Binding energy of the envelope of a star with an initial mass of $30 M_\odot$, as a function of the radius at RLOF. Solid lines indicate the results from our $\alpha_{CE} = 1$ grid with three different mass ratios $q = 0.17$, 0.29, and 0.45. Dotted lines show the binding energy at a given radius deduced from a single star model, including the results obtained with the fit of Xu & Li (2010), the fit of Claeyts et al. (2014), and assuming the bottom of the envelope to be at $X = 0.1$, $X = 0.2$, or to correspond to the bottom of the convective envelope. Symbols indicate whether the binary undergoes RLOF before or after core helium depletion.
4. Discussion

We have studied the outcomes from the interaction of low-metallicity massive stars with a companion BH using detailed binary evolution calculations performed with the MESA code. Our simulations, which consist of a 30 $M_\odot$ low-metallicity donor with a companion BH, show several features relevant to understand the formation of merging binary BHs through both stable MT and CE evolution. For these simulations, we have also updated the MT prescription developed by KR90, including the possibility of overflow from an outer Lagrangian point, and we also implemented a method to self consistently determine the core-envelope boundary in cases of CE evolution. The main conclusions we derive from our simulations are as follows:

1. Stable MT after the formation of the first BH in a binary can produce merging binary BHs for a broad range of orbital periods. The boundary between stable and unstable MT is located at a point that allows for orbital hardening just from stable MT, without requiring a CE phase (van den Heuvel et al. 2017). This includes cases where MT between the star and the first formed BH is initiated during the main sequence. We also find cases where the low density envelope of the donor would overflow its outer Lagrangian point. Using our model to quantify this outflow, we find that binaries can remain stable despite the high specific angular momentum of this outflow, resulting in more compact binaries after MT than if the process is ignored.

2. We do not find any case of CE evolution initiated with a star with a radiative envelope where the envelope is successfully ejected. Between the TAMS and the start of the rise in the Hayashi line, the binding energy of the envelope remains at $\sim 10^{50}$ erg, varying by just a factor of two despite an increase of more than an order of magnitude in radius. Before the formation of a deep convective envelope, we find that the prescriptions of Xu & Li (2010), as developed by Dominik et al. (2012) for $M > 20 M_\odot$, and Claeys et al. (2014) underestimate the binding energy of the envelope by up to an order of magnitude.

3. The parameter space where systems can survive CE is limited to a small range in the initial orbital period ($\sim 0.2$ dex) where RLOF happens after the formation of a deep convective envelope. Assuming that all internal and recombination energy as well as all the energy from inspiral during CE is used to eject the envelope, we find that the majority of these systems surviving CE would form binary BHs too wide to merge within a Hubble time.

4. As in Deloye & Taam (2010), we find that the core-envelope boundary is not uniquely defined, with different CE efficiencies (or similarly, different mass ratios) resulting in different binding energies for the envelope. In particular for the case discussed in Sect. 2.2, we computed binding energies that differ by over an order of magnitude depending on the efficiency of CE evolution. This is in contrast to the way binding energies are determined in population synthesis codes, where a star has a uniquely defined binding energy at each point of its evolution.

5. Under the simple assumption of a flat distribution in the mass ratio and a flat distribution in log $P$ after the formation of the first BH, we find that stable MT would dominate the formation rate of merging binary BHs. If we consider CE efficiency to be 100%, the ratio of binaries surviving CE evolution (independent of whether they form a compact or wide binary BH) to those that form merging binary BHs from stable MT is 0.35. This is further reduced to 0.017 if we take only the CE simulations that produce merging binary BHs. This is in significant contradiction with rapid population synthesis calculations that predict the vast majority (>90%) of BHs are formed via an initial phase of stable MT before the formation of the first BH, followed by a CE phase when the secondary fills its Roche lobe (cf. Table 4 of Dominik et al. 2012).

These results broadly agree with those of Klencki et al. (2020, 2021), who have used single star models with a fixed definition of the core-envelope boundary to propose that current population synthesis calculations may overestimate the production rate of merging binary BHs from CE evolution. Since our simulations are restricted to a 30 $M_\odot$ donor at $Z = Z_\odot/10$, it is necessary to further explore different masses and metallicities to understand how generally our conclusions apply (Gallegos-Garcia et al., in prep.). Recent investigations using rapid population synthesis...
codes have indicated that given current uncertainties in binary evolution such as the cosmic star formation rate (Neijssel et al. 2019) or the critical mass ratio for stable MT (Andrews et al. 2020), stable MT could be an important production channel for the merging binary BHs observed with gravitational waves. Our results indicate these rapid population synthesis calculations still significantly overestimate the number of systems surviving CE evolution at low metallicities, and accounting for this would further reduce the contribution from the CE evolution channel.

As we find that the boundary between stable and unstable MT is located right at a point that allows for the formation of merging binary BHs via stable MT, any uncertainties that can shift this boundary can have a large effect on the number and properties of expected GW sources. Among such uncertainties, we can point out that our simulations do not include stellar rotation or tidal coupling to the orbit. This additional sink of orbital angular momentum can potentially affect the stability of MT in our calculations. We have explicitly checked in our storable MT simulations that the moment of inertia of the star is too low to undergo the Darwin instability (Darwin 1879), which is the limit were tidal coupling leads to a runaway shrinkage of the orbit. Another important simplification is that we did not account for the effect of the companion in the momentum equation, which would make the adiabatic response of the star to mass loss dependent on the mass ratio of the system and its separation. It has also been argued that hydrodynamics need to be considered to assess the stability of MT for giant donors (Pavlovskii & Ivanova 2015), although a consistent treatment of hydrodynamics needs to consider the perturbation to the gravitational potential from the companion star as well. Additionally, a significant number of the systems we predict to form merging binary BHs also undergo overflow of the L2 and even the L1 equipotentials. Although for these large overflow cases, we have verified that merging binary BHs are formed even under extreme assumptions of mass and angular momentum outflows, three-dimensional hydrodynamic studies are required to accurately model outflows in such systems. All these uncertainties need to be studied in detail to determine if stable MT can indeed produce merging binary BHs.

Our simulations for the outcome of CE evolution also have important uncertainties. Although we do not have the core-envelope mass boundary as a free parameter, we still have the CE efficiency as an unknown variable. The efficiency can be reduced by radiative processes or also by a surplus of energy being removed as kinetic energy from the system (cf. Nandez & Ivanova 2016). Within the context of one-dimensional stellar models, it might be possible to constrain the efficiency of the CE process by means of simplified hydrodynamical simulations of the inspiral (Taam et al. 1978; Clayton et al. 2017; Fragos et al. 2019) or, ideally, with full three-dimensional simulations of the inspiral (Taam et al. 1978; Clayton et al. 2017; Andrews et al. 2020, 2019; Pavlovskii & Ivanova 2015), although a consistent treatment of hydrodynamics needs to consider the perturbation to the gravitational potential from the companion star as well. Additionally, a significant number of the systems we predict to form merging binary BHs also undergo overflow of the L2 and even the L1 equipotentials. Although for these large overflow cases, we have verified that merging binary BHs are formed even under extreme assumptions of mass and angular momentum outflows, three-dimensional hydrodynamic studies are required to accurately model outflows in such systems. All these uncertainties need to be studied in detail to determine if stable MT can indeed produce merging binary BHs.

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Appendix A: Computation of $d A/d\Phi$

The coefficients of the quadratic terms in the Taylor expansion of Eq. (7) are given by

$$C_1 = \frac{1}{2} \left( \frac{\partial^2 \Phi}{\partial y^2} \right)_{L_1} = \frac{GM_d}{2a^3} \left[ \frac{1}{|\hat{x}_{L1}|^3} + \frac{q}{|1 - \hat{x}_{L1}|^3} - (q + 1) \right],$$

$$C_2 = \frac{1}{2} \left( \frac{\partial^2 \Phi}{\partial z^2} \right)_{L_1} = \frac{GM_d}{2a^3} \left[ \frac{1}{|\hat{x}_{L1}|^3} + \frac{q}{|1 - \hat{x}_{L1}|^3} \right],$$

and the coefficients of the quartic terms are

$$C_3 = \frac{1}{24} \left( \frac{\partial^4 \Phi}{\partial y^4} \right)_{L_1} = -\frac{3GM_d}{8a^5} \left[ \frac{1}{|\hat{x}_{L1}|^5} + \frac{q}{|1 - \hat{x}_{L1}|^5} \right],$$

$$C_4 = \frac{1}{4} \left( \frac{\partial^4 \Phi}{\partial z^4} \right)_{L_1} = 2C_3,$$

$$C_5 = \frac{1}{24} \left( \frac{\partial^4 \Phi}{\partial z^4} \right)_{L_1} = C_3.$$ 

When we consider outflows from the outer Lagrangian point of the donor, $\hat{x}_{L1}$ is switched by the normalized $x$-coordinate of the outer Lagrangian point, $\hat{x}_{L_{out}}$, which, with our definition, has a negative value.

In order to estimate the area on the plane crossing $L_1$ for a given $\Delta \Phi$, it is useful to use cylindrical coordinates $y = \sigma \sin \theta$, $z = \sigma \cos \theta$, in which case

$$\sigma^2 = \frac{1}{\gamma} \Delta \Phi - \frac{C_1}{\gamma^2} (\Delta \Phi)^2 + O((\Delta \Phi)^3), \quad (A.1)$$

where $\gamma \equiv C_1 \sin^2 \theta + C_2 \cos^2 \theta$. This can be integrated to obtain the area $A(\Phi)$,

$$A = 2 \int_0^{\pi/2} \sigma^2 \, d\theta, \quad (A.2)$$

which after differentiation by $\Phi$ gives Eq. (8).

The relative importance of the constant and linear terms for $dA/d\Phi$ shown in Eq. (8) can be assessed by using the ratio of $dA/d\Phi(\Delta \Phi = 0)$ and $dA/d\Phi^2(\Delta \Phi = 0)$. This essentially quantifies how large $\Delta \Phi$ needs to be to produce a significant change in $dA/d\Phi$, with smaller values indicating a higher sensitivity to $\Delta \Phi$, and thus a larger importance of the higher-order term. Figure A.1 shows this value at different mass ratios both for the $L_1$ and $L_{out}$ points of a star. This value is always positive, indicating that the inclusion of the higher-order term results in an increased value of the computed MT rate. We also see that the value of the ratio is much smaller for the outer Lagrangian point, and in the case where the donor is the more massive component in the binary, a variation of $\Phi$ on the order of $0.1 \, GM_d/\alpha$ can modify $dA/d\Phi$ significantly. This points out that the inclusion of the higher-order term has a larger impact on the computation of outflows from the outer Lagrangian point than from $L_1$.

Appendix B: Dependence on numerical parameters of CE algorithm

As our algorithm to model CE evolution has various numerical parameters, it is important to evaluate in detail how our results depend upon these. Here we modify the three parameters described in Sect. 2.2: $M_{\text{high}}$, $M_{\text{low}}$, and $\delta$, for which in our default model we used $1 M_{\odot}$ $yr^{-1}$, $10^{-3} M_{\odot}$ $yr^{-1}$, and 0.02, respectively. Here, we consider the difference in the outcome of the CE simulations shown in Sect. 3.3 for variations in these three parameters, with the properties of the system post-CE being shown in Table B.1. For all parameters, we find only ~5% variations in the post-CE properties, except for variations of $M_{\text{high}}$ when using an efficiency $\alpha_{\text{CE}} = 1$. Detachment in these systems happens in the proximity of the mass coordinate from the bottom of the convective envelope at the onset of CE. Variations in $M_{\text{high}}$ modifies the mass of the star at the onset of CE, as is shown in Table B.1. This modifies the binding energy in the outermost layers, which is illustrated in Fig. B.1, making evolution sensitive to the exact point at which we switched between MT and CE evolution. This is not the case for the system with $\alpha_{\text{CE}} = 0.1$, as the core-envelope boundary is computed to be deeper in the star, in layers that are not affected by the exact point at which we switched between MT and CE evolution.
A comparison of our single metal-poor $30\, M_\odot$ model to similar stellar evolution tracks that are publicly available is shown in Fig. C.1. These include tracks by Meynet et al. (1994), Pols et al. (1998), Tang et al. (2014), and Klencki et al. (2020). Out of these, the only ones that use the same stellar evolution code are our own models and those of Klencki et al. (2020). From the models of Meynet et al. (1994), we took the $25\, M_\odot$ stellar track which is the closest one to $30\, M_\odot$. The track taken from Pols et al. (1998) corresponds to their models with overshooting, and the one from Klencki et al. (2020) is from their set of nonrotating stellar tracks. All tracks experience a long-lived phase as blue supergiants, halting their rapid expansion in the HG phase at $\log_{10}(T_{\text{eff}}/K) \sim 4.2$ (except the lower mass one from Meynet et al. 1994 that does so at a higher $T_{\text{eff}}$).

Given the formation of multiple intermediate convection zones in our models during the HG phase, we performed a resolution test to see if our results are significantly modified by increased spatial and temporal resolution. We made use of the `mesh_delta_coeff` and `time_delta_coeff` options that allow for an approximate scaling of all spatial and temporal resolution controls, and used models with approximately two, four, and eight times the resolution of our default setup. The resulting tracks in the HR diagram are shown in Fig. D.1. We see small variations with changes in resolution on the evolution during the HG phase, producing a variation of about 10% on the effective temperature at which the HG phase ends.

In particular, for this work we are interested in how the binding energy of the envelope of a star compares to prescriptions present in the literature. Figure D.2 shows the binding energies
obtained for models at increasing temporal and spatial resolution, as well as the results we obtain with the Xu & Li (2010) and Claeys et al. (2014) fits. Binding energies were computed from the stellar models taking the bottom of the envelope at the innermost mass coordinate where $X > 0.1$ and $X > 0.3$. Between the different models, we see variations in the computed binding energies at a given radius smaller than 12%. The larger variations happen at the end of the HG and are due to the slightly different radii achieved at the end of the HG. Ignoring the values between $\log_{10}(R/R_\odot) = 1.8 - 1.95$, we find that the differences are smaller than 6%.

**Appendix E: Application of $\lambda$ fits to stellar models**

The fits from Claeys et al. (2014) and Xu & Li (2010) depend on the mass of the star, its radius, and its evolutionary stage in terms of the star-types defined by Hurley et al. (2002). A model analogous to our $30 M_\odot$ model is assigned types "HG" during the HG, "CHeB" during core helium burning but before ascending the Hayashi line, and "EAGB" while on the Hayashi line. To distinguish between the CHeB and EAGB types in our models, we considered models with convective envelopes $>1 M_\odot$ to be of type EAGB, as we found this is close to the point where the star rises on the Hayashi line. As we are only concerned with these three assigned types, we only describe the fits associated with them.

**E.1. Claeys et al. (2014)**

The fits of Claeys et al. (2014) were first presented by Izzard (2004), who took them directly from the source code of the BSE code (Hurley et al. 2002). Claeys et al. (2014) do not specify the range of masses used to compute these fits. The fit for $\lambda$ including internal but not recombination energy is

$$\lambda = 2 \times \begin{cases} 
\lambda_2 & M_{\text{conv env}} = 0 \\
\lambda_1 + M_{\text{conv env}} \left( \lambda_1 - \lambda_2 \right) & 0 < M_{\text{conv env}} = M_\odot \\
\lambda_1 & M_{\text{conv env}} > M_\odot 
\end{cases} \quad (E.1)$$

where

$$\lambda_2 = 0.42 \left( \frac{R_{\text{ZAMS}}}{R} \right)^{0.4}, \quad (E.2)$$

with $R_{\text{ZAMS}}$ being the radius of the stellar model at the ZAMS. The value of $\lambda_1$ is given by

$$\lambda_1 = \min \{ 0.8, 1.25 - 0.15 \log_{10}(L/L_\odot), \lambda_3 \} \quad (E.3)$$

with

$$\lambda_3 = \min \{ 0.9, 0.58 + 0.75 \log_{10}(M/M_\odot) \} - 0.08 \log_{10}(L/L_\odot). \quad (E.4)$$

This fixes a typo (0.9 instead of $-0.9$ in the minimum function) from Claeys et al. (2014) and Izzard (2004) that is inconsistent with the implementation in the BSE code from which the fit was taken. Claeys et al. (2014) also describe variations in the $\lambda_1$ and $\lambda_3$ values for stellar types that are not relevant for our $30 M_\odot$ model, so we do not describe those.

**E.2. Xu & Li (2010)**

The $\lambda$ fits of Xu & Li (2010) cover up to a mass of $20 M_\odot$, while fits for higher masses were developed by Dominik et al. (2012). These fits are not published, but can be found in the source code of the COMPAS code. Fits are provided for the binding energy with and without thermal energy included, that is $\alpha_\text{th} = 0$ or 1 in Eq. (33). As we worked with $\alpha_\text{th} = 1$, we compared our results to the $\lambda$ fit including thermal energy. Fits are provided individually for each of the HG, CHeB, and EAGB phases, and the ones that correspond to our $30 M_\odot$ model at $Z_\odot/10$ are the Population II fits for masses $18 M_\odot < M < 35 M_\odot$. For HG stars $\lambda$ is computed as

$$\lambda_{\text{HG}} = \begin{cases} 
a_1 + a_2 \xi + a_3 \xi^2 + a_4 \xi^3 & R \leq 900 R_\odot \\
0.2 & R > 900 R_\odot \end{cases} \quad (E.5)$$

where $\xi = R/R_\odot$, and the value of the fitting coefficients is given in Table E.1. For CHeB stars we have

$$\lambda_{\text{CHeB}} = \begin{cases} 
a_1 + a_2 \xi + a_3 \xi^2 + a_4 \xi^3 & R \leq 230 R_\odot \\
0.1 & 230 R_\odot < R < 755 R_\odot \\
0.1 + 0.2(1 - b) & 755 R_\odot < R < 900 R_\odot \\
0.2 & R > 900 R_\odot \end{cases} \quad (E.6)$$

with the expression for the range $755 R_\odot < R < 900 R_\odot$ being a linear interpolation from 0.1 to 0.2 with $b = (900 - \xi)/(900 - 755)$. For the case of EAGB stars the fit is given by

$$\lambda_{\text{EAGB}} = \begin{cases} 
a_1 + a_2 \xi + a_3 \xi^2 + a_4 \xi^3 & R \leq 725 R_\odot \\
0.1 & 725 R_\odot < R < 850 R_\odot \\
0.1 + 0.2(1 - b) & 850 R_\odot < R < 900 R_\odot \\
0.2 & R > 900 R_\odot \end{cases} \quad (E.7)$$

where $b = (900 - \xi)/(900 - 850)$. All resulting values are capped between 0.05 and 1.5. The interpolation in the range $755 R_\odot - 900 R_\odot$ for CHeB stars is missing in the implementation of Dominik et al. (2012), and it erroneously applies the fitting polynomial in this range resulting in negative values, producing a positive value only because the final result is capped to a minimum of 0.05. Not including this correction and having $\lambda = 0.05$ in this range of radii does not change our conclusions, as the binding energy is still underestimated compared to our own models.
Table E.1. Fitting coefficients for the $\lambda$ fits of Xu & Li (2010) including thermal energy, as implemented by Dominik et al. (2012) for Population II stars with masses $18 M_\odot < M < 35 M_\odot$.

<table>
<thead>
<tr>
<th>Star type</th>
<th>$a_1$</th>
<th>$a_2$</th>
<th>$a_3$</th>
<th>$a_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>HG</td>
<td>1.27138</td>
<td>0.00538</td>
<td>-0.0012</td>
<td>$1.80776 \times 10^{-5}$</td>
</tr>
<tr>
<td>CHaB</td>
<td>0.69746</td>
<td>-0.0043</td>
<td>$8.97312 \times 10^{-6}$</td>
<td>$-5.83402 \times 10^{-9}$</td>
</tr>
<tr>
<td>EAGB</td>
<td>-436.00777</td>
<td>1.41375</td>
<td>-0.00153</td>
<td>$5.47573 \times 10^{-7}$</td>
</tr>
</tbody>
</table>

Notes. Values are taken from the source code of the COMPAS code. Exact numbers from the source code are included here, so the number of significant digits is variable.