Rotational equilibria by Lagrangian variational principle: toward multi-dimensional stellar evolutions

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ABSTRACT

We have developed a new formulation to obtain self-gravitating, axisymmetric configurations in permanent rotation. The formulation is based on the Lagrangian variational principle with a triangulated mesh. It treats not only barotropic but also baroclinic equations of state. We compare the various stellar equilibria obtained by our new scheme with those by Hachisu’s self-consistent field scheme for the barotropic case, and those by Fujisawa’s self-consistent field scheme for the baroclinic case. Included in these rotational configurations are those with shellular-type rotations, which are commonly assumed in the evolution calculation of rotating stars. Although radiation processes, convections and meridional flows have not been taken into account in this study, we have in mind the application of this method to the two-dimensional evolution calculations of rotating stars, for which the Lagrangian formulation is best suited.

Key words: stars: rotation — stars: evolution — stars: protostars

1 INTRODUCTION

Stellar evolution theory is well established especially for spherically symmetric stars, in which there is no rotation and no magnetic field. After the works of pioneers of the field (Burbidge et al. 1957; Cameron 1957; Havashi 1961\(^a\)\(b\)), Henyey proposed in his seminal papers a method, which later became the de facto standard for stellar evolution calculations (Henyey et al. 1964). The method is stable and capable of self-consistently incorporating various physical processes that occur in the stellar interior. It has been modified and extended a lot approximately in the last half century to accommodate rotation (Maeder & Meynet 2000; Woosley et al. 2002).

In stellar evolution calculations, we need to obtain stellar structures consistent with, nuclear reactions (and/or molecule formations), radiative transport of energy, convective and circular motions as well as a realistic equation of state. Since the time-scale of stellar evolution is normally much longer than the dynamical time-scale, it is well approximated by series of time-independent configurations in hydrostatic equilibrium. In the presence of rotation, this is not simple. One of the difficulties is to obtain rotational equilibria for a given angular momentum distribution. What is more, however, we do not know a priori what the distribution of angular momentum looks like in the stellar interior although the problem has been studied theoretically since the 19th century by Carl Jacobi, Richard Dedekind, Peter Lejeune Dirichlet, and Bernhard Riemann to mention a few (Chandrasekhar 1969). It was in the last century that some methods to obtain rotating hydrostatic equilibria were proposed, and it was mathematically shown that cylindrical distributions of specific angular momentum are realized for EOSs, in which matter pressure is a function of density alone. This type of EOS is referred to “barotrope” (Ostriker & Mark 1968; Hachisu 1986).

The EOS is not barotropic in the realistic stellar interior, but baroclinic in general, i.e., pressure depends on density, entropy and chemical compositions. Then the numerical construction of rotational equilibria becomes much more difficult compared with the case for barotropes, since the first integral of the hydrostatic equations that is available in the barotropic case no longer exists in the baroclinic case (Uryu & Eriguchi 1994; Uryu & Eriguchi 1997; Roxburgh 2006; Espinosa Lara & Rieutord 2007; Espinosa Lara & Rieutord 2013; Rieutord et al. 2016). Even if some rotational configurations are somehow obtained, it is a highly non-trivial issue how to make a sequence out of them that represents stellar evolutions appropriately. Note that in the presence of rotation fluid elements composing rotating stars could change their positions non-radially in complicated ways as the stars evolve in time even without convection. It would be highly difficult to describe such displacements of fluid elements with fluxes on a fixed numerical

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\(^1\) For compact stars such as white dwarfs and neutron stars, the EOS may be approximately barotropic, since the temperature is negligibly low.
mesh from a Eulerian point of view, since the motions are extremely slow. The Lagrangian formulation will be hence more appropriate just as in the spherically symmetric case. And this is exactly the idea in this paper. We introduced in this paper a triangulated mesh with each node representing a fluid element approximately. Starting from an arbitrary reference configuration, we seek for the Lagrangian displacement for each node that gives a rotational equilibrium as a result. In so doing, the variational principle is employed. The method has its own difficulty, though.

In this paper, we construct a couple of configurations in rotational equilibrium with different angular momentum distributions in order to demonstrate the capability of our new scheme. Included in them is the so-called shellular-type rotations. It has been argued that in the stellar interior turbulence is anisotropic, operating more strongly within each isobaric surface than in its normal direction, and that a uniform rotation should be realized in the isobar as a consequence. Such a rotation law is referred to as “shellular rotation” rather than cylindrical rotation (Zahn 1992; Meynet & Meynet 1997). It should be stressed, however, that no rotational equilibrium with the shellular rotation was constructed numerically, not to mention analytically. Very recently Fujisawa, one of the authors of this paper, has succeeded in producing some configurations with shellular-type rotation (Fujisawa 2015). It is a nice demonstration of performance of our new scheme to reconstruct these configurations.

The organization of the paper is as follows. In Section 2, we describe the formulation in detail: the Lagrangian variational principle, on which our scheme (referred to as the YFY scheme hereafter) is based, will be reviewed first; then the finite discretization on the triangulated grid will be explained and the handling of self-gravity is also mentioned; finally the minimization of the energy function with the Monte Carlo technique, which is analogous to those utilized, e.g., in nuclear physics, will be described. In Section 3, we demonstrate the nice performance of YFY scheme, constructing some rotational configurations and comparing them with the those obtained by other numerical schemes, including Fujisawa’s self-consistent field scheme mentioned above. In the last section, we summarize this study and discuss possible extensions of our scheme as future works.

2 FORMULATION

Rotational equilibria should satisfy the following equations:

(i) the force balance equation:

\[ \nabla P + \rho \nabla \phi - \frac{\rho \omega^2}{c^2} e_\phi = 0, \tag{1} \]

where \( P, \rho, \phi, j, \omega, e_\phi \) are pressure, density, gravitational potential, specific angular momentum, the distance from the rotation axis, and the unit vector perpendicular to the rotation axis, respectively;

(ii) the Poisson equation:

\[ \Delta \phi = 4\pi G \rho. \tag{2} \]

where \( G \) is the gravitational constant; these equations are supplemented with the equations of state (EOS).

Although equations (1), (2) look simple, especially for a very realistic EOS, numerically solving them may not be so easy. Most of previous works assumed a barotropic EOS, in which pressure is a function of density alone, and for which it is mathematically shown that rotation is uniform on concentric cylinders and, as a consequence, there is a first integral of equation (1). The realistic equation of state, however, is not barotropic but baroclinic, in which case pressure depends on density, entropy and chemical compositions. Then there is no first integral available any longer and the rotation law becomes unknown a priori. In this paper, we do not use the force balance equation, equation (1). Instead the variational principle is utilized as described in detail below.

2.1 Lagrangian variational principle

We first describe the Lagrangian variational principle, which is obeyed by rotational equilibria. We obtain the energy functional, which is minimized by a rotational equilibrium and will be discretized on a triangulated mesh later for numerical calculations. The variational principle itself is not our own invention but is actually given in a textbook (Tassoul 1978), although it is a core of the entire formulation proposed in this paper.

Suppose an arbitrary reference configuration with density, specific entropy and specific angular momentum distributions \((\rho(\tilde{r}), s(\tilde{r}), j(\tilde{r}))\), where \( \tilde{r} \) is a spatial coordinate and will also serve as a Lagrange coordinate as will become clear shortly. We, then, introduce a diffeomorphic mapping: \( r = f(\tilde{r}) \), which is identified with the Lagrangian displacement that deforms the reference configuration to another configuration. Then the following relation should hold if the mass were to be conserved in the deformation:

\[ \rho(\tilde{r}) \, dV(\tilde{r}) = \rho(r) \, dV(r), \tag{3} \]

where \( \rho(\tilde{r}) \) is a density at \( \tilde{r} \) and \( dV(\tilde{r}) \) is an infinitesimal volume around \( \tilde{r} \) in the reference configuration whereas \( \rho(r) \)

and \( dV(r) \) are the density and volume at the corresponding position in the mapped configuration. The volume element is transformed with the Jacobian \( J \) of the mapping as

\[ dV(\tilde{r}) = J(f^{-1}) \, dV(r), \tag{4} \]

the density after the displacement is expressed as

\[ \rho(r) = \rho(\tilde{r}) J(f^{-1}). \tag{5} \]

Introducing another diffeomorphism \( g \), we consider the consecutive mapping \( g \circ f \), which we also refer to as the product of maps \( g \) and \( f \) and interpret as a Lagrangian displacement:

\[ \tilde{r} \xrightarrow{f} r \xrightarrow{g} r', \tag{6} \]

where \( r' \) is the coordinate after the mapping \( g \).

Using the formula for the product of Jacobians,

\[ J((g \circ f)^{-1}) = J(f^{-1}) \cdot J(g^{-1}), \tag{7} \]
In this expression the integration domain is the stellar

\[ \rho(r') = \rho(\tilde{r})J((g \cdot f)^{-1}) = \rho(\tilde{r})J(f^{-1})J(g^{-1}) = \rho(r)J(g^{-1}). \]

This equation implies that the intermediate configuration

obtained after the mapping \( f \) served as the reference con-

figuration for the configuration after the mapping \( g \) and

one can forget about the original reference configuration.

Employing this fact, we consider infinitesimal displacements

alone in the following:

\[ \delta f : r \rightarrow r' = r + \xi \]

where \( \xi \) is the so-called Lagrangian displacement vector, an

infinitesimal vector that connects the original point with its

image by the map. The Jacobian for the (inverse) infinites-

imal mapping is given as

\[ J(\delta f^{-1}) = J^{-1}(\delta f) = 1 - \text{div}(\xi). \]

Then the Lagrangian variation of density \( \Delta \rho \) is obtained as

\[ \Delta \rho \equiv \rho(r') - \rho(r) = (J(\delta f^{-1}) - 1)\rho(r) = -\text{div}(\xi)\rho(r). \]

In this study, we consider the variational principle, in

which rotational equilibria are obtained. Supposing that

there is no Lagrangian variation in specific entropy and an-

gular momentum for the above infinitesimal displacement,

\[ \Delta s = 0, \]

\[ \Delta j = 0, \]

we consider the variation of the following functional, to

which we refer as the energy functional hereafter:

\[ E(\xi) = \int \varepsilon \rho \, dV + \frac{1}{2} \int \rho \phi \, dV + \int \frac{1}{2} \rho \left( \frac{j}{\omega} \right)^2 \, dV. \]

In this expression the integration domain is the stellar in-

terior, and \( \varepsilon \) and \( \phi \) denote the specific internal energy and

gravitational potential, respectively; \( \omega \) is the distance from

the rotational axis. We consider the variation of each term

on the right hand side of equation (12) in turn. The variation

of the first term is evaluated as follows:

\[ \delta \left( \int \varepsilon \rho \, dV \right) = \int \Delta \varepsilon \rho \, dV = \int \left( \frac{\partial \varepsilon}{\partial \rho} \right) \Delta \rho \, dV \]

\[ = \int \left( \frac{P}{\rho^2} \right) \Delta \rho \, dV = - \int P \text{div}(\xi) \, dV \]

\[ = \int \nabla P \cdot \xi \, dV. \]

Here \( \Delta \varepsilon \) is the Lagrangian variation of internal energy, which is

given by the Lagrangian variation of density upon using the

assumption that there is no entropy variation, equation (12). The final expression is obtained with the use of equation (12).

The second term is evaluated as follows:

\[ \delta \left( \frac{1}{2} \int \rho \phi \, dV \right) = \frac{1}{2} \int (\partial \rho/\partial \phi) \delta \rho \, dV \]

\[ = \frac{1}{2} \int \left( - \nabla (\rho \xi) \cdot \phi + \rho \cdot \delta \phi \right) \, dV \]

\[ = \frac{1}{2} \int \rho \cdot \phi \, dV \]

\[ = \frac{1}{2} \int \left( \rho \xi \cdot \nabla \phi + \rho \cdot \delta \phi \right) \, dV, \]

where the second equality is obtained with an employment of the following relation:

\[ \delta \rho = \Delta \rho - \xi \nabla \rho. \]

\[ \delta \phi \]

in the last term of the final expression is the Eulerian variation of the gravitational potential, which is evaluated from the Poisson equation, \( \Delta \phi = 4\pi G\rho \), and its solution with the Green’s function,

\[ \phi = - \int dV' \rho(r')/|r - r'|. \]

Then \( \int \rho \cdot \delta \phi \, dV \) is given as

\[ \int \rho \cdot \delta \phi \, dV = -G \int \delta \rho(r') \phi(r') \, dV' \]

\[ = \int \rho(r') \phi(r') \, dV' = \int \rho \xi \cdot \nabla \phi \, dV. \]

Equation (16) is hence expressed as

\[ \delta \left( \frac{1}{2} \int \rho \phi \, dV \right) = \int \rho \xi \cdot \nabla \phi \, dV. \]

The variation in the last term of equation (13) is calculated as

\[ \delta \left( \frac{1}{2} \int \rho \left( \frac{j}{\omega} \right)^2 \, dV \right) = \int \frac{1}{2} \rho \Delta \left( \frac{j}{\omega} \right)^2 \, dV \]

\[ = \int \frac{1}{2} \rho \left( \frac{j}{\omega} \right)^2 \cdot \nabla \rho \, dV \]

\[ = - \int \frac{1}{2} \rho \left( \frac{j}{\omega} \right)^2 \cdot \varepsilon \, dV. \]

In the second equality we employed the assumption that there is no Lagrangian variation in the specific angular momentum, equation (12).

We are now ready to derive the Euler-Lagrange equation for the energy functional in equation (13). All the variations given by equations (15), (20) and (21) are expressed with the Lagrangian displacement vector \( \xi \) and the Euler-Lagrange equation corresponds to its coefficient in \( \delta E \). The resultant equation is obviously equation (1). This implies that the rotational equilibrium is a stationary point of the functional and that it may be obtained not by solving equation (1) but by minimizing the energy functional somehow.

In the above formulation of the variational principle, we treated the gravitational potential \( \phi \) as a functional of density, formally solving the Poisson equation with the Green’s function. It is possible, however, to treat the gravitational potential also as an independent variational variable. The energy functional should be modified then to

\[ E(\xi) = \int \varepsilon \rho \, dV + \int \rho \phi \, dV + \int \frac{1}{2} \rho \left( \frac{j}{\omega} \right)^2 \, dV \]

\[ + \frac{1}{4\pi G} \int \frac{1}{2}(\nabla \phi)^2 \, dV. \]

\[ \text{The rotational equilibrium may not be a minimum point of the energy functional, which will be indeed the case if the configuration is convectively unstable. Possible treatments of such cases will be discussed elsewhere.} \]
In this case the Poisson equation for the gravitational potential itself is obtained as a Euler-Lagrange equation with respect to $\phi$. Note that the factor of 1/2 disappears from the second term, reflecting the fact that $\phi$ is an independent variable. The problem with this formula from a practical point of view is that the fourth term has a non-compact support, i.e., the integral region extends to infinity. This problem may be alleviated, however, by imposing an appropriate boundary condition at the stellar surface.

We employ the first formulation, in which the gravitational potential is treated as a functional of density through the energy functional given by equation (22), which is obtained once the node positions are determined appropriately. As for the gravitational potential $\phi$, equation (22) is solved numerically on the same triangulated mesh for the density obtained this way. It can be hence regarded also as a function of the coordinates of nodes.

2.3 Poisson equation for gravitational potential

As mentioned just now, in our formulation, the gravitational potential is regarded as a functional, or a function in the discretized version, of density by solving the Poisson equation. As also noted earlier in section 2.1, we employ the energy functional given by equation (22) to derive the discretized version of Poisson equation.

We first approximate the second and fourth terms on the right hand side of the equation on the triangulated mesh as

$$\sum_k \int_{\text{cell}} \left( \frac{1}{4\pi G} \frac{1}{2} \nabla \phi \cdot \nabla \phi + \phi \right) dV,$$

where the volume integral is done for each triangular cell and summed over all cells. The integrand is approximated linearly in each cell as

$$\phi = \alpha_1^\phi z + \alpha_2^\phi \varpi + \alpha_3^\phi,$$

where the coefficients, $\alpha$'s, are given in terms of the values of the gravitational potential at three nodes of the cell as

$$\alpha_a^\phi = T_{ab}^{-1} \phi_b.$$

In this expression, the subscripts $a$ and $b$ specify the node of the cell and the matrix $T_{ab}$ is determined by the geometry of the mesh alone and given in Appendix A. Then the derivative of $\phi$ in the integrand of equation (25) is approximated as

$$\nabla \phi = (\alpha_1^\phi, \alpha_2^\phi, 0).$$

Using this approximation, we evaluate the first term of equation (25) as

$$\sum_k \int_{\text{cell}} \nabla \phi \cdot \nabla \phi dV$$

$$= \sum_k \left( \alpha_1^\phi \alpha_1^\phi + \alpha_2^\phi \alpha_2^\phi \right) \int_{\text{cell}} dV$$

$$= \sum_k \sum_{a,b \in k} \left( T_{1a}^{-1} T_{1b}^{-1} + T_{2a}^{-1} T_{2b}^{-1} \right) \phi_a \phi_b V_{\Delta}.$$

The second term of equation (25) is, on the other hand, expressed as

$$\int_{\text{cell}} \phi \rho dV = \sum_i m_i,$$
nodes. Collecting the terms, we obtain the expression for equation (22) as
\[
\sum_{k} \sum_{a, b \in k} \frac{1}{2} \left( T_{1a}^{-1} T_{1b}^{-1} + T_{2a}^{-1} T_{2b}^{-1} \right) V_{\Delta} \cdot \phi_{a} \phi_{b} = \sum_{i} \phi_{i} m_{i}.
\]
(30)

The derivative of this expression with respective to \( \phi_{i} \) gives the discretized Poisson equation as
\[
\sum_{j} M_{ij} \phi_{j} = 4 \pi G m_{i}.
\]
Here we introduce a symmetric matrix \( M_{ij} \) defined as
\[
\sum_{i,j} M_{ij} \phi_{i} \phi_{j} = \sum_{k} \sum_{a, b \in k} \frac{1}{2} \left( T_{1a}^{-1} T_{1b}^{-1} + T_{2a}^{-1} T_{2b}^{-1} \right) V_{\Delta} \phi_{a} \phi_{b},
\]
in which the double sum on the left hand side runs over all nodes. Note that the matrix \( M_{ij} \) is determined by the mesh geometry alone. This approximate Poisson equation is solved algebraically as
\[
\phi_{i} = 4 \pi G M_{ij}^{-1} m_{j}.
\]
(32)

In so doing the boundary condition is imposed Dirichlet-type at the stellar surface with the values of the gravitational potential being calculated with the Green’s function (see below).

In order to validate the formulation, we compare it with another method. In fact, the Green’s function method, which is employed to calculate the boundary values of the potential, is actually applicable everywhere. Adopting the multipole expansion of the Green’s function, we write the potential as
\[
\phi_{i} = -2 \pi G \sum_{l=0}^{\infty} P_{l} (\cos \theta_{i}) \int_{0}^{\pi} \sin \theta' d\theta' P_{l} (\cos \theta')
\times \left( \int_{0}^{r_{i}} \left( \frac{r'}{r_{i}^{l}} \right)^{l+1} dr' + \int_{r_{i}}^{\infty} \frac{r'_{i}}{(r')^{l-1}} dr' \right) \rho_{i} (r', \theta'),
\]
(33)
where \( P_{l}' \)s are the Legendre polynomials. The integrals are evaluated numerically.

The comparison is done for a non-spherical density distribution shown in the left panel of Fig. 1. The triangulated mesh we deploy is displayed with the nodes and edges in the right panel. The mass \( m_{i} \) associated with node \( i \) is obtained from the density \( \rho_{i} \) at the node through equation (22). We evaluate the gravitational potential either via equation (32) or equation (33). The left and middle panels of Fig. 2 correspond to the numerical results for equations (32) and (33), respectively, which look almost identical to each other. As a matter of fact, the right panel of Fig. 2 demonstrates that the both results are in agreement within less than 1% errors. Here the error is defined as the absolute value of the ratio of the difference between the two results to the one obtained with equation (33).

2.4 Minimization of energy function

We employ a Monte Carlo technique to search for minimum energy function. The reason is the following. It is well known from the perturbation theory based on the Lagrangian displacement vector that a class of trivial Lagrangian dis-
We find that large deformations of triangular cells degrade accuracy of the energy function and in some cases generate a fictitious local minimum. To avoid such artifacts, we consider re-gridding as a solution as mentioned above. We apply a smoothing to the deformed portion of the grid, which has experienced more than 30% change in area by a single Monte Carlo sweep. We know empirically that such a large variation in area is commonly accompanied by the appearance of a very acute angle in the cell.

We introduce the residual in the virial relation defined as

\[ V_C = \left| \frac{2T + W + 3 \int PdV}{W} \right| , \tag{34} \]

in which \( T \) and \( W \) are the rotational and gravitational energies, respectively, and the third term is the volume integral of pressure. \( V_C \) is equal to zero in the exact theory and is a measure of numerical accuracy commonly used in the literature (e.g., Eriguchi & Mueller 1985), and we also adopt it together with energy convergence. In this paper, we have typically achieved \( V_C < 10^{-3} \) with ~500 grid points, which is comparable to previous studies where \( V_C \sim O(10^{-4}) \) was achieved (Eriguchi & Mueller 1985). It is possible to improve accuracy by deploying a larger number of nodes.

### 3 VALIDATION

In this section we apply our new method to some test cases: barotropic configurations in subsection 3.1 and baroclinic ones in subsection 3.2. We compare the resultant configurations with ones obtained by other Eulerian schemes. In subsection 3.3, we mention some technical issues such as the smoothing procedure.

#### 3.1 Comparison of barotropic equilibria

We begin with the barotropic case, in which pressure depends on density alone, \( P = P(\rho) \). We consider the isentropic configurations as a representative of barotropes. Here we compare our solutions with those given by Hachisu’s self-consistent field (HSCF) scheme (Hachisu 1986), a well-established numerical method for rapidly rotating barotropic configuration.

In this section, we consider two types of rotation laws. As repeatedly mentioned, the barotropic configuration rotates cylindrically, i.e., the iso-angular velocity surfaces are concentric cylinders.

- **(i) rigid rotation:**

  \[ \Omega^2(r) = \Omega_0^2, \tag{35} \]
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(ii) differential rotation:
\[ \Omega^2(r) = \frac{\Omega_0^2}{(\pi R_e)^2 + d^2}, \]

where \( \Omega(r) \) is an angular velocity at the position of \( r \). In equation (36), \( R_e \) is the equatorial radius, \( \Omega_0 \) is a parameter that specifies the overall magnitude of angular velocity, and \( d \) is a dimensionless parameter that determines the degree of differential rotation: as \( d \) increases, the angular velocity becomes uniform whereas in the opposite limit the rotational velocity becomes constant \( \) \cite{Eguchi & Mueller 1987}. We set \( d = 0.90 \) in the following.

It should be reminded that in our YFY scheme the distribution of angular velocity is not known a priori, since we assign a conserved specific angular momentum to each node of the triangulated mesh in the initial reference configuration, which is arbitrary, and the node position is changed later. This is not the case for the HSCF scheme, which is a Eulerian method and derives the density distribution on the fixed mesh, assuming from the beginning that the rotation is cylindrical. Note that even the cylindrical rotation is not guaranteed in the YFY scheme. It is hence a good diagnosis for our method. In order to make comparison between the two schemes, we take the following strategy: we first employ the HSCF scheme to obtain the rotational equilibrium for a given rotation law and cover the resultant configuration with a triangulated mesh and obtain the mass and specific angular momentum that should be assigned to each node; we then expand by 20 % in the radial direction and use it as the initial reference configuration for our YFY scheme. If it works properly, the mesh should return to the original shape after the minimization of the energy function. Incidentally we have confirmed that if the artificial expansion of the mesh is not administered, the mesh does not change by the minimization as it should.

One of such comparisons is shown in Fig. 3 where the density distributions obtained with the two schemes are presented for a rigidly rotation. Although HSCF scheme utilizes non-dimensional quantities are normalized with the maximum density \( \rho_0 \) and equatorial radius \( R_e \), we give here rather arbitrarily specific values \( \rho_0 \) and \( R_e \). As \( \rho_0 = 1.24 \times 10^8 \) g cm\(^{-3}\) and \( R_e = 2.57 \times 10^7 \) km in order to facilitate comparison. The ratio of poler radius \( R_p \) to the equatorial set to \( R_p/R_e = 0.80 \) in this model, which corresponds to the ratio of rotational energy to gravitational one of \( T/|W| = 3.7 \% \). As mentioned already, we assume a uniform entropy distribution in this model: the specific entropy is \( s = 14.7 \) K\(\text{g}^{-1}\) with \( K_B \) being the Boltzmann constant if the matter is composed of hydrogens alone.

The two configurations look quite similar to each other, which is also confirmed in the right panel in which relative differences are displayed as a contour color map: the density distributions agree with each other within an error of 5 % in most places. A closer inspection reveals, however, that the deviation gets larger near the surface, where the density itself is quite low. The vicinity of the rotation axis is another region, where the accuracy is compromised. This is due to the fact that our scheme is based on the variational principle and it is intrinsically difficult to determine the configuration of the regions that do not contribute much to the energy functional. Some multi-layer treatment may be needed to handle this problem.

In Fig. 4 we show how the energy function (left panel) and the virial residual (right panel) change as the Monte-Carlo sweep proceeds for the same model. The iteration process is terminated after 248 sweeps, at which point the value of the energy function is no longer lowered and, importantly, the virial residual is sufficiently small (\( < 10^{-3} \)) simultaneously. One recognizes that there are occasional glitches (three major ones and many minor ones) in the left panel, at which the value of the energy function rises discontinuously. There are corresponding discontinuities in the right panel. These irregularities are produced by what we call smoothing which is a reconfiguration of the mesh we conduct when the mesh gets distorted too much. Without such a measure, the search is trapped in local energy minimum as we describe later.

The corresponding change of the configuration is presented in Fig. 5. One can see a gradual shrink of the entire grid. Note that although the energy function looks settled to a minimum around 130 sweeps as shown in the left panel of Fig. 4 the configuration is far from rotational equilibrium. This is an example of the false local minimum just mentioned. The fact that this is a fake is reflected in the value of the virial residual given in the right panel of Fig. 5. This is the reason why we need to conduct the smoothing and the virial residual is an important diagnostic quantity to judge convergence.

Next, we present in Fig. 6 an example of the differential rotation given by equation (36). As in Fig. 5 the left panel shows the result by the YFY scheme, whereas the middle panel is for the HSCF scheme. We set again the central density and equatorial radius to \( \rho_0 = 1.24 \times 10^8 \) g cm\(^{-3}\) and \( R_e = 2.57 \times 10^7 \) km. The specific entropy is constant in space and the same as for the rigid rotation in the previous case. Then the total mass is also unchanged. The density distributions agree with each other within 5 % in most regions, but not near the surface again.

The behavior of the energy function (left panel) and virial residual (right panel) is qualitatively the same as what we saw in Fig. 4 for the rigid rotation. The iteration process is terminated after 248 sweeps. These panels are very similar to Fig. 6 though the rotation law is different.

In Fig. 7 the logs of the stellar structures for a differential rotation given by equation (36) are illustrated again. The structure changes from initial model—the results of HSCF scheme expanded by 20 % to the radial direction—to the hydro-static solution. The star continues to transform at 150 sweeps, although the energy function is converged in Fig. 7.

We also show the distributions of angular momentum in this model as \( \Omega_0 = 1.6 \times 10^{-3} \) rad s\(^{-1}\), and \( T/|W| \) is 4.3 %. As for the numerical accuracy, as we see some differences (~20%) between both schemes especially around rotational axis because of the division of zero or small values.

It is emphasized that this is not a trivial thing for YFY scheme. In fact, the rotation law is not known a priori in our formulation. The cylindrical rotation is not assumed but obtained as a result of computations. This is a strong vindication that our formulation works well indeed. We assume the other rotational laws including a rigid rotation in the reference configuration, and find in all cases cylindrical rotations in the outcomes.
3.2 Comparison of baroclinic equilibria

It is obvious that the EOS is generally baroclinic, i.e., pressure depends not only on density but also entropy and chemical abundance in stars. It is, then, necessary to handle such baroclinic equilibria for the study on the structures of rotating stars. Such investigations are not many in the literature, though (Uryu & Eriguchi 1994; Espinosa Lara & Rieutord 2002; Espinosa Lara & Rieutord 2013; Fujisawa 2013; Rieutord et al. 2016). In the stellar evolution calculation, the shellular rotation is commonly assumed as a rotation law. It is not obvious, however, whether there exist such hydrostatic equilibria or not, especially for rapid rotation. Recently, Fujisawa (2015), one of the authors of this paper, formulated a new self-consistent field scheme that is suitable for the construction of baroclinic equilibria. It is a Eulerian method and may be regarded as a (nontrivial) extension of the HSCF scheme we used in the previous subsection. It is probably superior to the formulation proposed in this paper in terms of accuracy. The advantage of the YFY scheme, however, is that it is a Lagrangian method and suitable to construct an evolutionary sequence out of an ensemble of rotational equilibria. Anyway, with two independent numerical schemes at hand to obtain baroclinic equilibria, it is a good opportunity for us to make comparisons and validate both of them simultaneously.

The baroclinicity is most easily introduced by an artificial modification of the entropy distribution. We employ the polytrope-like EOS, \( p = K \rho^n \) with \( K \) being position-dependent,

(i) spherical iso-\( K \) surfaces:

\[
K(r) = K_0 \left( 1 + \epsilon \frac{r^2}{R_e} \right),
\]

(ii) oblate iso-\( K \) surfaces:

\[
K(r) = K_0 \left( 1 + \epsilon_1 \left( 1 + \epsilon_2 P_2(\cos \theta) \right) \frac{r^2}{R_e} \right),
\]

where, the Legrendre polynomials are denoted by \( P_n \) with \( \theta \) being colattitude and employed to give non-sphericity to \( K \) distribution. Note that \( K \) corresponds to a (constant) specific entropy in the original polytropic EOS and, hence, the modifications above may be regarded as an introduction of entropy-dependence to pressure. Two dimensionless parameters \( \epsilon \) and \( \epsilon_1 \) specify radial non-uniformity whereas \( \epsilon_2 \) gives colatitudeal dependency. We set \( \epsilon = 0.35, \epsilon_1 = 0.45 \) and \( \epsilon_2 = 0.80 \) here. These models are highly artificial admittedly, and more realistic models, in which the entropy distributions are determined self-consistently with energy generations and...
Rotational equilibria by Lagrangian variational principle

Figure 5. (colour on-line). The change of configuration in the searching process for the solution given in Fig. 3.

Figure 6. (colour on-line). Same as Fig. 5 but for the differential rotation obeying the law given in equation (36). The values $\rho_0$, $R_e$, $s$, and $R_p/R_e$ are the same as those in Fig. 4.

Figure 7. (colour on-line). Same as Fig. 4 but for the differential rotation in Fig. 6. The iteration is terminated after 246 sweeps, at which point the energy functional is no longer lowered and the virial residual is sufficiently small $< 10^{-3}$.
radiative transfer, will be presented in the forthcoming papers.

In the FSCF scheme, the rotation law is given on the equatorial plane and the angular velocity distribution is solved self-consistently. We hence employ the rotation law given in equation (36) only on the equatorial plane. The resultant configurations are expanded radially by 20% and used as reference configurations for the YFY scheme just as in the previous sub section.

We first present the results for case (i) with the spherical isentropic surfaces given in equation (37). In Fig. 10, we show isopycnic color-contours, in which the left panel is for the YFY scheme and the middle panel for the FSCF scheme. Just as in the HSCF scheme, the FSCF scheme deals with non-dimensional quantities normalized with the maximum density $\rho_0$ and equatorial radius $R_e$. To facilitate the comparison with our scheme, we set $\rho_0 = 1.19 \times 10^7$ g cm$^{-3}$ and $R_e = 2.57 \times 10^5$ km. The values of $K_0$ and $\Omega_0$ chosen to be $5.49 \times 10^{13}$ in cgs unit and $1.9 \times 10^{-3}$ rad s$^{-1}$, respectively. This value of $\Omega_0$ is a few hundred times the solar angular velocity. We find again a reasonable agreement in most regions as is evident in the right panel of the figure, in which the relative difference is shown. Note that the number of the grid points employed in the FSCF method, $N_r = 513$ and $N_\theta = 257$, is much larger than that deployed in the YFY method, $N = 489$. It is also apparent from the same figure that the deviation gets larger in the vicinity of the surface for the reason mentioned earlier.

In Figs. 11 and 12 we present the values of the energy function and virial residual as a function of the number of sweeps and the corresponding change of the configuration, respectively, just as in the barotropic case. It is observed that the energy function is settled to a certain value after 130 sweeps. It is apparent from the values of virial residual
\( \sim 10^{-2} \) that this is a false minimum. The true one is reached at 261 sweeps after a couple of shuffles by the smoothings, which are marked by both major and minor spikes in the energy function shown in the left panel of Fig. 12. This demonstrates again that the virial residual is an important diagnostic measure for true minimum and the smoothing is an indispensable process to escape the trap of false minima.

It is well known that the stability of rotational equilibria against axisymmetric perturbations can be judged by the Solberg and Høiland criterion which is summarized as follows: stable configurations satisfy (i) the specific entropy \( s \), and (ii) the specific angular momentum \( j \) increases from the poles to the equator on the isentropic surface. The rotational configuration we obtained above satisfies these conditions as can be understood from Fig. 13, in which we display the iso-density contours; the arrows in the figure indicate the direction, in which the angular momentum increases.

There is another well-known rule we should care for: the Bjerkeness–Rosseland rule claims that if \( d\Omega/dz \) is positive at a given point, the isobaric surface is more oblate than the isopycnic one there (Tassoul 1978). This means in particular that if \( d\Omega/dz \) is positive everywhere, matter temperatures are lower at the poles than on the equator on each iso-baric surface (Tassoul 1978). The rule is also confirmed for the results presented above. The isobaric surfaces and iso-\( K \) (and, hence, iso-entropic) surfaces are shown in Fig. 14. As is evident from the figure, matter on each isobaric surface has indeed a smaller value of \( K \) at poles than on the equator. On the other hand, the gradient of specific angular momentum \( dj/dz \), and hence \( d\Omega/dz \) is also slightly positive as can be seen in Fig. 14.

We move on to case (ii), i.e., with oblate isentropic surfaces as specified in equation (38). What is important with this entropy distribution is that it leads to rotational equilibria with a shellular-type rotation law demonstrated by Fujisawa (2015). Figure 15 compares the isopycnic surfaces obtained with the YFY scheme (left panel) and for the FSCF scheme (middle panel). As quantified in the right panel, in which the relative difference is presented, the both schemes are in agreement within an error of \( \sim 5\% \) except near the surface as usual. The behavior of the energy function and virial residual as well as the configuration itself is shown in Figs. 11 and 12. It is quite similar to what we found for case (i) as well as for the baroclines. It is observed that the energy function is trapped erroneously to a local minimum around 130 sweeps as suggested by not-so-small values of the virial residual. The true minimum is reached only after 275 sweeps, where \( V_C < 10^{-3} \) is satisfied.

We show in Fig. 18 the angular velocity distributions obtained with the two methods for the same model. It is obvious that they are not cylindrical but of shellular-type. The relative difference between the results obtained with the two schemes is \( \sim 5\% \) except near the rotation axis as shown in the right panel. As should be apparent from the left panel, the numerical interpolation on the triangulated grid that is employed by a graphic software in plotting the contour certainly contributes to the discrepancy. The relative differences for the specific angular momentum and entropy are given in the lower panels of Fig. 18. The trends are quite similar to those for the angular velocity and density, respectively.

As mentioned repeatedly, the numerical construction of rotational equilibria with the shellular-type rotation law has important implications for stellar evolution theory, since most of the realistic calculations done so far have assumed shellular rotation (e.g. Meynet & Meynet 1997; Talon et al. 1994; Heger et al. 2000; Mathis et al. 2004; Potter et al. 2012). Talon et al. 1997; Heger et al. 2000; Mathis et al. 2004; Potter et al. 2012) are notable exceptions, their formulations are Eulerian, and will be unsuitable for stellar evolution, which could be a major break-through for the fully two-dimensional calculation of stellar evolution.

The entropy distributions and the iso-angular moment of the configuration presented in Fig. 15 are shown in Fig. 19. Obviously they satisfy the two conditions of the Solberg and Høiland criterion. In contrast to the previous case with the spherical isentropic surfaces, \( d\Omega/dz \) is negative everywhere in the current case with the oblate isentropic surfaces as found in Fig. 18. As expected and confirmed in Fig. 20, the values of \( K \) are larger at poles than on the equator this time. This model is hence also consistent with the Bjerkeness-Rosseland rule.
Figure 10. (colour on-line). Same as Fig. 3 but for a baroclinic configuration with the spherical isentropic surfaces given in equation (37). The maximum density and equatorial radius normalization factors $\rho_0$ and $R_e$ on FSCF scheme are set to $\rho_0 = 1.19 \times 10^2$ g cm$^{-3}$ and $R_e = 2.57 \times 10^5$ km, respectively, and the constant $K_0$ in equation (37) is chosen to be $5.49 \times 10^{13}$ in cgs unit.

Figure 11. (colour on-line). Same as Fig. 4 but for the baroclinic configuration presented in Fig. 10. The iteration is terminated at 261 sweeps.

Figure 12. (colour on-line). Same as Fig. 5 but for the baroclinic configuration presented in Figs. 10 and 11.
Figure 13. (colour on-line). (Upper panels) iso-angular momentum contours on top of the entropy color maps for the baroclinic configuration shown in Figs. [10][11]. The results for the YFY scheme and the FSCF scheme are displayed in the left and right panels, respectively. The arrows indicate the direction, in which the angular momentum increases. (Lower panels) the relative differences of specific angular momentum (left panel) and entropy (right panel) between the results obtained with the schemes.

Figure 14. (colour on-line). (Upper panels) iso-$K$ surfaces on top of the pressure color-contours for the baroclinic configuration in Figs. [10][13]. The arrows indicate the direction, in which $K$ increases. The left and right panels correspond to the results for the YFY and FSCF schemes, respectively. (Lower panels) the relative differences of pressure (left panel) and $K$ (right panel) for the results from the two schemes.
Figure 15. (colour on-line). Same as Fig. 10 but for the oblate isentropic surfaces given in equation (38). The maximum density and equatorial radius are set to $\rho_0 = 1.19 \times 10^2$ g cm$^{-3}$ and $R_e = 2.57 \times 10^5$ km, respectively.

Figure 16. (colour on-line). Same as Fig. 11 but for the configuration shown in Fig. 15. The iteration is terminated at 275 sweeps.

Figure 17. (colour on-line). Same as Fig. 12 but for configuration displayed in Figs. 15 and 16.
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3.3 Reference configuration and smoothing

In the application of the YFY scheme, we have always constructed the reference configurations from the configurations obtained with the HSCF or FSCF scheme, so far, expanding them radially by a certain factor. This may raise a concern that the reference configuration must be prepared in this particular way initially for our scheme. In order to demonstrate that this is not the case in fact, we try three more different constructions and see if the final configurations are the same or not.

To ensure that all models have the same angular momentum distribution in the final configuration, we still need to prepare the reference configurations by deforming this final configuration. Although it does not matter what configuration is employed for the latter, we adopt the model with the shellular-type presented in Figs. [15] [18] [19] and [20]. This time we do not expand but shrink the final configuration for this model by 30% in three different ways: (A) radially, (B) horizontally, and (C) vertically. To the three reference configurations thus obtained, we apply the YFY scheme and see if the same equilibrium, which should be also identical to the configuration shown in Fig. [17], is reached.

We show the changes of the configurations with the Monte Carlo sweeps in Figs. [21]-[23], which correspond to deformations (A)-(C) respectively. The upper left panels in these figures exhibit the reference configurations, which are clearly different from the final configurations shown in the lower right panels. In particular, the reference configuration is prolate for case (B) whereas the rotational equilibrium is oblate owing to centrifugal forces. It is evident from the comparison of the lower right panels in the figures that the

Figure 18. (colour on-line). The left and middle panels show the angular velocity distributions of shellular-type obtained for the configuration shown as in Figs. [15] - [17] with the YFY and FSCF schemes, respectively. The right panel exhibits the relative difference.

Figure 19. (colour on-line). Same as Fig. [18] but for the configuration presented in Figs. [15]-[18].
Figure 20. (colour on-line). Same as Fig. 14 but for the oblate isentropic surfaces given in equation (38).

resultant configurations after the optimizations look almost identical with each other as well as with the original configuration given in Fig. [14]. We, hence, conclude that our scheme is robust to the variation of reference configuration. It should be also noted that in the application to the stellar evolution calculation, we will be able to use the stellar structure obtained at the previous time step as the reference configuration, which should be hence pretty close to the true equilibrium.

It is one of the great advantages with the YFY scheme based on the Lagrange coordinates that we can record the history of the fluid element assigned to each node. This is true not only for the Monte Carlo sweep as has been demonstrated so far but also for the true evolution in time when the scheme is applied to the 2-dimensional calculation of stellar evolution. Since nuclear burnings occur locally in each fluid element, the chemical evolution is most easily treated on the Lagrange coordinates and hence by the YFY scheme. It should be also mentioned that since the specific entropy would be conserved for each fluid element in the absence of the generation and transport of heat, radiative transport of energy will be formulated most unambiguously on the Lagrange coordinates. This will be also the case for the transport of angular momentum. Convective transport of energy and angular momentum may be handled as well if it is justified to treat them approximately as diffusions.

The left three panels of Fig. 24 present the histories of the values of the energy function for the cases shown in Figs. 21-23. The values of the energy function decrease in most of the time in the Monte Carlo sweep. Sporadical small glitches are due to the smoothings, which are necessary, as repeatedly mentioned, to ensure the convergence to the true minimum, avoiding the trap of false minima.

Shown in the right panels of Fig. 24 are histories of the virial residuals for the same models. In the previous sections, we used the condition $V_C < 10^{-3}$ as one of the diagnostics to judge if the equilibrium is reached as described in section 2. This condition works well also in the current cases as can be seen from the comparison of the right panels in Fig. 24 with Figs. 21-23. The importance of the criterion is also understood from Fig. 16, in which the virial residual is still large, $V_C > 10^{-3}$, after $\sim 120$ iterations although the energy functional appears to have hit the minimum. The configuration is still out of equilibrium indeed, which can be confirmed in the corresponding structure shown in Fig. 17.

It should be noted, however, that the condition on the virial residual alone is not sufficient for the judgement, since the virial residual is an integrated quantity and may become small accidentally. For example, the configuration presented in the upper-middle panel in Fig. 23 for the 190th iteration has not yet reached the equilibrium although the virial residual satisfies $V_C < 10^{-3}$ as shown in Fig. 24. It is hence mandatory to take into account the convergence of energy function in addition to the condition on the virial residual. Considering the smoothing administered every $\sim 100$ sweeps (see the last paragraph of this section), we impose the condition that the energy function at the $i$-th iteration, $E(i)$, should not lower than $E(i - 100)$ after $i - 100$th step. Although this simple criterion seems to work well at least for the models considered in this paper, it may need improvement.

In this paper it is not our purpose to construct a se-
quence of rotational configurations with different angular
momenta up to the mass shedding limit. As a matter of
fact, the Monte Carlo method employed in this paper to
find a configuration that gives the minimum to the energy
functional is not suited very much for finding the limiting
configuration for two reasons. Firstly, in the Monte Carlo
method many (non-equilibrium) configurations are tried to
reach the true equilibrium and in so doing some trial con-
fignurations give energies lower than the true minimum er-
roneously owing to numerical errors more often than not,
which leads to a “numerical” mass shedding and no conver-
gence. Secondly, the equilibrium configuration may not be
unique for a given distribution of specific angular momen-
tum on the Lagrangian coordinates. It is well known in fact
that the angular momentum is not a monotonic function of
the ratio of the polar to equatorial radii, $R_p/R_e$, near the
mass shedding limit (see, for example, Table 3 in Fujisawa
(2015), in which the angular momentum increases initially
as the ratio decreases but it decreases after the peak that
appears at $R_p/R_e \sim 0.55$). In this regime we are afraid that
the Monte Carlo method may not work properly. As a mat-
ter of fact, we have confirmed for a model with $R_p/R_e = 0.5$
that this is indeed the case. We emphasize, however, that
our formulation can handle rapid rotators as demonstrated
in Fig. 25, in which we display a rotational configuration
with the axis ratio of $R_p/R_e = 0.60$, which we find is es-
sentially the same as the result obtained with the FSCF
method. Incidentally, this model is in the regime, where the
solution is unique in the above sense.

At the end of section, we would like to discuss the im-
portance of the smoothing. We employ the variational prin-
ciple to obtain the hydrostatic equilibria. The idea is sim-
ple, but the implementation is not indeed. Without an ap-
propriate smoothing, Monte Carlo sweeps tend to make too
deformed triangles, which lead the generation of a false local
energy minimum. In Fig. 26 we show the result of the calcu-
lation without the smoothing of acute-angled triangulations.
There is no glitch in the panels of the energy function nor
of the virial residual, since we do not take into account the
smoothing. The energy function is settled after 1000 sweeps,
although the virial residual does not satisfy $V_C < 10^{-3}$. This
result suggests that the search of minimum energy is trapped
in a local minimum. The resultant structure is far from the
final results shown in Figs. 21-23. This is the reason why the
smoothing process is necessary.

There will be many possible ways to alleviate highly
acute triangles, which could be the origin of false local min-
ima. Followings are the simple measures we employed in this
study:

1. if there is a triangular cell that experiences more than
30% of change in area after a shift of a node during
the Monte Carlo sweep, we move this particular node of
the triangle to the center of the polygon formed with its
nearest neighboring nodes.

2. since the local shifts of some nodes in the first step some-
times produce zigzag distributions of the nodes that are
initially located on one of the concentric surfaces (see
the construction of the triangulated mesh in the refer-
ence configuration), we make them somewhat smoother
by moving the grid on the top or bottom of the zigzag dis-
tribution toward the middle of the two neighboring nodes;
the first step is always followed by the second step.

3. in rare cases, the order of the grid points on the axis
or on the equator is changed by their large shifts in the
Monte Carlo sweep; then we restore the original order
by pulling back the grids in trouble toward the original
positions appropriately; this measure is take only when
it is needed.

4. even if we do not find any large areal-change, we admin-
ister the smoothing prescribed in (1) to all nodes period-
ically; from our experience it should be done every $\sim 100$
($\sim$ the number of grid points / 5) sweeps; although this
is mainly meant to be a preventive measure, it is also
effective for the escape from a local minimum.

The above prescriptions are admittedly empirical and de-
pend on the construction of the triangulated mesh as well as
the direction of Monte Carlo sweep. Although there should
be more sophisticated and systematic ways to treat the false
local minimum, we defer a further improvement to the fu-
ture work, since the measures given above appear to work
well at least for the models considered in this paper.
Figure 21. (colour on-line). Same as Fig. [17] but for the reference configuration given by the deformation for case (A).

Figure 22. (colour on-line). Same as Fig. [21] but for case (B).
4 SUMMARY AND DISCUSSIONS

We have developed a new formulation to obtain rotational equilibria numerically. The scheme can handle not only barotropic but also baroclinic configurations, which are critically important for the application to realistic stars. Such an achievement is itself a major break through to the status quo, in which previous works are not many and, more importantly, they are all based on the Eulerian description (Uryu & Eriguchi 1994; Uryu & Eriguchi 1995; Espinosa Lara & Rieutord 2007; Espinosa Lara & Rieutord 2013; Rieutord et al. 2016). Our method, on the other hand, employs the Lagrangian description just as in one-dimensional counterparts for non-rotating stars and hence stellar evolution calculations, since we can trace the potentially complex movements of each fluid element. Our formulation is based on the variational principle, in which rotational equilibria are obtained as the configurations that optimize the energy functional for given distributions of mass, specific entropy, and angular momentum on the Lagrangian coordinates. In this paper all physical quantities are discretized on triangulated grids.

In order to validate our scheme, we compare the configurations obtained by our scheme with those by other Eulerian schemes: the HSCF scheme developed by Hachisu (Hachisu 1984) for the barotropic configuration, and the FSCF scheme conceived by Fujisawa (Fujisawa 2013) more recently for the baroclinic one. We have confirmed that all the configurations including the ones with shellular-type rotations obtained in this paper are linearly stable against axisymmetric perturbations and comply with the Bjorkness-Rosseland rule.

In this comparison, we have reproduced the equilibria a shellular-type rotation, which were obtained only recently by Fujisawa although the shellular rotation is commonly assumed in the one-dimensional stellar evolution calculation with rotation being taken into account only approximately. It is found that the result obtained with the YFY scheme agree with those derived by other schemes with an error of $\sim 5 \sim 10\%$, which we think is reasonable, considering the relatively small number of nodes (489) we used in this paper. It is then a legitimate question to ask how the accuracy is improved with the number of nodes. In order to address this issue, we have conducted additional computations, changing the node number from $\sim 200$ to $\sim 1000$, the latter of which is the maximum node number we can afford at present, since the numerical cost is proportional to $N^2$ in our currently unparallelized code, where $N$ is the node number.

We find that the higher resolutions with $\sim 800$ and $\sim 1000$ nodes do not improve the accuracy at a recognizable level as shown in the leftmost panel of Fig.27. As demonstrated for the highest resolution model in the middle and rightmost panels of this figure, as the final configuration is approached, the values of energy function and virial residual repeat a sudden jump followed by a settlement to some values which satisfy our convergence criteria. This is due to the smoothing that we administer regularly to avoid the trapping by false minima (see the last part of the last section). Because of the probabilistic nature of the Monte Carlo method, the final value of the virial residual after each repetition fluctuates around the mean value. In the left panel of Fig.27 we present these mean values and variances as dots and bars, respectively, for different node numbers.

The reason why the precision is not improved is the following: in our formulation we search for the rotational configuration that gives the lowest value to the energy functional for given mass, specific entropy and angular momentum; it is not surprising then that some nodes contribute more than others; in particular, the nodes close either to
Figure 24. (colour on-line). Same as Fig. 16 but for the reference configurations given by the deformation for cases (A)-(C).

Figure 25. (colour on-line). The density distribution (left panel), and the edges and nodes (right panel) for shellular-type configuration for $R_p/R_e = 0.60$ obtained with the YFY scheme. The result is essentially the same as the one derived with the FSCF scheme. The unit of density is given in cgs. The central density, equatorial radius are $\rho_0 = 1.51 \times 10^2$ g cm$^{-3}$, $R_e = 2.57 \times 10^5$ km, respectively.
See also Kiuchi et al. (Kiuchi et al. 2010) combined with the spectral method, achieves very high accuracy.

The global equilibrium, we need to find an equilibrium serving as fixed gravitational potentials. In order to obtain be applied to each region individually, with other regions regions. Then the minor contributions are no longer minor. If the multi-layer treatment is employed, these regions, whose contributions to the energy functional are minor. If the multi-layer treatment is employed, these minor regions can be treated separately from the major regions, since they have either small densities or volumes; this means in turn that their positions are very difficult to obtain accurately; this may be understood if one scrutinizes the right panels of Figs. 2, 3, 6, 9, 10, 15, 18 as well as the lower panels of Figs. 14, 19, 20; on the hand, these nodes give contributions of the order of $10^{-4}$ to the virial residual.

In principle, if the number of nodes is sufficiently large particularly in the regions of our concern, then the virial equality should be improved. The current resolutions we can afford are way too small, though, as can be understood from the figures showing the node configurations. This may be remedied if we implement the multi-layer treatment.

Multi-layer treatment may be useful to improve the accuracy, which will be also required in applying the new scheme to rotating stars in advanced evolutionary stages. As explained earlier, the numerical error comes mainly from the regions, whose contributions to the energy functional are minor. If the multi-layer treatment is employed, these minor regions can be treated separately from the major regions. Then the minor contributions are no longer minor. The important thing is that the variational principle can be applied to each region individually, with other regions serving as fixed gravitational potentials. In order to obtain the global equilibrium, we need to find an equilibrium configuration in each region consecutively and iteratively until all these configurations do not change any longer simultaneously. In addition to the minor regions mentioned above, the central region may also need to be treated with a special care, since the nodes are rather sparse there as can be seen, e.g., in the right panel of Fig. 1.

We have demonstrated that our scheme is robust, obtaining the same final configuration in equilibrium irrespective of the reference configurations assumed initially. We have also observed, on the other hand, that the Monte Carlo sweeps to get to the minimum of the energy functional are prone to be trapped by false local minima, which are generated by deformations of the numerical grid and that the smoothing should be properly administered to escape them. In order to distinguish the true minimum from a false one, the virial residual is found to be a good diagnostic. Although we have not made any attempt in this paper to reconstruct a triangulated mesh when it becomes too deformed, such re-gridding will be necessary when applying the YFY scheme to the evolution of rotating stars.

The next step toward the application to the stellar evolution calculation will be to incorporate radiation transport in rotational equilibria. Unlike the previous works (Uryu & Eriguchi 1994; Uryu & Eriguchi 1995; Roxburgh 2006; Espinosa Lara & Rieutord 2007; Espinosa Lara & Rieutord 2013; Rieutord et al. 2016), we will solve time-dependent diffusion equation on the same triangulated grid, which will not be a difficult task. The meridional circulation is another important ingredient (Zahn 1989; 1992) when considering the application of our scheme to actual rotating stars, since it is supposed to play a major role.

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**Figure 26.** (colour on-line). The values of the energy function (left panel) and of the virial residual (middle panel) with initial model (A), but without the smoothing against acute-angled triangles. The right panel corresponds to the resultant structure at 1000-th iteration number.

**Figure 27.** (colour on-line). The range of converged virial residual for each resolution (left panel). The middle and the right panels show an example of the energy functional and the virial residual with 1073 nodes.

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\[ \text{the surface or to the rotational axis are the least contributors, since they have either small densities or volumes; this means in turn that their positions are very difficult to obtain accurately; this may be understood if one scrutinizes the right panels of Figs. 2, 3, 6, 9, 10, 15, 18 as well as the lower panels of Figs. 14, 19, 20; on the hand, these nodes give contributions of the order of } 10^{-4} \text{ to the virial residual.} \]

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\[ \text{Multi-layer treatment may be useful to improve the accuracy, which will be also required in applying the new scheme to rotating stars in advanced evolutionary stages. As explained earlier, the numerical error comes mainly from the regions, whose contributions to the energy functional are minor. If the multi-layer treatment is employed, these minor regions can be treated separately from the major regions. Then the minor contributions are no longer minor. The important thing is that the variational principle can be applied to each region individually, with other regions serving as fixed gravitational potentials. In order to obtain the global equilibrium, we need to find an equilibrium configuration in each region consecutively and iteratively until all these configurations do not change any longer simultaneously. In addition to the minor regions mentioned above, the central region may also need to be treated with a special care, since the nodes are rather sparse there as can be seen, e.g., in the right panel of Fig. 1.} \]

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\[ \text{The next step toward the application to the stellar evolution calculation will be to incorporate radiation transport in rotational equilibria. Unlike the previous works (Uryu & Eriguchi 1994; Uryu & Eriguchi 1995; Roxburgh 2006; Espinosa Lara & Rieutord 2007; Espinosa Lara & Rieutord 2013; Rieutord et al. 2016), we will solve time-dependent diffusion equation on the same triangulated grid, which will not be a difficult task. The meridional circulation is another important ingredient (Zahn 1989; 1992) when considering the application of our scheme to actual rotating stars, since it is supposed to play a major role.} \]
in the re-distribution of angular momentum and elements (e.g. Mathis [2000]. Although such motions cannot be handled directly with our scheme, they may be incorporated either as advections or as diffusions as just as in current 1D calculation.

Convection is even more difficult to treat. The redistributions of angular momentum and elements as well as entropy in this case may be approximated as diffusions or advections like the meridional circulation but the real difficulty with the convection is the fact that the convectively unstable configurations do not correspond to the minimum of the energy functional but to a saddle point. A couple of ideas are currently being tested and will be reported elsewhere.

Since our scheme is applicable only to axisymmetric configurations, in which the specific angular momentum is conserved in each fluid element, extension of our formalism to 3D configurations such as triaxial equilibria (Tassoul 1978) is much beyond the scope of this paper.

Possible applications of our scheme are not limited to the ordinary stars, though. They will be extended to e.g., compact stars, proto-stars and proto-planets to mention a few. Incorporation of magnetic fields and/or general relativity should be considered in due course.

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APPENDIX A: TRANSFORMATION MATRIX

In this appendix we give the transformation matrix $T_{lm}$ that is used in the expression of the discretized energy functional, equation (22). It is actually introduced in association with the interpolation of physical quantities in each triangular cell.

Suppose that a quantity $X$ is given by on all nodes and we interpolate them in each triangular cell as

$$X = \alpha_1^X z + \alpha_2^X \zeta + \alpha_3^X,$$ (A1)

in which $\zeta$ and $z$ are the cylindrical coordinates of an arbitrary point in the cell and $\alpha$’s are the expansion coefficients to be determined. Evaluating the above equation at three apexes (labeled with $l$) of the $n$-th triangular cell, one obtains the following equations:

$$X_l(n) = T_{lm}(n)\alpha_m^X,$$ (A2)

where $l, m = 1, 2, 3$. The matrix $T(n)$ in this expression is referred to the transformation matrix in this paper, in which defined as

$$T(n) = \begin{pmatrix} z_1(n) & \zeta_1(n) & 1 \\ z_2(n) & \zeta_2(n) & 1 \\ z_3(n) & \zeta_3(n) & 1 \end{pmatrix}.$$ (A3)

This matrix is useful in the finite element method employed here. For example, the volume of the triangular cell $V_\Delta$ is evaluated as

$$V_\Delta = \int_{\text{cell}} 2\pi \zeta d\zeta dz = 2\pi \frac{|J|}{2} \frac{\zeta_1 + \zeta_2 + \zeta_3}{3}. \quad (A4)$$

Here $|J|$ is the Jacobian for the diffeomorphic mapping $f$ of an isosceles right triangle constructed by the unit fundamental vectors $e_x, e_y$ of the two-dimensional Cartesian coordinates to the triangle with three apexes $X_1X_2X_3$ as shown in Fig. A1. The Jacobian is then equal to the determinant $|T(n)|$. This is obtained from the following relation between two forms: Here, the areal element under the mapping is shown as

$$du \wedge dv = J(f)dx \wedge dy,$$ (A5)

where $u$ and $v$ are the vectors $\vec{X}_1\vec{X}_2$ and $\vec{X}_1\vec{X}_3$, respectively. $|T(n)|/2$ in equation (A4) hence is the area of the triangular-cell whereas $(\zeta_1 + \zeta_2 + \zeta_3)/3$ is the $\zeta$-coordinate of the geometrical centre of the triangle.

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