

Marginally-Stable Thermal Equilibria of Rayleigh-Bénard Convection

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Natural convection is ubiquitous throughout the physical sciences and engineering, yet many of its important properties remain elusive. In this investigation, we derive and solve a quasilinear form of the Rayleigh-Bénard problem by representing the perturbations in terms of marginally stable eigenmodes. The amplitude of each eigenmode is determined by requiring that the background state maintains marginal stability. The background temperature profile evolves due to the advective flux of each eigenmode, as well as diffusion. The entire calculation is one-dimensional, and can be run on a workstation. We find the background temperature field evolves to an equilibrium state, where the advective flux from the marginally-stable eigenmodes and the diffusive flux sum to a constant. These marginally-stable thermal equilibria (MSTE) are exact solutions of the quasilinear equations. The mean temperature profile has thinner boundary layers and larger Nusselt numbers than thermally-equilibrated 2D and 3D simulations of the full nonlinear equations. We find the Nusselt number scales like $Nu \sim Ra^{1/3}$. When an MSTE is used as initial conditions for a 2D simulation, we find that Nu quickly equilibrates without the burst of turbulence often induced by purely conductive initial conditions, but we also find that the kinetic energy is too large and viscously attenuates on a long viscous time-scale.

I. INTRODUCTION

Rayleigh-Bénard convection plays a foundational role in astrophysical and geophysical settings. The resulting buoyancy-driven flows regulate heat transfer and generate large-scale vortices [1]. Turbulent convection, which is associated with large Rayleigh numbers Ra , is difficult to simulate. State of the art simulations performed by [2] have reached $Ra \sim 10^{14}$ but estimates for the sun's convective zone and earth's interior are $Ra \sim 10^{16} - 10^{20}$ and $Ra \sim 10^{20} - 10^{30}$ respectively [3, 4]. The scaling behavior of the Nusselt number $Nu \sim Ra^\beta$ in the asymptotic ultimate regime is of particular interest. There is a substantial body of work pertaining to this specific topic with no general agreement between numerical simulations and asymptotic theories [5–11].

Absent solid evidence from direct numerical simulation, other methods have been developed to try to infer large- Ra behavior or otherwise gain insight. In the presence of other physical effects (e.g., rotation, magnetic fields), one can sometimes derive an asymptotically consistent set of reduced equations [12, 13]. Reduced models are useful because they allow us to study the problem with less expensive computations. Another approach relates to unstable exact coherent states (ECS) which are steady solutions to the full nonlinear problem [14–17]. Simulations and analyses performed by [18, 19] suggest that chaotic solution trajectories might “visit” these ECS. Should that be the case, it is crucial that we discover and classify such equilibria.

Others have turned to studying quasilinear systems. The quasilinear approximation starts with a decomposition of all variables into a background and perturbations about this background. This approximation neglects the nonlinear interactions between perturbations which mod-

ify the perturbations themselves [20]. This renders the perturbation equations linear. Although the quasilinear approximation greatly simplifies the problem, an additional condition must be imposed to determine the amplitude of the perturbations. In ref. [21], researchers compute ECS in parallel shear flows by deriving and solving a quasilinear formulation of the Navier-Stokes equations via multi-scale asymptotic arguments. They assume the background velocity evolves on a slow time-scale, and, to determine the perturbation amplitudes, they require marginal stability at each timestep. A similar strategy is employed by ref. [22] to study acoustic streaming. In that work, an analytic expression for the first-order perturbation's amplitude is found by deriving a solvability condition.

In this paper we derive and solve a quasilinear form of the Rayleigh-Bénard convection problem using a similar time-scale separation argument. In section II we recall the underlying equations, and in section III we describe how we evolve the background temperature profile while maintaining marginal stability. Section IV describes the properties of the marginally-stable thermal equilibria, in particular how the Nusselt number and characteristic wavenumbers vary with the Rayleigh number. Finally, we describe the results of simulations initialized with marginally-stable thermal equilibria in section V, and conclude in section VI.

II. MODEL SETUP

We begin with the Boussinesq approximation for Rayleigh-Bénard Convection, nondimensionalized on the freefall time-scale. The domain \mathcal{D} is 2-dimensional, rectangular, and horizontally periodic with cartesian spatial

coordinates $0 \leq x < 4$ and $-1/2 < z < 1/2$. The fluid of interest is constrained between two flat boundaries at $z = -1/2$ and $z = 1/2$ with fixed temperatures $1/2$ and $-1/2$ respectively. At both boundaries we specify impenetrable, no-slip conditions, such that the velocity $\mathbf{u} = u\hat{x} + w\hat{z} = \mathbf{0}$ at $z = \pm 1/2$, where \hat{x}, \hat{z} are the unit vectors in the x and z directions. The equations of motion are then given by

$$\nabla \cdot \mathbf{u} = 0 \quad (1)$$

$$\frac{\partial \mathbf{u}}{\partial t} + \mathbf{u} \cdot \nabla \mathbf{u} = -\nabla p + T\hat{z} + \mathcal{R}\nabla^2 \mathbf{u} \quad (2)$$

$$\frac{\partial T}{\partial t} + \mathbf{u} \cdot \nabla T = \mathcal{P}\nabla^2 T \quad (3)$$

where p is pressure and T is temperature. Any system of this form can be characterized by its dimensionless Rayleigh number $\text{Ra} = \frac{g\alpha L^3 \Delta T}{\nu\kappa}$ and Prandtl number $\text{Pr} = \frac{\nu}{\kappa}$, where $g, \alpha, L, \Delta T, \nu, \kappa$ are the gravitational acceleration, coefficient of thermal expansion, domain height, opposed temperature difference, kinematic viscosity, and thermal diffusivity respectively. In this paper, we fix $\text{Pr} = 1$. For convenience, we define

$$\mathcal{R} = \sqrt{\frac{\text{Pr}}{\text{Ra}}}, \quad \mathcal{P} = \frac{1}{\sqrt{\text{Pr}\text{Ra}}}. \quad (4)$$

To derive the quasilinear form, we posit that an arbitrary field f can be represented as the sum of a mean profile (denoted by \bar{f}) and a perturbation function (denoted by f').

$$\mathbf{u}(x, z, t) = \mathbf{u}'(x, z, t) \quad (5)$$

$$= u'(x, z, t)\hat{x} + w'(x, z, t)\hat{z} \quad (6)$$

$$T(x, z, t) = \bar{T}(z, t) + T'(x, z, t) \quad (7)$$

$$p(x, z, t) = \bar{p}(z, t) + p'(x, z, t). \quad (8)$$

where the mean-velocity components vanish due to incompressibility and symmetry. Perturbations are defined to have no horizontal-average

$$\langle f'(x, z, t) \rangle_x \equiv \int_0^4 f'(x, z, t) dx = 0. \quad (9)$$

Substituting (7) into (3) and taking the horizontal-average reduces the system to a simple initial value problem (IVP) for \bar{T}

$$\frac{\partial \bar{T}}{\partial t} + \frac{\partial}{\partial z} \langle w'T' \rangle_x = \mathcal{P} \frac{\partial^2 \bar{T}}{\partial z^2}, \quad (10)$$

with associated boundary conditions $\bar{T}(-1/2, t) = 1/2$ and $\bar{T}(1/2, t) = -1/2$. It should be noted that we could obtain a similar IVP for u by breaking symmetry and considering some nontrivial mean horizontal flow $\bar{u}(z, t)$. However we must have $\bar{w}(z, t) = 0$ due to incompressibility.

Substituting (8) into (2) and taking the horizontal average reveals that the mean pressure field $\bar{p}(z)$ must satisfy

$$0 = -\frac{\partial \bar{p}}{\partial z} + \bar{T} \quad (11)$$

To solve (10) numerically, we need an expression for the perturbations so we can calculate the advective heat flux. Here we will make the quasilinear approximation, dropping the $\mathbf{u}' \cdot \nabla \mathbf{u}'$ and $\mathbf{u}' \cdot \nabla T'$ terms from the evolution equations for the perturbations. Substituting (5)–(8) into (1)–(3) followed by subtracting (10) & (11) from the resulting temperature and \hat{z} momentum equations gives

$$\nabla \cdot \mathbf{u}' = 0 \quad (12)$$

$$\frac{\partial \mathbf{u}'}{\partial t} = -\nabla p' + T'\hat{z} + \mathcal{R}\nabla^2 \mathbf{u}' \quad (13)$$

$$\frac{\partial T'}{\partial t} + \frac{\partial \bar{T}}{\partial z} w' = \mathcal{P}\nabla^2 T' \quad (14)$$

with Dirichlet boundary conditions

$$T'|_{z=\pm\frac{1}{2}} = 0, \quad \mathbf{u}'|_{z=\pm\frac{1}{2}} = 0, \quad p'|_{z=\pm\frac{1}{2}} = 0. \quad (15)$$

This is now a linear problem in \mathbf{u}' and T' which can be solved as an eigenvalue problem.

In his groundbreaking report [23], Lord Rayleigh observed that (12)–(14) can be manipulated into a separable form with generalized solutions

$$w'(x, z, t) = A \Re \left[W(z) e^{i(k_x x - st)} \right], \quad (16)$$

$$u'(x, z, t) = A \Re \left[U(z) e^{i(k_x x - st)} \right], \quad (17)$$

$$T'(x, z, t) = A \Re \left[\theta(z) e^{i(k_x x - st)} \right], \quad (18)$$

$$p'(x, z, t) = A \Re \left[P(z) e^{i(k_x x - st)} \right], \quad (19)$$

where A is the (undetermined) mode amplitude, $s = \omega + i\sigma$ and k_x is constrained, by periodicity, to the countably infinite set (spectrum) of wavenumbers

$$k_x \in \left\{ \frac{n\pi}{2} \mid n \in \mathbb{N} \right\}. \quad (20)$$

We normalize the eigenmodes to have

$$\langle |\theta|^2 \rangle_{\mathcal{D}} = 1 \quad (21)$$

where $\langle \cdot \rangle_{\mathcal{D}}$ denotes the spatial mean over the entire domain.

For each k_x , we can assess the stability of the perturbations by solving for the eigenvalue s , whose imaginary component σ plays the role of an exponential growth rate. Positive eigenvalues indicate that the system is unstable to small disturbances of wavenumber k_x , while negative eigenvalues indicate stability. A complete linear stability analysis requires solution over the full spectrum of

wavenumbers. The prototypical case is used to demonstrate that the critical Rayleigh number $Ra_c = 1708$ when $\frac{\partial \bar{T}}{\partial z} = -1$.

To calculate the advective heat flux in equation 10, we can sum $\langle w'T' \rangle_x$ from each horizontal wavenumber individually. In this way, the heat flux from the perturbations influence the evolution of \bar{T} . But the evolution of \bar{T} also influences the perturbations, as equation 14 depends on $\partial_z \bar{T}$. Thus, the mean temperature and perturbation fields are coupled.

III. PERTURBATION EVOLUTION

The linearized system (12)–(14) does not constrain the amplitude of the eigenmodes, A . However, the advective heat flux is proportional to A^2 , so we need to specify the amplitude in order to solve equation 10. To evolve \bar{T} , we assume the perturbations evolve on a much faster time-scale than the mean temperature, as in [22]. Stable modes ($\sigma < 0$) decay away rapidly. Unstable modes ($\sigma > 0$) will not persist on the slow time-scale because the advective term $\langle w'T' \rangle_x$ tends to stabilize \bar{T} , thereby creating a negative feedback loop. Only marginally stable modes can be maintained on the slow time-scale. Therefore the amplitude A must satisfy

$$\max_{k_x} \{\sigma\} = 0. \quad (22)$$

For various Ra and fixed $Pr = 1$, we seek marginally-stable thermal equilibria (MSTE) satisfying $\frac{\partial \bar{T}}{\partial t} = 0$ according to (10). We employ the *Dedalus* pseudo-spectral python framework¹ [24] to solve the eigenvalue problem outlined in Section II as well as the time evolution equations (10). We represent each field with Chebyshev polynomials and use the 3/2 dealiasing rule to calculate the advective heat flux. The necessary number of basis functions increases with Ra as the eigenfunctions include increasingly small-scale features (see Appendix C). We use the *Eigentools* package² [25] to manipulate the eigenfunctions and calculate the advective heat flux $\langle w'T' \rangle_x$.

To initialize our calculations, we construct a marginally-stable initial temperature profile $\bar{T}(z, t = 0)$ whose equation is given in appendix A. We use (10) to evolve $\bar{T}(z, t_0)$ into a new marginally-stable profile $\bar{T}(z, t_0 + \Delta t)$. We use a second-order, two-stage IMEX Runge-Kutta method. The eigenfunctions and amplitudes are assumed to be constant over the timestep. We symmetrize \bar{T} by setting the coefficients of its even Chebyshev basis functions to zero at the start of each

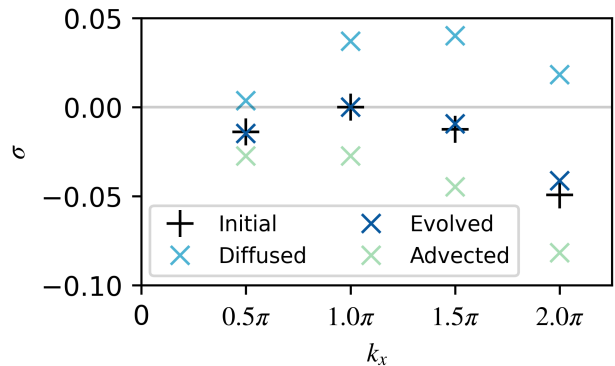


FIG. 1. Eigenvalue spectra for $Ra = 10^5$. The spectrum of an “initial” marginally-stable mean temperature profile $\bar{T}(z, t_0)$ has a maximum eigenvalue of 0. Given a small fixed timestep Δt , diffusion destabilizes the system, increasing σ . Advection tends to stabilize the system, decreasing σ . We find the eigenfunction amplitude A^2 such that the combination of diffusion and advection yields a new, “evolved,” marginally-stable mean temperature profile $\bar{T}(z, t_0 + \Delta t)$. In this case $A^2 \approx 1$ due to approximately equal magnitudes of the diffused and advected eigenvalues.

timestep. This ensures $\bar{T}(z) = -\bar{T}(-z)$. On every timestep, we calculate the marginally stable eigenfunctions and their amplitude A . It is essential that we pick the correct eigenfunction amplitude when calculating the advective term to maintain marginal stability. We will now illustrate our method of finding the appropriate A through an example.

Consider a marginally-stable temperature profile $\bar{T}(z, t_0)$. By definition, its maximum growth rate is 0. Diffusing $\bar{T}(z, t_0)$ tends to increase its growth rates while ignoring the diffusive term and evolving according to advection tends to decrease its growth rates. A must be selected such that these two influences are equal and opposite. We can measure the effects of diffusion and advection on the maximum eigenvalue by solving two new IVPs

$$\frac{\partial \bar{T}_{\text{diff}}}{\partial t} = \mathcal{P} \frac{\partial^2 \bar{T}_{\text{diff}}}{\partial z^2} \quad (23)$$

$$\frac{\partial \bar{T}_{\text{adv}}}{\partial t} + \frac{\partial}{\partial z} \langle w'T' \rangle_x = 0 \quad (24)$$

where \bar{T}_{diff} and \bar{T}_{adv} denote to the diffused and advected temperature profiles. We initialize \bar{T}_{diff} and \bar{T}_{adv} with $\bar{T}(z, t_0)$, and evolve the equations to $t = t_0 + \Delta t$. Suppose k_0 is the wavenumber of the marginally-stable mode. Then, using the diffused and advected profiles, we compute the eigenvalues σ_{diff} and σ_{adv} at $k_x = k_0$ and $t = t_0 + \Delta t$. A preliminary approximation Λ_0^2 of the appropriate amplitude A^2 is given by

$$A^2 \approx \Lambda_0^2 \equiv -\frac{\sigma_{\text{diff}}}{\sigma_{\text{adv}}} \quad (25)$$

where the square in A^2 and Λ_0^2 is due to the quadratic

¹ We employ *Dedalus* version v2.2006
<https://github.com/DedalusProject/dedalus/releases>

² We employ *Eigentools* commit
 f457afc3193c32c16fdff94cdad962d07edea56b
<https://github.com/jsoishi/eigentools>

nonlinearity of the eigenfunctions in the advective term and the minus sign is due to the fact that $\sigma_{\text{adv}} < 0$ and $\sigma_{\text{diff}} > 0$ in general. We illustrate these trends in Figure 1. The subscript 0 denotes a preliminary estimate which we later refine.

Using the approximate amplitude Λ_0^2 , the background temperature $\bar{T}(z, t_0)$ is then evolved to $t = t_0 + \Delta t$ according to (10) and another eigenvalue solve is performed. This background temperature is typically close to, but not exactly, marginally stable. We use Newton's method to find an amplitude A^2 such that the maximum growth rate is zero to within 10^{-9} . Marginally-stable modes do not oscillate in time, i.e. $\sigma = 0$ implies $\omega = 0$. This agrees with the conventional notion of exchange of stabilities [26]. Crucially, we do not assume the k_x of the marginally-stable mode is fixed. In section III A we specify procedures for the treatment of multiple simultaneously marginal modes.

A. Treatment of Multiple Marginally-Stable Modes

In most cases, we encounter eigenvalue spectra with multiple marginal modes. To accommodate this we generalize the advective term in (10) to accommodate N simultaneously marginal modes

$$\langle w'T' \rangle_x = \sum_{n=1}^N 2A_n^2 \Re[W_n \theta_n^*] \quad (26)$$

where W_n and θ_n are the marginally stable eigenmodes associated with different values of k_x . There are now N modes, each with their own amplitude to solve for and eigenvalue to keep marginally-stable. Given a small fixed time step Δt , let \mathbf{A}^2 be the amplitude vector and $\boldsymbol{\sigma}(\mathbf{A}^2)$ be the eigenvalues of the N modes. We are searching for $\tilde{\mathbf{A}}^2$ that satisfies of $\boldsymbol{\sigma}(\tilde{\mathbf{A}}^2) = 0$. We approximate the amplitudes with Λ_0^2 by generalizing (25):

$$\tilde{\mathbf{A}}^2 \approx \Lambda_0^2 = -\Sigma_{\text{adv}}^{-1} \boldsymbol{\sigma}_{\text{diff}}. \quad (27)$$

Here $\boldsymbol{\sigma}_{\text{diff}} = \boldsymbol{\sigma}(\mathbf{0})$ refers to the eigenvalues of the background temperature after evolving under only diffusion (equation 23) for a time interval Δt . We account for the influence of N advection terms on N eigenvalues (one for each mode) by constructing an eigenvalue matrix Σ_{adv} . To calculate element of Σ_{adv} at row i and column j , we evolve the background temperature under only advection by mode j for a time interval Δt . Then the i, j component of Σ_{adv} is given by the eigenvalue of the i th mode.

Our estimate Λ_0^2 does not give a background temperature for which all N modes are exactly marginally stable. We refine our vector of amplitudes via Newton's method. This requires calculating the Jacobian matrix

$$J = \begin{bmatrix} \nabla \sigma_1(A_1, A_2, \dots, A_N) \\ \nabla \sigma_2(A_1, A_2, \dots, A_N) \\ \vdots \\ \nabla \sigma_N(A_1, A_2, \dots, A_N) \end{bmatrix}. \quad (28)$$

We approximate J via first-order finite differences, which requires an additional N^2 sparse eigenvalue solves. We iterate Newton's method until all marginally stable modes have eigenvalues within 10^{-9} of zero. We find $\boldsymbol{\sigma}(\mathbf{A}^2)$ does indeed have a unique root provided the time step is not too large and there are no numerical instabilities, as outlined in appendix B. Presumably this is due to the coupling of $\langle w'T' \rangle_x$ with \bar{T} . Over the course of a large time step, \bar{T} evolves according to (10) and eventually the original eigenfunctions cease to provide a stabilizing influence. This limits the timestep size we can take.

Difficulty arises when transitioning between different numbers of marginal modes. We facilitate these transitions by defining an adjustable candidate tolerance $\varepsilon_{\text{cand}} \in [10^{-6}, 10^{-8}]$. A mode which meets the candidate tolerance can be included as a candidate marginal mode. Candidate modes are rejected when the root-finding algorithm converges on a negative amplitude $A^2 < 0$. Otherwise the mode becomes marginally-stable. In a similar manner, a marginally-stable mode need not remain marginally-stable. If its amplitude converges to some $A^2 < 0$ as before, then that mode is discarded and the iteration is repeated.

IV. PROPERTIES OF THERMALLY EQUILIBRATED STATES

We evolve \bar{T} as described above until $\max |\partial_t \bar{T}| < 10^{-5}$. In this marginally-stable thermal equilibrium, \bar{T} does not evolve in time, and the perturbations also do not evolve in time, as they are marginally stable. Thus, these configurations are exact solutions to the quasilinear equations (equations 10–14). They differ from the usual ECS in that ECS are fixed points of the full nonlinear problem (1)–(3). Such definitions are not mutually exclusive, but in general we can assume that MSTE and ECS are not steady with respect to their counterparts' equations. We compute symmetric MSTE for Ra in the range $10^5 - 10^9$.

Figure 2 gives temperature profiles for Ra = 10^8 , where the initial profile is outlined in appendix A. We ran a direct numerical simulation (DNS) of (1)–(3) with *Dedalus*, and also plot the horizontal- and time- averaged temperature profile. The temperature boundary layers in the DNS are wider than in the MSTE, which is in turn wider than the boundary layer of the initial temperature profile. Performing an eigenvalue solve by setting \bar{T} equal to the DNS profile yields unstable eigenvalues. This suggests that MSTE might maximize boundary layer thickness, subject to the marginal stability constraint.

The most resilient and unexpected feature of MSTE temperature profiles are the pronounced dips adjacent to the boundary layers. These dips appear in every solution, regardless of Ra. Physically, they correspond to thin layers in which the mean temperature gradient reverses, contradicting an important hypothesis of [5, 6]. This counter-diffusion, which opposes overall heat trans-

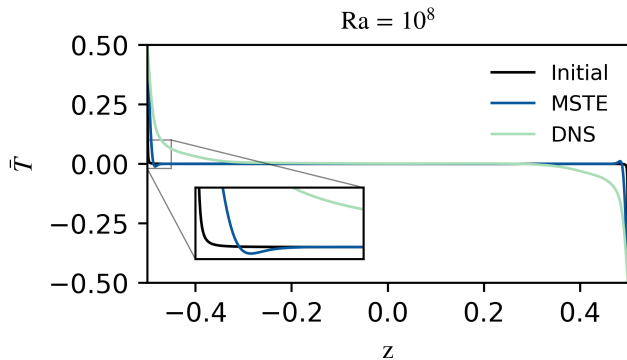


FIG. 2. Mean temperature profiles \bar{T} for $Ra = 10^8$. The initial profile is given by (A1). We evolve this background temperature profile according to section III until we reach a marginally-stable thermal equilibrium (MSTE). The DNS curve is obtained from a 2D nonlinear simulation of (1)–(3) with *Dedalus*. DNS temperature data are horizontally- and time-averaged. The initial profile has the narrowest boundary layer, while the DNS profile has the widest boundary layer. The MSTE profile exhibits prominent dips, nested alongside the boundary regions.

fer, is overcome by the coinciding advective flux, shown in Figure 3. We do not understand the source of these dips, but similar temperature gradient reversals were reported by [17] along the midlines of 2D convective cellular solutions at $Ra \sim 10^6$. In that case, the reversals were due to nonlinear advection, which is not present in our quasilinear model.

In Figure 3, we give heat flux profiles and eigenvalue spectra for two cases: $Ra = 2 \times 10^5$ (top) and $Ra = 10^9$ (bottom). For $Ra = 2 \times 10^5$, there is a single marginal mode at $k_x = 1.5\pi$ whose advective flux occupies the bulk of the domain. These states have wide boundary layers, with diffusive fluxes which gradually subside as advection becomes the dominant flux component. Transitional regions occur over a smaller length scale for $Ra = 10^9$ where the shift from diffusion to advection is sharp. At $Ra = 10^9$ we find five marginally-stable modes are necessary to reach an MSTE. Thin advection profiles, belonging to high-wavenumber modes with $k_x = 23.5\pi, 24\pi$, hug the boundary layer. Closer to the bulk of the domain, we see wider advection profiles corresponding to modes in a second group of marginal modes $k_x = 6\pi, 6.5\pi$. The $k_x = 1.5\pi$ mode forms the large-scale convective cell structures observed in DNS, again occupying the bulk of the domain. The pairs of modes $k_x = 6\pi, 6.5\pi$ and $k_x = 23.5\pi, 24\pi$ are each associated with a single maximum in our plots of growth rate σ as a function of k_x (lower right panel of Figure 3). If we allowed wavenumbers to vary continuously, there would be an unstable mode between these pairs of wavenumbers. However, since we have fixed the horizontal size of our domain, we are left with pairs of discrete marginal modes.

MSTE for large Ra tend to have a diverse combina-

tion of marginal modes. In every case, the $k_x = 1.5\pi$ mode is included. In Figure 4 we give the wavenumbers k_x of marginal modes. For some values of Ra we obtain states with pairs of marginally-stable modes, with wavenumbers separated by $\pi/2$. Like $k_x = 6\pi, 6.5\pi$ and $k_x = 23.5\pi, 24\pi$ for the $Ra = 10^9$ MSTE, these are due to the discretization of wavenumbers from our domain of width 4. We think of the pairs of modes as acting together as part of a single maximum of the growth rate as a function of the wavenumber. When wavenumbers are adjacent, we plot them in the same color and denote the larger mode with an x and the smaller with a +. For $Ra \geq 10^6$, a second branch of marginal modes is shown in light green. Least-squares regression gives $\max\{k_x\} \propto Ra^{0.300}$ with $R^2 = 0.998$ for this maximum branch. For large Ra , the advective fluxes of this maximum branch compensates for the strongly peaked diffusive flux in the thin boundary layers. At $Ra \geq 10^8$, a third branch appears (shown in blue), splitting the widening gap between the other two. For these points regression gives $k_x \propto Ra^{0.155}$ with $R^2 = 0.988$. The blue branch is associated with moderately wide advection profiles, filling a niche in the total flux by uniting the thin profiles of the maximum branch with those of the bulk-domain-oriented minimum branch.

The largest marginal wavenumber $\max\{k_x\}$ (represented by the light green branch in Figure 4) serves as an inverse minimum length-scale in the x direction. The finest vertical structures in $\bar{T}(z)$ appear near the boundaries, requiring more basis functions (resolution) at large Ra . Naturally, this provides a complementary minimum length-scale for z . We define the boundary layer width δ as the distance from the boundary where $\partial_z T$ equals zero, i.e.,

$$\left. \frac{\partial \bar{T}}{\partial z} \right|_{z=-\frac{1}{2}+\delta} = 0, \quad 0 < \delta < 1/2. \quad (29)$$

This height corresponds to the local extrema of the MSTE temperature profile, e.g., in Figure 2. In Figure 5, we show $\max\{k_x\}$ is proportional to δ^{-1} over our range of Ra . Least-squares regression gives $\delta^{-1} = 1.71 \max\{k_x\} - 2.13$ with $R^2 = 0.996$.

We find the MSTE can be characterized by their boundary layer height. In Figure 6, we illustrate the scaling behavior of the boundary layer height $\delta = Ra^{-1/3}$. This is consistent with Malkus' classical marginal-stability theory, a scaling argument which perceives the boundary regions as subdomains which are themselves marginally-stable [5].

The Nusselt number, which measures convective performance is given by

$$Nu = \frac{\langle \langle w'T' \rangle_x - \mathcal{P} \frac{\partial \bar{T}}{\partial z} \rangle_z}{\langle -\mathcal{P} \frac{\partial \bar{T}}{\partial z} \rangle_z}. \quad (30)$$

There is no general consensus surrounding the scaling behavior of Nu for high Ra systems, which are of par-

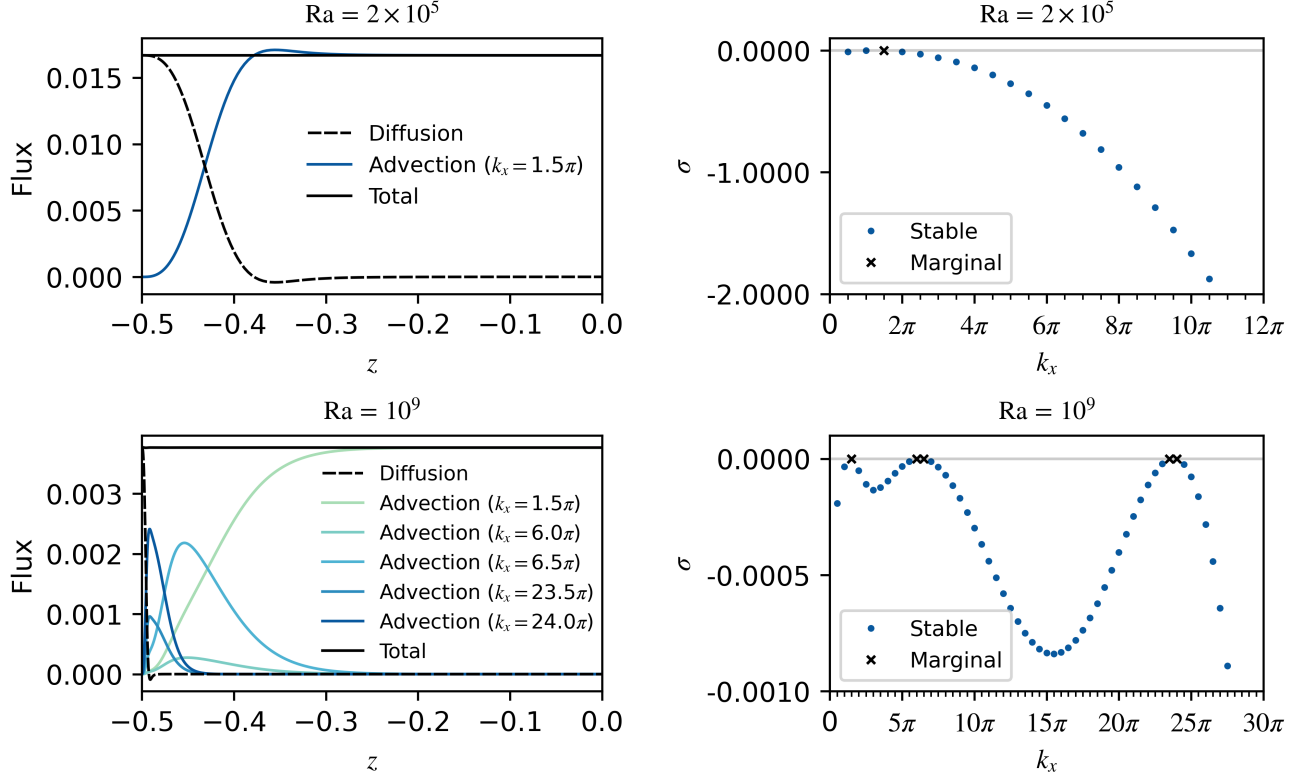


FIG. 3. Heat fluxes (left) and eigenvalue spectra (right) of equilibrated states with $Ra = 2 \times 10^5$ (top) and $Ra = 10^9$ (bottom). The heat flux profiles are symmetric about $z = 0$, so we only plot for $z < 0$. We plot the advection profiles for marginally-stable modes. At low Ra , a single mode with $k_x = 1.5\pi$ is sufficient to oppose boundary layer diffusion and facilitate heat flux throughout the bulk of the domain. At large Ra , high-wavenumber modes contribute pronounced small-scale advection profiles which tightly hug the thin boundary layers. A combination of progressively wider advection profiles is necessary to transition to bulk advection provided by the $k_x = 1.5\pi$ mode.

ticular importance in astrophysical and geophysical systems. In Figure 7 we report Nu for MSTE, “steady rolls” ECS [16], and DNS [27, 28]. We find that MSTE satisfy $Nu \sim Ra^{1/3}$, consistent with our finding that the boundary layer height scales like $\delta \sim Ra^{1/3}$. The Nusselt numbers of the ECS are somewhat lower and the DNS Nusselt numbers are yet lower still. In both cases, the Ra dependence appear slightly more shallow than for the MSTE. [16] hypothesized that the Nu of all ECS which admit classical Malkus scaling must always exceed the Nu of turbulent convection. If we generalize this notion to include quasilinear equilibria, our findings agree; MSTE have larger Nu than 2D and 3D DNS. This might be due to the chaotic transitions among the unstable periodic orbits outlined by [18, 19] inhibiting heat flux. We might also anticipate the existence of similar equilibria with smaller Nu , occupying complementary nodes in the Markov chain whose behavior agrees with DNS.

V. SIMULATIONS WITH THERMALLY EQUILIBRATED INITIAL CONDITIONS

This investigation is partially motivated by the prospect of decreasing DNS runtimes by employing MSTE as initial conditions. One common choice of initial conditions for DNS of equations (1)–(3) are

$$\begin{aligned} T(x, z)|_{t=0} &= 0.5 - z + \mathcal{N} \\ \mathbf{u}(x, z)|_{t=0} &= \mathbf{0} \\ p(x, z)|_{t=0} &= 0 \end{aligned} \quad (31)$$

where \mathcal{N} is low-amplitude random noise. Here we instead initialize using the MSTE,

$$\begin{aligned} T(x, z)|_{t=0} &= \bar{T}(z) + \sum_{n=1}^N A_n \Re[\theta_n(z) e^{ik_{x_n} x}] + \mathcal{N} \\ \mathbf{u}(x, z)|_{t=0} &= \sum_{n=1}^N A_n \Re[(U_n(z) \hat{x} + W_n(z) \hat{z}) e^{ik_{x_n} x}] \\ p(x, z)|_{t=0} &= \sum_{n=1}^N A_n \Re[P_n(z) e^{ik_{x_n} x}] \end{aligned} \quad (32)$$

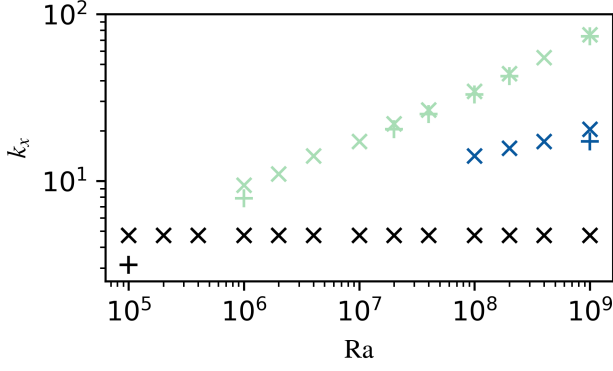


FIG. 4. Wavenumbers of marginally-stable modes in thermally equilibrated states. Marginal modes often appear in adjacent pairs, which we denote with a common color. Within each of these pairs, the largest and smallest k_x are denoted with an x and a $+$ respectively. For example, the spectrum corresponding to $Ra = 10^5$ has adjacent marginal wavenumbers $k_x = \pi, 1.5\pi$. The $Ra = 10^9$ spectrum, shown in lower right corner of Figure 3, has three groups of maxima, with a single marginal mode in the first group ($k_x = 1.5\pi$), two adjacent marginal modes in the second group ($k_x = 6\pi, 6.5\pi$), and two adjacent marginal modes in the third group ($k_x = 23.5\pi, 24\pi$). The largest wavenumbers of the green and blue branches depend on Ra according to power laws. For the green branch, $\max\{k_x\} \propto Ra^{0.300}$ with $R^2 = 0.998$ while for the blue branch, $k_x \propto Ra^{0.155}$ with $R^2 = 0.988$.

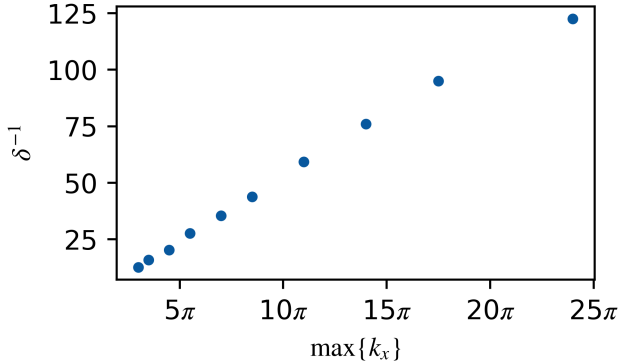


FIG. 5. For $Ra \geq 10^6$, the maximum marginally-stable wavenumber (corresponding to the green x markers in Figure 4) are inversely related to the boundary layer height δ . $(\max\{k_x\})^{-1}$ gives a minimum x length scale for the perturbations, and consequently, the advection. For large Ra , the boundary layers admit small scale features, requiring more vertical basis functions (higher resolution, see Appendix C for specifications). The boundary layer width gives an estimate of the minimum vertical length-scale in the problem. This suggests that the minimum horizontal and vertical length-scales are proportional to each other over a wide range of Ra .

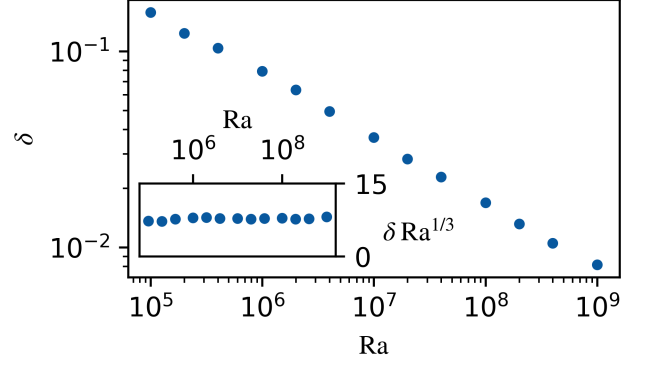


FIG. 6. Boundary layer height δ of MSTE. We define the boundary layer height based off the location where $\frac{\partial T}{\partial z} = 0$ (see equation 29). Plotting on a log-log scale, we find that δ and Ra obey a power-law relationship. We also demonstrate that $Ra^{1/3}\delta$ is approximately constant with respect to Ra which is consistent with [5]

where $\theta_n(z), U_n(z), W_n(z), P_n(z); A_n$; and k_{x_n} refer to the complex eigenfunctions, amplitude, and wavenumber of the n th marginal mode respectively. Note that although the MSTE is an equilibrium of the quasilinear equations, it is not an equilibrium of the full nonlinear equations. Accordingly the simulation state would evolve on initialization absent a random noise term. Here we include noise as a source of asymmetries. We also perform a simulation with MSTE absent any initial velocity ($\mathbf{u}(x, z)|_{t=0} = \mathbf{0}$). This state is not in equilibrium but we refer to it as “MSTE No Flow” for clarity.

Simulations initialized with the conductive equilibrium plus low-amplitude thermal noise have a large peak in Nu early on in their evolution (Figure 8). This is due to a burst of turbulence which occurs when the convective motions first become nonlinear. The MSTE no flow simulation undergoes a similar transient period, albeit less pronounced as illustrated by the smaller peak in the green curve. A simulation initialized with the MSTE, however, does not exhibit this transient burst of turbulence, as the large-scale anatomy of convective cells exists on initialization. Simulation of this transitional period is prohibitive [29]. For high Ra experiments, researchers often “bootstrap” data by initializing simulations with the results of similar Ra runs [27, 30]. MSTE can be perceived as a set of initial conditions, designed for avoiding the simulation of transient high Reynolds number flows. MSTE absent velocity achieve the same goal but give way to seemingly-persistent oscillations in Nu .

MSTE are laminar, lacking the small-scale structures associated with moderate- to high- Ra experiments. This is an apparent consequence of the quasilinear assumptions. If we perceive MSTE as background states, DNS suggest that plumes, vortex sheets, and other unstable turbulent features inhibit total heat transfer. This perspective agrees with conventional models of transitions to

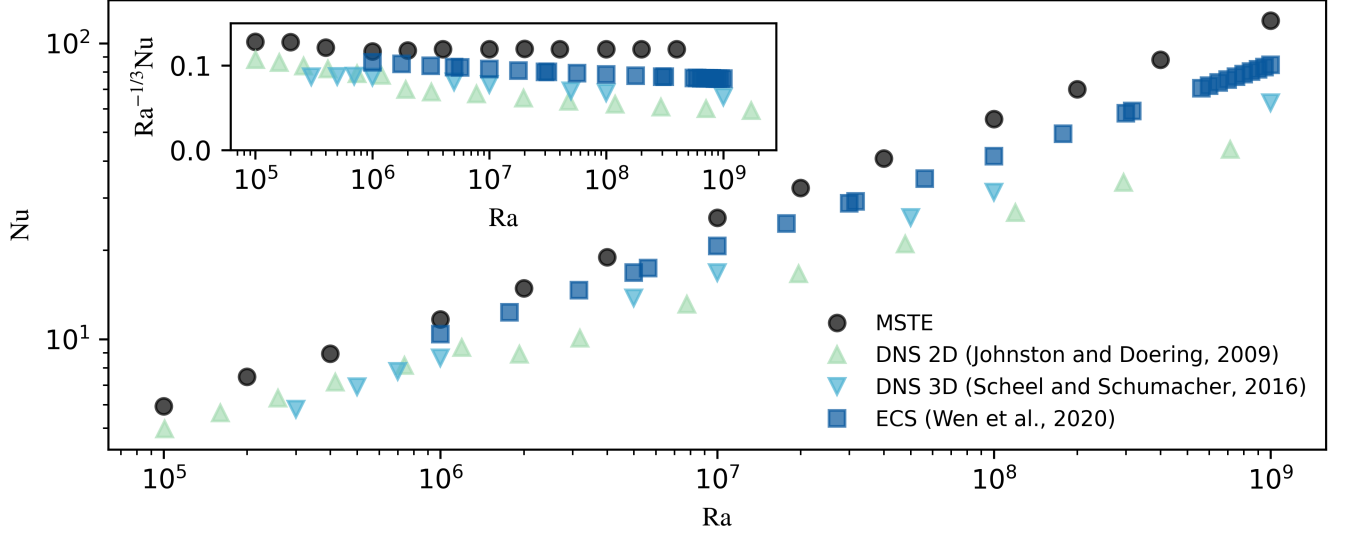


FIG. 7. Nusselt numbers are shown for MSTE, aspect-ratio-optimized “steady rolls” ECS [16], as well as statistically-steady 2D and 3D DNS [27, 28]. All datasets obey power-law relationships, with the MSTE and ECS scaling like $Nu \sim Ra^{1/3}$. MSTE have greater Nu than the ECS, which in turn, have greater Nu than the DNS. This can be explained by the contrasting boundary layer geometries shown in Figure 2.

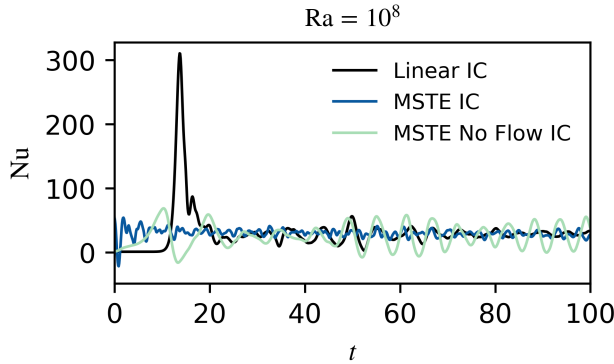


FIG. 8. Nusselt numbers of simulations performed at $Ra = 10^8$, initialized with the conductive equilibrium plus thermal noise (black), the MSTE (blue), and the MSTE absent initial flow (green). The MSTE simulation does not undergo a convective-transient period because the characteristic large-scale convective cell structure exists on initialization. The MSTE no flow simulation exhibits quasi-periodic oscillations in Nu which appear to persist for long times. The oscillations are great enough in amplitude that Nu is quasi-periodically negative, indicating brief periods of average heat transfer reversal.

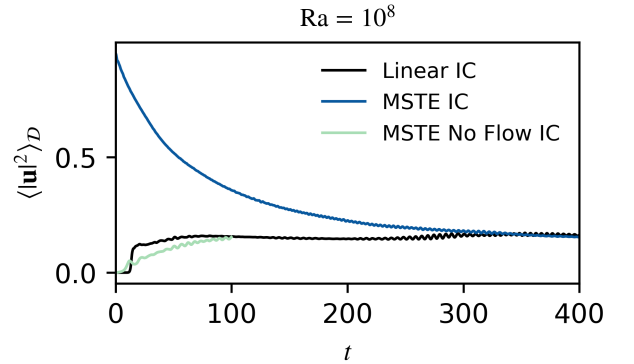


FIG. 9. Average kinetic energies are reported for the same simulations illustrated in Figure 8 ($Ra = 10^8$). The eigenfunctions belonging to MSTE have significantly more kinetic energy than the statistically-steady state. Kinetic equilibrium is achieved on a long viscous time-scale.

turbulent flows, such as Boussinesq’s turbulent-viscosity hypothesis [31]. The emergence of small-scale velocity structures is indicative of nonlinear energy transfer to small scales where energy is lost due to viscosity. Buoyancy driven flows are therefore impeded and the advection in the bulk of the domain decreases in magnitude [26, 32]. We could also attribute the diffuse DNS temperature profile in Figure 2 with unsteady boundary-layer penetration and mixing that MSTE do not exhibit.

We also find that the average kinetic energy of the simulation initialized with MSTE is significantly larger than those found in either of the other cases, as shown

Figure 9. This is because the MSTE contain strong large-scale flows, which decay on a long viscous time-scale t_ν . Fluid in the other cases is initialized at rest. Two viscous time-scales can be derived for this problem: one for the boundary layer

$$t_{\nu,\delta} = \frac{\delta^2}{\mathcal{P}} = 2.86$$

and another for the entire domain

$$t_{\nu,L} = \frac{L_z^2}{\mathcal{P}} = 10^4.$$

Empirically, we find $t_\nu \approx 200$ suggesting the true kinetic energy decay time-scale is based on a length scale $\ell \approx 0.14$ which is longer than the boundary layer height δ but shorter than the domain height L_z . Consequently, MSTE initial conditions do not reduce the simulation time required to achieve a statistically-steady state—rather they increase it considerably! This is alleviated by using no-flow MSTE initial conditions. The MSTE temperature state is a promising candidate initial condition for high-Rayleigh number convection simulations.

VI. DISCUSSION

In this paper we describe a new way to study Rayleigh–Bénard convection. We compute marginally-stable thermal equilibria (MSTE), which are equilibria of the quasilinear equations. To compute MSTE, we construct a marginally-stable mean temperature profile and evolve it according to the advective flux of its marginally-stable eigenfunctions, and its own diffusion. We assume that at least some modes are always in a marginally-stable configuration. The marginal stability constraint then fixes the ratio between advection and diffusion (eigenfunction amplitude A^2). We use standard root-finding algorithms to solve for the appropriate A^2 at each timestep until the combination of diffusive and advective flux sum to a constant. The MSTE calculation is a one-dimensional problem, combining eigenvalue solves, and the time evolution of the one-dimensional mean temperature profile. Thus, MSTE can be calculated on a single workstation.

The MSTE retain several key features that are prevalent in experiments and simulations: $\text{Nu} \sim \text{Ra}^{1/3}$ scaling, large-scale convective cell structures, and minimum length scale agreement. They also exhibit unique and unexpected features: mean temperature gradient-reversals/dips, high kinetic energy flows, and a larger Nu than other time-invariant solutions. When initializing with different mean temperature profiles, we find the same MSTE, suggesting these equilibria might be unique.

Simulations initialized with the MSTE (32) do not undergo an early convective transient period, but have faster flows when compared with DNS. From a dynamical systems perspective, unstable orbits depart from MSTE

and approach the global attractor on a viscous time-scale. This requires more computational effort to achieve relaxation when compared to the conventional conductive initial condition (31). Initializing with only the temperature state of the MSTE avoids an early convective transient, but does not have excessive large-scale flows. These state could be excellent initial conditions for high-Rayleigh number DNS.

Using the mean temperature in a statistically-steady DNS as a background state for an eigenvalue problem yields positive eigenvalues: the system is in a perpetual state of instability. Unstable modes tend to stabilize the system rapidly, creating a negative feedback loop whose average state is linearly unstable. Future work could adjust the marginal-stability criterion to allow for moderately unstable modes. Should the fast and slow time scales not be entirely separate, these modes might persist for long times. This might lead to greater agreement with DNS.

ACKNOWLEDGMENTS

The authors thank Geoff Vasil, Greg Chini, Kyle Augustson, and Emma Kaufman for their valuable feedback and suggestions. We also thank the *Dedalus* and *Eigentools* development teams. Computations were conducted with support by the NASA High End Computing (HEC) Program through the NASA Advanced Supercomputing (NAS) Division at Ames Research Center on Pleiades with allocation GIDs s2276.

Appendix A: Initial Buoyancy Profile

We initialize the thermal-equilibration algorithm with an analytical thermal boundary layer equation, derived by [33]

$$\begin{aligned} \bar{T}_0(\xi) &= \frac{\sqrt{3}}{4\pi} \log \frac{(1+a\xi)^3}{1+(a\xi)^3} + \frac{3}{2\pi} \arctan \left(\frac{4\pi}{9} \xi - \frac{1}{\sqrt{3}} \right) + \frac{1}{4} \\ \xi = \frac{z+1/2}{\delta_0} &\geq 0, \quad a = \frac{2\pi}{3\sqrt{3}} \end{aligned} \quad (\text{A1})$$

where δ_0 is a new measure of the boundary layer height ($\delta_0 \neq \delta$ in general). This function is meant to describe the temperature near $z = -1/2$ so it does not pass through the origin. An appropriate initial mean temperature profile \bar{T}_0 must be odd-symmetric, i.e. $\bar{T}_0(-z) = -\bar{T}_0(z)$. Due to continuity, this implies $\bar{T}_0(0) = 0$. Accordingly, we construct $\bar{T}_0(z)$ by translating $\bar{T}_0(\xi)$ vertically to pass through the origin. We then take its odd-extension and include a unique scaling coefficient to satisfy the boundary conditions $\bar{T}_0(-1/2) = 1/2$ and $\bar{T}_0(1/2) = -1/2$. The initial mean temperature profile is

therefore given by

$$\bar{T}_0(z) = \frac{1}{2} \begin{cases} 1 - \frac{\bar{\tau}_0\left(\frac{z+1/2}{\delta_0}\right)}{\bar{\tau}_0\left(\frac{1}{2\delta_0}\right)} & -1/2 \leq z \leq 0 \\ -1 + \frac{\bar{\tau}_0\left(\frac{1/2-z}{\delta_0}\right)}{\bar{\tau}_0\left(\frac{1}{2\delta_0}\right)} & 0 < z \leq 1/2. \end{cases}$$

We expect each Ra to be associated with a unique δ_0 for which $\bar{T}_0(z)$ is marginally-stable. It should be noted that when experimenting with various initial profiles (tanh, erf, etc.), we obtain indistinguishable equilibrated states. Therefore these initial states lie in the MSTE basin of attraction with respect to the quasilinear system. This might also suggest that solutions are unique. An example of (A1) is given by the blue curve in Figure 2.

Appendix B: Timestep Management

To find MSTE, we advance \bar{T} by timesteps of size Δt . For large Δt , we find the advective flux terms fail to provide a stabilizing influence. This is due to coupling between the eigenfunctions $W(z)$, $\Theta(z)$ and the mean temperature profile \bar{T} . In the context of our algorithm, this effectively deletes the sought-after root of the max-

imum eigenvalue function $\sigma_{\max}(A^2)$, causing the root-finding methods to fail. This is not a numerical instability, rather, it is an inherent limitation of our timestepping algorithm. To curtail this, we halt the root-finding algorithms after 20 successive approximations, and reduce the timestep by a factor of 1.1, and try again.

The timestep must be also reduced to avoid a numerical instability. We find that for large Δt , after several hundred iterations, highly concave features develop in the advective flux term $\langle w'T' \rangle_x$ near $z = 0$. Such features are undesired, as they are uncommon in similar calculations [5] and we do not believe they accurately represent a physical process. If ignored, the concave features grow in magnitude until they affect \bar{T} on a readily apparent scale. Eventually \bar{T} develops oscillations near $z = 0$ and the timestep must be reduced. Once these oscillations reach some amplitude, they cannot be eliminated via timestep reduction and the roots of $\sigma_{\max}(A^2)$ vanish. To avoid this, we measure the curvature of the advective flux $|\frac{\partial^2}{\partial z^2} \langle w'T' \rangle_x|_{z=0}$ of the dominant $k_x = 1.5\pi$ mode. We find that reducing the timestep Δt by the same factor of 1.1 whenever this curvature measure exceeds 10^{-6} curtails the problem. Both of these issues appear to become more prominent as the boundary layers diffuse over the course of the algorithm's implementation (see Figure 2). Thus there is never a practical opportunity to increase the timestep.

Appendix C: MSTE Parameters and Results

Ra	N_z	Nu	δ_0	marginal k_x
10^5	256	5.9343	0.15746	$1\pi, 1.5\pi$
2×10^5	256	7.45986	0.12341	1.5π
4×10^5	256	8.92597	0.10395	1.5π
10^6	256	11.6689	0.07922	$1.5\pi, 2.5\pi, 3\pi$
2×10^6	256	14.8463	0.06345	$1.5\pi, 3.5\pi$
4×10^6	256	18.9433	0.04933	$1.5\pi, 4.5\pi$
10^7	256	25.6821	0.03632	$1.5\pi, 5.5\pi$
2×10^7	256	32.4531	0.02820	$1.5\pi, 6.5\pi, 7\pi$
4×10^7	512	40.7925	0.02289	$1.5\pi, 8\pi, 8.5\pi$
10^8	512	55.4383	0.01690	$1.5\pi, 4.5\pi, 10.5\pi, 11\pi$
2×10^8	512	69.8349	0.01318	$1.5\pi, 5\pi, 13.5\pi, 14\pi$
4×10^8	512	87.8525	0.01053	$1.5\pi, 5.5\pi, 17.5\pi$
10^9	768	119.318	0.00817	$1.5\pi, 6.0\pi, 6.5\pi, 23.5\pi, 24\pi$

TABLE I: Control parameters and results are given for the MSTE timestepping algorithm. N_z denotes the number of Chebyshev basis functions employed. This, along with Ra are specified on initialization. The remaining quantities are computed directly from the MSTE. The low Ra = 10^5 case requires ~ 12 hours while the high Ra = 10^9 case requires ~ 72 hours. Eigenvalue solves are performed for various k_x simultaneously using 28 cores.

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