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—particularly in the large Rayleigh number regime. In this investigation, we derive and solve a quasilinear form of the Rayleigh-Benard 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 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 initializing a 2D simulation with a marginally-stable thermal equilibrium, we find the simulation does not exhibit the initial burst of turbulence typical of simulations initialized with the conductive background state plus noise. Although the Nu quickly equilibrates in these simulations, the kinetic energy evolves on a viscous timescale as domain-filling flywheel flows viscously attenuate.

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]. Aggressive 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 stars are Ra $\sim 10^{20}$ [3]. The scaling behavior of the Nusselt number Nu $\sim Ra^{\beta}$ in the asymptotic ultimate regime $Ra \to \infty$ is of particular interest. There exists a substantial body of work pertaining to this specific topic with no general consensus [4–10].

Reduced models are particularly useful in this context because they allow us to study the problem with relatively low expense. Much of the published work on this topic relates to unstable exact coherent states (ECS) [11– 14]. Simulations and analysis performed by [15, 16] suggest that chaotic solutions to the full nonlinear problem can be represented by a Markov chain of unstable equilibria. Should that be the case, it is crucial that we discover and classify such states.

In [17], researchers compute ECS in parallel shear flows by deriving and solving a reduced form of the Navier-Stokes equations via multi-scale asymptotic arguments. This is realized by defining a mean state which, hypothetically, evolves on a slow time-scale. Marginal stability is required at each successive timestep to constrain the perturbation amplitudes. A similar strategy is employed by [18], where an analytic expression for the first-order perturbations' amplitude is found by deriving a solvability condition of the second-order perturbations and integrating over the domain spatially. We seek to solve the

Boussinesq Rayleigh-Bénard convection equations using an analogous strategy. The application of this procedure yields time-invariant solutions to the quasilinear problem with a range of expected and unexpected features.

MODEL SETUP

We begin with the non-dimensionalized Boussinesa approximation for Rayleigh-Bénard Convection. The domain \mathcal{D} is 2D rectangular and horizontally periodic with spatial dimensions 0 < 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 and 0 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 $z = \pm 1/2$. The equations of motion are then given by

$$\nabla \cdot \mathbf{u} = 0 \tag{1}$$

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

$$\nabla \cdot \mathbf{u} = 0 \tag{1}$$

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

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

where p is pressure and T is temperature. For completeness, we specify a final boundary condition $p = p_0$ at $z = \pm 1/2$. Any system of this form can be characterized by its dimensionless Rayleigh number Ra = $\frac{g\alpha L^3\Delta T}{\nu\kappa}$ and Prandtl number 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.

For convenience, we define

$$\mathcal{R} = \sqrt{\frac{\Pr}{\operatorname{Ra}}}, \qquad \mathcal{P} = \frac{1}{\sqrt{\Pr \operatorname{Ra}}}.$$
 (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').

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

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

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

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

where A(t) is some unknown real amplitude and the mean-velocity components vanish due to incompressibility and symmetry. Perturbations must admit no horizontal-average

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

Assuming the existance of nontrivial solutions, we also fix the volume-average over the entire domain

$$\langle T^{\prime 2} \rangle_{\mathcal{D}} = 1 \tag{10}$$

thereby alleviating any ambiguity in A(t). Substituting Eq. (7) into Eq. (3) and taking the horizontal-average reduces the system to a simple initial value problem (IVP) for \bar{T}

$$\frac{\partial \bar{T}}{\partial t} + A^2 \langle w'T' \rangle_x = \mathcal{P} \frac{\partial^2 \bar{T}}{\partial z^2},\tag{11}$$

with associated boundary conditions $\bar{T}(-1/2,t)=1$ and $\bar{T}(1/2,t)=0$. 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.

To solve Eq. (11) numerically, we need the perturbations and their associated amplitude function A(t). The former is obtained by solving the linearized system. Substituting Eqs. (5)–(8) into Eqs. (1)–(3) and subtracting Eq. (11) from the resulting temperature equation gives

$$\nabla \cdot \mathbf{u}' = 0 \tag{12}$$

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

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

with Dirichlet boundary conditions

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

In his famous report [19], Lord Rayleigh observed that Eqs. (12) & (14) can be manipulated into a separable

form with generalized solutions

$$w'(x, z, t) = W(z) e^{i(k_x x - st)}$$
 (16)

$$u'(x,z,t) = U(z) e^{i(k_x x - st)}$$

$$\tag{17}$$

$$T'(x,z,t) = \theta(z) e^{i(k_x x - st)}$$
(18)

$$p'(x, z, t) = P(z) e^{i(k_x x - st)}$$
 (19)

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

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

For each k_x , we can assess the stability of the perturbations by solving for the eigenvalue s, whose imaginary component ω plays the role of an exponential growth rate. Solution yields a finite set of eigenvalues, among which, that with maximum ω is assumed to be dominant. 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 $\mathrm{Ra}_c=1708$ when $\frac{\partial \bar{T}}{\partial z}=-1$.

At some arbitrary time t_0 , the forcing term $\langle w'T'\rangle_x$ can be found by solving an eigenvalue problem (EVP) which itself involves $\frac{\partial T}{\partial z}$.

III. PERTURBATION EVOLUTION

Solving the linearized system Eqs. (12)–(14) does not yield A(t). To evolve \bar{T} , we assume the perturbations and mean quantities evolve according to different time-scales, as in [18]. Stable modes decay away rapidly. Unstable modes 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 traverse this hypothetical time-scale separation. Therefore the amplitude function A(t) must satisfy

$$\max_{k_{-}}\{\omega\} = 0. \tag{21}$$

For various Ra and fixed Pr = 1, we seek marginally stable thermal equilibria (MSTE) satisfying $\frac{\partial \bar{T}}{\partial t} = 0$ according to Eq. (11). We employ the Dedalus pseudospectral python framework to solve the EVP outlined in Section II as well as the IVP Eq. (11) by representing each field with a dealiased Chebyshev polynomial basis. The necessary number of basis functions varies, as eigenfunctions and temperature profiles associated with large Ra admit increasingly small-scale features. We supplement Dedalus with the Eigentools package to compute $\langle w'T' \rangle_x$.

At t=0, we begin with a marginally-stable initial profile $\bar{T}(z,0)$ whose construction is outlined in Section VI A. At some arbitrary time $t=t_0$, we seek to evolve $\bar{T}(z,t_0)$

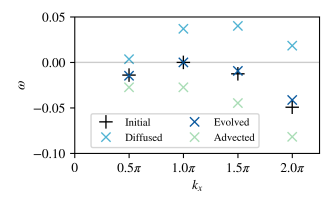


FIG. 1. Eigenvalue spectra for Ra = 10^5 . The spectrum of an "initial" marginally-stable mean temperature profile $\bar{T}(z,t_0)$ has a maximum eigenvalue of 0. Given a small fixed timestep Δt , diffusion destabilizes the system, increasing its eigenvalues. Advection tends to stabilize the system, decreasing its eigenvalues. It follows that there exists some proportion $A^2(t_0)$ of these two terms which yields a new marginally-stable mean temperature profile $\bar{T}(z,t_0+\Delta t)$, whose spectrum is "evolved".

into a new marginally stable profile $\bar{T}(z, t_0 + \Delta t)$ according to Eq. (11) via forward-Euler. This generally involves a unique and unknown value $A^2(t_0) > 0$. Our method of finding $A^2(t_0)$ is best illustrated through example.

An iteration is performed as follows: diffusing a marginally-stable temperature profile $\bar{T}(z,t_0)$ tends to increase its eigenvalues. Ignoring the diffusive term and evolving according to advection tends the stabilize the system. The appropriate amplitude $A^2(t_0)$ can then be approximated by

$$A^2(t_0) \sim -\frac{\omega_{\text{diff}}}{\omega_{\text{adv}}}$$
 (22)

where ω_{diff} and ω_{adv} refer to the diffused and advected eigenvalues of the initially marginal mode. These quantities correspond to the light blue and green points at $k_x = \pi$ in Figure 1). $\bar{T}(z,t)$ is then evolved according to Eq. (11) and another eigenvalue solve is performed. Given a fixed timestep Δt , we assume the dominant eigenvalue can be described by a continuous function $\omega_{\text{max}}(A^2)$ which is locally differentiable. We use Newton's method to find an amplitude which satisfies our marginal stability tolerance criterion. The dominant mode is not fixed; an iteration can facilitate the transfer of marginal-stability from one mode to another. 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 eventually encounter eigenvalue spectra with multiple simultaneously marginal modes. If ignored, we are forced to reduce the timestep and allow

the modes to alternate. Instead, we generalize the advective term in Eq. (11) to accommodate N simultaneously marginal modes

$$A^{2}\langle w'T'\rangle_{x} = \sum_{n=1}^{N} A_{n}^{2}\langle w'T'\rangle_{x,n}$$
 (23)

where $\langle w'T'\rangle_{x,\,n}$ is composed of perturbations associated with $k_x=\frac{n\pi}{2}$. There are now N amplitudes to solve for and N eigenvalues to keep marginally-stable. In general, we expect a function $\omega:\mathbb{R}^N\to\mathbb{R}^N$ to have isolated roots (should they exist). We employ Broyden's method for root-finding in multi-dimensional functions whose derivatives are not explicitly known. Difficulty arises when transitioning between different numbers of marginal modes. Here we rely on $A^2>0$ by asserting that a mode which is close enough to marginal-stability can be included in the iteration provided that its respective amplitude is positive. Should we converge upon a negative amplitude, that mode is discarded and the iteration is repeated. In general we find that no other obvious course of action yields equilibria.

IV. PROPERTIES OF THERMALLY EQUILIBRATED STATES

MSTE are fixed points of the quasilinear form derived in section II, differing from ECS in that ECS are fixed points of the full nonlinear problem Eqs. (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' problems. We compute symmetric MSTE for Ra in the range $10^5 - 10^9$.

Figure 2 gives temperature profiles for ${\rm Ra}=10^8$ where the initial profile, whose construction is outlined in , is employed at iteration 0; the MSTE profile corresponds to the thermally relaxed state; and the DNS (direct numerical simulations) profile is obtained by solving Eqs. (1)–(3) with <code>Dedalus</code>, followed by horizontal- and time- averaging. The DNS curve is more diffuse than the MSTE curve, which in turn, is more diffuse than the initial curve. Performing an eigenvalue solve by setting \bar{T} equal to the DNS profile yields positive eigenvalues. Comprehensively, this suggests that MSTE might maximize boundary layer thickness, while 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 [4]. This counter-diffusion, which opposes overall heat transfer, is overcome by the coinciding advective flux, shown in Figure 3. We don't understand the source of these dips, but similar temperature gradient reversals were reported by [14] along the midlines of 2D convective cellular solutions at Ra $\sim 10^6$.

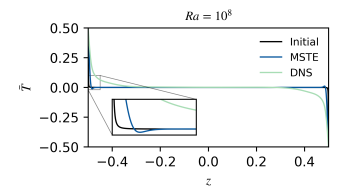


FIG. 2. Mean temperature profiles \bar{T} for Ra = 10^8 . The initial profile is given by Eq. (26). MSTE refers to the mean temperature profile of the marginally-stable thermally-equilibrated state. DNS was obtained by performing a 2D nonlinear simulation of Eqs. (1)–(3) with Dedalus until thermal and kinetic relaxation. DNS temperature data are horizontally- and time-averaged. The initial, MSTE, and DNS profiles have increasingly relaxed boundary regions respectively. The MSTE profile exhibits prominent dips, nested alongside the boundary regions. The source of this feature is not well understood, but similar temperature gradient reversal regions were found by [14] along the midlines of 2D convective cellular solutions at Ra $\sim 10^6$.

Equilibrated states exhibit distinct behaviors for large and small Ra. This contrast is illustrated in Figure 3, where we give heat flux profiles and eigenvalue spectra for two cases: Ra = 2×10^5 and Ra = 10^9 . For $Ra = 2 \times 10^5$, there is a single presumed local maximum in the eigenvalue spectrum, whose adjacent modes' advection profiles occupy the bulk of the domain. These states have relaxed boundary layers which gradually subside as advection becomes the dominant flux component. Such transitional regions do not appear for $Ra = 10^9$, where the shift from diffusion to advection is sharp, requiring pronounced, small-scale advection profiles belonging to modes in the third local maxima pair of the eigenvalue spectrum. Traversing towards the midline of the domain, we see increasingly thicker advection profiles corresponding to modes in the second local maxima pair, eventually culminating in the dominant $k_x = 1.5\pi$ mode.

Thermally equilibrated states with large Ra tend to have diverse combinations of marginal modes. In every case, the $k_x=1.5\pi$ mode is included and is often unaccompanied by adjacent modes. In Figure 4 we give the wavenumbers k_x of marginal modes, color-coded according to their adjacencies. Marginal modes often appear in neighboring pairs, where the larger mode is denoted with an x and the smaller mode is denoted with a +. In these cases we infer the existence of a local maximum eigenvalue between the two adjacent marginally-stable modes. For Ra $\geq 10^6$, a second branch of marginal modes is introduced as shown in light green, obeying some power-law with respect to Ra. The introduction of this largest branch is likely required to oppose the diffusion of the

thinning boundary layers. For Ra $\geq 10^8$, a third branch appears (shown in light blue), splitting the widening gap between the other branches. This development is associated with moderately thick advection profiles, filling a niche in the total flux by uniting the sharp profiles of the largest branch and those of the bulk-domain-oriented smallest branch.

The largest marginal wavenumber $\max\{k_x\}$ (represented by the light green branch in Figure 4) serves as a inverse minimum length-scale in the x direction. The finest structures in $\bar{T}(z)$ appear near the boundaries, providing us with a complementary minimum length-scale for z. We define the interior boundary layer threshold as the z-coordinate at which $\frac{\partial \bar{T}}{\partial z} = 0$. In Figure 5, we illustrate the constant ratio between these two quantities for various Ra. The quasilinear model cannot describe small-scale behaviors in either direction. Discretization of the normal modes allows us to collapse the problem into a single dimension, but this comes at a cost.

In general, MSTE can be characterized by their boundary layer height, from which, the rest of their properties follow. In Figure 6, we illustrate the scaling behavior of the boundary layer height $\delta = \mathrm{Ra}^{-1/3}$. This is consistent with Malkus' classical marginal-stability theory, a scaling argument which perceives the boundary regions as subdomains which are themselves marginally-stable [4].

The Nusselt number, which measures convective performance, reduces to

$$Nu = \frac{\langle A^2 \langle w'T' \rangle_x - \mathcal{P} \frac{\partial \bar{T}}{\partial z} \rangle_z}{\langle -\mathcal{P} \frac{\partial \bar{T}}{\partial z} \rangle_z}.$$

There is no general consensus surrounding the scaling behavior of Nu for high Ra systems, which are of particular importance in astrophysical systems. In Figure 7 we report Nu for MSTE, "steady rolls" ECS [13], and DNS [20, 21]. All states obey Nu $\sim \text{Ra}^{\beta}$ with classical scaling $\beta = 1/3$ in the ECS and MSTE. [13] hypothesized that Nu of ECS which admit classical Malkus scaling must always exceed 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 [15, 16], inhibiting heat flux. We might also anticipate the existence of similar equilibria, having smaller Nu and occupying complementary nodes in the Markov chain whose long-term behavior agrees with DNS.

V. SIMULATIONS WITH THERMALLY EQUILIBRATED INITIAL CONDITIONS

This investigation is partially motivated by the prospect of decreasing DNS times by employing MSTE initial conditions. Conventionally, Eqs. (1)–(3) are solved

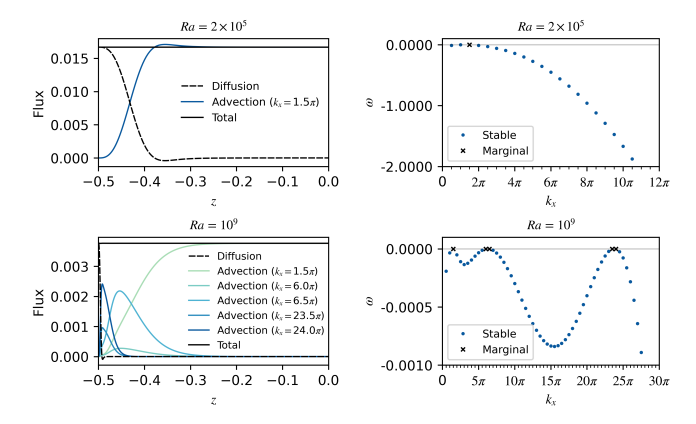


FIG. 3. Heat fluxes (left) and eigenvalue spectra (right) of equilibrated states $Ra = 2 \times 10^5$ (top) and $Ra = 10^9$ (bottom). Advection profiles belong to marginally-stable modes. For low Ra, a single mode with $k_x = 1.5\pi$ is sufficient to oppose boundary layer diffusion and facilitate heat flux throughout the bulk of the domain. For large Ra, high-wavenumber modes contribute pronounced small-scale advection profiles which tightly hug the thin boundary layers. A combination of of progressively wider advection profiles is necessary to transition to the $k_x = 1.5\pi$ mode.

via DNS with the linear initial conditions

$$T(x,z)\big|_{t=0} = 0.5 - z$$

 $\mathbf{u}(x,z)\big|_{t=0} = \mathbf{0}$
 $p(x,z)\big|_{t=0} = 0.$ (24)

t in this section now refers to the simulation time as opposed to the thermal equilibration time parameter used previously. We define the MSTE initial conditions

$$T(x,z)\big|_{t=0} = \bar{T}(z) + \sqrt{2} \sum_{n=1}^{N} A_n \operatorname{Re} \Big[\theta_n(z) e^{ik_{x_n} x} \Big]$$

$$\mathbf{u}(x,z)\big|_{t=0} = \sqrt{2} \sum_{n=1}^{N} A_n \operatorname{Re} \Big[\Big(U_n(z) \hat{x} + W_n(z) \hat{z} \Big) e^{ik_{x_n} x} \Big]$$

$$p(x,z)\big|_{t=0} = \sqrt{2} \sum_{n=1}^{N} A_n \operatorname{Re} \Big[P_n(z) e^{ik_{x_n} x} \Big]$$
(25)

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 at the *n*th marginal mode respectively. The $\sqrt{2}$ factor is necessary for eigenfunction normalization.

Simulations with MSTE initial conditions do not exhibit a convective-transient period, as the large-scale anatomy of convective cells exists on initialization. This is illustrated in Figure 8, where the sharp spike in Nu is associated with a turbulent transitional period of upwelling until the distinctive plume-like structure is established. Simulation of this transitional period is prohibitive [22]. For high Ra experiments, researchers often "bootstrap" data by initializing simulations with the results of similar Ra runs [20, 23]. MSTE can be perceived as a set of initial conditions, designed for avoiding the simulation of transitional high Reynolds number flows.

MSTE are laminar, lacking the small-scale structures associated with moderate to high Ra experiments. The absence of chaotic dynamics is an apparent consequence of the quasilinear assumptions. If we treat 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 turbulent flows, such as Boussinesq's turbulent-viscosity hypothesis [24], where the emergence of small-scale velocity structures tends to increase total shear [25–27], thereby impeding buoyancy-driven flows and decreasing advection in the bulk of the

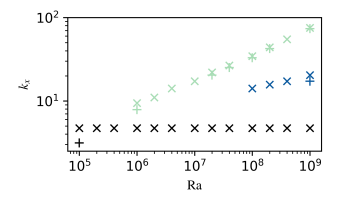


FIG. 4. Wavenumbers of marginally-stable modes of thermally equilibrated states. Markers are color-coded according to their adjacent local maxima index in the eigenvalue spectrum. For example, the spectrum corresponding to Ra = 10^5 has adjacent marginal wavenumbers $k_x = \pi, 1.5\pi$. The Ra = 10^9 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 branch obey a power-law relationship with Ra

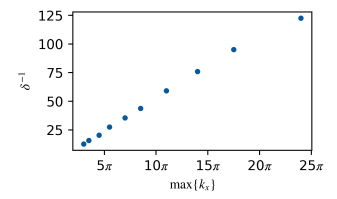


FIG. 5. For Ra $\geq 10^6$, the maximum marginally-stable wavenumbers (corresponding to the green X markers in Figure 4) are inversely related to the boundary layer heights δ . (max $\{k_x\}$)⁻¹ gives a minimum x length scale for the perturbations, and consequently, the advection. We expect the minimum z length scale to agree with the boundary layer height δ because otherwise the boundary layer would continue to diffuse. This is illustrated in the lower left corner of Figure 3, where the advection profile for $k_x = 24\pi$ is tightly flanked by the surrounding boundary layer. This suggests that the minimum x and z length scales obey some constant ratio over various Ra.

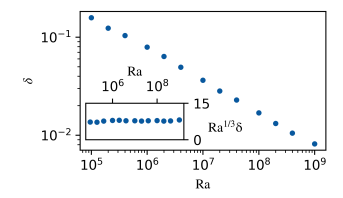


FIG. 6. Boundary layer height δ of marginally-stable thermal equilibria. We define the threshold of each boundary layer as the z-coordinate at which $\frac{\partial \bar{T}}{\partial z} = 0$, corresponding to the local extrema of the equilibrated curve in Figure 2. Plotting on a log-log scale, we find that δ and Ra obey a power-law relationship. We also demonstrate that $\mathrm{Ra}^{1/3}\delta$ is approximately constant with respect to Ra which is consistent with [4, 13]

domain. We could also attribute the diffuse DNS temperature profile in Figure 2 with unsteady boundary-layer penetration and mixing that MSTE do not exhibit.

The average kinetic energies of MSTE are significantly larger than those associated with the statistically-steady state, as shown Figure 9. This is postpones kinetic relaxation because the large-scale flows decay on a viscous time-scale

$$t_{\nu} \sim \sqrt{\frac{\mathrm{Ra}}{\mathrm{Pr}}}.$$

Consequently, MSTE initial conditions do not reduce the simulation time required to achieve a statistically-steady state – instead they increase it considerably. This suggests the MSTE background state perspective is partially flawed, as an appropriate background state would approximate $\langle |\mathbf{u}|^2 \rangle_{\mathcal{D}}$ with more fidelity.

VI. DISCUSSION

In general, the reduced model we study in this paper can be perceived as a unique way to describe Rayleigh–Bénard convection. To compute MSTE, we construct a marginally-stable mean temperature profile and develop the quasilinear model via independent time-scales. Solutions retain several key features that are ubiquitous in experiments and simulations: Nu $\sim \mathrm{Ra}^{1/3}$ scaling, large-scale convective cell structures, and agreement with classical minimum length-scale approximations [4]. They also exhibit unique and unexpected features: mean temperature gradient-reversals/dips, kinetic exaggeration, and greater Nu than other time-invariant solutions. We obtain indistinguishable solutions despite initializing

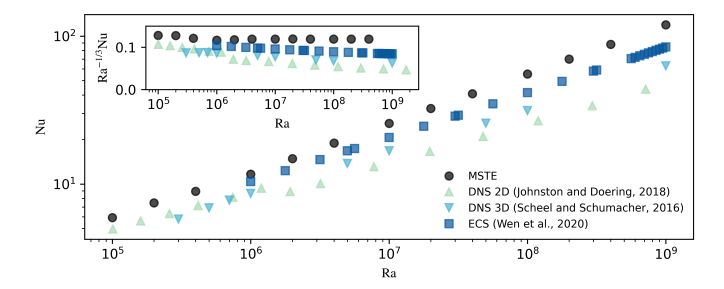
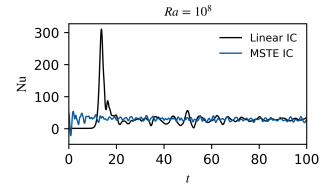


FIG. 7. Nusselt numbers are shown for MSTE, aspect-ratio-optimized "steady rolls" ECS [13], as well as statistically-steady 2D and 3D DNS [20, 21]. All datasets obey power-law relationships, with the MSTE and ECS admitting classical scaling $\text{Nu} \sim \text{Ra}^{1/3}$. MSTE have greater Nu than 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.



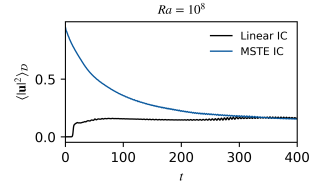


FIG. 8. Nusselt numbers of simulations performed at $\rm Ra=10^8$ with the conventional initial conditions (black) and MSTE initial conditions (blue). MSTE simulations do not undergo a convective-transient period because the characteristic large-scale convective cell structure exists on initialization.

FIG. 9. Average kinetic energies are reported for the simulations performed in Figure 8 at Ra = 10^8 . The combination of eigenfunctions associated with MSTE have significantly more kinetic energy than the statistically-steady state. Kinetic equilibrium is achieved according to the viscous timescale $t_{\nu} \sim \sqrt{{\rm Ra}/{\rm Pr}}$.

with various mean temperature profiles $\bar{T}(z,0)$. Thus solutions might be unique, but we cannot demonstrate this rigorously.

Simulations performed with MSTE initial conditions Eq. (25) do not undergo an early convective transient period, but have faster flows and greater Nu when compared with DNS. From a dynamical systems perspective, unstable orbits depart from MSTE and approach the global attractor (the compact manifold of states whose weighted-average agrees with experiments) on a viscous time-scale. This requires more CPU-hr to achieve relaxation as compared to the conventional linear initial

condition Eq. (24).

As previously noted, solving the EVP for statisticallysteady DNS data 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. We might curtail the disagreement between MSTE and DNS by loosening our marginal-stability criterion. Should the time-scales not be entirely separate, we might anticipate the long-term persistance of moderately unstable modes. Such an instability cannot be described by our model independently (we do not know, a priori, how unstable MUTE should be) but it might prove fruitful to investigate moderately unstable thermal equilibria (MUTE). More precisely, instead of traversing the manifold of marginally-stable states toward MSTE, we could consider the manifold of states whose maximum eigenvalue(s) $\omega_{\rm max}$ agree with some other stability constraint. Note that under this definition, ECS are themselves MUTE, because they are thermally-equilibrated and generally unstable, though they need not satisfy the quasilinear form.

We cannot use our reduced quasilinear model to predict Nu $\sim \mathrm{Ra}^{\beta}$ for large Ra without some complementary stability constraint. The eigenvalues of DNS data might provide an informative stability constraint (we could find MUTE and compare them with DNS data), but this would be self-defeating in general. Instead we seek to combine reduced models to approximate β without DNS. Under the marginal-stability constraint, we found that MSTE exhibit classical Malkus scaling, having marginally-stable boundary layers with $\beta = 1/3$. We could instead require that the eigenvalues of MUTE agree with the eigenvalues of the analytic 1D boundary layer predictions of [28, 29] (which remain in development), allowing us to refine and expand these estimates into 2D

Ra	\Pr	$k = 2\pi/\Gamma$
$10^{13/4}$	1	π
1.9×10^{3}	1	π
2×10^3	1	π
2.25×10^{3}	1	π
2.5×10^3	1	π
2.75×10^3	1	π
3×10^3	1	π
$10^{14/4}$	1	π
3.5×10^3	1	π
4×10^3	1	π
4.5×10^3	1	π
5×10^3	1	π
$10^{15/4}$	1	π
8×10^3	1	π
10^{4}	1	π
$10^{17/4}$	1	π
$10^{18/4}$	1	π
$10^{19/4}$	1	π
10^{5}	1	π
$10^{21/4}$	1	π

MUTE. Accessible MUTE states would not rely on DNS, satisfying only the quasilinear problem and stability constraint. They ought to agree with DNS data more than MSTE, ideally having slower flows This is reasonable to expect because accurate boundary layer approximations and DNS data should have similar stability characteristics. By definition, MUTE are unstable and would therefore admit more diffuse boundary layer structures, requiring smaller $A^2(t)$ (less advection) to satisfy the stability constraint and thereby reducing Nu, $\langle |\mathbf{u}|^2 \rangle_{\mathcal{D}}$ and possibly β .

APPENDIX

A. Initial buoyancy profile

We initialize the thermal-equilibration algorithm with [28]'s analytical thermal boundary layer equation for turbulent Rayleigh-Bénard convection, given by

$$\bar{T}_0(\xi) = \frac{\sqrt{3}}{4\pi} \log \frac{(1+a\xi)^3}{1+(a\xi)^3} + \frac{3}{2\pi} \arctan\left(\frac{4\pi}{9}\xi - \frac{1}{\sqrt{3}}\right) + \frac{1}{4}$$

$$\xi = \frac{z}{\delta_0}, \qquad a = \frac{2\pi}{3\sqrt{3}}$$
(26)

where δ_0 is the boundary layer height. We expect that each Ra is 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, implying that solutions may be unique. An example of Eq. (26) is given by the blue curve in Figure 2.

B. MSTE Metrics

$N_x \times N_z$	Nu	Re
128×65	1.056590	1.617336
128×65	1.145807	2.682155
128×65	1.212037	3.317190
128×65	1.355410	4.550975
128×65	1.474455	5.537770
128×65	1.575599	6.391812
128×65	1.663162	7.159844
128×65	1.714193	7.624400
128×65	1.808754	8.526064
128×65	1.926775	9.740578
128×65	2.025985	10.85141
128×65	2.111714	11.88534
128×65	2.204811	13.09152
128×65	2.476330	17.05494
128×65	2.648664	20.07400
128×65	3.122843	29.50047
128×65	3.665041	42.29585
128×65	4.287042	59.56858
128×65	4.994322	82.84462
128×65	5.795869	114.2355

$10^{22/4}$	1	π	128×97	6.703915	156.5252
$10^{23/4}$	1	π	128×97	7.732236	213.4031
10^{6}	1	π	128×97	8.896615	289.7982
$10^{25/4}$	1	π	256×97	10.21546	392.2837
$10^{26/4}$	1	π	256×129	11.71065	529.6372
$10^{27/4}$	1	π	256×129	13.40898	713.6005
10^{7}	1	π	512×129	15.34493	959.9367
1.35×10^{7}	1	π	512×129	16.46456	1119.932
1.5×10^{7}	1	π	512×129	16.87881	1182.172
1.6×10^{7}	1	π	512×193	17.13944	1222.005
1.65×10^{7}	1	π	512×193	17.26636	1241.484
1.7×10^{7}	1	π	512×193	17.39216	1260.709
1.736×10^{7}	1	π	512×193	17.48282	1274.414
1.76×10^{7}	1	π	512×193	17.54351	1283.493
$10^{29/4}$	1	π	512×193	17.58987	1290.377
1.786×10^{7}	1	π	512×193	17.60946	1293.277
1.8×10^{7}	1	π	512×193	17.64500	1298.522
1.85×10^{7}	1	π	512×193	17.77160	1317.122
1.9×10^{7}	1	π	512×193	17.89693	1335.501
1.95×10^{7}	1	π	512×193	18.02044	1353.657
2×10^7	1	π	512×193	18.14193	1371.593
2.4×10^{7}	1	π	512×193	19.04221	1507.890
2.8×10^{7}	1	π	512×193	19.83413	1633.418
$10^{30/4}$	1	π	512×193	20.47798	1739.647
4.5×10^{7}	1	π	512×193	22.44036	2087.182
$10^{31/4}$	1	π	512×193	23.76002	2340.822
10^{8}	1	π	768×257	27.50669	3144.931
$10^{33/4}$	1	π	768×257	31.81154	4220.616
2.15×10^{8}	1	π	768×257	33.36657	4649.541
$10^{34/4}$	1	π	768×257	36.75427	5658.648
4.64×10^{8}	1	π	896×321	40.44652	6875.704
$10^{35/4}$	1	π	896×321	42.42917	7580.057
10^{9}	1	π	1024×321	48.94284	10145.79
$10^{37/4}$	1	π	1024×321	56.42926	13571.10
2.15×10^{9}	1	π	1024×321	59.13988	14935.81
2.5×10^{9}	1	π	1024×321	61.38714	16116.94
$10^{38/4}$	1	π	1024×321	65.06338	18145.61
$10^{153/16}$	1	π	1024×321	67.42466	19511.79
4×10^9	1	π	1024×321	68.96487	20429.24
4.64×10^{9}	1	π	1024×321	71.58241	22019.93
$10^{39/4}$	1	π	1024×321	75.09725	24262.22
10^{10}	1	π	1024×321	86.68318	32430.06
$10^{41/4}$	1	π	1024×321	100.0909	43339.02
2×10^{10}	1	π	1024×321	103.0738	45995.60
2.15×10^{10}	1	π	1024×321	104.9517	47705.08

Table S1: Details for numerical solutions with Pr = 1 and $\Gamma = 2$. N_x and N_z represent the numbers of Fourier and Chebyshev modes, respectively.

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