Lagrangian Statistics of Heat Transfer in Homogeneous Turbulence Driven by Boussinesq Convection

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Abstract: The movement of heat in a convecting system is typically described by the nondimensional Nusselt number, which involves an average over both space and time. In direct numerical simulations of turbulent flows, there is considerable variation in the contributions to the Nusselt number, both because of local spatial variations due to plumes and because of intermittency in time. We develop a statistical approach to more completely describe the structure of heat transfer, using an exit-distance extracted from Lagrangian tracer particles, which we call the Lagrangian heat structure. In a comparison between simulations of homogeneous turbulence driven by Boussinesq convection, the Lagrangian heat structure reveals significant non-Gaussian character, as well as a clear trend with Prandtl number and Rayleigh number. This has encouraging implications for simulations performed with the goal of understanding turbulent convection in natural settings such as Earth's atmosphere and oceans, as well as planetary and stellar dynamos.

Keywords: Lagrangian transport; geophysical fluid flows; convection; turbulence; statistical methods

1. Introduction

Convection occurs in many natural settings, including the oceans, the atmosphere, and the interior of the Earth and the stars. These convective flows are not typically constrained by solid boundaries, and therefore have some characteristics that differ from experimental set-ups like Rayleigh-Bénard convection. Convection-driven flow in natural systems is often turbulent, but may be unaffected by boundary layers, large-scale circulations, or regular convection rolls. These large-scale flow features may be absent, or they may be so large that they do not influence the local characteristics of the turbulence. Thus convective turbulence in the interior of an atmospheric layer or the interior of a stellar convection zone appears homogeneous on scales smaller than the pressure scale-height. These are the length scales accessible to study with direct numerical simulations.

Early studies of convection in natural settings focused heavily on the development of Lagrangian statistics to describe diffusion and dispersion, e.g., Gifford [1], Roberts [2], Sutton [3], Frenkiel and Katz [4], Kellogg [5]. Lagrangian tracer particles continue to be widely regarded as an ideal way to describe turbulent diffusion and transport (see for example [6,7]). Recent work using Lagrangian statistics to examine fundamental properties of turbulent convection is delivering a wealth of new insight into these processes [8–13]. Exit-time and exit-distance statistics, sometimes also called first-passage times, can be produced with Lagrangian particles, and have been shown to provide useful information that minimizes the effect of intermittency. Popular examples of exit-time and
exit-distance statistics are the doubling-time [14], finite-size Lyapunov exponents (FSLE) [15–18], finite-time Lyapunov exponents (FTLE), and inverse structure functions [19,20].

The heat transfer is one of the most fundamental aspects of convection. Measuring the evolving temperature, and how it correlates with velocity, is a problem of active interest for oceanographers and atmospheric scientists (e.g., as discussed by [21]). Experimental investigations of convection [8,13,22] have recently engineered smart particles that can sense and record the temperature that they experience in a turbulent flow. This experimental development also increases our need to define clearer theoretical measures to understand heat transfer in turbulent convection.

Heat transfer has typically been approached using a dimensionless heat flux averaged over the simulation time and volume, called the Nusselt number (Nu). However, in many cases the Nusselt number describes the heat transfer in a limited way. This is particularly true in simulations of homogeneous turbulent convection, where the most relevant dynamical length scale is not necessarily associated with the imposed temperature gradient, and the dynamical length scale of the flow may actively change and evolve. A wide variation of the Nusselt number in both space and time results. Intermittency can be observed for different diagnostics in a wide range of turbulent flows [23], including stratified turbulence relevant to geophysical fluids (e.g., [24]). For comparing different realizations of homogeneous turbulent convection, a more detailed description of the heat flux is needed. In this work we therefore discuss the development of a new higher order statistic related to the heat transfer.

We build this new statistic as a Lagrangian exit-distance statistic to characterize how heat is transported over distances in a turbulent flow. The use of an exit-distance statistic has the advantage, documented in Lagrangian descriptions of turbulence [18], that it permits us to compare fluid particles that have transported the same amount in temperature, rather than particles that have diffused for the same length of time. This removes contamination due to crossover from other temperature scales, a quality that has helped to clarify intermittent behavior in hydrodynamic turbulence (e.g., as discussed in [25]). Examination of this statistic provides higher-order information about the heat transfer from a different perspective, supplementing a straightforward global average like the Nusselt number.

Heat transfer and heat flux are highly dependent on a fluid’s Prandtl number, the ratio of viscosity to thermal diffusivity (e.g., [26–28]). The fundamental idea is that low Pr flows can support small-scale velocity fluctuations while the corresponding small-scale temperature fluctuations are smoothed out by diffusion; the opposite situation occurs at large Pr. In the oceans, viscosity dominates and the Prandtl number ranges between approximately 7 (warm sea water) to 13 (sea water near freezing). Atmospheric convection occurs in the air, where the Prandtl number can range between \(O(10^{-1})\) and \(O(10^1)\) with a typical value of 0.72 for air. In the solar convection zone thermal diffusivity dominates, and Prandtl numbers on the order of \(O(10^{-5})\) to \(O(10^{-7})\) are estimated. Current direct numerical studies of convection typically are able to describe a range of Prandtl numbers between \(O(10^{-2}) < Pr < O(10^2)\) (e.g., [29,30]), while dynamo studies often focus on \(Pr = 1\). A more detailed theoretical understanding of how the dynamics of thermal fluctuations change when different Prandtl numbers are considered is key to effectively modeling convection in these different settings. Using our new Lagrangian exit-distance statistic, the Lagrangian heat structure, we compare simulations of turbulent convection over the range of Prandtl numbers \(0.1 < Pr < 10\). This comparison illustrates the utility of the Lagrangian heat structure to expose differences in the transport of heat.

This work is structured as follows. In Section 2 we describe the physical and numerical models used to produce simulations of homogeneous turbulent Boussinesq convection at different Prandtl numbers. In Section 3 we discuss the Nusselt number and its variation during our simulations. In Section 4 we derive a Lagrangian exit-distance statistic to provide more detail about the heat transfer than can be extracted from the Nusselt number. We present the results of this new diagnostic for our simulations. In Section 5 we discuss the implications of these results.
2. Simulations

We perform direct numerical simulations of homogeneous turbulence driven by Boussinesq convection. In each simulation, the dynamical equations are solved using a pseudospectral method in a cubic simulation volume with a side of length $2\pi$. The nondimensional equations for Navier-Stokes turbulence driven by Boussinesq convection are

$$\frac{\partial \omega}{\partial t} - \nabla \times (v \times \omega) = \nu \nabla^2 \omega - \nabla \theta \times g_0, \quad (1)$$

$$\frac{\partial \theta}{\partial t} + (v \cdot \nabla) \theta = \kappa \nabla^2 \theta - (v \cdot \nabla) T_0, \quad (2)$$

$$\nabla \cdot v = 0. \quad (3)$$

These equations include the solenoidal velocity field $v$, the vorticity $\omega = \nabla \times v$, and the thermal fluctuation $\theta$ about the mean temperature $T_0$. In our fluid model, the total temperature is expressed as the sum of mean and fluctuating quantities: $T = T_0 + \theta$. The turbulent flow is driven solely by the fixed, unitary, mean temperature gradient in the vertical direction, opposing the direction of gravity; the gradient of the mean temperature can be expressed fully by the $z$-component $\partial_z T_0$. In Equation (1), $g_0$ is a unit vector in the direction of gravity; the term that includes $g_0$ is commonly known as the buoyancy force. The simulation volume and physical set-up are indicated alongside a visualization of thermal energy in Figure 1. Two dimensionless parameters appear in the equations: $\hat{\nu}$ and $\hat{\kappa}$. They derive from the kinematic viscosity $\nu$ and the thermal diffusivity $\kappa$, and are defined so that the Prandtl number $Pr \equiv \nu / \kappa = \hat{\nu} / \hat{\kappa}$. The thermal diffusivity $\kappa$ relates to the nondimensional quantity by $\hat{\kappa} = \kappa / \sqrt{g_0 \alpha \Delta T L^2}$. A low-storage third-order Runge-Kutta method [31] is used for the time-integration of Equations (1)-(3).

![Figure 1](image.png)

**Figure 1.** Visualization of thermal energy in simulation HC4 described in Table 1. The physical set-up of the simulations is also indicated.

Boundary conditions strongly influence the structure and dynamics of a flow. For convection simulations the choice of fully periodic boundary conditions (sometimes called homogeneous Rayleigh-Bénard boundary conditions) allows a macroscopic elevator instability to form [32], which manifests as an exponentially growing mean velocity component along the direction of gravity. The elevator instability rapidly destroys the turbulent flow field, making it impossible to collect long-time steady-state statistical data. To avoid this, the simulation volume considered in this work is confined by quasi-periodic boundary conditions (as in [33–35]). Quasi-periodic boundary conditions explicitly suppress the mean flow parallel to gravity, which is removed at each time step. Our simulations are pseudospectral, and therefore a vertical mean flow is straightforwardly isolated in
Fourier space. These boundary conditions are also periodic, like the homogeneous Rayleigh-Bénard boundary conditions. Quasi-periodic boundary conditions combine the conceptual simplicity of statistical homogeneity with a physically natural convective driving of the turbulent flow. With these boundary conditions, a large-scale structuring of the turbulent flow, such as the convection-cell pattern caused by Rayleigh-Bénard boundary conditions does not occur. Unlike Rayleigh-Bénard convection, our simulations therefore produce homogeneous anisotropic convection.

Table 1. Parameters of homogeneous hydrodynamic convection simulations. The simulations are defined by the value of the Rayleigh number, calculated according to Section 2.2, and the Prandtl number. The Nusselt number is calculated according to Equation (7). The Grashof number is defined \( Gr = Ra/Pr \). The nondimensionalized kinematic viscosity \( \hat{\nu} \) and the thermal diffusivity \( \hat{\kappa} \) are also provided.

| Simulation | HC1  | HC2  | HC3  | HC4  |
|-----------|------|------|------|------|
| grid size | 1024³ | 512³ | 1024³ | 1024³ |
| simulation time (\( \tau_\eta \)) | 484  | 1605 | 416  | 427  |
| \( n_p \) (10⁶) | 4    | 4    | 4    | 8    |
| Prandtl number | 0.1  | 1.0  | 10   | 1.0  |
| Rayleigh number Ra | \( 2.5 \times 10^5 \) | \( 2.5 \times 10^5 \) | \( 2.5 \times 10^5 \) | \( 2.5 \times 10^6 \) |
| Grashof number Gr | \( 2.5 \times 10^6 \) | \( 2.5 \times 10^5 \) | \( 2.5 \times 10^4 \) | \( 2.5 \times 10^6 \) |
| Reynolds number Re(\( L_{\text{conv}} \)) | 3230 | 1350 | 570  | 4180 |
| Nusselt no. NuL | 582  | 2025 | 6980 | 6503 |
| time maximum of NuL | 651  | 3200 | 8873 | 7903 |
| \( \eta_{\text{kol}} \) (10⁻³) | 3.43 | 7.60 | 17.0 | 3.23 |
| \( \tau_\eta \) (10⁻²) | 1.85 | 5.03 | 4.57 | 1.64 |
| \( \eta_{\text{therm}} \) (10⁻³) | 1.92 | 7.57 | 3.01 | 3.24 |
| \( \tau_{\text{therm}} \) (10⁻²) | 10.18 | 2.21 | 2.52 | 2.86 |
| \( \hat{\nu} \) | \( 6.3 \times 10^{-4} \) | \( 2 \times 10^{-3} \) | \( 6.3 \times 10^{-3} \) | \( 6.3 \times 10^{-4} \) |
| \( \hat{\kappa} \) | \( 6.3 \times 10^{-3} \) | \( 2 \times 10^{-3} \) | \( 6.3 \times 10^{-4} \) | \( 6.3 \times 10^{-4} \) |

A summary of the fundamental parameters describing each simulation is given in Table 1. In this table, we define the Reynolds number to be \( Re = \langle E_v^{1/2} L \rangle / \hat{\nu} \), where \( E_v = v^2 / 2 \) is the kinetic energy, and the brackets indicate a time-average. For statistically homogeneous turbulent convection the characteristic length scale is the instantaneous temperature gradient length scale \( L \equiv L_{\text{conv}} = T_0 / \partial_x T \), where \( T_0 \) is the root-mean-square of thermal fluctuations [36]. We measure length in units of the Kolmogorov microscale \( \eta_{\text{kol}} \equiv (\hat{\nu}^3 / \epsilon_v)^{1/4} \) and time in units of the Kolmogorov time-scale \( \tau_\eta \equiv (\hat{\nu} / \epsilon_v)^{1/2} \), where \( \epsilon_v \) is the dissipation of kinetic energy; the Kolmogorov scales are the smallest length and time scales that characterize turbulent flows. The Kolmogorov microscale multiplied by \( k_{\text{max}} \), the highest wavenumber in the simulation, is often used to test whether a simulation is adequately resolved on small spatial scales. In this work all of the simulations fulfill the standard criterion used for hydrodynamic turbulence due to Pope [37] for adequate spatial resolution of the velocity field: \( k_{\text{max}} \eta_{\text{kol}} > 1.5 \). The grid size used and \( \eta_{\text{kol}} \) are given in Table 1.

Because this work treats convection at different Prandtl numbers, an independent requirement is necessary to be sure that the field of thermal fluctuations \( \theta \) is also adequately resolved. This question has been closely examined for Rayleigh-Bénard convection (e.g., [38,39]), and the Nusselt number has been found to be particularly sensitive to the resolution. In contrast to Rayleigh-Bénard convection, for homogeneous convection we expect that the buoyancy force will be significant, the flow is potentially strongly anisotropic, and strong velocity and temperature gradients can occur with equal
probability throughout the flow. For this reason, a resolution criterion for the thermal fluctuations equivalent to that used for the velocity field is desirable in our simulations. By dimensional argument, the equivalent to Pope’s resolution criterion [37] can be straightforwardly extended to the field of thermal fluctuations. The dimension of kinematic viscosity is length$^2$/time and the dimension of the kinetic energy dissipation rate $\epsilon_v$ is length$^2$/time$^3$. The only combination of these dimensions that has the dimension of time is the Kolmogorov time-scale $\tau_\eta$. The thermal diffusivity $\kappa$ also has units of length$^2$/time, and the thermal energy dissipation rate $\epsilon_\theta$ has the same units as the kinetic energy dissipation rate. Thus we define the thermal microscale $\eta_{therm} = (\hat{\kappa}^3/\epsilon_\theta)^{1/4}$. We consider the field of thermal energy to be adequately resolved in space if the thermal microscale fulfills the requirement $k_{max}/\eta_{therm} > 1.5$. Each of our simulations also fulfills this criterion. The satisfactory nature of this criterion is clear from a comparison of the average spectra of kinetic and thermal energy for simulations HC1, HC2, HC3, and HC4 (see Figure 2). For simulation HC2 ($Pr = 1$) the kinetic energy and thermal energy experience a similar level of damping. For simulation HC1 ($Pr = 0.1$) it is clear that the thermal energy spectrum experiences more damping than the kinetic energy spectrum; the reverse is true for simulation HC3 ($Pr = 10$). For simulation HC4 ($Pr = 1$), the spectra are similar to HC2, but exhibit a wider inertial range.

![Figure 2. Kinetic energy spectrum $E_v$ (a) and thermal energy spectrum $E_\theta$ (b) for simulations HC1, HC2, HC3, and HC4.](image)

2.1. Lagrangian Tracer Particles

Each simulation is launched with random Eulerian fields of unitary magnitude. After a statistically stationary steady state is attained by the convective driving, i.e., when the global kinetic energy has established a well-defined average value in time, the simulation is judged to have produced a physically realistic initial state to begin the calculation of Lagrangian statistics. This period of relaxation lasts more than $10^3\tau_\eta$ in each simulation. The positions of Lagrangian tracer particles are then initialized in a homogeneous random distribution. The total number of particles $n_p$ is given in Table 1. This is a standard density of tracer particles for such simulations, and the Lagrangian statistics produced have been tested and found to be well-resolved in space. The Eulerian time step for each simulation is much less than the limit of the CFL number, and chosen so that $\tau_\eta/\delta t \approx 100$; the same time step is used when integrating the trajectories of tracer particles. Lagrangian data is recorded every four Eulerian time steps for post-processing, so that $\tau_\eta/\delta t_L \approx 25$. At each time step the particle velocities, thermal fluctuations, and gradients of these quantities are interpolated from the instantaneous Eulerian velocity field using a tricubic (see for example [40]) polynomial interpolation scheme. Particle positions are calculated by numerical integration of the equation of motion using a third order Runge-Kutta
method. Each simulation is run for at least 400\(\tau_\eta\), and the total simulation time is given in Table 1. Our Lagrangian statistics are validated by examining Lagrangian single-particle diffusion, the average square distance a given particle has moved from its initial position. The Lagrangian single-particle diffusion in directions perpendicular and parallel to gravity is shown in Figure 3. In the figure, triangular brackets indicate an average over all Lagrangian particles in the simulation. The diffusion curves perpendicular to gravity exhibit the ballistic scaling with \(t^2\) for early times and the diffusive scaling with \(t\) for long times. The diffusion curves parallel to gravity are slightly superdiffusive for long times due to the anisotropy introduced by the convective driving.

For the purpose of further validation of our Lagrangian statistics, we examine the Lagrangian autocorrelation functions shown in Figure 4. These have the typical shape known from hydrodynamic turbulence (e.g., [6]); they are initially one, and decay smoothly to zero at long times without ever becoming negative. Velocity autocorrelation functions have been examined more recently to determine the anisotropy of magnetohydrodynamic (e.g., [41]), and rotating stratified systems (e.g., [42]). In Figure 4, the decay of the autocorrelation function of \(\theta\) takes place over a slightly longer time than the autocorrelation of velocity in the direction parallel to gravity. The velocities in the plane perpendicular to gravity have a much shorter correlation time, due to the anisotropy that convection introduces to the system. For simulation HC1 (\(Pr = 0.1\)) this decay time is more than twice as long as simulation HC3 (\(Pr = 10\)). A longer decay time is to be expected for simulation HC1 than for HC3, since the lower Prandtl number is achieved by using a lower viscosity.

### 2.2 Rayleigh Number

The Rayleigh number is a dimensionless measure of the convective driving: the product of the Prandtl number with the Grashof number, the ratio of the buoyancy to viscous force acting on a fluid. In comparisons of convecting systems, the Rayleigh number is typically held constant when the Prandtl number is varied so that results are compared from systems with identical convective driving. The Rayleigh number is typically defined by \(Ra = g\alpha\Delta TL^3/\nu\kappa\), where \(\Delta T\) is the temperature difference across the fluid layer of height \(L\). Because our system is periodic, the typical temperature difference across any height of fluid is defined only by the mean temperature gradient that we impose. This contrasts with the boundary conditions that are typically used in Rayleigh-Bénard convection, and has consequences for several nondimensional parameters that characterize a convecting system. Our mean temperature gradient is unitary, and therefore \(\Delta T = 1\) for \(L = 1\). Adopting \(L = 1\) for
the length scale of the fluid layer is a sensible choice in our nondimensional, periodic system; this length scale should be fixed to allow for the Rayleigh number to be well-defined. The quantities $g$ and $\alpha$ are also one in our nondimensional units. We therefore calculate the Rayleigh number for our homogeneous simulations in nondimensional units using 

$$Ra = \frac{1}{\hat{\nu} \hat{\kappa}}.$$ 

This Rayleigh number is held fixed at $Ra = 2.5 \times 10^5$ for the three simulations HC1, HC2, and HC3 described in Table 1. This is the highest Rayleigh number achievable using a grid size of $1024^3$ while also maintaining our criterion for adequate spatial resolution of the turbulent velocity and thermal fluctuations described above for the three simulations at different Prandtl numbers. The three simulations HC1, HC2, and HC3 provide a comparison of different Prandtl number flows at fixed Rayleigh number; simulations HC1 and HC4 provide a useful comparison of different Prandtl number flows at fixed Grashof number (e.g., [29]). Simulation HC4 uses a Rayleigh number that is ten times higher than simulation HC2, to allow for a comparison between results at different Rayleigh numbers for fixed Prandtl number.

**Figure 4.** Lagrangian autocorrelation function of (a) the velocity aligned with gravity $v_z$, and (b) the thermal fluctuations $\theta$. In (b) a dotted grey line is provided in the background as a reference for the autocorrelation function of $v_z$ in the $Pr = 1$ simulation HC2. Time is given in units of the Kolmogorov time-scale $\tau_\eta$. Triangular brackets indicate an average over Lagrangian particles.

### 3. Results: Nusselt Number

The Nusselt number, which describes heat transport, is standardly defined as (e.g., [43,44])

$$Nu = \frac{\langle v_z T \rangle_{V,t} - \kappa \partial_z \langle T \rangle_{V,t}}{\kappa \Delta T} \frac{L}{\kappa \Delta T}. \quad (4)$$

This definition is constructed using the total heat flux $Q = \langle v_z T \rangle_{V,t} - \kappa \partial_z \langle T \rangle_{V,t}$, where $T$ is the total temperature in dimensionless units. The triangular brackets $\langle \rangle_{V,t}$ indicate an averaging over the simulation volume $V$ and the simulation time $t$. The change in temperature $\Delta T$ and length scale $L$ are positive quantities, identical to the quantities used in the definition of the Rayleigh number. In our system the mean temperature gradient is constant and linear, so that $\partial_z \langle T \rangle_{V,t} \equiv \partial_z T_0 = -\Delta T/L = -1$. Using these definitions, the nondimensional thermal diffusivity can be seen to be identical to the thermal diffusivity $\kappa \equiv \kappa$. The formula for the Nusselt number can then be reduced to

$$Nu = \frac{\langle v_z T \rangle_{V,t}}{\kappa} + 1. \quad (5)$$

This formula is often used for Rayleigh-Bénard systems [8,9,22].
For Lagrangian particles the volume average can be performed as an average over Lagrangian fluid particles, since the particles sample the volume at high resolution. Equation (4) can be re-written

$$\text{Nu}_L = (\langle v_z T \rangle_p - \hat{k} \partial_z \langle T \rangle_p) \frac{1}{\hat{R}}. \tag{6}$$

Here \(v_z\) and \(T\) are Lagrangian measurements rather than Eulerian measurements and the brackets now indicate an average over all tracer particles \(p\) and the simulation time \(t\). We have used the fact that \(\Delta T = L = 1\) for all of our simulations, to eliminate non-Lagrangian measurements from this equation. Although Equation (6) is averaged in time, it is also useful to examine an instantaneous Nusselt number, which is not averaged in time:

$$\text{Nu}_L(t) = (\langle v_z T \rangle_p - \hat{k} \partial_z \langle T \rangle_p) \frac{1}{\hat{R}}. \tag{7}$$

For our linear temperature gradient, this reduces to

$$\text{Nu}_L(t) = \frac{\langle v_z T \rangle_p}{\hat{R}} + 1. \tag{8}$$

Obtaining identical results for Equations (7) and (8) provides a verification that the resolution of Lagrangian fluid particles is adequate. These definitions produce the same result for all of our simulations.

Nusselt numbers in simulations of turbulent flows are typically on the order of \(O(10^{1})\) to \(O(10^{3})\); the Nusselt numbers we calculate are in the high end of this range, and are given in Table 1. Several theoretical predictions for scaling of the Nusselt number with Prandtl number exist (e.g., [45,46]). Niemela [43] discuss that if the vertical heat transport were entirely by turbulent fluctuations, then the Nusselt number should scale as \(\text{Nu} \sim (RaPr)^{1/2}\). For the simulations we study in this work this scaling proves remarkably accurate, as shown in Figure 5. In this figure, there is a slight dislocation between simulation HC1 with \(Pr = 0.1\), and the simulations at \(Pr = 1\) and 10. Because simulations HC1 and HC4 have a significantly larger Reynolds numbers than HC2 and HC3, and more energy in small-scale motions, the heat transfer in these simulations is likely due more completely to turbulence. Laminar convective features are likely to account for the slightly higher Nusselt numbers at \(Pr = 1\) and 10. For the highest point on the horizontal axis of Figure 5, we can compare the simulations HC3 and HC4; the Nusselt number in HC3 is larger than in HC4, providing a calibration of how much turbulence affects this scaling relation.

One reason for the inaccuracies in this scaling stems from the use of an average Nusselt number. We find that the instantaneous Nusselt number can change rapidly and vary over a wide range during a simulation of steady state convection, so that neither the average Nusselt number nor its standard deviation characterizes the heat transfer particularly well. Other works have found a similar large time-variation in the Nusselt number, for example see Figure 1 of Schmalzl et al. [28]. To quantify this variation, we examine the probability density functions (PDFs) of the instantaneous Nusselt number from Equation (7) (see Figure 6). In this figure, the PDFs are all multi-modal; two of the four PDFs also feature a heavy tail. To some extent, this may be attributed to the fact that, in our homogeneous system intended to model convection in natural settings, local temperature gradients can become more significant than the imposed \(\Delta T\) and \(L\). Local temperature gradients can also change in time with the changing structure of the flow.

The Nusselt number also varies strongly in space (see Figure 7). Higher values of the Nusselt numbers cluster within coherent flow structures that can be associated with convective plumes. The wide range of Nusselt numbers obtained in these controlled direct numerical simulations makes the definition of an average Nusselt number to characterize the whole flow questionable. We therefore propose a more detailed diagnostic to provide additional information about the heat transport, which we called the Lagrangian heat structure.
Figure 5. The scaling of the Nusselt number with the product of Prandtl and Rayleigh numbers. The shaded region indicates one standard deviation of the instantaneous Nusselt number above and below the time-averaged value.

Figure 6. Probability density functions of the instantaneous Nusselt number for (a) simulation HC1 (Pr = 0.1), (b) simulation HC2 (Pr = 1.0), (c) simulation HC3 (Pr = 10), and (d) simulation HC4 (Pr = 1.0).
Figure 7. Visualization of particles in a slice of the simulation volume determined by the first three grid spaces in the z-direction in simulation HC1: two-dimensional visualization at the initial time point (left), and three dimensional visualization as the same particles trace the evolution of the evolving convective flow structures (right). Particles are colored by the instantaneous local Nusselt number that they experience between zero and $10^4$, with dark colors indicating a low Nusselt number and bright colors indicating a high Nusselt number.

4. Results: Lagrangian Heat Structure

4.1. Definition of the Lagrangian Heat Structure

To understand in better detail the transport of heat in convective turbulence and to supplement the information from the average Nusselt number, we define the Lagrangian heat structure. We develop this as an exit-distance statistic, similar in spirit to the finite-time Lyapunov exponent (FTLE) which is used to calculate Lagrangian coherent structures (LCS) [47]. Lagrangian coherent structures have recently been used to clarify length scales for the dynamo [48–50]. To calculate the FTLE, a certain finite time $\tau$ is chosen, and the separation $\delta$ of particle pairs is measured after that time is reached. The FTLE of each particle pair is then

$$\lambda_{\text{ftle}} = \frac{1}{\tau} \ln \left( \frac{\delta}{\delta_0} \right). \quad (9)$$

In this definition $\delta_0$ is the separation of the particle pair at the initial time. Rearranging this definition we see that

$$\exp (\tau \lambda_{\text{ftle}}) = \frac{\delta}{\delta_0}. \quad (10)$$

The FTLE is thus designed to supply a local rate associated with the integrated pair separation. It is this definition of a rate that can be a useful concept when we adapt it to examine changes in temperature. Instead of a time rate of change, we want to measure an analogous quantity that has units of inverse temperature, and is associated with the diffusion of Lagrangian tracer particles. Rather than emphasize particle pairs, as the FTLE does, we develop a single-particle Lagrangian diagnostic. A possible weakness of employing a single-particle diagnostic rather than a pair diagnostic is that the effect of macroscopic flows can dominate over the small scales. However, since larger scale motions contribute most strongly to heat transport, a single-particle diagnostic seems appropriate to the problem, and has the advantage of also permitting simpler interpretation. In addition, single-particle diagnostics can be averaged over many initial conditions using the conventional
methods described in Dubbeldam et al. [51]. This allows these diagnostics to be calculated in a way that is insensitive to the initial conditions of the flow. An investigation of heat transfer might be extended to pair diagnostics in the future.

We construct an equation for the heat in a similar form to Equation (10) so that

$$\exp(\Delta T \lambda_{hs}) = \Delta r.$$  \hspace{1cm} (11)

We choose a separation in temperature $\Delta T = |T - T_i|$ where $T_i$ is the initial temperature that a tracer particle feels. This temperature separation $\Delta T$ is nondimensional, expressed in units of the root-mean-square of temperature in the system. In Equation (11) we measure the distance that a single particle must diffuse before it achieves that change in temperature in units of the Kolmogorov microscale

$$\Delta r = \sqrt{(x - x_0)^2 + (y - y_0)^2 + (z - z_0)^2 / \eta_{kol}}.$$  \hspace{1cm} (12)

Here $(x_0, y_0, z_0)$ represents the initial position of the particle. This is a nondimensional construction of distance similar to that used for the diffusion curves in Figure 3. We thus define the Lagrangian heat structure as the ratio

$$\lambda_{hs} = \frac{1}{\Delta T} \ln(\Delta r).$$  \hspace{1cm} (13)

The formulation of Equation (13) as a natural logarithm is useful because it permits the particles that have not yet separated as far as a Kolmogorov microscale to be shown as negative values of $\lambda_{hs}$.

The Lagrangian heat structure reveals in a local and visual way the positions of particles that experience the fastest change in temperature. We further refine the Lagrangian heat structure in order to isolate only the distance along the mean temperature gradient,

$$\lambda_{hsz} = \frac{1}{\Delta T} \ln(\Delta z).$$  \hspace{1cm} (14)

Here the distance used is the distance that the particle moves against gravity, so that $\Delta z = |z - z_0| / \eta_{kol}$. This provides a Lagrangian exit-distance statistic for the transport of heat in the vertical direction. In Figure 8, visualizations show the spatial variation of $\lambda_{hsz}$ in a horizontal plane of simulation HC1. The color scale is chosen so that areas of extremely high vertical transport of heat, corresponding to low $\lambda_{hsz}$, are dark. Although the flow is turbulent, clear structures associated with high and low vertical heat transfer are identifiable, and are consistently positioned for different temperature separations. These visualizations may be compared to the visualization of the Nusselt number in Figure 7. The initial time for the calculation of $\lambda_{hsz}$ and the horizontal plane are the same as that shown in Figure 7, but the size and position of structures of high vertical transport of heat differ between these figures. In contrast to the instantaneous and rapidly changing Nusselt number, the Lagrangian heat structure represents how changes in heat are connected to an integrated diffusion process. The visualizations in Figure 8 thus reveal more information about the size of structures related to heat transfer in the flow.
Figure 8. Visualizations of the Lagrangian vertical heat structure $\lambda_{hsz}$ in the same slice of the simulation volume shown in Figure 7, in simulation HC1 ($Pr = 0.1$) for temperature separations $\Delta T = 0.1$ (left), $\Delta T = 0.5$ (middle), and $\Delta T = 1.0$ (right). Darker color indicates a lower value of $\lambda_{hsz}$.

4.2. Prandtl Number Dependence of Heat Transfer in Hydrodynamic Convection

For a chosen temperature difference, we calculate the probability density function $P$ and the cumulative distribution function (CDF) $C$ of $\lambda_{hsz}$ (see Figure 9). These distributions are calculated for 6 statistically independent flow realizations in simulation HC1. The probability density function deviates from a Gaussian distribution mildly because of the larger lower wing, which is displayed in greater detail by the CDF. We find that this distribution has a shape that changes only slightly for these statistically independent initial points; it does not exhibit the large time intermittency that we observe in the Nusselt number. A similar result is found for each of the other simulations in Table 1. Thus by examining distributions and moments of the Lagrangian vertical heat structure $\lambda_{hsz}$ calculated based on all available time data for each simulation, we characterize universally how heat changes in each simulation.

Figure 9. Distribution functions of the Lagrangian vertical heat structure $\lambda_{hsz}$ for $\Delta T = 0.1$, constructed from approximately 4 million Lagrangian tracer particles in simulation HC1 ($Pr = 0.1$). (a) Probability density function constructed from the data for 6 different statistically independent flow realizations. (b) Cumulative distribution function. Six dashed lines show distributions constructed from data at different independent flow realizations; the solid line represents a distribution constructed from the data at all 6 times. A dotted line marks the point where $C(\lambda_{hsz}) = 0.5$.

A sample of the cumulative distribution functions $C(\lambda_{hsz})$ for simulations HC1, HC2, and HC3 and several temperature separations $\Delta T$ are shown in Figure 10. The CDFs we produce are broader
for higher Prandtl number, indicating that as the Prandtl number is increased, for more particles, the distance that a particle must diffuse to reach a certain temperature difference is shorter. This result agrees with an intriguing finding of Schmalzl et al. [28] for Rayleigh-Bénard convection. In Figure 8 of that work, the surface area of an arbitrary temperature iso-surface is shown to become smaller with increasing Prandtl number. Since we find that a generally shorter diffusion distance is required to reach a certain temperature difference, our results agree with the finding that the surface area of a temperature iso-surface should be typically smaller in a simulation with larger Prandtl number.

![Figure 10. Cumulative distribution function of the Lagrangian vertical heat structure $\lambda_{hz}$ for (a) a temperature separation of $\Delta T = 0.1$ in simulations HC1 ($Pr = 0.1$), HC2 ($Pr = 1.0$), and HC3 ($Pr = 10$), and (b) for a range of temperature separations in simulation HC1. A dotted line marks the point where $C(\lambda_{hz}) = 0.5$. Distributions are constructed from approximately 4 million Lagrangian tracer particles, sampled for 6 different statistically independent flow realizations.](image1)

The mean value of the Lagrangian vertical heat structure $\lambda_{hz}$ is shown in panel (a) of Figure 11. When a difference in temperature of $\Delta T < 0.1$ is considered, $\lambda_{hz}$ is on average negative for HC3 ($Pr = 10$); this indicates that this separation in temperature is reached before a particle moves as far as the Kolmogorov microscale. The temperature difference that requires a particle must diffuse further than a Kolmogorov microscale is smaller for simulation HC2 ($Pr = 1.0$) than HC3 ($Pr = 10$), and smaller still for simulation HC1 ($Pr = 0.1$). This is in agreement with general ideas about flows at different Prandtl number.

![Figure 11a.](image2)

As larger differences in temperature are considered, the mean value of $\lambda_{hz}$ appears to approach an asymptotic value that is similar between these simulations. The ordering of the means for different Prandtl numbers agrees with the result of Schmalzl et al. [28] for each $\Delta T$. At $\Delta T = 2$, the ratio of the mean $\lambda_{hz}$ between HC1 and HC3 is 1.86; between HC2 and HC3 it is 1.27. For these three simulations, for moderately small $\Delta T$, the mean Lagrangian vertical heat structure gains a progressively sharper and higher peak as the Prandtl number is decreased. In addition, a comparison between simulations HC2 and HC4 reveals that as the Rayleigh number is increased for fixed Prandtl number, this peak becomes sharper and higher. A comparison between simulations HC1 and HC4 shows that for fixed Grashof number and increasing Prandtl number, the peak becomes less sharp and lower. The results of Figure 11a imply that dynamo simulations that are produced using $Pr = 1.0$ may represent a transfer of heat that is characteristically similar to the lower Prandtl number settings that they represent, and more accurate for larger heat changes. The standard deviation of $\lambda_{hz}$ is shown in Figure 11b. Surprisingly, unlike the mean, the standard deviation of $\lambda_{hz}$ is nearly identical for all
simulations, despite the different Prandtl numbers. The data indicate that this standard deviation decreases approximately as the inverse of the temperature separation.

The skewness is defined

\[
\text{skewness}(x) = \frac{E[(x - m)^3]}{E[(x - m)^2]^{3/2}}.
\]  

Here \(E\) indicates the expected, or average, value of the quantity in brackets, and \(m\) is the mean of \(x\). For a perfectly symmetric distribution the skewness is zero. The form of the kurtosis that we calculate is

\[
\text{kurtosis}(x) = \frac{E[(x - m)^4]}{E[(x - m)^2]^2} - 3.
\]  

This is sometimes also called the excess kurtosis \([52]\), or flatness, and is zero for all univariate Gaussian distributions. The skewness provides a measure of the asymmetry of a distribution, while the kurtosis measures the heaviness of its tails. The skewness and kurtosis of \(\lambda_{\text{hsz}}\) as a function of the temperature separation \(\Delta T\) are shown in Figure 12. The skewness is negative for all temperature separations we calculate, indicating that the left side of the distribution function of \(\lambda_{\text{hsz}}\) is always fatter. This means small values of \(\lambda_{\text{hsz}}\) are more likely than large values, for all \(\Delta T\) obtained in our simulations. For all simulations the skewness shows a characteristic shape with a peak for small values of \(\Delta T\), followed by a long decay for large values of \(\Delta T\). For simulation HC1 (\(Pr = 0.1\)) this peak is the sharpest, followed by simulation HC2 (\(Pr = 1.0\)), and then simulation HC3 (\(Pr = 10\)). The peak for simulation HC4 (\(Pr = 1.0\)) is similar to simulation HC2 (\(Pr = 1.0\)). As larger \(\Delta T\) are considered, the skewness becomes more negative, indicating that the distribution of \(\lambda_{\text{hsz}}\) becomes more asymmetric, and small values of \(\lambda_{\text{hsz}}\) become more likely. All three simulations obtain similar values of the skewness for larger temperature separations \(\Delta T\).

For all simulations the kurtosis of \(\lambda_{\text{hsz}}\) drops sharply for small temperature separations \(\Delta T\) (see Figure 12b). It then grows slowly with increasing \(\Delta T\). The initial drop is sharpest for HC1 (\(Pr = 0.1\)), followed by simulation HC2 (\(Pr = 1.0\)), and then HC3 (\(Pr = 10\)), showing again a clear trend in Prandtl number at fixed Rayleigh number. The drop for simulation HC4 (\(Pr = 1.0\)) is slightly more than simulation HC2 (\(Pr = 1.0\)), and slightly less than simulation HC1 (\(Pr = 0.1\)), indicating that
the Rayleigh number also plays a role. The higher the kurtosis, the higher the tails of the distribution of $\lambda_{h_{sz}}$ relative to the tails of a Gaussian distribution. For moderate and large $\Delta T$, the tails of the distributions become more super-Gaussian. However for larger temperature separations $\Delta T$, any trend in Prandtl number is lost.

Figure 12 shows that the non-Gaussian aspects of the Lagrangian vertical heat transfer are most dependent on the Prandtl number for small temperature separations $\Delta T$. The characteristic shapes produced for the skewness and kurtosis of $\lambda_{h_{sz}}$ are universal for all simulations, suggesting that aspects of heat transfer in a low Prandtl number simulation could possibly be captured at moderately higher Prandtl number. The non-zero values of the skewness and kurtosis indicate that any purely Gaussian theory is unlikely to capture important aspects of heat transfer. This supports the similar finding of Emran and Schumacher [53] that PDFs of temperature and its fluctuations are strongly non-Gaussian. Unlike Emran and Schumacher [53], there is no height dependence to the skewness and kurtosis because we examine a homogeneous system.

5. Discussion and Conclusions

After studying the intermittency of the Nusselt number in our simulations of homogeneous turbulent convection, we develop an exit-distance statistic, the Lagrangian heat structure, to quantify properties of heat transfer in a way that is less susceptible to noise and intermittency effects. The Lagrangian heat structure is inspired by the finite-time Lyapunov exponent and the mathematical framework surrounding Lagrangian coherent structures. Local spatial deviations as well as variance in time are easily extracted from the Lagrangian heat structure. This ability to accurately classify extreme variations in a convecting flow is broadly desirable for the study of natural systems. Our new statistic therefore provides a useful supplement to the information provided by the Nusselt number.

We use the Lagrangian vertical heat structure to compare several direct numerical simulations of homogeneous turbulent convection at Prandtl numbers 0.1, 1.0, and 10. Different regimes of heat transfer, based on different temperature separations $\Delta T$, are revealed by the moments of the distribution of the Lagrangian vertical heat structure. The behavior of the mean, standard deviation, skewness, and kurtosis are characteristically similar for the range of Prandtl numbers studied here. This suggests that a simulation with $Pr = 1.0$ may be able to reasonably capture qualities of the heat transfer at somewhat larger or smaller Prandtl numbers. This is particularly true if large temperature separations are of primary importance.
This is an encouraging insight for simulations performed with the goal of understanding turbulent convection in natural settings such as Earth’s atmosphere and oceans, as well as planetary and stellar dynamos. Because many numerical studies of the dynamo have been performed at $Pr = 1$, this observation lends support to the practice of applying these simulation results to realistic natural settings such as a planetary or stellar interior. Our results also indicate that a convecting fluid at room temperature has similarities in the shape of the Lagrangian heat structure to convection in the arctic oceans. If such simulations are performed as large eddy simulations where the smallest scales are modeled rather than resolved, the differences that we note between different Prandtl number simulations at the smallest scales of temperature separation may have even less impact on the simulation results.

This work expands on the findings of several earlier studies of Rayleigh-Bénard convection. Building on the pioneering work of Schumacher [10] to describe convective turbulence from the Lagrangian point of view, we have shown that Lagrangian tracer particles can be used for an exploration of heat transport. Our perspective on heat transport adds to the larger work of Emran and Schumacher [53] that examines the Rayleigh number dependence of higher-order Eulerian statistics of the thermal gradient during turbulent convection. Our development of Lagrangian statistics produces a detailed confirmation of the Eulerian results of Schmalzl et al. [28], particularly the dependence of the iso-surfaces of temperature on the Prandtl number.

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