

Direct numerical simulations of turbulent Rayleigh-Bénard convection with polymer additives

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(Received 13 January 2025; revised 28 April 2025; accepted 14 May 2025)

We present results of three-dimensional direct numerical simulations of turbulent Rayleigh-Bénard convection of dilute polymeric solutions for Rayleigh number (Ra) ranging from 10^6 to 10^{10} , and Prandtl number Pr = 4.3. The viscoelastic flow is simulated by solving the incompressible Navier-Stokes equations under the Boussinesq approximation coupled with the finitely extensible nonlinear elastic Peterlin constitutive model. The Weissenberg number (Wi) is either Wi = 5 or Wi = 10, with the maximum chain extensibility parameter L = 50, corresponding to moderate fluid elasticity. Our results demonstrate that both heat transport and momentum transport are reduced by the presence of polymer additives in the studied parameter range. Remarkably, the specific parameters used in the current numerical study give similar heat transfer reduction values as observed in experiments. We demonstrate that polymers have different effects in different regions of the flow. The presence of polymers stabilises the boundary layer, which is found to be the primary cause of the overall heat transfer reduction. In the bulk region, the presence of polymers slows down the flow by increasing the effective viscosity, enhances the coherency of thermal plumes, and suppresses the small-scale turbulent fluctuations. For small Ra, the heat transfer reduction in the bulk region is associated with plume velocity reduction, while for larger Ra, it is caused by the competing effects of suppressed turbulent fluctuations and enhanced plume coherency.

Key words: Bénard convection, turbulence simulation, viscoelasticity

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1. Introduction

It is well-known that adding a small amount of polymer additives in turbulent flows can lead to a significant drag reduction (Toms 1948). Extensive studies have been performed to address this phenomenon in wall-bounded flows through theoretical, experimental and numerical approaches (Procaccia, L'vov & Benzi 2008; White & Mungal 2008). However, the effects of polymer additives on heat transport in turbulent thermal convection remain less understood despite many processes involving heat transfer in nature, e.g. in oils, melts and molten magma, as well as those prevalent in industrial processes such as polymer solutions and emulsions, concerning non-Newtonian flows (Denn 2004). These flows exhibit distinctive rheological properties, like viscoelasticity, plasticity, shear thinning and shear thickening. Among them, the viscoelastic behaviour of non-Newtonian fluids determines surprising and complex rheological properties and flow dynamics, which can lead to different unexpected fluid-flow phenomena, such as elasto-inertial turbulence (Samanta et al. 2013; Li, Sureshkumar & Khomami 2015; Sid, Terrapon & Dubief 2018; Song et al. 2021; Dubief, Terrapon & Hof 2023) and elastic turbulence (Groisman & Steinberg 2000; Steinberg 2021; Datta et al. 2022; Song et al. 2022, 2023a,b). Therefore, studying the thermal convection in viscoelastic fluids can deepen our understanding of the interplay between the fluid viscoelasticity and convective processes. Such insights are crucial for various applications, ranging from geophysical and astrophysical phenomena such as mantle convection, to industrial processes such as polymer processing and thermal management systems.

As a canonical system to study thermal convection, Rayleigh–Bénard (RB) convection of Newtonian fluids has been studied extensively over the past decades (Bodenschatz, Pesch & Ahlers 2000; Ahlers, Grossmann & Lohse 2009; Lohse & Xia 2010; Chillà & Schumacher 2012; Xia 2013; Shishkina 2021; Lohse & Shishkina 2024). In RB convection, the flow is driven by buoyancy forces due to the temperature difference Δ between a hot bottom plate and a cold top plate. The main dimensionless control parameters in RB convection are the Rayleigh number *Ra*, Prandtl number *Pr*, and aspect ratio of the container Γ , defined as

$$Ra = \frac{g\alpha\Delta H^3}{\kappa\nu}, \quad Pr = \frac{\nu}{\kappa}, \quad \Gamma = \frac{D}{H}.$$
 (1.1)

Here, g denotes the gravitational acceleration, α the thermal expansion coefficient, v the kinematic viscosity, and κ the thermal diffusivity, and H and D are the height and width of the container, respectively. Studies of this system have focused mainly on: (i) the flow structures, identifying kinetic and thermal boundary layers (BLs), and an approximately homogeneous bulk flow (Grossmann & Lohse 2000; Ahlers *et al.* 2009); and (ii) the dependence of the global heat transport, quantified by the Nusselt number Nu, and momentum transport, characterised by the Reynolds number Re, on the different control parameters mentioned above. The Nusselt number is defined as the ratio of the total heat flux in the flow to the molecular heat conduction, which for an incompressible flow is

$$Nu = \frac{\langle u_z \theta \rangle_{A,t} - \kappa \,\partial_z \,\langle \theta \rangle_{A,t}}{\kappa \,\Delta/H}.$$
(1.2)

Here, u_z is the vertical velocity, and θ is the temperature, while $\langle \cdots \rangle_{A,t}$ denotes the horizontal-plane and time average at any distance between the heated and cooled plates.

As regards complex fluids, some studies also considered RB convection of viscoelastic fluids, focusing on the effects of fluid elasticity on two critical Rayleigh numbers: Ra_{c1} for the transition from conduction to convective flow, and Ra_{c2} for the onset of chaotic flows

	Ex	perime	ents					
	Ra	Pr	c (ppm)	Effect				
Ahlers & Nikolaenko (2010)	$5 \times 10^9 \sim 10^{11}$	4.38	$0 \sim 120$		HTR HTR in smooth cell, HTE in rough cell			
Wei <i>et al.</i> (2012)	$\sim 10^9$	4.34	0~180					
Xie <i>et al.</i> (2015)	6.18×10^9	4.3	0~180		HTE for large <i>c</i> in rough cell, HTR for small <i>c</i> in rough cell			
Cai et al. (2019)	$\sim 10^9$	5.43	0~400		HTR			
Xu et al. (2025)	$5 \times 10^8 \sim 7 \times 10^9$	4.34	0~20		HTR			
	Si	mulatio	ons					
	Ra	Pr	β	Wi	Effect			
Benzi et al. (2010)	$10^{11} \sim 10^{13}$	1		[0,100]	HTE for $Wi \leq 1$,			
(homogeneous)					HTR for $Wi > 1$			
Benzi et al. (2016a)	2.07×10^{5}	7	0.9	[0,50]	HTE			
Cheng et al. (2017)	108	0.7	0.9	[0,10]	HTR			
Dubief & Terrapon (2020)	10 ⁵	7	0.9	[0,45]	HTE for small L, HTR for larger L			
Wang <i>et al.</i> (2023)	$0 \sim 10^5$	7	0.9	[0,50]	HTR, HTE for specific cases			

Table 1. Parameters in previous studies of turbulent RB convection with polymer additives. The fourth column gives the polymer concentration c in the experiments or the viscosity ratio β in the simulations. The Weissenberg number Wi is given only for simulations (except for Benzi *et al.* (2010), which defines Wi using the r.m.s. velocity and a large length scale of the flow in the absence of polymers, other data are adapted to the same definition using the free-fall velocity and domain height), since to measure it in experiments is challenging. In the last column, HTE means heat transfer enhancement, HTR means heat transfer reduction, and L is the extensibility parameter. Here, we provide only a list of literature specifically discussing the modification of the heat transfer in turbulent RB convection with polymers, while omitting numerous studies focused on pattern formation, onset of convection and other issues.

from stationary conditions. Moreover, pattern formations at small and moderate *Ra* have been studied through linear (Green 1968; Vest & Arpaci 1969; Sokolov & Tanner 1972) or nonlinear stability analysis (Park & Ryu 2001), or via numerical simulations (Wang *et al.* 2023; Zheng *et al.* 2023).

The effect of polymer additives on turbulent RB convection has received attention only in recent years. Table 1 lists the studies of turbulent RB convection with polymer additives, and the corresponding range of parameters examined. Here, the Weissenberg number Wi, which is commonly used in viscoelastic flow to quantify the elastic effects of polymer solutions, is defined as

$$Wi = \frac{\lambda U}{H},\tag{1.3}$$

where λ is the relaxation time scale of the polymers, and U is the relevant flow velocity. In terms of experimental studies, Ahlers & Nikolaenko (2010) did the first experiments at Ra spanning from 5×10^9 to 10^{11} to investigate the turbulent heat transport modifications by polymers. They found heat transport reduction (HTR), and that HTR amount increases with polymer concentration c up to 10%. Later, RB convection with polyethylene oxide (PEO) solutions was studied for $Ra \sim 10^9$ and $Pr \approx 4.34$ for both smooth and rough plates (Wei, Ni & Xia 2012). In the smooth cell, HTR was observed, whereas heat transport enhancement (HTE) was found in the rough cell. Moreover, a maximum HTR saturation state was observed as polymer concentration *c* increases. Based on these observations, Xie *et al.* (2015) further analysed the HTE mechanism within the rough cell. Their analysis revealed that the polymer additives can enhance coherent heat fluxes while suppressing incoherent heat fluxes within the bulk region, leading to the HTE. In the BL region, the plume emission rate is strongly suppressed. Thus whether HTE or HTR takes place depends on the competition between these two effects. Cai *et al.* (2019) found HTR in their experiments at similar $Ra \sim 10^9$, and concluded that the reduction is related to the smaller size of large-scale circulation (LSC) in polymer solutions. In a recent study, Xu *et al.* (2025) also found an HTR in a cylindrical cell, and revealed that the addition of polymers re-establishes the cylindrical symmetry of the large-scale structures, leading to a larger flow coherence.

As mentioned above, the HTR saturate state was observed in high-Ra experiments (Wei et al. 2012: Cai et al. 2019). This behaviour is reminiscent of the maximum drag reduction (MDR) asymptote caused by polymer additives in other wall-bounded turbulent flows, such as channel, pipe and plane Couette flows (Toms 1948; Lumley 1969; Virk 1975). The cornerstone work by Samanta *et al.* (2013) proposed that the dynamics of MDR is driven by an elasto-inertial instability that can even eliminate the Newtonian turbulence, and the MDR asymptote flow state can be interpreted as a self-sustained elasto-inertial turbulence, where the polymer stretching acts to suppress the small-scale vortices and turbulent fluctuations. This interpretation has gained traction, while it is not yet conclusive that the MDR asymptote is solely governed by elasto-inertial turbulence (Dubief et al. 2023). In addition, the kinetic energy spectrum in the elasto-inertial turbulence has a scaling exponent approximately -14/3, which is distinctly different from the Kolmogorov scaling -5/3 for inertial turbulence, but close to the scaling for elastic turbulence -3.5; see Groisman & Steinberg (2000), Fouxon & Lebedev (2003), Dubief, Terrapon & Soria (2013) and Steinberg (2019). However, this spectral universality and associated turbulent dynamics of elasto-inertial turbulence have not been examined in natural thermal convection systems so far.

Regarding the numerical studies, polymer chains in turbulent flows are generally modelled as dumbbells composed of two beads connected by the elastic Hookean linear spring Oldrold-B model with an infinite extensibility or a nonlinear elastic entropic spring with finite extensibility, which is known as the finitely extensible nonlinear elastic-Peterlin (FENE-P) model (Bird *et al.* 1987). The FENE-P model can overcome several limitations of the Oldroyd-B model, including: (i) the possibility of singular extensional viscosity or stress in extensional flows, due to the absence of an upper bound on the polymer extensibility; (ii) the inability to reproduce the shear-thinning behaviour commonly observed in most polymer solutions (Dubief *et al.* 2023).

In the pioneering work of Benzi, Ching & De Angelis (2010), homogeneous turbulent thermal convection with polymer additives was studied using direct numerical simulations (DNS) based on the FENE-P model, and then a simplified shell model was used to understand the phenomenon. The DNS were performed under periodic boundary conditions driven by a constant temperature gradient, which mimics the bulk region of RB convection, and the HTE were observed for Wi < 1. With shell model caculations, a transition from HTE to HTR at Wi > 1 was obtained with increasing Wi. These authors speculated that the addition of polymers would increase the drag within the BL, leading to a decrease in Nu. The overall modulation of the heat transfer would then depend on the balance between the contribution from the BLs and bulk region. Moreover, they highlighted three intriguing and promising approaches to observe HTE: (i) increasing Ra, where heat transfer becomes dominated by the bulk region; (ii) artificially reducing the energy dissipation contribution from the BL, which is the approach explored in

the experimental work of Xie *et al.* (2015) and already validated by their studies; and (iii) utilising a slender domain to shift the the main transport towards the bulk region.

Later, Benzi, Ching & De Angelis (2016a) performed simulations over a range of Wi from 0 to 50, at $Ra = 2.07 \times 10^5$, Pr = 7, by using the Oldroyd-B model. They observed HTE, and that the HTE amount first increases and then decreases with increasing W_i , which was attributed to the combined effects of decreasing viscous energy dissipation and increasing energy dissipation due to the polymers. Cheng et al. (2017) explored a range of Wi from 0 to 10 within a two-dimensional enclosed RB cell with Oldroyd-B polymers at $Ra = 10^8$ and Pr = 0.7. These authors observed that the HTR behaves nonmonotonically with Wi (first increasing and then decreasing). They concluded that this non-monotonic behaviour results primarily from the contribution of elastic stresses to the turbulent kinetic energy, which absorbs energy from the turbulence for small W_i , while releasing stored energy to small-scale motions for relatively large Wi. Dubief & Terrapon (2020) investigated the effects of the FENE-P polymer chain extensibility parameter L. In their simulations, Wi ranges from 0 to 45, and L spans from 10 to 100 at $Ra = 10^5$, Pr = 7. They reported that polymers with small L lead to HTE, whereas polymers with larger $L \ (L \gtrsim 50)$ result in HTR. The mechanism underlying HTE involves an increase in the number of convection cells, while in the case of HTR, the LSC slows down simultaneously. Recently, Wang *et al.* (2023) found similar HTR for additional configurations at lower Ra, except for some particular parameters where multiple pairs of LSC exist, leading to HTE.

In all these studies, based on simulations and experiments, researchers speculated that the effects of polymers on the BL and bulk regions differ significantly. Whether the total heat transfer is enhanced or reduced depends mainly on the contributions of these two regions. To elucidate the effects of polymers on a laminar BL flow, Benzi, Ching & Chu (2012) generalised the classical Prandtl-Blasius-Pohlhausen theory to Oldroyd-B viscoelatic flows at moderate Ra. Their analysis demonstrated that the stretching of polymers induces a space-dependent effective viscosity, increasing near the plate, and vanishing away from it. Such an increase leads to an enhancement of friction drag, and thus results in the reduction of the heat transport in the BL region. Moreover, the effective viscosity increases with W_i and viscosity ratio γ , therefore the amount of the HTR and drag enhancement increases as well. Here, γ is the ratio of polymer viscosity to zeroshear-rate viscosity of polymer solutions ($\gamma = 1 - \beta$, where β is used in our simulation, as defined below.) Later, Benzi et al. (2016b) extended their theoretical framework to FENE-P modelled polymer solutions. They suggested that HTR occurs for polymers with large extensibility parameter L, while HTE should be observed for small L. Moreover, they concluded that heat transport is enhanced when the region of substantial polymer stretching moves outside the thermal BL. These conclusions have been confirmed by Dubief & Terrapon (2020) at low Rayleigh number $Ra = 10^5$, where they observed that in cases of HTE, the first normal stress difference is predominantly concentrated inside the plumes within the bulk region, while in cases of HTR, the magnitudes of the first normal stress difference inside the plumes and BL are similar. However, we have not found any evidence of this phenomenon at high *Ra* in the literature.

As summarised in table 1, previous experimental investigations have focused primarily on high-Ra (10⁹~10¹⁰) situations, whereas simulations in the existing literature have been limited predominantly to lower Ra values, typically smaller than 10⁶, due to the difficulties associated with the numerical simulations of viscoelastic flows (Alves, Oliveira & Pinho 2021). This leads to a gap in Ra values of several orders between experimental and numerical studies. Moreover, previous numerical studies on turbulent RB convection with polymer additives focused mainly on the dependence on Wi, viscosity ratio β , or extensibility parameter L, while the specific effects of Ra on the heat and momentum



Figure 1. Sketch of viscoelastic RB convection in a parallelepiped geometry.

transfer modifications, flow structures and turbulent velocity, temperature and elastic stress statistics remain unclear. To fill this gap, we have therefore performed three-dimensional DNS over a broad range of Ra spanning from 10⁶ to 10¹⁰ at Pr = 4.3 while keeping moderate Wi, namely, Wi = 5 and 10. This work seeks to provide understanding of the role played by the Rayleigh number in turbulent RB convection of viscoelastic fluids, with particular interest in the heat-transport modifications, changes of thermal plumes and BLs, and kinetic and thermal energy dissipation rates.

The paper is organised as follows. The physical problem, governing equations, numerical methods and computational details are described in § 2. The flow structures and statistic profiles are shown in § 3. The polymer effects on heat flux, dissipation rate and energy spectrum are discussed separately in § 4. Finally, the main conclusions are summarised in § 5.

2. Physical problem and computational details

2.1. Governing equations

A sketch of RB convection with polymer additives in a parallelepiped geometry with $\Gamma = 1$, i.e. a cube, is shown in figure 1. The FENE-P viscoelastic constitutive model is used to model the contribution of polymer additives to the total stress. The orientation and extension of polymer molecules are described by the end-to-end vectors q_i , and represented in the statistical model by the phase-averaged configuration tensor denoted by $C_{ij} = \langle q_i q_j \rangle$. The FENE-P model is frequently used in DNS of viscoelastic flows due to its physical connection with real viscoelastic liquids, specifically dilute solutions of high molecular weight, finitely extensible flexible polymers in a theta solvent. In this model, interactions between polymer chains are not considered. Although it is not easy to compare the relaxation time in the FENE-P model against real experiments, it can qualitatively capture the relevant experimental results (White & Mungal 2008; Larson & Desai 2015; Xi 2019; Ching 2024; Serafini et al. 2024). The FENE-P model is closed by three control parameters: (i) the maximum polymer chain extensibility parameter L; (ii) the Weissenberg number Wi, here defined by $Wi \equiv \lambda U_{ff}/H$, where $U_{ff} \equiv \sqrt{g\alpha\Delta H}$ is the characteristic free-fall velocity; and (iii) the viscosity ratio $\beta = v_s/(v_s + v_p)$, which is the ratio of the solvent viscosity to the total zero-shear rate viscosity of the polymer solution (subscripts s and p represent solvent and polymer, respectively). The maximum polymer chain extensibility remains constant throughout all simulations. Thus polymer chain scission is not captured in our simulations. We choose a commonly used value L = 50, which implies moderate extensibility. Viscosity ratio can be regarded as a measure of polymer concentration by assuming that the viscosity depends only on the polymer concentration. In our DNS, β is fixed at 0.9 for viscoelastic fluids ($\beta = 1$ for Newtonian), which denotes dilute polymer solutions.

The chosen reference scales for the non-dimensionalisation are the height of the domain H, the temperature difference between the plates Δ , the characteristic free-fall velocity U_{ff} , pressure ρU_{ff}^2 , and stress $\mu_p U_{ff}/H$. Non-dimensional temperature θ , velocity u_i (where i = x, y, z represent the three coordinate directions), pressure p, polymer stress τ_{ij} and time t are obtained by using these scales. The dimensionless governing equations for the incompressible FENE-P fluid are as follows:

$$\frac{\partial u_i}{\partial x_i} = 0, \tag{2.1}$$

$$\frac{\partial u_i}{\partial t} + u_j \frac{\partial u_i}{\partial x_j} = -\frac{\partial P}{\partial x_i} + \beta \sqrt{\frac{Pr}{Ra}} \frac{\partial^2 u_i}{\partial x_j \partial x_j} + (1 - \beta) \sqrt{\frac{Pr}{Ra}} \frac{\partial \tau_{ij}}{\partial x_j} + \theta \delta_{iz}, \qquad (2.2)$$

$$\frac{\partial C_{ij}}{\partial t} + u_k \frac{\partial C_{ij}}{\partial x_k} = C_{ik} \frac{\partial u_j}{\partial x_k} + C_{kj} \frac{\partial u_i}{\partial x_k} - \tau_{ij}, \qquad (2.3)$$

$$\frac{\partial\theta}{\partial t} + u_j \frac{\partial\theta}{\partial x_j} = \frac{1}{\sqrt{Ra Pr}} \frac{\partial^2\theta}{\partial x_j \partial x_j},$$
(2.4)

where the polymer stress τ_{ij} is related to the conformation tensor C_{ij} via

$$\tau_{ij} = \frac{f(C_{kk}) C_{ij} - \delta_{ij}}{Wi}.$$
(2.5)

The function $f(C_{kk})$, known as the Peterlin function, is defined as

$$f(C_{kk}) = \frac{L^2 - 3}{L^2 - C_{kk}},$$
(2.6)

and δ_{ij} in (2.2) and (2.5) denotes the Kronecker symbol.

In the above equations, the trace of the conformation tensor C_{kk} represents the square of the average polymer-chain extension. It should be noted that for dilute polymer concentrations, the effect of polymer additives on the thermodynamic properties can be ignored, except for the fluid viscosity (Rubinstein & Colby 2003). No-slip boundary and constant temperature conditions are applied at the bottom and top plates, whereas periodic boundary conditions are applied in the wall-parallel horizontal directions.

2.2. Numerical method and validations

The equations introduced above are solved via an in-house viscoelastic solver based on the open source AFiD code (Verzicco & Orlandi 1996; van der Poel *et al.* 2015*b*). The AFiD code is an energy-conserving second-order finite difference code that has been proven to be a versatile Navier–Stokes solver for wall-bounded turbulent flows (Verzicco & Orlandi 1996). Moreover, the code is parallelised using a two-dimensional pencil domain decomposition strategy, allowing it to effectively handle large-scale computations (van der Poel *et al.* 2015*b*). The code uses a staggered grid for the spatial discretization, with velocities located on the cell faces, and the other variables (pressure, conformation tensor and elastic stress) at the cell centre, except temperature, which is collocated with the vertical velocity u_z . In the new version of the code, all spatial derivatives are



Figure 2. Comparison of statistics profiles from our simulations with the results of Dubief & Terrapon (2020) for the Newtonian flow and non-Newtonian flows with an HTE flow state at Wi = 10, L = 25 and an HTR flow state at Wi = 40, L = 100. The other control parameters are fixed at $Ra = 10^5$, Pr = 7, $\beta = 0.9$ and $\Gamma = 16$. Root mean square (r.m.s.) of (a) horizontal velocity $u'_{h,rms}$, (b) vertical velocity $u'_{z,rms}$, (c) temperature fluctuations θ'_{rms} . (d) Mean heat flux $\langle u_z \theta \rangle$ across the half-gap.

approximated by using the second-order central finite differences except for the advection term in (2.3), where a Kurganov-Tadmor scheme (Vaithianathan et al. 2006; Lin et al. 2022) is used; indeed, the hyperbolic nature of this equation requires a special treatment of the convective term to ensure numerical convergence, especially at high Wi (Min, Yoo & Choi 2001). Specifically, the Kurganov–Tadmor scheme is a modified central difference scheme that ensures the symmetric-positive-definite property of C_{ii} based on flux limiters. The tensor C_{ij} at cell interfaces is constructed from the second-order, piecewise, linear approximations based on three potential candidates for approximating the gradients at cell centre: first-order forward and backward schemes, and a second-order central scheme (Vaithianathan et al. 2006). In addition, a fractional-step third-order Runge-Kutta (RK3) scheme combined with a Crank-Nicolson scheme for the implicit terms is used for all the time advancements. Specifically, in (2.3), the linear stress relaxation term is treated implicitly to strictly enforce the chain finite maximum extension limit (Vaithianathan & Collins 2003; Dubief et al. 2005; Song et al. 2021). To this end, this algorithm is numerically stable and preserves the positive definiteness as well as the boundedness of the polymer conformation tensor ($0 < C_{ii} < L^2$).

To validate the code, we compare our DNS results with the three-dimensional numerical simulations of Dubief & Terrapon (2020) at $Ra = 10^5$, Pr = 7, $\beta = 0.9$ and $\Gamma = 16$ using the same grid resolutions. The results of the statistical profiles for velocities, temperature and heat flux are shown in figure 2, which are in good agreement with their results. Additionally, we compared the heat transfer modification (HTM), defined

as $HTM = (Nu^V - Nu^N)/Nu^N \times 100$ %. In our simulations, the HTM is 14.4 % for the HTE flow state, and -31.4 % for the HTR flow state, closely matching their results 11 % and -31 %, respectively. The small quantitative differences may be attributed to the different numerical schemes, specifically the inclusion of an artificial diffusion term in their code. Moreover, the present DNS results of the heat transport modification for highly turbulent thermal convection with polymer additives agree quantitatively with previous experiments in a similar parameter range. Specifically, our DNS at $10^9 \le Ra \le 10^{10}$, Pr = 4.3 give an HTR approximately 9 %~15 %, which is quite close to the 10 %~12 % measured in the experiments by Ahlers & Nikolaenko (2010), Wei *et al.* (2012) and Cai *et al.* (2019) for similar parameters $10^9 \le Ra \le 7 \times 10^{10}$, Pr = 4.3. For further validation cases and details of the code, we refer to Song, Xu & Shishkina (2025).

2.3. Simulation parameters

In our simulations, we explore the range of Ra spanning from 10^6 to 10^{10} at fixed Pr = 4.3, corresponding to pure water at 40 °C. We consider a moderate level of elasticity: Wi = 5 and Wi = 10, with Wi = 0 representing Newtonian flow.

In classical Newtonian RB convection, the mesh resolution should be adequate to resolve the Kolmogorov $\eta_K^* \equiv (\nu^3/\varepsilon_u^*)^{1/4}$ and Batchelor $\eta_B^* \equiv (\nu\kappa^2/\varepsilon_u^*)^{1/4}$ length scales, where ε_u^* represents the kinetic energy dissipation rate, with * indicating dimensional quantities. The dimensionless forms of these two length scales are calculated by

$$\eta_K(\boldsymbol{x},t) = \left(\frac{Pr}{Ra}\right)^{3/8} \varepsilon_u(\boldsymbol{x},t)^{-1/4}, \quad \eta_B(\boldsymbol{x},t) = \frac{\eta_K}{\sqrt{Pr}}, \quad (2.7)$$

where the dimensionless kinetic energy dissipation rate is given by $\varepsilon_u(\mathbf{x}, t) = \sqrt{Pr/Ra} [\partial_i u_j(\mathbf{x}, t)]^2$ (Scheel, Emran & Schumacher 2013). We ensure adequate grid resolution in all directions by requiring that the maximal value of the ratio of the mesh size to the local Kolmogorov h_{max}/η_K and Batchelor h_{max}/η_B microscales is smaller than 3, which was found empirically to be acceptable (Verzicco 2003; Shishkina *et al.* 2010), where $h_{max} = \max(\Delta x, \Delta y, \Delta z)$. For all the cases, the mesh width fulfils these requirements (see table 2, h_{max}/η_K and h_{max}/η_B). Notably, according to Stevens, Verzicco & Lohse (2010), higher resolution in the horizontal direction is necessary to resolve the plume dynamics in the BL. Further, we check the number of mesh points inside the thermal and viscous BLs (see table 2, N_{θ} and N_v) to ensure adequate resolutions inside the BL. The resolutions in our DNS fully satisfy the requirements for minimal mesh points suggested by Shishkina *et al.* (2010).

In a viscoelastic flow, the typical length is similar or even larger than those in a Newtonian fluid (Benzi *et al.* 2010; Wei *et al.* 2012; Cheng *et al.* 2017), therefore, for viscoelastic flows we use the same mesh resolutions as in the corresponding Newtonian flows. In table 2, we similarly provide h_{max}/η_K , h_{max}/η_B and the mesh points inside the BL with the minimal mesh points according to Shishkina *et al.* (2010). The Kolmogorov and Batchelor length scales are defined using the Newtonian solvent dissipation rate $\varepsilon_u^{[s]}$, excluding polymer dissipation $\varepsilon_u^{[p]}$, as $\eta_K^* = (v_s^3/\varepsilon_u^{[s]^*})^{1/4}$ and $\eta_B^* = (v_s \kappa^2/\varepsilon_u^{[s]^*})^{1/4}$. Their dimensionless forms are

$$\eta_K(\mathbf{x},t) = \left(\frac{Pr}{Ra}\right)^{3/8} \beta^{3/4} \varepsilon_u^{[s]}(\mathbf{x},t)^{-1/4}, \quad \eta_B(\mathbf{x},t) = \frac{\eta_K(\mathbf{x},t)}{\sqrt{\beta Pr}},$$
(2.8)

Ra	$N_x \times N_y \times N_z$	Nu	Nu_h	Max-diff	Re	h_{max}/η_K	h_{max}/η_B	$N_{ heta}$	N_v	t_{avg}			
Newtonian flow $Wi = 0$													
1×10^{6}	$192 \times 192 \times 128$	10.96	10.96	0.13 %	62.30	0.25	0.51	11/2.2	14/3.6	300			
3×10^{6}	$192 \times 192 \times 128$	13.83	13.77	0.24~%	103.84	0.35	0.73	9/2.5	12/4.1	300			
7×10^{6}	$256\times 256\times 192$	16.71	16.82	0.39 %	155.98	0.35	0.73	14/2.7	19/4.5	300			
1×10^{7}	$256\times 256\times 192$	17.93	18.08	0.46%	183.54	0.38	0.78	13/2.8	19/4.7	300			
3×10^{7}	$256\times 256\times 192$	23.64	23.73	0.44~%	310.95	0.54	1.13	11/3.2	17/5.3	300			
7×10^{7}	$384\times 384\times 256$	29.94	29.98	0.17 %	467.64	0.55	1.14	13/3.6	23/6.9	300			
1×10^{8}	$384\times 384\times 256$	32.75	32.69	0.19 %	538.69	0.63	1.30	12/3.8	23/6.3	300			
3×10^{8}	$384\times 384\times 256$	44.86	44.95	0.39 %	931.59	0.90	1.86	10/4.4	21/7.4	300			
7×10^{8}	$512 \times 512 \times 384$	57.17	57.17	0.17~%	1396.45	0.81	1.67	14/5.0	33/8.3	300			
1×10^{9}	$512 \times 512 \times 384$	63.44	63.63	0.46~%	1622.95	0.90	1.88	13/5.3	33/8.7	400			
3×10^{9}	$576 \times 576 \times 432$	88.98	89.08	0.19 %	2736.58	1.16	2.40	12/6.3	35/10.4	400			
1×10^{10}	$768\times768\times512$	129.21	128.43	0.28~%	4762.14	1.45	3.01	12/7.5	38/12.5	600			
Viscoelastic flow $Wi = 5$													
1×10^{6}	$192 \times 192 \times 128$	9.91	9.92	0.11~%	49.82	0.24	0.47	11/2.1	15/3.5	300			
3×10^{6}	$192\times192\times128$	12.92	12.93	0.17~%	86.28	0.35	0.68	10/2.4	13/ 3.9	300			
7×10^{6}	$256\times 256\times 192$	15.69	15.68	0.09~%	127.49	0.36	0.71	15/2.6	19/ 4.4	300			
1×10^7	$256\times 256\times 192$	16.82	16.84	0.34~%	143.95	0.40	0.78	14/2.7	19/4.5	300			
3×10^{7}	$256\times 256\times 192$	22.41	22.38	0.21~%	232.08	0.58	1.14	11/3.1	17/5.2	300			
7×10^{7}	$384\times 384\times 256$	27.85	27.80	0.17~%	340.24	0.52	1.13	14/3.5	24/ 5.8	300			
1×10^{8}	$384\times 384\times 256$	30.51	30.62	0.18~%	394.24	0.59	1.17	13/3.7	23/6.1	300			
3×10^{8}	$384\times 384\times 256$	41.28	41.24	0.25~%	647.78	0.85	1.67	10/4.3	21/ 7.1	300			
7×10^{8}	$512 \times 512 \times 384$	52.89	52.90	0.20%	975.66	0.89	1.76	15/4.8	33/8.0	300			
1×10^{9}	$512 \times 512 \times 384$	58.38	58.50	0.94~%	1148.39	1.00	1.96	14/5.1	32/8.4	400			
3×10^{9}	$576 \times 576 \times 432$	78.78	78.72	0.35 %	1877.12	1.29	2.53	13/5.9	32/9.7	400			
1×10^{10}	$768\times768\times512$	109.12	108.82	0.18~%	3293.28	1.45	2.85	13/6.9	35/11.5	500			
			Visc	coelastic fl	ow $Wi =$	10							
1×10^{6}	$192 \times 192 \times 128$	9.43	9.43	0.10 %	46.56	0.24	0.47	13/2.0	15/3.4	300			
3×10^{6}	$192 \times 192 \times 128$	11.97	11.99	0.20%	77.77	0.32	0.63	10/2.3	13/3.8	300			
7×10^{6}	$256\times 256\times 192$	14.61	14.60	0.26%	112.96	0.33	0.65	15/2.5	20/4.2	300			
1×10^{7}	$256\times 256\times 192$	16.41	16.49	0.11~%	137.09	0.39	0.76	14/2.7	20/4.4	300			
3×10^{7}	$256\times 256\times 192$	21.72	21.89	0.32%	222.47	0.57	1.11	11/3.1	18/5.1	300			
7×10^{7}	$384\times 384\times 256$	26.92	26.72	0.41 %	322.68	0.51	1.01	14/3.4	26/5.7	300			
1×10^8	$384\times 384\times 256$	29.77	29.86	0.28~%	380.02	0.58	1.15	13/3.6	25/6.0	300			
3×10^{8}	$384 \times 384 \times 256$	40.85	40.92	0.53 %	610.02	0.83	1.64	10/4.2	21/7.0	300			
7×10^8	$512 \times 512 \times 384$	52.42	52.49	0.26%	934.52	0.84	1.73	15/4.8	34/8.0	300			
1×10^9	$512 \times 512 \times 384$	58.23	58.42	0.88~%	1077.57	1.00	1.97	14/5.1	32/8.4	400			
3×10^9	$576 \times 576 \times 432$	78.72	78.37	0.39 %	1814.40	1.30	2.56	13/5.9	32/9.7	400			
1×10^{10}	$768\times768\times512$	108.75	109.92	0.16 %	3141.61	1.35	2.79	13/6.9	35/11.5	600			

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Table 2. Simulation parameters. Columns from left to right indicate Ra, grid resolution, spatiotemporal averaged Nusselt number (Nu) determined by averaging the results from four methods, Nusselt number Nu_h computed by using half of the statistical time (Stevens *et al.* 2010), maximum difference between four methods (Max-diff), Reynolds number Re, the maximum value of the ratio of the local mesh size $h_{max} = \max(\Delta x, \Delta y, \Delta z)$ to the local Kolmogorov η_K and Batchelor η_B length scales, the number of grid points inside the thermal BL, N_{θ} , and viscous BL, N_v (actual resolution/requirement; Shishkina *et al.* 2010), and the statistical averaging time t_{avg} in the free-fall time units.

where

$$\varepsilon_{u}^{[s]} = \beta \sqrt{\frac{Pr}{Ra}} \left[\partial_{i} u_{j}(\boldsymbol{x}, t) \right]^{2}, \quad \varepsilon_{u}^{[p]} = -(1 - \beta) \sqrt{\frac{Pr}{Ra}} u_{i} \partial_{j} \tau_{ij}.$$
(2.9)

To start the simulation of a Newtonian flow, the temperature field is initialised by a linear profile, and the velocity field is initialised with random fluctuation of maximum amplitude 0.001 to trigger the transition to the chaotic turbulent motion. In the viscoelastic cases, the temperature and velocity flow fields are initialised with those of the corresponding Newtonian flows in the statistically stationary state, while the initial conformation tensor is set to the unit tensor, corresponding to the equilibrium state. Statistical data are collected after the flow reaches a statistically stationary state, determined by the convergence of Nu and volume-averaged turbulent kinetic energy, which typically happens after at least 500 dimensionless time units for all cases. Table 2 provides the time span t_{avg} for the statistical data sampling in free-fall time units. In this paper, the statistic profiles presented in § 3.3 are further averaged over horizontal planes (*xy*-plane), with a prime indicating fluctuations around this average.

We use four different methods to calculate the Nusselt number Nu. One is the time and volume averaged Nu: $Nu = 1 + \sqrt{Ra Pr} \langle u_z \theta \rangle$. The second is derived from the exact relation for the thermal dissipation rate: $Nu = \sqrt{Ra Pr} \langle \varepsilon_{\theta} \rangle$ (Shraiman & Siggia 1990; Grossmann & Lohse 2000). The third and fourth values are obtained from the averages of the temperature gradients at the bottom and top plates, respectively. In table 2, the column Nu shows the average value obtained from these four methods, and the column headed 'Max-diff' shows the maximum difference between the results from these four methods. Notably, all differences are within 1 %, indicating the convergence of our simulations. We have also calculated the average Nu_h using half of the statistical averaging time. We observe consistency between Nu and Nu_h for all cases, confirming that our simulations are indeed well converged, and the averaging times are sufficient. Finally, the Reynolds number Re is defined as $Re = U_{rms}\sqrt{Ra/Pr}$, where $U_{rms} = \sqrt{\langle u_x^2 + u_y^2 + u_z^2 \rangle}$ is the r.m.s. of the velocity averaged over time and the entire domain.

3. Results

3.1. Nusselt number and Reynolds number

We start by examining the polymer effects on the macroscopic heat and momentum transfer. Figure 3(a) presents the Ra dependence of the Nusselt number Nu compensated with $Ra^{1/3}$, and figure 3(b) presents the Reynolds number Re compensated with $Ra^{1/2}$, where these two specific scaling exponents are suggested by Grossmann & Lohse (2000) for the classical regime of turbulent RB convection at medium Ra and $Pr \gtrsim 1$. It is evident that within our parameter ranges, both Nu and Re are drastically reduced with the addition of polymer additives, indicating a reduction of the heat and momentum transport. Moreover, comparing to the polymer solutions with small Wi (Wi = 5), we note that the amount of HTR and momentum transport reduction (MTR) increases at larger Wi (Wi = 10). This effect is more pronounced at the lower Ra under investigation, while for $Ra > 10^8$, Nu is nearly the same at Wi = 5 and Wi = 10, likely indicating the onset of a saturation regime. A fit to the data of Newtonian flows yields $Nu \sim Ra^{0.27}$ and $Re \sim Ra^{0.47}$. In general, the data follow the theoretical scaling of the Grossmann and Lohse theory (Grossmann & Lohse 2000, 2001); see figure 3(a). With polymer additives, the scalings remain similar: $Nu \sim Ra^{0.26}$ and $Re \sim Ra^{0.45}$ at Wi = 5, and $Nu \sim Ra^{0.27}$ and $Re \sim Ra^{0.46}$ at Wi = 10, but the absolute values of Nu and Re are smaller in the viscoelastic case compared to the Newtonian case. These results are consistent with previous experiments of Wei et al. (2012) and Cai et al. (2019); these authors found that the scalings of Nu versus Ra, and Re versus Ra, remain almost unchanged in their parameter ranges (*Ra* span from 10^9 to 10^{10} at $Pr \approx 4.3$).



Figure 3. (a) Nusselt number Nu - 1 compensated with $Ra^{-1/3}$. (b) Reynolds number Re compensated with $Ra^{-1/2}$. Ratios of (c) Nu and (d) Re for a viscoelastic flow to those for the corresponding Newtonian flow; N and V represent Newtonian and viscoelastic flows, respectively. The solid line in (a) is the result of the Grossmann–Lohse (Grossmann & Lohse 2000, 2001) fit for Pr = 4.38 from Ahlers *et al.* (2022).

Next, we directly compare Nu and Re of viscoelastic flows with those of Newtonian flows, as shown in figure 3(c,d). The ratio Nu^V/Nu^N between Newtonian and viscoelastic flows, lower than 1 for the quenching effect of polymer additives, exhibits a non-monotonic dependence on Ra. For the Nusselt number ratio (Nu^V/Nu^N) , we observe first an increase between $Ra = 10^6$ and $Ra = 10^7$, followed by a slight decrease as Ra rises further from $Ra = 10^7$ to $Ra = 10^9$, with a more pronounced decrease at large $Ra \ge 10^9$. This nonmonotonic behaviour suggests the existence of different heat transfer mechanisms in the low-Ra and large-Ra regions. Regarding the ratio of the Reynolds numbers (Re^V/Re^N) , a different trend is observed: a gradual decrease is noted for $Ra < 10^9$, after which it eventually reaches a plateau. It can be inferred that the polymer-induced drag enhancement becomes more pronounced with increasing Ra, and the plateau at higher Ra indicates that the maximum drag enhancement due to the polymer additives has been achieved. Although there are some fluctuations in the Nu and Re ratios at the low-Ra regime ($Ra \le 10^7$), which should be attributed to the sensitivity of these quantities due to the pseudo-laminar flow state, the overall trends remain clear and consistent.

In order to compare our results with existing experiments, we note that Benzi *et al.* (2016*b*) provided a reference for extracting the dependence of the viscosity ratio in the FENE-P model on the polymer concentration *c*, based on the experimental measurements of the viscosity of the PEO solution; cf. figure 7 of their paper for details. Since we have relatively large polymer extensibility, when ignoring the shear-thinning behaviour, the kinetic viscosity of the polymer solution $v_{sol}(c)$ for different polymer concentrations *c* can be simplified as the zero-shear viscosity $v_s + v_p$ in the model. Thus we have a

reasonable approximation of the relation between c and β . In our DNS, $\beta = 0.9$ would therefore correspond to the polymer concentration between 90 (ppm) and 120 (ppm). In our DNS, the ratio of Nu in a viscoelastic flow to that in the corresponding Newtonian flow at $10^9 \leq Ra \leq 10^{10}$ is approximately 0.85-0.91, very close to the findings of Wei *et al.* (2012), who observed ratio 0.88 for concentration c = 120 (ppm) at $Ra = 8.1 \times 10^9$, to the results reported by Ahlers & Nikolaenko (2010), who found ratio 0.90 for concentration c = 120 (ppm) at $Ra = 6.74 \times 10^{10}$, and also to the value 0.92 for concentration c = 100(ppm) at $Ra = 2 \times 10^9$ reported in Cai *et al.* (2019).

In what follows, we will focus on two key questions: first, why do the heat and momentum transport decrease, and second, why are the trends non-monotonic? Given that the results for Wi = 5 and Wi = 10 yield similar trends, our primary focus will be on the flow cases with Wi = 10.

3.2. Flow structure

The typical flow structures for both Newtonian and viscoelastic thermal convection are shown in figure 4. At low Ra ($Ra = 10^6$), the Newtonian and viscoelastic flows exhibit an ascending hot plume and a descending cold plume, which generate an LSC in the bulk region (see figure 4a,d). These plumes show quasi-periodic wave-like motions (see the supplementary movies 1 and 2) without visible horizontal sweeping. These oscillations may be attributed to plume-vortex interactions. The flow structures of the polymeric flow are similar to those of the Newtonian case, while the plumes become less wavy. With an increase of Ra to $Ra = 10^8$ (see figure 4b), the thermal BL becomes thinner, and more plumes are ejected from the BL. The comparison between Newtonian and viscoelastic flows reveals that the polymer additives change the structure of the plumes: they become thinner, while their stems become elongated and their caps almost reach the opposite plate; see figure 4(b,e). The waviness of these plumes is substantially weakened. This is reminiscent of the flow structures characterising drag reduction in turbulent channel flows, where the near-wall streamwise streaks at the core of the turbulence self-sustaining cycle become more stable in the presence of polymer additives, a calmer phase also denoted as 'hybernation' (Xi & Graham 2012).

At an even higher $Ra = 10^{10}$ (see figure $4c_s f$), the polymer additives reduce small-scale structures, making the plumes more coherent. This polymer-induced stabilisation has been reported in homogeneous turbulence (De Angelis *et al.* 2005), channel and pipe flows (Dubief *et al.* 2023), plane Couette flow (Teng *et al.* 2018), Rayleigh–Taylor turbulence (Boffetta *et al.* 2010) and Taylor–Couette turbulence (Song *et al.* 2023*b*).

Finally, the bottom panels in figure 4(g,h,i) display the trace of the polymer conformation tensor, which represents the spatial distribution of the elongation of the polymer chains. We can see that at relatively low Ra, most of the elongation concentrates inside the plumes, where the velocity gradient is higher, leading to maximum stretching of the polymer chains; these stretched molecules reach the opposite wall and form mushroom-like structures when interacting with the thermal BL (see the supplementary movie 3). As Ra increases, the magnitude of the polymer extension intensifies, and the region with highly stretched polymers becomes wider. Moreover, we also note that near the bottom/top wall, the polymers have stronger horizontal stretching at $Ra = 10^8$. For the largest $Ra = 10^{10}$ case, highly stretched polymers are even more dispersed, and they can be found throughout the entire flow field.

To continue our analysis of the main flow features, vertical and horizontal crosssections (mid-plane) of the temperature and trace fields are shown in figures 5 and 6, respectively. The temperature fields pertaining to the polymeric flows (second row) are



Figure 4. Instantaneous temperature fields of (a,b,c) Newtonian flows and (d,e,f) viscoelastic flows, and (g,h,i) the corresponding trace fields of viscoelastic flows. The columns from left to right represent $Ra = 10^6$, $Ra = 10^8$ and $Ra = 10^{10}$, respectively. (The viewing angle of trace fields is rotated 30° along the horizontal x-axis for a better visualisation.)

directly compared with the corresponding Newtonian flow (top row), whereas the images in the bottom row depict the trace of the polymer conformation tensor. The figures clearly show that with polymer additives, the thermal BLs become thicker, and the plumes become longer, wider and less fragmented. In the core region, the polymers suppress the small-scale structures, and the temperature is less efficiently mixed, leading to more distinct plume boundaries, especially at higher Ra. As concerns the polymer stretching, we note that with increasing Ra, the well-identified slim regions of high polymer stretching



Figure 5. Instantaneous vertical cross-sections (*xz*-plane) of the temperature fields for (*a,b,c,d*) Newtonian flows, (*e,f,g,h*) the corresponding viscoelastic flows, and (*i,j,k,l*) the trace of the conformation tensor in the same cross-section. The columns from left to right represent $Ra = 10^6$, $Ra = 10^8$, $Ra = 3 \times 10^9$ and $Ra = 10^{10}$, respectively.

gradually become wider and more chaotic. They are seen to connect with each other and spread over the whole domain with higher intensity and finer spatial scales. The reader is also referred to the supplementary movies to better appreciate the differences in flow structures due to polymer additives.

Figure 7 presents space-time plots of the temperature in Newtonian and viscoelastic flows along a horizontal line in the mid-height plane. The images are organised to show data for increasing Ra in each column. In the left-hand column (figure 7a, c, e, g), we first see that the Newtonian flow exhibits regular and periodic patterns with a stable and coherent structure. As Ra increases, the fluctuations increase, and the flow eventually exhibits a strongly chaotic behaviour. In contrast, the right-hand column (figure 7b, d, f, h) shows that the viscoelastic flows maintain more stable and coherent structures compared to the Newtonian flows, with less variation over time. Although the turbulence intensity increases for the larger $Ra > 10^8$, the viscoelastic flows retain a higher coherence compared to their Newtonian counterpart. This suggests that the viscoelastic properties of the fluid help to maintain the flow stability. At an even higher $Ra = 10^{10}$, the viscoelastic flow continues to demonstrate a stronger coherence, with reduced chaotic behaviour. Overall, the flow structures of the viscoelastic solutions exhibit a larger degree of coherence in their spatiotemporal structures compared to the corresponding Newtonian flows, as evidenced by more pronounced plume structures and less turbulent flow patterns.

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Figure 6. Instantaneous mid-height horizontal cross-sections (*xy*-plane) of the temperature fields for (*a,b,c,d*) Newtonian flows, (*e,f,g,h*) the corresponding viscoelastic flows, and (*i,j,k,l*) the trace of the conformation tensor in the same cross-section. The columns from left to right represent $Ra = 10^6$, $Ra = 10^8$, $Ra = 3 \times 10^9$ and $Ra = 10^{10}$, respectively.

3.3. Turbulent and polymer statistics

In this subsection, we discuss the turbulent velocity and temperature statistics, polymeric deformation, and BL thicknesses. Figure 8 depicts the wall-normal profiles of the r.m.s. vertical $u'_{z,rms}$ and horizontal $u'_{h,rms}$ velocity fluctuations for three representative cases. Both the vertical and horizontal velocity fluctuations are attenuated in the presence of polymer additives, indicating a reduction of turbulence intensity and an increase in flow drag. This observation is consistent with the HTR cases reported by Dubief & Terrapon (2020) at $Ra = 10^5$, where the addition of polymer additives resulted in a slowdown of the LSC and thus a reduction in heat transfer.

The profiles of the mean temperature θ and r.m.s. temperature fluctuation θ'_{rms} for the same representative cases as in figure 8 are shown in figure 9. As *Ra* increases, an enhanced convection leads to the contraction of the thermal BL region, resulting in a steeper temperature gradient near the wall, and more efficient mixing within the bulk region, for both Newtonian and viscoelastic flows. From figure 9(*a*), we observe that the differences of mean temperature between polymeric (dashed lines) and Newtonian flows (solid lines) are small in the bulk region, where the temperature remains almost constant. Within the thermal BL, polymeric flows exhibit a slightly smoother increase of the mean temperature compared to Newtonian flows, suggesting that the addition of polymers affects the stability and structure of the thermal BL. More pronounced differences are seen in the temperature fluctuation data: in viscoelastic flows, the r.m.s. values are smaller than



Figure 7. Temperature space-time plots for (a,c,e,g) Newtonian and (b,d,f,h) viscoelastic flows along a horizontal line $x \in [0, 1]$, y = 0.5 in the mid-height plane for (a,b) $Ra = 10^6$, (c,d) $Ra = 10^8$, (e,f) $Ra = 3 \times 10^9$, (g,h) $Ra = 10^{10}$.



Figure 8. Intensity of the velocity fluctuations: r.m.s. profiles of (a) vertical $u'_{z,rms}$ and (b) horizontal $u'_{h,rms}$ velocity fluctuations for Newtonian (Wi = 0) and viscoelastic flows with Wi = 10 for different values of Ra.

in Newtonian flows near the wall, and larger than those in Newtonian flows in the bulk region. The peaks of θ_{rms} are lower in viscoelastic flows and located further away from the wall (which supports the idea of a thicker BL in polymer solutions). This behaviour is consistent with previous studies on polymer solutions in homogeneous turbulence (De Angelis *et al.* 2005), where the polymer-induced suppression of small-scale motions is found to lead to increased fluctuations at larger scales. The reduction in temperature fluctuation near the wall implies an enhanced stability of the thermal BL, which could potentially reduce the frequency of the plume release.



Figure 9. Profiles of (a) the average temperature θ and (b) the temperature fluctuations θ'_{rms} for different values of Ra. The inset shows the profile in the near-wall region.



Figure 10. Profiles of (*a*) mean trace $\langle C_{ii} \rangle$ normalised by L^2 , and (*b*) first normal stress difference N_1 of viscoelastic flows at different Ra for Wi = 10.

Next, we explore the impact of polymer additives on the deformation of polymer molecules. Figure 10(a) presents wall-normal profiles of the mean trace of the polymer conformation tensor, representing the average polymer length, normalised by the square of the polymer extensibility parameter, $\langle C_{ii} \rangle / L^2$. The data in the figure reveal that the polymer extension increases with Ra, indicating stronger polymer-flow interactions due to the enhanced turbulent intensity. Moreover, as expected, the maximum values of the trace are observed near the wall, whereas in the bulk region, especially far from the wall, the polymer stretching remains relatively uniform. A classic measure of the non-Newtonian character of a suspension is the first normal stress difference, defined as $N_1 = (1 - \beta) \sqrt{Pr/Ra} (\langle \tau_{zz} \rangle - \langle \tau_{xx} \rangle)$, the normalised difference between two components of the normal stresses. Note that with our definition of N_1 , we consider z the principal flow direction (which is the case in the bulk but not in the thermal BLs). The wall-normal profiles of N_1 are depicted in figure 10(b). Near the wall, N_1 is negative except for case $Ra = 10^6$, whereas in the bulk region, it becomes positive, with positive values smaller than the absolute values of the negative ones. This validates the conclusion in Benzi et al. (2016b), who showed that when the stretching of polymers occurs predominantly within the BL, HTR could be observed. Also it implies that near the wall, τ_{xx} dominates and the polymer undergoes greater stretching in the horizontal direction, while in the bulk, τ_{zz} dominates and the stretching is more pronounced in the vertical



Figure 11. Visualisation of the definitions of (a) thermal and (b) viscous BL thicknesses using the example of the flow for $Ra = 10^8$ and Wi = 10.

direction. This observation is intuitive, as the polymer chains align with the primary flow direction. Indeed, a positive normal stress difference indicates that polymer chains are becoming oriented in the direction of the flow, in agreement with the visualisations shown above. For the case $Ra = 10^6$, the polymer stretching is mainly in the vertical direction, which is expected since the flow is laminar, and the plumes become less wavy with polymers, resulting in reduced horizontal motion. Furthermore, an increase in Raresults in a monotonic decrease of the magnitude of N_1 . Since the flows become more chaotic and turbulent as Ra increases, the polymers are not able to maintain significant stretching in one specific direction due to the increasingly isotropic and fluctuating nature of the flow. To conclude, we note that from a dynamics perspective, we therefore retrieve a positive value of N_1 for the average flow direction, as typically observed in shear flows. This indicates the presence of elasticity, polymer molecules trying to return to their coiled state.

Next, we investigate the effects of polymer additives on the BL thickness. We adopt the frequently used slope method to define the edge of the thermal BL. That is, the thermal BL is defined as the depth where the linear fit to the mean temperature profile θ near the bottom (or top) wall intersects the linear fit to the mean temperature profile at midheight, as illustrated in figure 11(a). The viscous BL, instead, is defined as the distance between the wall and the maximum value of the horizontal velocity fluctuations, $u'_{h,rms}$; see figure 11(b). The thermal and viscous BL thicknesses and their scaling behaviours are presented in figure 12. Figure 12(a) shows that the thermal BL thickness increases when adding polymer additives, suggesting an enhanced thermal BL stability. For smaller Ra, this increase is more pronounced with higher Wi = 10. On the contrary, at larger Ra, the thermal BL thicknesses remain nearly constant for both Wi = 5 and Wi = 10, indicating that the thermal BL thickness becomes less sensitive to polymer extensibility as Ra increases. However, the polymer additives do not modify the scaling of λ_{θ} with respect to *Ra*. Moreover, all the data collapse on the line $0.5Nu^{-1}$ (see figure 12b), fitting well with the theoretical prediction: $\lambda_{\theta} = H/(2Nu)$ for turbulent thermal convection. Interestingly, the viscous BL thickness exhibits a different behaviour. Specifically, the viscous BL thickness scales differently in flows with polymers than in Newtonian flows, displaying a steeper decrease of λ_u with increasing Ra. The intersection of the λ_u curves in Newtonian and viscoelastic flows happens at approximately $Ra = 10^8$. The scaling of λ_u with Reyields $\lambda_{\mu} \sim Re^{-0.24}$ for Newtonian flows, as found previously in both simulations (Breuer et al. 2004; King, Stellmach & Buffett 2013) and experiments (Lam et al. 2002). For the



Figure 12. The thermal BL thickness λ_{θ} versus (a) the Rayleigh number Ra and (b) the Nusselt number Nu. The viscous BL thickness λ_{u} versus (c) Ra and (d) Re. The dashed lines depict a polynomial fitting to the data. For better visualisation, in (a) and (c) we present only the fitting lines corresponding to Wi = 0 and Wi = 10.

viscoelastic cases, the scaling becomes steeper, $\lambda_u \sim Re^{-0.30}$, indicating a modification of the kinetic BL dynamics.

4. Discussion

4.1. Effective viscosity

It is generally argued in the literature that HTR is related to the enhanced stability of the thermal BL, which can be attributed to two primary factors (Xie *et al.* 2015): (i) the reduced bulk velocity fluctuations that constantly perturb the BL, as depicted in figure 8; (ii) the increased effective viscosity of the polymer solutions that reduces the convection velocity. In order to verify this, we examine the relative effective viscosity of the polymer solutions. According to Benzi *et al.* (2016*a*), the relation between the space-dependent effective viscosity v_{eff} and the solvent viscosity v_s can be expressed as

$$\frac{v_{eff}}{v_s} = \frac{E_p}{\varepsilon_u^{[s]}},\tag{4.1}$$

where E_p is the rate of energy transferred to polymer elastic energy, calculated by $E_p = (1/2)(1 - \beta/Re)\tau_{ii}$ (Min *et al.* 2003), and $\varepsilon_u^{[s]}$ is the solvent viscous dissipation rate defined in (2.9).

Figure 13 depicts the wall-normal profiles of the effective viscosity v_{eff} normalised by the solvent viscosity v_s for the polymeric flows. The data show that the normalised effective viscosity exhibits a local maximum in the near-wall region, with a smaller value



Figure 13. Wall-normal profiles of the effective viscosity v_{eff} normalised by the solvent viscosity v_s for different values of Ra.

compared to the bulk region. As one moves away from the wall, the effective viscosity increases (more dramatically at lower Ra) and then remains almost constant inside the bulk region. This indicates that the interaction between polymers and the turbulent flow is more pronounced in the bulk, leading to a more pronounced enhancement of viscosity in that region. The non-monotonic behaviour observed near the wall is difficult to explain only from the definition in (4.1) and the complex interactions between turbulence and viscoelastic effects. The local maxima near the wall may be related to the peak of the polymer elastic energy (profiles of E_p), and are similar to the trace $\langle C_{ii} \rangle$ presented in figure 10(*a*).

More importantly, we observe that v_{eff} is correlated with the ratio of Reynolds numbers Re^V/Re^N shown in figure 3(d). With increasing Ra, the effective viscosity ratio increases, leading to a decrease in Re up to $Ra = 10^9$. Beyond this threshold, the effective viscosity remains nearly unchanged, resulting in an almost constant Re. We then can conclude that for $Ra < 10^9$, the increased viscosity causes a slower circulation, as corroborated by the supplementary movies, thus decreasing the heat transfer rate. Further, the effective viscosity ratios are almost equal for higher $Ra \ge 10^9$, meaning that there may exist a maximum drag enhancement. The additional HTR for $Ra > 10^9$ is therefore caused by other factors, which we discuss below.

4.2. Heat flux from plume and turbulence background in the mid-height plane

In this subsection, we aim to investigate why the heat transport is reduced and why the HTR shows a non-monotonic trend with Ra. Since plumes are the primary carrier of heat in turbulent thermal convection, we consider the contribution from the plumes and the turbulent background separately. First, we calculate the probability density functions (PDFs) of the vertical heat flux $(u_z\theta')$ at the mid-height plane. It should be noted that $Nu_{centre} = \sqrt{Ra Pr} \langle u_z\theta' \rangle_{A,t}$ (Kaczorowski & Xia 2013). The negative part of the vertical heat flux PDF is due to turbulent background fluctuations, known as incoherent heat flux, while the positive part contains both incoherent heat flux and coherent heat flux arising from plumes (Shang *et al.* 2003; Xie *et al.* 2015). The large positive value of $u_z\theta'$ should be related to plumes. For each case presented in figure 14, the dataset includes flow field information for at least 300 dimensionless time units. Intriguingly, the PDFs exhibit different characteristics for small $Ra (Ra < 10^9)$ and relatively high $Ra (Ra \ge 10^9)$ regimes.



Figure 14. Probability density functions (PDFs) of the vertical heat flux $u_z \theta'$ at the mid-height plane for (a) $Ra < 10^9$ and (b) $Ra \ge 10^9$.

In the low $Ra < 10^9$ regime, as depicted in figure 14(*a*), the addition of polymer additives leads to a significant suppression of both negative and positive tails, particularly for very large values of the positive vertical heat flux - those related to the plumes, suggesting a suppression of both incoherent turbulent background fluctuations and coherent heat transport. This effect is more pronounced for smaller Ra. The suppression of incoherent heat flux is confirmed by the significant decrease in horizontal turbulent fluctuations in the central plane discussed above (see figure 8). This substantial suppression of the coherent flux is attributed to a slower circulation speed, which is more evident for $Ra = 10^6$, where we observe a pair of convection cells of lower LSC speed (see supplementary movies 1 and 2). Since the relative effective viscosity ratio increases significantly with Ra, one would intuitively expect a stronger suppression of the coherent motions at higher Ra. However, the figure shows the opposite trend, which may indicate that polymers have a twofold action. For higher $Ra \ge 10^9$ (see figure 14b), our observations align with the Xie *et al.* (2015) finding that the negative part of the PDF is suppressed, and the probability of large positive values is significantly higher for the polymer solutions. Given the slight increase in temperature fluctuations (see figure 9) and the halving of the velocity fluctuations at mid-height (see figure 8), this enhancement is most likely due to the increased correlation between velocity and temperature, in other words, to the increasing coherence of the thermal plumes. In summary, the data suggest that in the laminar or weakly turbulent regime, $Ra < 10^9$, polymers reduce the plume intensity, whereas at larger $Ra \ge 10^9$, they enhance the coherent transport by the plumes by reducing the effect of the background fluctuations.

To assess more quantitatively the effect of polymers additive on plumes, it is crucial to detect plumes and then separate them from the background fluctuations. In previous studies, various methods have been employed to extract plumes based on their distinct characteristics, such as high temperature (Zhou & Xia 2002), high thermal dissipation rate (Shishkina & Wagner 2006, 2008), high vertical heat flux (Ching *et al.* 2004), and negative horizontal divergence of the horizontal velocity field (Shevkar *et al.* 2022). Here, we adopt the criteria that combine both temperature and heat flux conditions (Huang *et al.* 2013; van der Poel *et al.* 2015*a*). That is, in a certain horizontal cross-section, the plume is the area satisfying $\pm [\theta(x, y) - \langle \theta \rangle_{xy}] > c \langle \theta_{rms} \rangle_{xy}$ and $\sqrt{Ra Pr} u_z(x, y) \theta'(x, y) > c(Nu - 1)$, where '+' indicates a hot plume, and '-' indicates a cold one.

Note that we use the factor (Nu - 1) to exclude conductive heat transfer. The remaining part of the cross-section is defined as background. We choose the same empirical constant



Figure 15. Plumes extraction at (a,b,e,f) the edge of the thermal BL (z = 0.5H/Nu) and (c,d,g,h) the middleheight xy-plane for (a,b,c,d) Newtonian and (e,f,g,h) viscoelastic (Wi = 10) flows at $Ra = 10^8$, Pr = 4.3. The temperature shown in (a,c,e,g) has the same colour bar as in figure 6, and the red and blue colours in (b,d,f,h) denote the extracted hot and cold plumes, respectively.

c = 1.2 as in van der Poel *et al.* (2015*a*). The choice of *c* influences the absolute value of the plume area and plume mean velocity, but the trends presented below remain independent of *c*. The results of this plume-extraction procedure are displayed in figure 15 for both Newtonian and viscoelastic flows at $Ra = 10^8$. One can see that this criterion successfully extracts both hot and cold plumes in the thermal BL and bulk regions. We use this procedure to calculate the mean temperature and vertical velocity of plumes, as well as the heat flux contributions from both plumes and background, in the middle-height plane by averaging over 300 free-fall time units.

Figure 16(a,b) show the mean temperature and mean vertical velocity differences between viscoelastic and Newtonian flows for hot and cold plumes. These are defined as $\Delta \theta_p = \langle \theta \rangle_{p,t}^V - \langle \theta \rangle_{p,t}^N$ and $\Delta u_{z,p} = \langle u_z \rangle_{p,t}^V - \langle u_z \rangle_{p,t}^N$. Interestingly, we observe that, compared to the Newtonian flow, the hot plumes are less hot, and the cold plumes are less cold, in viscoelastic flows at small $Ra < 10^8$. However, for $Ra \ge 10^8$, this trend reverses, with hot plumes becoming hotter, and cold plumes becoming colder. Moreover, the magnitude of this difference increases with Ra. The vertical velocity of the plumes in viscoelastic flows is lower than in Newtonian flows, and the magnitude of this decrease becomes smaller with increasing Ra until $Ra = 10^9$, beyond which it reaches a plateau, which aligns with the trend for Re shown in figure 3, and the effective viscosity shown in figure 13. The data seem to confirm our interpretation above: at small Ra, the addition of polymers dampens the intensity of thermal plumes by increasing the effective viscosity of the flow. Consequently, the effective thermal diffusivity increases, which leads to a decreased temperature of the hot plumes, and increased temperature of the cold ones. Conversely, at high Ra, the stabilised BL in the viscoelastic flows makes the released plumes more coherent. Additionally, the presence of polymer additives reduces the turbulence intensity, leading to weaker mixing. This reduced mixing slows down the horizontal heat transport, causing heat to remain more concentrated inside the plumes.



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Figure 16. Results of plume extraction: difference in (a) plume mean temperature $\Delta \theta_p = \langle \theta \rangle_{p,t}^V - \langle \theta \rangle_{p,t}^N$ and (b) plume vertical velocity $\Delta u_{z,p} = \langle u_z \rangle_{p,t}^V - \langle u_z \rangle_{p,t}^N$ between Newtonian flows and viscoelastic flows at Wi = 10. (c) Contributions to the total Nu from the plumes and background chaotic flow. (d) Difference $\Delta Nu = Nu^V - Nu^N$ between Newtonian and viscoelastic flows at Wi = 10, considering the heat flux in plumes, background, and the total value.

To further investigate the heat transfer mechanisms, we now discuss the heat flux contributions from the plumes and the background chaotic flow, still focusing on the midplane. Figure 16(c) displays the total $Nu = \sqrt{Ra Pr} \langle u_z \theta' \rangle$, as well as the heat flux from the turbulent background and plumes. We note that the heat flux from both turbulent background and plumes increases with increasing Ra, and plumes serve as the main heat carriers for both Newtonian and viscoelastic flows. Focusing on the differences, figure 16(d) presents the HTM between viscoelastic and Newtonian flows $\Delta Nu = Nu^V - Nu^N$, considering the total flux, and the plume and background contributions. First, we observe that the heat flux reduction increases with the Rayleigh number, and that this difference can be ascribed to the plumes for $Ra \leq 10^7$, when the flow is in a laminar-like regime. Further increasing Ra, instead, the reduction is mainly due to the reduction of the background turbulence. The contributions from the plumes actually increase at the largest values of the Rayleigh number under consideration. Nevertheless, the increased heat flux associated with more coherent plumes is not enough to balance the reduction associated with the background fluctuations.

The results therefore confirm the observations made when examining the PDFs. That is, the heat flux in the mid-plane from the background generally decreases with polymer additives, while the heat flux from the plumes first decreases for $Ra < 10^9$, and then increases. While at small Ra the reduction in the total heat flux arises mainly from the plumes, at large Ra, even though the heat flux from the plumes region increases, the significant decrease in the heat flux from the turbulent background determines the total HTR.



Figure 17. Difference between the Nusselt numbers of Newtonian and viscoelastic flows, $\Delta Nu = Nu^V - Nu^N$, in different regions: BLs, plumes and background.

Overall, the polymer additives affect the turbulent background and plumes in different ways. That is, polymers act to suppress turbulence in the background flow, leading to lower turbulence intensity. As concerns the plumes, the net effect of polymer additives varies with the Rayleigh number: (i) they reduce the vertical velocity of strong plumes, thereby leading to the suppression of strong heat transport events, and this effect is more pronounced at lower Ra when the background flow is laminar-like; (ii) they enhance the coherence of the plumes, thus increasing the heat flux. These two effects are in competition, with the latter becoming dominant as Ra increases. In the following, we investigate how the flow in the BL is affected by the polymers, and how this reflects in the heat fluxes.

4.3. Volume-averaged heat flux from plumes, BLs and background

To further elucidate the impact of viscoelasticity on the HTM of the entire system, we examine the contributions of the BLs, plumes and turbulent background to the volumeaveraged heat flux derived by the exact relation between Nu and the thermal dissipation rate. The decomposition is based on the criterion for plume extraction, as mentioned previously. The BLs are defined as the regions within distance 0.5H/Nu from the top and bottom walls exclusion plume region. The remaining area is classified as turbulent background. Figure 17 displays the differences between the Nusselt numbers of Newtonian and viscoelastic flows across these regions, with all results averaged over 100 free-fall time units. We observe that the total $\Delta N u$ shows a non-monotonic trend as the HTR: also, the heat flux contributions from the plumes and background follow trends similar to those observed in the mid-plane, i.e. the total heat transferred by the plumes first decrease and then increase with Ra in the viscoelastic flows while it decreases in the background flow, with the latter decrease being more pronounced. Most interestingly, we note that the largest part of the reduction in viscoelastic flows is associated with the heat flux in the BLs, and it increases with Ra. In the experiments by Xie et al. (2015) at similar $Ra = 6.18 \times 10^9$, Pr = 4.3, it was found that polymer additives enhance heat transport in the turbulent thermal convection bulk flow by increasing the coherency of thermal plumes and suppressing turbulent background fluctuations. On the other hand, the polymer additives stabilise the thermal BLs and reduce plume release. Xie et al. (2015) explained the observed HTE in their experiments by saying that the effect of enhanced plume coherency overcomes the effect of reduced plume emission. Their observations on the effects of polymers are consistent with our study. The qualitative difference in heat



Figure 18. (a) Viscous (circle) and elastic (triangle) energy dissipation rates in Newtonian and viscoelastic (Wi = 10) flows versus Ra. (b) The fraction of viscous (green circle) and elastic (yellow triangle) contributions to the total kinetic dissipation rate in viscoelastic flows.

transport modification – i.e. HTE in their experiments and HTR in our simulations – is attributed to rough walls in their experiments, which makes both kinetic energy and thermal dissipation rates bulk-dominated.

4.4. Kinetic energy dissipation rate and energy spectra

In this subsection, we focus on the contribution from turbulence and polymers to the kinetic energy dissipation rate. In viscoelastic flows, the fluid kinetic energy is partly dissipated by the viscous stresses at small scales, and partly transferred to the polymers, which is called the elastic dissipation rate. By taking the dot product of the momentum equation (2.2) with the velocity u_i , and averaging over time and over the entire volume (Benzi *et al.* 2016*a*), the exact balance between energy production rate by buoyancy, viscous dissipation rate in the Newtonian solvent $\varepsilon_u^{[s]}$ and elastic dissipation rate due to the polymer additives $\varepsilon_u^{[p]}$ is given as

$$\frac{\nu^3}{H^4} Ra Pr^{-2} (Nu - 1) = \varepsilon_u^{[s]*} + \varepsilon_u^{[p]*}.$$
(4.2)

After normalisation using U_{ff} and H, this can be rewritten as

$$\frac{Nu-1}{\sqrt{Ra\ Pr}} = \varepsilon_u^{[s]} + \varepsilon_u^{[p]} = \beta \sqrt{\frac{Pr}{Ra}} \left[\langle \partial_i u_j(\mathbf{x}, t) \rangle_{V,t} \right]^2 - (1-\beta) \sqrt{\frac{Pr}{Ra}} \langle u_i \ \partial_j \tau_{ij} \rangle_{V,t}.$$
(4.3)

The value of Nu can thus be directly related to $\varepsilon_u^{[s]}$ and $\varepsilon_u^{[p]}$.

The dissipation rates $\varepsilon_u^{[s]}$, $\varepsilon_u^{[p]}$ and total dissipation rate are displayed in figure 18(*a*). As expected, both $\varepsilon_u^{[s]}$ and $\varepsilon_u^{[p]}$ increase with *Ra*, and the total energy dissipation rates $\varepsilon_u = \varepsilon_u^{[s]} + \varepsilon_u^{[p]}$ of polymer solutions decrease compared to Newtonian flows, leading to the HTR. Here, the increasing $\varepsilon_u^{[p]}$ is related to the amount of the polymer stretching, as indicated by the trace shown in figure 10(*a*).

The relative contributions of the viscous and elastic dissipation to the total energy dissipation rate of polymer solutions are reported in figure 18(*b*). The data in the figure reveal that when increasing *Ra*, the contribution of the elastic part increases monotonically and becomes dominant over the viscous dissipation at $Ra \ge 10^7$, which indicates the growing importance of elastic stresses in the flow. In addition, at higher *Ra*, more of



Figure 19 (a) Turbulent kinetic energy spectra normalised by $u_i^2/2$, $\overline{E_u}(k)$, and (b) temperature energy spectra normalised by $\theta^2/2$, $\overline{E_{\theta}}(k)$, sampled at the mid-height of the domain, and averaged over the plane and over time. The spectra are plotted as functions of normalised wavenumber k for different Ra, comparing Newtonian flows (solid lines) with viscoelastic flows at Wi = 10 (dashed lines). The solid and dashed black lines show the scaling laws -5/3 and -14/3, respectively.

the kinetic energy is transferred into elastic potential energy as the polymers are stretched by the turbulent flow. This elastic energy is released back to the flow once the polymer stretching reduces. The effect of this mechanism in thermal convection, and how the kinetic energy redistributes at different spatial scales, deserve further investigation.

To shed some light on these energy transfer mechanisms, we consider the onedimensional turbulent kinetic energy $\overline{E}_{\mu}(k)$ and temperature energy spectra $\overline{E}_{\theta}(k)$. The one-dimensional energy spectra are calculated by the fluctuations from the three velocity components and temperature at the middle-height plane. The spectra in figure 19 are obtained by first computing the Fourier transform in the x (or y) direction, then averaging in the orthogonal horizontal direction, i.e. y (or x). Finally, averages are performed over the two horizontal directions and over at least 300 dimensionless time units. As depicted in figure 19(a), with increasing Ra, the classical inertial scaling $k^{-5/3}$ is gradually reached in the Newtonian flows in the intermediate wavenumber range. When polymer additives are added, the large-scale content decreases (energy in the low wavenumber range), while the small-scale energy content increases (high wavenumber range). Moreover, the spectra tend to follow the power-law decay with exponent nearly -14/3 at high $Ra \ge 10^8$, as also reported in other flow configurations with elasto-inertial turbulence (Dubief et al. 2013; Sid et al. 2018; Song et al. 2021). The suppression of large-scale motions and the activation of small-scale fluctuations observed here may be related to polymer stretching (see enhanced polymer dissipation in figure 18), and is consistent with the typical features of elasto-inertial turbulence, where the small-scale motions are sustained by a significant polymer stretching (Sid et al. 2018). Note that a similar activation of small-scale motions is also observed in other multiphase systems, for example, in emulsions, where the small-scale motions are induced by surface tension releasing energy to the flow upon relaxation and coalescence (Crialesi-Esposito, Chibbaro & Brandt 2023; Moradi Bilondi et al. 2024).

Finally, examining the temperature fluctuation spectra of polymeric flows in figure 19(*b*), we note, as for the temperature fluctuations, that the large-scale content increases slightly, while small-scale content decreases in the presence of polymer additives. It is also worth noticing that the polymeric temperature spectra do not follow the same scaling law $k^{-14/3}$ observed for turbulent kinetic energy spectra, indicating a lower correlation between velocity and temperature fluctuations in the polymeric turbulent thermal convection.

5. Conclusions

In this work, three-dimensional DNS are performed to explore the effects of polymer additives on turbulent RB convection by using the FENE-P constitutive model with fixed polymer extensibility parameter L = 50 and viscosity ratio $\beta = 0.9$. The Prandtl number is set to 4.3, and two moderate Weissenberg numbers (Wi = 5 and 10) are considered. The simulations are conducted for a broad range of Ra from 10^6 to 10^{10} , which is much broader than any of the existing numerical simulations of RB convection with polymer additives. This parameter range is encompassing the Ra range explored in experiments.

Our results show that within our parameter range, the heat and momentum transfer are reduced by polymer additives. The effects of polymers are different in different flow regions. Specifically: (i) within the BL region, polymers stabilise the thermal BL, leading to an apparent HTR, which serves as the primary mechanism for HTR in all cases of our study; (ii) in the bulk region, the polymers slow down the flow velocity by increasing the effective viscosity, enhance the plume coherency, and suppress the small-scale turbulent fluctuations. We observed a non-monotonic behaviour of heat transfer by plumes in the range $10^8 \leq Ra \leq 10^9$, where it initially decreases and then increases. By separating the plumes from the turbulent background in the mid-height plane, we find that at small Ra, in a laminar-like regime, the HTR is due mainly to the decreased plume velocity. As Raincreases, the effect of enhanced plume coherency gradually overcomes the slowdown in velocity, allowing plumes to transport more heat. The reduction in heat transfer in the bulk region is thus the result of the competition between suppressed background fluctuations and enhanced plume coherency.

Comparing to Newtonian flows, the main features of RB convection of polymeric flows can be summarised as follows: (i) a significant reduction in both vertical and horizontal turbulent velocity fluctuations; (ii) an increase in temperature fluctuations within the bulk region, and a decrease in temperature fluctuations within the BL; (iii) an enhancement of the thermal BL thickness, which still follows the classical scaling; (iv) an increase ($Ra < 10^8$) and then decrease ($Ra > 10^8$) of the viscous BL thickness, which yields a steeper scaling relation with Re; (v) a decrease in the turbulent kinetic energy content at the large energy-containing scales, while increasing at the small scales, in agreement with other multiphase flows, e.g. emulsions and particle-laden flows.

Our simulations support the experimental findings of Xie et al. (2015): the increased coherence of the thermal plumes, the suppression of turbulent fluctuations, and the enhanced BL stability. The differences in heat transport modulation between their experiments and our simulations are attributed to the side walls and rough surfaces of the the container used in their study, which alter the BL stability. While our study provides a more detailed view of the flow structures modifications in RB convection with polymer additives, further investigations are required to explore the effect of rough walls or of modifications of the boundary conditions on the heat and momentum transfer in the convection cell. In particular, it remains to investigate why the viscous BL thickness decreases at high $Ra > 10^8$ despite the turbulence intensity being reduced by the polymers. Moreover, it would be interesting to better understand how the stored elastic energy is released to the flow at the different length scales by a scale-by-scale analysis. In conclusion, this study has started to fill the gap between previous experimental and numerical studies on turbulent thermal convection with polymers, and shed some insights on the flow physics and heat transport mechanisms, focusing on the coherent structure modification of turbulent convection in complex fluids.

Supplementary movies. Supplementary movies are available at https://doi.org/10.1017/jfm.2025.10286.

Acknowledgements. We thank Professor D. Lohse for his valuable suggestions. C.X. gratefully acknowledges support from the China Scholarship Council (grant 202206340009), J.S. acknowledges support from the Alexander von Humboldt Foundation and European Research Council (ERC) advanced grant no. 101094492 MultiMelt, and O.S. acknowledges support from the German Research Foundation (DFG). The calculations were completed on the HoreKa supercomputer and the HPC systems of the Max Planck Computing and Data Facility (MPCDF).

Declaration of interests. The authors report no conflict of interest.

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