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Elastic turbulence in curvilinear flows of polymer solutions

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Abstract. Following our first report (A Groisman and V Steinberg 2000 Nature 405 53), we present an extended account of experimental observations of elasticity-induced turbulence in three different systems: a swirling flow between two plates, a Couette-Taylor (CT) flow between two cylinders, and a flow in a curvilinear channel (Dean flow). All three set-ups had a high ratio of the width of the region available for flow to the radius of curvature of the streamlines. The experiments were carried out with dilute solutions of high-molecular-weight polyacrylamide in concentrated sugar syrups. High polymer relaxation time and solution viscosity ensured prevalence of non-linear elastic effects over inertial non-linearity, and development of purely elastic instabilities at low Reynolds number (Re) in all three flows. Above the elastic instability threshold, flows in all three systems exhibit features of developed turbulence. They include: (i) randomly fluctuating fluid motion excited in a broad range of spatial and temporal scales and (ii) significant increase in the rates of momentum and mass transfer (compared with those expected for a steady flow with a smooth velocity profile). Phenomenology, driving mechanisms and parameter dependence of the elastic turbulence are compared with those of the conventional high-Re hydrodynamic turbulence in Newtonian fluids. Some similarities as well as multiple principal differences were found. In two out of three systems (swirling flow between two plates and flow in the curvilinear channel), power spectra of velocity fluctuations decayed rather quickly, following power laws with exponents of about -3.5. It suggests that, being random in time, the flow is rather smooth in space, in the sense that the main contribution to deformation and mixing (and, possibly, elastic energy) is coming from flow at the largest scale of the system. This situation, random in time and smooth in space, is analogous to flows at small scales (below the Kolmogorov dissipation scale) in high-Re turbulence.

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Notation

С	passive tracer (dye) concentration
<i>c</i>	average concentration
c_0	initial concentration of dye
C _{rms}	root-mean-square of deviations of dye concentration from its average,
	divided by the average
d	gap width
d/R	gap ratio
f	frequency
k	wave number
Κ	parameter of elastic non-linearity for normal stress driven instabilities
	in curvilinear flows $(=(d/R)(\eta_p/\eta)Wi^2)$
L	system size
M_w	average molecular weight

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Ν	number of half-rings in a curvilinear channel
N_1	first normal stress difference
р	pressure
P	power of fluctuations
Pe	Peclet number
Q	flow discharge rate
r, ϕ, z	cylindrical co-ordinates (radial, azimuthal, axial)
R	radius of either a plate in rotating flow between two plates, or inner
	cylinder in Couette–Taylor flow
R_{g}	radius of gyration of a polymer
Re	Reynolds number $(=VL/v)$
t_{vd}	viscous diffusion time $(=d^2/\nu)$
Т	torque
T _{lam}	torque in polymer solution flow in a laminar regime
T_s	torque measured in a solvent flow
Та	Taylor number $(=(d/R)Re^2)$
V_r	radial (spanwise) velocity component, D-diffusion coefficient
V_{ϕ}	azimuthal (longitudinal) velocity component
Wi	Weissenberg number
Wi _c	critical Wi of the elastic instability onset
z_b	characteristic thickness of boundary layer
Ϋ́	shear rate
λ	polymer relaxation time
η	viscosity
η_s	viscosity of Newtonian solvent
η_p	part of solution viscosity due to polymer molecules
η'	in-phase component of viscosity in an oscillatory test
$\eta^{\prime\prime}$	out-of-phase component of viscosity in an oscillatory test
ν	kinematic viscosity
ρ	density
τ	stress tensor
$ au_s$	stress tensor due to Newtonian solvent
$ au_p$	polymer stress tensor
$ au_w$	shear stress near a wall
$ au_w^{lam}$	shear stress near a wall in a laminar flow
Ω	rotation (angular) velocity

1. Introduction

Turbulence is one of the most fascinating phenomena in nature and one of the biggest challenges for modern physics. It is common knowledge that the flow of a simple, low molecular, Newtonian fluid is likely to be turbulent, when the velocity, V, is high, kinematic viscosity of the fluid, ν , is low and size of the tank, L, is large. All of these factors contribute to a high value of the Reynolds number, $Re = VL/\nu$. Motion of the Newtonian fluids is governed by the Navier–Stokes equation [1]

$$\frac{\partial \vec{\mathbf{V}}}{\partial t} + (\vec{\mathbf{V}}\vec{\nabla})\vec{\mathbf{V}} = -\nabla p/\rho + \nu\Delta\vec{\mathbf{V}},\tag{1}$$

where p is the pressure and ρ the density of the fluid. The equation has a non-linear term, $(\vec{V}\vec{\nabla})\vec{V}$, which is inertial in nature. *Re* defines the ratio of this non-linear term to the viscous dissipative term, $\nu\Delta\vec{V}$. When *Re* is high, the non-linear effects are strong and the flow is likely to be turbulent. Therefore, turbulence in fluids at high *Re* is a paradigm for a strongly non-linear phenomenon in spatially extended systems [1, 2].

Solutions of flexible high-molecular-weight polymers are viscoelastic liquids, and they differ from Newtonian fluids in many aspects [3]. The most striking elastic property of the polymer solutions is, probably, the dependence of mechanical stresses in flow on the history of the flow. So, the stresses do not immediately become zero when fluid motion stops, but rather decay with some characteristic relaxation time, λ , which can be well above 1 s. When a polymer solution is sufficiently dilute, its stress tensor, τ , can be divided into two parts, $\tau = \tau_s + \tau_p$. The first part, τ_{s_2} is defined by viscosity of the Newtonian solvent, η_s , and rate of strain in the flow, $\tau_s = -\eta_s [\nabla \vec{V} + (\nabla \vec{V})^T]$. Therefore, the equation of motion for a dilute polymer solution becomes

$$\frac{\partial \vec{\mathbf{V}}}{\partial t} + (\vec{\mathbf{V}}\vec{\nabla})\vec{\mathbf{V}} = -\vec{\nabla}p/\rho + (\eta_s/\rho)\Delta\vec{\mathbf{V}} - \nabla\tau_p/\rho.$$
⁽²⁾

Here the elastic stress tensor, τ_p , is due to the polymer molecules, which are stretched in the flow, and it depends on the history of the flow. One can see that τ_p enters the equation of motion linearly. So the degree of non-linearity of the equation of motion can still be defined by $Re = VL\rho/\eta_s$.

The simplest model incorporating the elastic nature of the polymer stress tensor, τ_p , is a Maxwell-type constitutive equation [3] with a single relaxation time, λ ,

$$\tau_p + \lambda \frac{\mathrm{D}\tau_p}{\mathrm{D}t} = -\eta_p [\vec{\nabla}\vec{\mathrm{V}} + (\vec{\nabla}\vec{\mathrm{V}})^{\mathrm{T}}].$$
(3)

Here $D\tau_p/Dt$ is a material time derivative of the polymer stress. An appropriate expression for the time derivative $D\tau_p/Dt$ has to take into account that the stress is carried by fluid elements, which move, rotate and deform in the flow. The translational motion implies an advection term $(\vec{V}\vec{\nabla}) \cdot \tau_p$ in an appropriate expression for $D\tau_p/Dt$, while the rotation and deformation of the fluid particles should lead to contributions like $(\vec{\nabla}\vec{V}) \cdot \tau_p$ or $\tau_p \cdot (\vec{\nabla}\vec{V})$ [3]. Therefore, along with terms linear in τ_p and \vec{V} , some non-linear terms, in which τ_p is coupled to \vec{V} , should appear in the constitutive relation. A simple model equation for $D\tau_p/Dt$, which is commonly used for description of dilute polymer solutions, is the upper convected time derivative,

$$\frac{\mathrm{D}\tau_p}{\mathrm{D}t} = \frac{\partial\tau_p}{\partial t} + (\vec{\mathbf{V}}\vec{\nabla}) \cdot \tau_p - (\vec{\nabla}\vec{\mathbf{V}})^{\mathrm{T}} \cdot \tau_p - \tau_p \cdot (\vec{\nabla}\vec{\mathbf{V}}).$$
(4)

Equations (3) and (4) together with the expression for τ_s constitute the Oldroyd-B model of polymer solution rheology [3]. One can see that non-linear terms in the constitutive equations (3) and (4) are all of the order $\lambda(V/L)\tau_p$. The ratio of those non-linear terms to the linear

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relaxation term, τ_p , is given by a dimensionless expression $\lambda(V/L)$, which is usually called the Weissenberg number, *Wi*. (The relaxation term τ_p is somewhat analogous to the dissipation term in the Navier–Stokes equation.)

One can expect mechanical properties of the polymer solutions to become notably nonlinear at sufficiently large Weissenberg numbers. Indeed, quite a few effects originating from the non-linear polymer stresses have been known for a long time [3]. So, in a simple shear flow of a polymer solution there is a difference between normal stresses along the directions of flow and velocity gradient. At low shear rates this normal stress difference, N_1 , is proportional to the square of shear rate. When flow lines are curvilinear, it gives rise to a volume force acting on the liquid in the direction of the curvature, the 'hoop stress'. Therefore, if a rotating rod is inserted in an open vessel with a highly elastic polymer solution, the liquid starts to climb up on the rod, instead of being pushed outwards [4]. This phenomenon is known as 'rod climbing', or the 'Weissenberg effect'. Furthermore, in a purely extensional flow, resistance of a polymer solution depends on the rate of extension in a strongly non-linear fashion. There is a sharp growth in the elastic stresses, when the rate of extension exceeds $1/(2\lambda)$, that is at Wi > 1/2. As a result, apparent viscosity of a dilute polymer solution can increase by up to three orders of magnitude [5]. Both the Weissenberg effect and the growth of the extensional flow resistance have been most clearly observed in very viscous polymer solutions and in flows with quite low *Re*, when non-linear inertial effects were insignificant.

A natural question arising here is whether there may exist some kind of turbulent flow produced by the non-linear polymer stresses alone, in the absence of any significant inertial effects, at low *Re*. An important step in this direction was made about a decade and a half ago when purely elastic instabilities were experimentally identified in a rotational flow between two plates [6] and in the Couette–Taylor (CT) flow between two cylinders [7]. The experiments were carried out with a Boger fluid [8], a dilute solution of high-molecular-weight polymers in a viscous Newtonian solvent. The Boger fluids are almost universally used as model viscoelastic fluids. Their relaxation times can be quite large, reaching seconds or even minutes, while their rheological properties are semi-quantitatively described by the simple Oldroyd-B model. The purely elastic instabilities occurred at *Wi* of order unity and vanishingly small *Re*. As a result of the instabilities, secondary vortex flows developed [7], and flow resistance somewhat increased [6]. The problem of linear stability of the CT flow was also treated theoretically using the Oldroyd-B model [9]. The analysis showed that the non-linear mechanical properties of the polymer solution can indeed lead to a flow instability, and a simple mechanism of this purely elastic instability was proposed.

During the past decade the purely elastic instabilities in the Boger fluids have been the subject of many theoretical and experimental studies, which are partially reviewed in [10, 11]. After the pioneering work by Larson and co-workers [7, 9], purely elastic instabilities were also found in other shear flows with curvilinear streamlines. They included the flow between a rotating cone and a plate and the Taylor–Dean flow [10, 11]. The original theoretical analysis of Larson *et al* [9] was refined, and more elaborate experiments were carried out. Some new mechanisms of flow instability driven by non-linear elastic stresses were suggested for cone-and-plate and for Taylor–Dean flows. The original mechanism proposed in [7] was verified experimentally in [12].

Some flow patterns observed above the purely elastic instability threshold in the curvilinear flows had rather disordered appearance and exhibited chaotic spectra of velocities. So, it was reasonable to suggest that, under certain conditions, a truly turbulent flow might be excited by

elastic stresses at vanishingly small *Re*. This idea was explicitly stated in [13], where the analogy between elastic and inertial flow transitions was discussed. In fact, irregular flow patterns and growth of flow resistance in elastic polymer solutions at low *Re* were observed even before the purely elastic instabilities were identified [14]. Such flow phenomena were also sometimes called 'elastic turbulence'. In all such cases, however, the term 'turbulence' was used in a rather loose fashion, without being given a proper definition. More importantly, no quantitative data on either flow velocity field or spatial and temporal velocity spectra in those irregular flows were ever presented.

Although the notion of turbulence is widely used in scientific and technical literature, there is no unique commonly accepted definition of it. Therefore, turbulent flow is usually identified by its main features. Turbulence implies fluid motion in a broad range of temporal and spatial scales, so that many degrees of freedom are excited in the system. There are no characteristic scales of time and space in the flow, except for those restricting the excited temporal and spatial domains from above and below. Turbulent flow is also usually accompanied by a significant increase in momentum and mass transfer. That is, flow resistance and rate of mixing in a turbulent flow become much higher than they would be in an imaginary laminar flow with the same Re.

In a recent publication [15] we showed how the first three of these features of turbulence appeared in the flow of a highly elastic polymer solution at low *Re*. The experiments were performed in a swirling flow between two plates with a wide gap, and the phenomenon was called elastic turbulence. In the present paper, we give a more complete account of our experiments on the elasticity-driven turbulent flow. We present additional velocity spectra, and distributions of probabilities of flow velocities and their derivatives. We also show that the elastic turbulence leads to quite efficient mixing in the flow. Furthermore, we present results of our experiments on Couette–Taylor (CT) flow and on an open flow in a curvilinear channel (Dean flow), where we also observed the elasticity-driven turbulence.

The paper is organized as follows. In the next section we discuss some practical problems concerning the experiment and the polymer solution. In sections 3, 4 and 5 the results on the swirling flow between two plates, the CT flow and the flow in the curvilinear channel, respectively, are presented. The results are briefly summarized and discussed in section 6.

2. Experimental considerations

As was suggested in [13], where the CT flow was discussed, there is some analogy between flow transitions driven by elasticity and inertia. So, the inertially driven Taylor instability occurs at constant Taylor number [1, 2], $Ta = (d/R)Re^2$, while the elastic instability is controlled by a parameter $K = (\eta_p/\eta)(d/R)Wi^2$ [9, 12], where R is the radius of the inner cylinder, d the width of the gap between the cylinders in the CT set-up, η the total solution viscosity, and η_p the polymer contribution to the viscosity, $\eta_p \equiv \eta - \eta_s$. The Weissenberg number is defined here as $Wi = \lambda \Omega R/d$, where Ω is the angular velocity of the rotating inner cylinder. (It was referred to as the Deborah number in some previous studies [9, 12, 13].) The Weissenberg number, which reflects the non-linearity of the constitutive equations (3) and (4), appears to be analogous to Re, which expresses non-linearity of the equation of motion. The geometric parameter determining curvature, the gap ratio, d/R, enters the expressions for both Ta and K. (There is also a specific term η_p/η , however, which shows the polymer contribution to the solution viscosity and naturally appears only in the expression for the parameter K for the elastic instability.) Scales of time and velocity for the purely elastic flow transition are given by λ and d/λ , respectively. As was shown in [13] they are analogous to t_{vd} and d/t_{vd} , which define scales of time and velocity for the inertially driven flow transitions. Here t_{vd} is the viscous diffusion time defined as $t_{vd} = d^2/v$.

Nevertheless, along with all those analogies, there are still some important differences between flow transitions driven by inertia and elasticity. Hence, it is an experimental fact that *any* laminar flow of a Newtonian fluid becomes unstable at sufficiently high *Re*, and all high-*Re* flows are turbulent. It includes rectilinear shear flows, such as Poiseille flow in a circular pipe and plane Couette flow, which are supposed to be linearly stable at any *Re*. In contrast to it, purely elastic flow instabilities in shear flows have been observed so far only in systems with curvilinear stream lines. All these instabilities are supposed to be driven by the hoop stress, which originates from the normal stress differences. Elastic instabilities also occur in flows of polymer solutions with open surfaces and in extensional flows through contractions in channels [10, 16]. A particular feature of the latter flows is that the rate of strain changes along the flow lines, so that even the basic flow states are not stationary in the Langrangian sense. Instabilities in those flows are not discussed in this paper.

This difference between the inertial and elastic instabilities may originate, of course, from the distinct governing equations. There are, however, some purely practical factors that can explain rather well the lack of observations of purely elastic flow transitions in rectilinear shear flows. Inertial instabilities in rectilinear shear flows of Newtonian fluids occur at quite high *Res*. Those are typically about two orders of magnitude higher than *Re* at which curvilinear shear flows with large gap ratios, d/R, become unstable. One may suggest, *a priori*, that rectilinear and curvilinear shear flows would have a similar relation between *Wi* at thresholds of the purely elastic flow instabilities as well. The problem is, however, that while it is rather easy to generate high-*Re* flows with low viscosity Newtonian fluids, it is usually impossible to reach the corresponding high values of *Wi* in shear flows of elastic polymer solutions. That is, there always exist rather severe practical limitations restricting non-linearity in elastic polymer stresses in shear flows. Their molecular mechanisms have been recently elucidated in a seminal paper by Chu's group [17].

Polymer molecules have finite extensibility, and their relaxation time decreases when they get stretched in a shear flow. This thinning of the relaxation time at high *Wi* is usually quite a strong and well-recognized effect. In fact, different variations of the basic Oldroyd-B model have been specially developed to take into account the shear thinning [3]. Significant decrease of λ with shear rate, $\dot{\gamma}$, renders growth of the Weissenberg number, $Wi = \lambda(\dot{\gamma})\dot{\gamma}$, much slower than linear in $\dot{\gamma}$. Substantial stretching of the polymer molecules in the primary shear flow also reduces their ability for further extension and susceptibility of the elastic stresses to flow perturbations, which is necessary for the generation of the flow instabilities and secondary vortex flows [9]. Finally, high shear rates cause mechanical degradation of the polymer molecules. It leads to permanent reduction of elasticity during experimental runs and decay of λ that can be very fast at high *Wi*. For all these reasons, it was very difficult or even impossible to observe elastic instabilities, when expected values of *Wi* at the instability threshold were high. This was the case in curvilinear flows with small gap ratios, d/R [18], and small viscosity ratios η_p/η [12].

Therefore, in order to maximize the non-linear elastic effects and to get a better opportunity to observe the elastic turbulence, we had to choose experimental conditions quite carefully [15].

First, it was important to obtain an elastic instability at a suitably low critical Weissenberg number, Wi_c . For that purpose, the gap ratio and the viscosity ratio had to be correspondingly large. Therefore, we used a polymer solution with a rather large η_p/η of about 1/4. (Further increase of the polymer concentration and η_p/η was not very efficient, and would also complicate the solution rheology, including large shear thinning of the solution viscosity.) To have a large gap ratio with the possibility of variation, we carried out our experiments in a swirling flow between two plates. The ratios d/R which we used were 0.263 and 0.526. In fact, historically we first tested a CT flow with d/R = 1/2, but growth of flow resistance in it was significantly smaller than in the swirling flow set-up, which we studied later (see sections 3 and 4).

Secondly, an appropriate polymer sample for the solution had to be chosen, which would not suffer major mechanical degradation under experimental flow conditions. Furthermore, the limit for extensibility of the polymer molecules had to be as high as possible compared with their typical conformations at the instability threshold. We used polyacrylamide (PAAm) with a large average weight $M_w = 18\,000\,000$ dalton and a broad molecular weight distribution (and low concentration of polyacrylic acid monomers) supplied by Polysciences (Warrington, USA). This commercial polymer sample proved to be remarkably stable with respect to mechanical degradation, which allowed us to reach high values of Wi and to explore strongly non-linear flow regimes. The high molecular weight of the polymers resulted in a large characteristic relaxation time, λ , even with solvents of moderate viscosity and a small characteristic stress, $\tau_0 = \eta_s / \lambda$. One can learn from the equations (3) and (4) that τ_0 sets a scale of the polymer stress, at which its non-linearity becomes significant. Therefore, the value of τ_0 together with η_p/η and d/R determine the polymer stress τ_p in the primary shear flow at the instability threshold. It is rather natural to suggest that when polymer molecules transduce less stress, they are also less subjected to mechanical degradation. Furthermore, molecular interpretation of the Weissenberg number in a shear flow is the degree of deformation of polymer molecules from their relaxed random coil conformations. So, Wi = 1 can be regarded as a characteristic value at which extension of polymer molecules becomes considerably larger than the size of relaxed coils. By the same token, extension of a polymer molecule at Wi_c (the elastic instability threshold) is supposed to be a fixed multiple of the relaxed coil size (in the former, linear approximation valid at moderate Wi_c [17]). High molecular weight and flexibility of a polymer suggests a large number of Kuhn segments [3, 19] in the polymer chain, and a high ratio between its contour length (size, when fully extended) and the size (radius of gyration, R_g) of a relaxed coil. (For a polymer molecule in a good solvent, with $R_g \sim M_w^{3/5}$, this ratio should increase as $M_w^{2/5}$ [19]. We would like to point out here that addition of 1% of NaCl to the solution significantly reduced R_g .) Therefore, using a higher M_w polymer we increase the 'reserve' of extensibility starting from the typical conformation at Wi_c . That opens a way to reach flow dynamics above the elastic instability threshold.

3. Swirling flow between two plates

3.1. Experimental set-up and procedure

The experimental apparatus is schematically shown in figure 1. The polymer solution was held in a stationary cylindrical cup with a flat bottom (lower plate). A co-axial rotating upper plate was just touching the surface of the fluid. The cup was mounted on top of a commercial rheometer, AR-1000 of TA-instruments. The upper plate was attached to the shaft of the rheometer, which



Figure 1. Schematic drawing of the experimental set-up. The set-up consists of a stationary cylindrical cup with a flat bottom (the lower plate), which is concentric with the rotating upper plate. A special cover is put above to minimize evaporation of the liquid.

allowed precise control of its rotation velocity, Ω , and measurements of the torque, T. The average shear stress at the upper plate, τ_w , was calculated using the equation $T = \tau_w \int r \, dS$, that gave $\tau_w \equiv 3T/(2\pi R^3)$. (The integration is over the upper plate surface.)

Sidewalls of the cup were machined from a single piece of perspex, which was optically clear. The cup was circular from the inside and square from the outside in horizontal crosssection. That allowed measurements of flow velocity in the horizontal plane by a laser Doppler velocimeter (LDV) with two crossing frequency-shifted beams. By appropriate positioning and orientation of the beam crossing region, azimuthal (longitudinal) and radial (spanwise) velocity components, V_{ϕ} and V_r , respectively, could be measured at different r and z. Here (r, ϕ, z) are cylindrical co-ordinates. The bottom of the cup was machined of stainless steel and the temperature was stabilized at 12 °C by circulation of water below the bottom plate.

A slightly modified version of the set-up was designed to photograph the flow from below and to observe mixing in the flow. A special cup of the same shape but with a transparent bottom (lower plate, made of perspex) was attached to the rheometer concentrically with the shaft but above the rheometer base, and a mirror tilted by 45° was placed under the cup, as is schematically shown in figure 1. The mirror was used both to illuminate the fluid by diffuse light and to relay images of the flow to a CCD camera. The camera was equipped with a regular video lens and mounted horizontally near the rheometer (figure 1). The source of the diffuse light was an illuminated white screen around the camera. The images were digitized by an 8 bit 512×512 frame grabber. In order to provide thermal stabilization, in this case the whole rheometer was placed in a thermally isolated box with through flow of temperature controlled air.

In the basic standard set-up, the radii of the upper plate and of the cup were R = 38 mm and $R_2 = 43.6$ mm, respectively, and the distance between the plates was d = 10 mm. The configuration was similar to the devices with rotating flow between two plates used in experiments on purely elastic instability [6, 11, 18, 20]. Its gap ratio, d/R, was significantly higher, though. In order to study dependence of flow conditions on the size of the system, two smaller set-ups, a half-size and a quarter-size, with all the dimensions reduced by factors of two and four, respectively, were used. Whenever the dimensions of an experimental set-up are non-standard, it is stated explicitly in the text.

We prepared a stock solution of PAAm ($M_w = 18\,000\,000$ dalton by Polysciences) and used it for the entire course of the experiments. First we dissolved 0.9 g of PAAm powder and 3 g of NaCl in 275 ml of deionized water by gentle shaking. (Addition of NaCl was necessary to improve solubility of PAAm.) Next the solution was mixed for 3 h in a commercial mixer with a propeller at a moderate speed. The idea behind the procedure was to cause mechanical degradation of PAAm molecules with the highest weights, and to 'cut' a high M tail of the broad molecular weight distribution of the PAAm sample. In a solution with a broad distribution of polymer molecular weights, the heaviest molecules, that are most vulnerable to mechanical degradation, may also make a major contribution to the solution's elasticity. A possible negative effect of that is a significant degradation of elasticity during experimental runs, and inconsistency in the results of the experiments. We found empirically that the procedure of pre-degradation in the mixer leads to substantial reduction of the degradation during the experiments and to substantial improvement of their consistency [21]. Finally, 9 g of isopropanol was added to the solution (to preserve it from ageing) and water was added up to 300 g. The final concentrations of PAAm, NaCl and isopropanol in the stock solution were 3000 ppm, 1% and 3%, respectively.

As a viscous Newtonian solvent for PAAm, we used a solution of 65% sugar (saccharose) and 1% NaCl in water. The salt was added to fix the ionic contents. Viscosity and relaxation time were measured with the same AR-1000 rheometer in a temperature-controlled narrowgap Mooney–Ewart geometry. The solvent viscosity was $\eta_s = 0.324$ Pa s at 12 °C. The PAAm concentration was 80 ppm, see figure 2. Solution viscosity, $\eta(\dot{\gamma})$, was carefully measured in a broad range of shear rates, $\dot{\gamma}$ (figure 2). Viscosity, η , slowly decreased with $\dot{\gamma}$, so that its shear thinning was about 7% per decade of $\dot{\gamma}$. At a shear rate of $\dot{\gamma} = 1 \text{ s}^{-1}$, corresponding to the onset of a purely elastic instability in the standard configuration (see below), η was 0.424 Pa s, and the viscosity ratio was $\eta_p/\eta = 0.235$. Polymer relaxation time, λ , was measured in oscillatory tests with a shear rate amplitude of 1 s^{-1} . Components of the apparent viscosity of the solution in-phase and out-of-phase with the applied shear, η' and η'' , respectively, were measured in long series of oscillations in a range of angular frequencies, ω . (Individual series were typically repeated about 10 times, and average viscosity values were calculated.) The same procedure was applied then to the solvent, and its viscosity components, η'_s and η''_s , were calculated as well. (The latter was nearly zero and it was an important test of virtual absence of inertial effects and of general applicability of the procedure.) The values for the polymer in-phase and out-of-phase



Figure 2. Upper curve: dependence of the apparent viscosity, $\eta(\dot{\gamma})$, on shear rate for an 80 ppm PAAm solution in 65% sucrose and 1% NaCl in water at 12 °C (semi-logarithmic co-ordinates). Applied shear rate was gradually increased during the test. Lower curve shows a similar run for the pure solvent. *Inset:* apparent relaxation time, λ , as a function of angular frequency, ω , measured for shear rate oscillations with an amplitude of 1 s⁻¹.

viscosity were calculated then as $\eta'_p = \eta' - \eta'_s$ and $\eta''_p = \eta'' - \eta''_s$. The (frequency-dependent) relaxation time was calculated as $\lambda(\omega) = \tan^{-1}(\eta''_p/\eta'_p)/\omega$, and $\lambda(\omega)$ at $\omega \to 0$, estimated as 3.4 s, was chosen as a representative relaxation time, λ , see inset in figure 2. The diffusion coefficient for the saccharose molecules was about $D = 8.5 \times 10^{-7} \text{ cm}^2 \text{ s}^{-1}$ [22].

In a swirling flow between two plates shear rate is quite inhomogeneous over the fluid bulk, even when the flow is laminar. So, in the simplest case of a narrow gap, $d/R \ll 1$, when the shear rate is constant along the z-direction, it grows along the radius, from zero at r = 0to the maximal value of $\Omega R/d$ at r = R. Shear rate averaged over the surface of the upper plate (and over the fluid volume) is then $2\Omega R/(3d)$. In our case, when the gap is quite wide (d/R = 0.263 in the standard configuration) and the sidewalls are close to the upper plate, the situation is even more complicated. The shear rate becomes strongly non-homogeneous along z in a large region of space corresponding to r beyond R - d. So, measurements of the ratio T/Ω in a laminar flow of a viscous Newtonian liquid gave a value 1.68 times larger than a value calculated for homogeneous shear along z. (The latter suggestion is not realistic for the large gap ratio set-up, of course.) This discrepancy is obviously due to a high shear rate layer near the upper plate at large r. One can see that choice of a representative shear rate becomes somewhat arbitrary under these circumstances. We decided to consider the simple expression $\Omega R/d$ as a characteristic shear rate, and to define the Weissenberg number as $Wi = \lambda \Omega R/d$. The Reynolds number was defined as $Re = \Omega R d\rho/\eta$.

In order to evaluate growth of flow resistance due to elastic instability and irregular secondary flow in the system, the average shear stress near the upper plate, τ_w^{lam} , in an imaginary

laminar shear flow at the same Ω had to be estimated. The stress, τ_w^{lam} , would depend on an average shear rate, $\dot{\gamma}_{av}$, at the upper plate, and on viscosity of the polymer solution, η , at this shear rate. An appropriate expression for it is $\tau_w^{lam} = \eta(\dot{\gamma}_{av})\dot{\gamma}_{av}$. The average shear rate, $\dot{\gamma}_{av}$, was estimated from measurements of the ratio τ_w/η_0 in laminar flow of a Newtonian fluid with a large viscosity, η_0 , at low *Re*. The shear rate in the laminar flow, calculated as $\dot{\gamma}_{av} = \tau_w/\eta_0$, was proportional to Ω , being $\dot{\gamma}_{av} = 1.12\Omega R/d$ in the standard configuration, and it gave a properly weighted average over the surface of the upper plate.

A suitable parameter characterizing growth of the flow resistance is the ratio of the average stress at the upper plate, τ_w (defined as $\tau_w \equiv 3T/(2\pi R^3)$ again), to the stress τ_w^{lam} in the laminar flow at the same Ω . In Newtonian fluids this ratio generally increases with Re as the flow becomes increasingly irregular, and the magnitude of τ_w/τ_w^{lam} can be considered as a measure of strength of turbulence and of the turbulent resistance. In our standard set-up, when a low viscosity Newtonian fluid is used, τ_w becomes 30% higher than τ_w^{lam} at Re = 70. Therefore, Re = 70 can be regarded as a characteristic value at which inertial effects in the flow become considerable.

3.2. Observation of elastic turbulence

3.2.1. Flow resistance. The first indication of a strongly non-linear state in the swirling flow was significant growth of the flow resistance above the elastic instability threshold. Dependence of τ_w/τ_w^{lam} on the shear rate, $\dot{\gamma} = \Omega R/d$, is shown in figure 3. The shear rate was gradually increased in time, very slowly (by about 10% h⁻¹) in the transition region, $\dot{\gamma} = 0.8-1.1 \text{ s}^{-1}$, and faster below and above it. One can see a sharp transition in flow of the polymer solution (curve 1, black line), which occurs at $\dot{\gamma}$ of about 1 s⁻¹ (corresponding to Wi = 3.5), and it manifests in a significant increase in τ_w compared to the laminar flow. Re at the transition point is about 0.3, so that the inertial effects are quite negligible there. The transition has pronounced hysteresis (grey line, corresponding to slow reduction of $\dot{\gamma}$), which is rather typical for the purely elastic flow instability [12]. The ratio τ_w/τ_w^{lam} keeps growing with the shear rate and at the highest $\dot{\gamma}$ that has been reached the flow resistance is about 12 times larger than it would be in a laminar flow. In the same range of shear flow rates of the pure solvent, curve 3, the flow resistance is completely laminar and the ratio τ_w/τ_w^{lam} is unity within resolution of the rheometer (about 1%).

Growth of the resistance in the polymer solution flow becomes even larger, when the size of the gap is increased to 20 mm (figure 3, curve 2), and the gap ratio becomes 0.526. Then the ratio τ_w/τ_w^{lam} reaches a value of 19. For Newtonian fluids in the same flow geometry such a growth of the flow resistance is found at Re of about 2×10^4 . For flow in a circular pipe this value of τ_w/τ_w^{lam} is reached at $Re \simeq 10^5$, which is usually considered as a region of rather developed turbulence [1, 2].

Mechanical degradation of the polymers was quite small at shear rates below 1.5 and 1 s⁻¹, for d = 10 and 20 mm, respectively. The dependence of τ_w/τ_w^{lam} on $\dot{\gamma}$ is therefore reproducible in consecutive runs within about 1%. At higher shear rates, degradation effects became appreciable. Thus, to ensure consistency of the results, time of measurements at high shear rates had to be made as short as possible. So, for $\dot{\gamma}$ above 1.5 s^{-1} (curve 1), the shear rate was raised by about 7% min⁻¹ and the measurement time was not sufficient to average out fluctuations of τ_w produced by the flow. Irregular undulations in curves 1 and 2 in figure 3 at high $\dot{\gamma}$ are due to those fluctuations and the short time of averaging.



Figure 3. The ratio of the average stress at the upper plate, τ_w , measured in the flow, to the stress, τ_w^{lam} , in imaginary laminar shear flow with the same boundary conditions, as a function of the shear rate, $\dot{\gamma}$. The curves 1 and 2 are for the polymer solution flow with the gap d = 10 and 20 mm, respectively. The shear rate was gradually varied in time. Thin black lines represent increasing $\dot{\gamma}$; thick grey lines represent decreasing $\dot{\gamma}$. Curve 3 is for the pure solvent.

In spite of the short time of the experiments, the solution relaxation time, λ , typically decreased by up to 10% after runs of the kind shown by the curves 1 and 2. Nevertheless, a major part of the degradation occurred at the highest shear rates, above 5 s^{-1} and above 3.5 s^{-1} for the curves 1 and 2, respectively. Therefore, the results at lower shear rates can be regarded as quite consistent.

Resistance is an integral characteristic of a flow, and its fluctuations give some information about flow events on the scale of the whole system. To get information about characteristic frequencies of the flow on this integral scale, we measured spectra of fluctuations of angular velocity of the upper plate at constant applied torque *T*. (AR-1000 is a stress-controlled rheometer, so it was more feasible to apply a constant torque and monitor rotation velocity, than to do it the other way around.) Spectra of fluctuations of Ω at a few *T* corresponding to different average shear rates (all above the transition point, $\dot{\gamma} \simeq 1 \text{ s}^{-1}$) are presented in figure 4. Rms of fluctuations of the angular velocity, Ω_{rms} , were about 4% of its average values, $\bar{\Omega}$, for all *T*. Because of the low rotation rates and low oscillation frequencies, inertial effects associated with acceleration of the shaft of the rheometer and of the liquid were always quite small. $(I\dot{\Omega}_{rms}$ was less than 1% of $(T/\bar{\Omega})\Omega_{rms}$, where *I* is the moment of inertia of the system, and $\dot{\Omega}_{rms}$ is the rms of the time derivative of Ω .) So, the spectra in figure 4 should reflect time scales of fluctuations of viscous and elastic stresses in the flow.

As the average shear rate increases, the power of fluctuations, P, increases and characteristic frequencies become higher (figure 4). The general form of the spectra remains very much the same, however. There are no distinct peaks, except for instrumental peaks at f, which are multiples of the average frequencies of the upper plate rotation, $\overline{\Omega}/(2\pi)$. Furthermore, each spectrum has a broad region, spanning about a decade in frequencies, where dependence of P on



Figure 4. Power spectra of fluctuations of the angular velocity, Ω , of the upper plate at different applied torques, *T*. Curves 1–5 correspond to average shear rates of 1.25, 1.85, 2.7, 4 and 5.9 s⁻¹, respectively (all above the transition point $\dot{\gamma} \simeq 1 \text{ s}^{-1}$, see figure 3). The power, *P*, of fluctuations is fitted by a power law, $P \sim f^{-4.3}$, for $\dot{\gamma} = 4 \text{ s}^{-1}$ over about a decade in frequencies, *f*. Curve 6, taken with the pure solvent at a shear rate of 4 s^{-1} , shows instrumental noise.

f is close to a power law, $P \sim f^{-\alpha}$, with $\alpha \simeq 4.3$. (Some increase in the power of fluctuations at higher f is due to instrumental noise, and it was also measured in laminar flow of the pure solvent, curve 6 in figure 4.) The power-law dependence implies that flow events on the integral scale of the system occur in this whole region of frequencies. As was argued above, excitation of fluid motion in a broad continuous range of frequencies is one of the characteristic features of turbulence.

3.2.2. Temporal and spatial spectra. Time spectra of fluctuations of flow velocity in a point at different constant shear rates (all above the elastic instability threshold) are shown in figure 5. One component of the velocity in the horizontal plane was measured with LDV in the centre of the set-up, where its average value was zero. The shear rates, $\dot{\gamma}$, for curves 1–5 were the same as the average shear rates for curves 1–5, respectively, in figure 4. The spectra in figure 5 have the same general features as those in figure 4. Power of the fluctuations and their characteristic frequencies increase with $\dot{\gamma}$, but the spectra remain very similar in their appearance. In particular, the spectra do not have distinct peaks, and do have broad regions of frequencies, where the power of fluctuations decays according to a power law $P \sim f^{-\alpha}$. (Flattening of the curves at high f is due to instrumental noise; see also discussion of figure 16 below.) Again, for each spectrum, the power-law-decay region spans about an order of magnitude in f, which implies excitation of the fluid motion in the whole range of the corresponding temporal scales. One can see that characteristic frequencies of fluctuations in a point and at the integral scale (cf figures 4 and 5) are rather close. The exponent, α , is about 3.5, which is rather close to the



Figure 5. Power spectra of velocity fluctuations in the standard set-up at different shear rates, $\dot{\gamma}$. The fluid velocity was measured by LDV at the centre of the flow. Curves 1–5 correspond to $\dot{\gamma} = 1.25$, 1.85, 2.7, 4 and 5.9 s⁻¹, respectively (all above the transition point $\dot{\gamma} \simeq 1$, see figure 3). The power, P, of fluctuations is fitted by a power law, $P \sim f^{-3.5}$, for $\dot{\gamma} = 4 \text{ s}^{-1}$ over about a decade in frequencies, f.

exponent of 4.3 in figure 4, and much larger than the Kolmogorov exponent of 5/3, found for velocity spectra of high *Re* inertial turbulence [1, 2].

A few representative snapshots of the flow viewed from below are shown in figure 6. In order to visualize the flow, the fluid was seeded with light reflecting flakes (1% of Kalliroscope liquid). The upper plate was black. So, the bright regions are those where the flakes are oriented parallel to the upper plate. The patterns of the polymer solution flow above the transition, at Wi = 6.5 (figures 6(a) and 6(b)) and at Wi = 13 (figures 6(c)-6(e)), look quite irregular and exhibit structures of different sizes. One can see, however, that the structures tend to have a spiral-like form, which is probably imposed by the average azimuthal flow and circular symmetry of the set-up. Furthermore, there is a dark spot in the middle, which appears in most of the snapshots. It corresponds to the centre of a big persistent toroidal vortex, which has dimensions of the whole set-up (see also below). Direction of the vortical motion was inwards near the upper plate, downwards near the centre and outwards near the lower plate. Average flow velocity along the radial direction was measured by LDV in a few points, and the results agreed quite well with the presence of the big persistent toroidal vortex. Flow of the pure solvent at the same shear rate was completely laminar as can be seen from the snapshot in figure 6(f). It looks quite uniform up to some variation along the radial direction due to varying shear rate and inhomogeneity along the circumference due to a somewhat uneven (and directed) illumination.

The visual impression of spatial disorder in the flow at high Wi is confirmed by a more careful analysis. Figure 7 shows average Fourier spectra of the brightness profiles at Wi = 13 along the diameter and along the circumference at r = 2d. The spectra were averaged over long series of flow pattern snapshots taken at consecutive moments of time. The wavelength is measured in units of d, so that the wave number, k, of unity corresponds to a length of $2\pi d$. The amplitude of fluctuations of the brightness along the circumference does not have any



Figure 6. Representative snapshots of the flow taken from below. The field of view corresponds to the upper plate area. The flow was visualized by seeding the fluid with light reflecting flakes. (a, b) Polymer solution at Wi = 6.5, Re = 0.35; (c–e) polymer solution at Wi = 13, Re = 0.7; (f) pure solvent at Re = 1.

peak and it exhibits a power-law decay, $A \sim k^{-1}$, over almost a decade in the wave-number domain. This implies that there are no selected wave numbers in the azimuthal direction and the fluid motion is excited in a broad range of spatial scales. (The plateau at high k may be due to restricted spatial resolution of the method of visualization that was used.)

The spectrum taken along the diameter shows a general tendency of power-law decay in even broader range of wave numbers. However, it apparently differs from the azimuthal spectrum by a series of broad peaks, which may be due to the persistent toroidal vortex discussed above. The average flow shear rate and the local gap ratio, d/r, vary along the radius, so that the radial direction is not neutral. Thus, the flow is not structureless and homogeneous along the radial direction. The visualization of the flow with the light reflecting flakes does not provide direct information about the fluid velocity field. Therefore, the specific value of the exponent in the power-law fit in figure 7 does not have any special meaning.

In order to obtain quantitative information about the spatial structure of the fluctuating velocity field, we explored velocity spectra in various off-centre points, with non-zero average azimuthal velocity, \bar{V}_{ϕ} . Spectra of fluctuations of the radial component of velocity, V_r , at $\dot{\gamma} = 4 \text{ s}^{-1}$ (Wi = 13.5) at four different radii are shown in figure 8. All of them were measured at z = d/2, where z is the distance from the upper plate. Rms of the fluctuations in all four points were rather close, varying between 0.88 and 0.99 mm s⁻¹. The spectra in figure 8 bear the same general features of turbulence as the spectra in figures 4, 5 and 7 discussed above. They



Figure 7. Average Fourier spectra of the brightness profiles taken along the diameter (thin black line) and along the circumference at a radius of 2d (thick grey line). The averaging was made over about 20 min of the flow time and about 2000 snapshots.



Figure 8. Power spectra of fluctuations of radial velocity, V_r , at $\dot{\gamma} = 4 \text{ s}^{-1}$ measured at z/d = 0.5 at different radii. Curves 1–4 correspond to r = 0, d/2, 2d and 3d, respectively. The average flow velocities (V_{ϕ}, V_r) (in mm s⁻¹) were (0, 0), (0.13, 0.19), (3.81, 1.17), (6.99, 0.89) for the curves 1–4, respectively.

do not exhibit distinct peaks, and have broad regions of f, where the power of fluctuations, P decreases according to $P \sim f^{-\alpha}$.

One can learn from figure 8 that as the point of measurement is moved away from the centre, characteristic frequencies of the fluctuations become higher. The most reasonable explanation for that will be growth of the average azimuthal velocity, \bar{V}_{ϕ} , which was 3.81 mm s⁻¹ and 6.99 mm s⁻¹, at r = 2d and 3d, respectively. So, the fluctuations of the velocity with time at these two points are mainly due to fluctuations in space, which are advected by the large mean flow velocity, \bar{V}_{ϕ} . Applying the Taylor hypothesis, we can view the spectra in time as spectra in space, with the relation between the frequency and the wave number given by $k = 2\pi f/\bar{V}_{\phi}$. Then the power-law-decay regions in curves 3 and 4 imply that the fluid motion is excited in the whole corresponding ranges of k. The ranges of the spatial scales, where the motion is excited, span about an order of magnitude for both the curves. The exponents α in the power laws $P \sim f^{-\alpha}$ (and, so, $P \sim k^{-\alpha}$) are again quite large, about 3.6 for r = 2d and about 3.3 for r = 3d. These large values of α imply that the power of fluctuations decays very quickly as the size of eddies decreases. The main contribution to fluctuations decays very quickly and velocity gradients (the power of the latter should scale as $k^{-(\alpha-2)}$) should therefore be due to the largest eddies.

3.2.3. Mixing in the flow. Mixing in the flow was observed using a droplet of black ink added to the working fluid before rotation of the upper plate was started (figure 9). Using a micropipette, the droplet was carefully placed near the centre (at r = 0) at about a half of the fluid depth. The ink was dissolved in a concentrated sugar syrup, to match the density of the droplet with the density of the working fluid. We used the half-size set-up with $R_1 = 19$ mm, $R_2 = 21.9$ mm and d = 5 mm. The volume of the working fluid was about 9 ml, and the amount of the ink was 50 μ l. The upper plate was coloured white and the fluid was viewed from below. So, regions without the ink appeared bright, whereas dark regions corresponded to positions in the (r, ϕ) plane, where the total amount of the ink across the fluid layer was large.

Consecutive stages of mixing in the polymer solution are shown in figure 9. Rotation of the upper plate was abruptly started at t = 0 at a rate of 1 turn per 4.26 s, which corresponded to a shear rate $\dot{\gamma} = 5.6 \,\mathrm{s}^{-1}$. It took about 1 min for the irregular flow to develop, after the rotation of the upper plate was started. (Development of the irregular flow was judged by growth of the flow resistance, which was saturated after about 1 min, cf [23].) So, no significant changes in the ink distribution occurred during the first 15 s (figure 9). After 30 s, the ink spread over the surface of the lower plate by the big toroidal vortex discussed above. In the snapshots taken at later moments in time (60, 90 and 120 s in figure 9) one can see many fine scale structures. Those can be either due to excitation of the fluid motion at small spatial scales or significant stretching of fluid elements along their Lagrangian trajectories by randomly fluctuating large-scale eddies. The contrast of the patterns gradually decreases with time, which indicates progress in mixing. The pattern in the last snapshot, taken 8 min after the flow has been started, looks completely homogeneous. From the appearance of the mixing patterns in figure 9, characteristic time of mixing can be estimated as 120s, corresponding to about 30 full turns of the upper plate. The time required for mixing by molecular diffusion without macroscopic flow can be estimated as dR/D. Substituting for D the diffusion coefficient for the saccharose molecules, $D = 8.5 \times 10^{-7} \,\mathrm{cm}^2 \,\mathrm{s}^{-1}$, we obtain a diffusion time of about 10⁶ s, which is four orders of magnitude larger than the mixing time in the flow. (Diffusion time for suspended solid particles of the black ink could be significantly larger.)



Figure 9. Snapshots of consecutive stages of mixing of a droplet of ink in the polymer solution in the half-size set-up, view from below. The area of the photographs corresponds to the area of the white upper plate. Rotation of the upper plate at $\Omega = 1.47 \text{ s}^{-1}(\dot{\gamma} = 5.6 \text{ s}^{-1})$ was suddenly started at t = 0.

Mixing in the flow of the pure solvent at the same shear rate is shown for comparison in figure 10. One can see that distribution of the ink remained inhomogeneous even after 9 h, although the ink became somewhat spread with the time. *Re* was about 0.5, and there were some non-vanishing inertial effects in the flow. In particular, centrifugal force gave rise to a slow fluid motion out of the $r-\phi$ plane. This motion had the form of a big toroidal vortex with outflow near the upper plate and inflow near the lower plate. (So, this vortex spun in a direction opposite to that of the toroidal vortex produced by the elastic stresses.) The vortical motion stirred the fluid but it did not really mix it. So, when the set-up was viewed from a side, one could see that the ink did not get to the toroidal core of the vortex.

Summarizing the experimental results, we conclude that the flow of the elastic polymer solution at sufficiently high *Wi* has, indeed, all the main features of developed turbulence that





Figure 10. Snapshots of consecutive stages of mixing of a droplet of ink in the pure solvent in the half-size set-up, view from below. Rotation of the upper plate at $\Omega = 1.47 \text{ s}^{-1}$ was suddenly started at t = 0.

were stated above. The fluid motion is excited in broad ranges of frequencies and wave numbers, both spanning about an order of magnitude. The flow is accompanied by a dramatic increase in the rate of transfer of momentum and mass. By the strength of the turbulent resistance and by the span of scales in space and time, where the fluid motion is excited, the observed flow can be compared to the turbulence of a Newtonian fluid in a pipe at Re of about 10^5 . This apparent turbulent fluid motion in the swirling flow between two plates arises at very low Re, where inertial effects are negligible, solely because of the non-linear mechanical properties of the elastic polymer solution. We therefore call the phenomenon *elastic turbulence*. Distinctions between the elastic turbulence and the usual inertial turbulence, which is observed in Newtonian fluids at high Re, are discussed in section 3.4 below.

3.3. Further properties of the flow

3.3.1. Flow structure and velocity profiles. To learn more about the velocity field generated by elastic turbulence, we measured the average velocity and rms of the velocity fluctuations at different points. Profiles in the z-direction of the average azimuthal velocity, \bar{V} , and the rms of its fluctuations, V_{rms} , at different flow conditions are shown in figure 11. (It was the standard set-up.) The measurements were taken at r = 2d, which is rather far from the edge of the upper plate (R - r = 1.8d). Profile of \bar{V} in a low *Re* flow of the pure solvent is almost a straight line



Figure 11. Curves 1–3: average azimuthal velocity, \bar{V} along the *y*-axis on the left; and curve 4: rms of fluctuations of the azimuthal velocity, V_{rms} along the *y*-axis on the right, as functions of the distance, *z*, from the upper plate. The measurements were taken at r = 2d. The average velocities are divided by the upper plate velocity at r = 2d. Curve 1: polymer solution at $\dot{\gamma} = 2.7 \text{ s}^{-1}$; curves 2, 4: polymer solution at $\dot{\gamma} = 4 \text{ s}^{-1}$ (see figures 3–5 and 8); curve 3: pure solvent at $\dot{\gamma} = 4 \text{ s}^{-1}$, $Re \simeq 1.2$. The rms of velocity fluctuations in the polymer solution (curve 4) is multiplied by λ/d to make it dimensionless.

(figure 11, curve 3). The elasticity-driven turbulent flow significantly changes the distribution of \bar{V} . It produces a high-shear-rate layer near the upper plate (curve 1), and a low-shear-rate region near the middle of the gap (at z/d = 0.5). Such a distribution of \bar{V} is reminiscent of average velocity profiles in usual high *Re* turbulence. The perturbation of the \bar{V} profile becomes stronger when $\dot{\gamma}$ is increased (curve 2). In particular, the slope of the \bar{V} -curve at small z/d becomes larger, which obviously corresponds to growth of τ_w/τ_w^{lam} with $\dot{\gamma}$ (figure 3).

Fluctuations of the azimuthal velocity (figure 11, curve 4) are small near the upper plate, reach a maximum at $z/d \simeq 0.25$, and start to decrease at larger z. Again, such a distribution of V_{rms} along z is reminiscent of velocity fluctuations in turbulent flows of Newtonian fluids [1, 2]. Rms of fluctuations reaches a value of about $0.5d/\lambda$, so that the rate of deformations produced by the fluctuating velocity field is of the order of $1/\lambda$. That implies an essentially non-linear relation between the rate of deformations and the fluctuating elastic stress. (This is discussed in more detail in the next section.) The maximal rms of the velocity fluctuations at $\dot{\gamma} = 4 \, \text{s}^{-1}$ was about 1.55 mm s⁻¹, it is about 7.5% of the upper plate velocity (V_{max}) at r = 2d and about 25% of the local value of \bar{V} .

As one can learn from curves 1 and 2 in figure 11, there is significant asymmetry in the distribution of the average shear rate, $\partial \bar{V}/\partial z$, along z. The main decrease in \bar{V} occurs near the upper plate. So, at $\dot{\gamma} = 4 \text{ s}^{-1}$, \bar{V}/V_{max} becomes 0.5 at z/d = 0.12 and at z/d = 0.5the average velocity is only $0.18V_{max}$. (The velocity profile near the lower plate could not be measured because of instrumental restrictions.) The uneven distribution of the shear rate along the z-direction should create an essentially inhomogeneous profile of N_1 along z. Indeed, in a laminar shear flow, we have $N_1 = 2\eta_p \lambda \dot{\gamma}^2 = 2\tau_p \lambda \dot{\gamma}$ (where τ_p is the polymer contribution to the shear stress). Therefore, the high shear rate near the upper plate should generate large N_1 and large hoop stress there. A result of this large unbalanced hoop stress would be inward motion of the fluid near the upper plate in the radial direction, which is quite consistent with our observation of the permanent toroidal vortex discussed above (see figure 6). In fact, the shape of the $\bar{V}(z)$ profiles in figure 11, which is strongly asymmetric with respect to the mid-plane (z = d/2), is also consistent with the picture of a *single* large-scale toroidal vortex. Indeed, even a possible increase in the slope of the $\bar{V}(z)$ profile near the lower plate would not match the high shear at small z, and the vortical motion in the *rz*-plane should be mainly driven by the high hoop stress near the upper plate. In contrast, a symmetric $\bar{V}(z)$ profile would imply equally high hoop stress near the upper and the lower plate (and low hoop stress in the middle) that would produce a pattern of two toroidal vortices on top of each other.

In fact, the turbulent $\overline{V}(z)$ -profiles of the kind shown by curves 1 and 2 in figure 11 were not always stable, especially at larger gap ratios. So, in a system with d/R = 0.563 at $\dot{\gamma} = 2.5 \text{ s}^{-1}$ the high shear rate layer randomly moved between the vicinity of the upper plate and the vicinity of the lower plate with a typical residence time of a few tens of minutes and a transition time of about a minute. These fluctuations in the location of the high shear rate layer were accompanied by inversions of the spinning direction of the big toroidal vortex. When the main drop of \overline{V} occurred near the lower plate, it was at its vicinity where the radial flow was directed inwards. Therefore, the asymmetry of the $\overline{V}(z)$ profile with respect to the z = d/2 plane (figure 11) is indeed directly related to the large-scale vortical flow.

Big toroidal vortices driven by the hoop stress are actually quite well known to appear in swirling flows of elastic fluids [3, 24], and inhomogeneity of the shear-rate profile in the primary laminar flow has long been recognized as their common origin. In our system, this vortex first arises as a stationary structure at low-shear rates. As can be seen in figure 3 (curve 1 at $\dot{\gamma} < 0.75 \,\mathrm{s}^{-1}$ and curve 2 at $\dot{\gamma} < 0.4 \,\mathrm{s}^{-1}$), the toroidal vortex leads to some increase in flow resistance even before the transition to the elastic turbulence. As suggested by the first three snapshots in figure 9, the big toroidal vortex is also the first vortical motion to arise upon sudden inception of a shear rate, which is above the elastic instability threshold level. (Further redistribution of the average azimuthal velocity due to development of the turbulent flow should enhance this large-scale vortical motion.) Hence, we can conclude that transition to the elastic turbulence in the swirling flow between two plates is mediated by this vortex. The vortex is always present in the flow, and it probably suffers permanent instability and is permanently fluctuating, which may be an important mechanism for the production of fluid motion at smaller scales. The toroidal vortex is providing a smooth, large-scale velocity field, which is randomly fluctuating in time, and in which the liquid and the stress tensor imbedded in it are chaotically advected. This type of advection can create variations of stress in a range of smaller scales, which may cause small-scale fluid motion [25, 26]. (This would be analogous to generation of small-scale concentration variations in chaotic mixing by large fluctuating vortices.)

The large-scale vortical flow is a probable reason for low Wi at the elastic instability threshold, and for the very strong increase of resistance above the transition to elastic turbulence in the flow between two plates. Indeed, growth of the flow resistance in the CT flow at comparable Wi (see figure 18) is significantly smaller. This situation is somewhat similar to inertial turbulence in Newtonian fluids in the same systems. In the swirling flow between two plates, a big toroidal vortex, which is driven by centrifugal force, causes transition to turbulence at quite low Re and



Figure 12. O, PDF of flow velocity measured at r = 2d, z = d/4 and $\dot{\gamma} = 4 \text{ s}^{-1}$. Solid lines represent fits by Gaussians with some skewness; (a) azimuthal velocity, V_{ϕ} and (b) radial velocity, V_r .

provides a route of efficient energy transfer to the turbulent flow. Many instructive pictures of the toroidal vortex, its evolution and instabilities in inertial, inertia–elastic, and elasticity-dominated flow regimes in a swirling flow set-up with larger gap ratios can be found in [24].

3.3.2. Velocity statistics in a point. Characteristic probability distribution functions (PDF) of azimuthal and radial components of the flow velocity in the regime of elastic turbulence are shown in figure 12. These distributions were obtained from LDV measurements taken at r = 2d, z = 0.25d at a shear rate of 4 s^{-1} . PDFs of the azimuthal and radial velocity can be fitted quite well by the functions $A_1 \exp[-(V-\bar{V}_{\phi})^2/(2a_1^2) + 0.032(V-\bar{V}_{\phi})^3/a_1^3]$ and $A_2 \exp[-(V-\bar{V}_r)^2/(2a_2^2) - 0.017(V-\bar{V}_r)^3/a_2^3]$, respectively. Here $\bar{V}_{\phi} = 6.34 \text{ mm s}^{-1}$ and $\bar{V}_r = -1.84 \text{ mm s}^{-1}$ are the average velocities; $a_1 = 1.55 \text{ mm s}^{-1}$ and $a_2 = 0.97 \text{ mm s}^{-1}$ are the rms of fluctuations for the azimuthal and radial components, respectively. One can see that both the distributions have only minor skewness and are very close to a Gaussian. (Skewness is probably due to the non-zero average velocity.)

Probability distribution functions for gradients of velocity (rates of deformation) in the longitudinal and transversal directions, obtained from the same LDV time series, are shown in figures 13(a) and 13(b), respectively. The rates of deformation are multiplied by the relaxation time, λ , to make them dimensionless. The velocity gradients were estimated by using the Taylor hypothesis, with smoothing over a distance of about 1.45 mm. The difference, ΔV , between consecutive velocity readings with even time intervals of $\Delta t = 0.22$ s was divided by Δt and by the average velocity $\bar{V} = (\bar{V}_{\phi}^2 + \bar{V}_r^2)^{1/2} = 6.6 \text{ mm s}^{-1}$. One can see that the distributions of $\partial V_{\phi}/r \,\partial \phi$ and $\partial V_r/r \,\partial \phi$ cannot be fitted by Gaussians. In contrast with the velocity distributions in figure 12, both the PDFs in figure 13 have well-pronounced exponential tails, which imply significant intermittency of the velocity gradients. The situation of nearly Gaussian statistics of velocities and essentially non-Gaussian, strongly intermittent distributions of velocity gradients is actually quite typical for high-*Re* inertial turbulence [27]. Hence, the elastic turbulence resembles high-*Re* inertial turbulent flows in this respect.



Figure 13. O, PDF of velocity gradients measured at r = 2d, z = d/4 and $\dot{\gamma} = 4 \text{ s}^{-1}$. The velocity gradients are made dimensionless by multiplication by the relaxation time, λ . Solid lines represent Gaussian fits; (a) longitudinal gradient, $\partial V_{\phi}/r \partial \phi$; (b) transversal gradient, $\partial V_r/r \partial \phi$.

3.4. Comparison with inertial turbulence at high Re

3.4.1. Role of the elastic stresses. Along with the apparent similarity in phenomenology between the elastic and inertial turbulence, there are also quite a few very important differences. An obvious reason being the difference in the physical mechanisms, which underlie the two kinds of turbulent motion. Turbulence in Newtonian fluids is observed at high Re, when the equation of motion (1) is highly non-linear. The non-linear term in equation (1) is connected with the kinematics of *translational motion* in a continuous medium and is due to fluid inertia. Elastic turbulence occurs at low Re, when non-linearity of the equation of motion (equation (2)) is negligible. The origin of the elastic turbulence is nonlinear dependence of the elastic stresses on the rate of deformation in flow. (This non-linear dependence is seen very well in the model constitutive relations (3) and (4).) The latter non-linearity is connected with kinematics of *deformation* in a continuous medium, and it is due to the finite relaxation time of polymer molecules. Therefore, one can suggest that, in the case of the elastic turbulence, it would be more relevant to study the field of stresses and rates of deformations rather than the velocity field.

The products of the randomly varying rates of deformation $(\partial V_{\phi}/r \partial \phi$ and $\partial V_r/r \partial \phi$ in figure 13) and λ can actually be viewed as different estimates for Weissenberg number, Wi', of the fluctuating velocity field. The distributions in figure 13 show that Wi' is rather often above unity, and the rms of its fluctuations estimated from figure 13(a) is about 0.83. Fluctuations of the rate of deformation are directly connected with fluctuations of the elastic stresses by the constitutive relation (see equations (3) and (4)). Since the fluctuating Weissenberg number is rather high, this connection should be essentially non-linear. Therefore, one can suggest that the field of τ_p in the elastic turbulence is highly complex and strongly intermittent. It would certainly be quite instructive to explore spatial structure and temporal distribution of the elastic stresses, but there is currently no technique for local measurements of τ_p in a turbulent flow. On the

other hand, large-scale properties of the τ_p field can be inferred from measurements of the total flow resistance.

As is well known, the high flow resistance in high Re inertial turbulence is due to large Reynolds stresses. The Reynolds stress tensor is defined by the average values of $\rho \langle V_i V_j \rangle$, where V_i and V_j are different components of flow velocity. In the case of elastic turbulence, Reynolds stresses are quite small, since Re is low. Indeed, $\langle V_i V_j \rangle$ can be estimated as $2V_{rms}^2$. Taking $V_{rms} = 1.55 \text{ mm s}^{-1}$, which is the maximal value for rms of the azimuthal velocity fluctuations at r = 2d and $\dot{\gamma} = 4 \text{ s}^{-1}$, we obtain a Reynolds stress of 6×10^{-3} Pa, while τ_w at the same conditions is about 16 Pa. Therefore, the whole increase of flow resistance in the elastic turbulence is caused by growth of average elastic stresses [15, 23]. By the same token, the irregular fluctuations of the total flow resistance in the elastic turbulence, which were discussed above (figure 4), are due to fluctuations in the τ_p field at large spatial scales.

Average growth of the polymer stresses can be estimated from the curves in figure 3. Indeed, by momentum conservation, the stress near the wall should be equal to its average value in the bulk, $\tau_w = \tau_p + \tau_s$, $\tau_w^{lam} = \tau_p^{lam} + \tau_s^{lam}$ (we omitted signs of averaging for brevity). The viscous stress due to the Newtonian solvent, τ_s , is linearly proportional to the shear rate, and its average value depends only on the boundary conditions and cannot change as a result of a turbulent flow [23]. Therefore, we can take $\tau_s = \tau_s^{lam} = (\eta_s/(\eta - \eta_s))\tau_p^{lam}$, where η is the viscosity of the polymer solution measured in a laminar shear flow at a corresponding shear rate. Then for the ratio of the average polymer stresses in turbulent and laminar flow we obtain a relation $\tau_p/\tau_p^{lam} = (\eta/(\eta - \eta_s))[(\tau_w/\tau_w^{lam}) - (\eta_s/\eta)]$. After plugging in numeric values from figure 3 (curve 2) and from measurements of $\eta(\dot{\gamma})$ (figure 2) we get ratios of τ_p/τ_p^{lam} as high as 65 at high shear rates. For a set-up with larger d/R and a solution with a smaller polymer concentration, the ratio of the polymer stresses in turbulent and laminar flows estimated this way reached a factor of 170 [23].

In molecular theory of polymer dynamics the tensor τ_p is found to be proportional to the polymer concentration and to the average tensor of inertia of the polymer molecules, $\tau_{p,ij} \sim k \langle R_i R_j \rangle$ (where k is an elasticity coefficient, which remains constant in a linear elasticity regime). Since elastic energy density of the stretched polymer molecules is given by a quadratic form of a similar kind, one can suggest linear dependence of the elastic energy on τ_p . Therefore, the density of the elastic energy should increase in the turbulent flow by about the same factor as τ_p . The density of kinetic energy is proportional to the Reynolds stress and although it certainly increases with development of the elastic turbulence, it always remains quite small compared with the elastic energy density. So, we can suggest that the elastic stress, τ_p , should be the object of primary importance and interest in the elasticity-driven turbulent flow, and it may be appropriate to view elastic turbulence as turbulence of the τ_p field.

The growth of the elastic stresses is also an evidence of significant extension of the polymer molecules in the flow. The relation $\tau_{p,ij} \sim k \langle R_i R_j \rangle$ suggests that polymer extension should grow as $\sqrt{\tau_p}$. Taken together with the data on τ_p / τ_p^{lam} , it implies an eight-fold stretching of the polymer molecules in the turbulent flow beyond their extension in a laminar flow with the same average shear rate. (For $\tau_p / \tau_p^{lam} = 170$, the corresponding extension factor is 13.) Significant extension of the flexible polymer molecules in the turbulent flow resistance, has been recently confirmed by direct visualization of individual polymer molecules in flow [28].

The behaviour of polymer molecules in chaotic and turbulent flows has been a subject of controversy for long [29], and it deserves special discussion. Dynamics of polymer liquids have

been most well studied in basic rheometric flows, such as simple shear and extension. In the former case, τ_p is usually found to increase slower than linearly with the rate of deformation $\dot{\gamma}$ (shear thinning). In the latter case, τ_p grows faster than linearly with the rate of deformation, $\dot{\varepsilon}$, and at $\dot{\varepsilon} > 1/(2\lambda)$, τ_p can grow by a few orders of magnitude as the total deformation (measured by the Henky strain, $\int \dot{\varepsilon} dt$) increases. The molecular basis for this behaviour, stretching and tumbling in the case of a shear flow, and a coil–stretch transition in the case of an extensional flow, has been clearly demonstrated in the recent experiments by Chu's group [17, 30]. A turbulent flow involves a complex three-dimensional velocity field, with the rate-of-deformation tensor randomly varying in time and space. Therefore, an attempt to fit it directly into the context of rheometric flows may lead to frustration and confusion.

Being stochastic phenomena, chaotic and turbulent flows are most appropriately treated statistically. An object of particular interest for polymer dynamics is statistics of deformations of the fluid elements, in which the polymer molecules reside. Since the size of the polymer molecules is almost always much smaller than the size of the smallest turbulent eddies, the velocity field is smooth on the scale of the molecules. At every moment in time it is defined by some average velocity and a tensor of rate of deformation, $V(\vec{r}) = V_0 + (\partial V_i / \partial r_i)(\vec{r} - \vec{r}_0)$. Statistics of deformations of a fluid element can be derived from statistics of the tensor $\partial V_i / \partial r_i$ in the Lagrangian reference frame moving with the fluid element. The tensor $\partial V_i / \partial r_i$ is threedimensional and in all non-degenerate cases it has a positive eigenvalue, corresponding to extensional flow along some axis. Rate of the extension and direction of the axis vary randomly at a time scale, connected with the characteristic time of variations in the flow. A detailed statistical analysis was first made by Lumley [31] in the early 1970s and was revised and refined recently [32, 33]. Under quite general assumptions about statistics of $\partial V_i / \partial r_i$, it was shown that a fluid element, which is initially a sphere, becomes an ellipsoid with its longest axis growing on average as $a = a_0 \exp(\alpha t)$. The coefficient α is a Lyapunov exponent of divergence of two closely spaced points in the flow, and the value of α is usually of the order of $\langle (\partial V_i / \partial r_i)^2 \rangle^{1/2}$. A somewhat surprising conclusion of the theory is that a generic random flow is on average an extensional flow in every point, with the rate of extension $\dot{\varepsilon} = \alpha$ and unlimited Henky strain. Therefore, one should expect the polymer molecules to become strongly stretched by the random (turbulent) flow, when the Lyapunov exponent α , as given by the statistics of $\partial V_i / \partial r_i$, becomes larger than $1/(2\lambda)$. (More refined theory suggests that a significant fraction of the molecules should become substantially stretched even at lower α due to intermittency in the statistics [32, 33].) One can see that the extension of the polymer molecules to the regime of elastic turbulence, taken together with the estimate $Wi' \simeq 0.8$, is in good agreement with the theoretical predictions [32, 33].

Based on the discussion above, one can suggest the following scenario of development of elastic turbulence. The polymer molecules are stretched in the primary shear flow that leads to large elastic stresses, τ_p . The elastic stresses render the primary shear flow unstable and cause an irregular secondary flow. The flow acts back on the polymer molecules, stretching them further, and raising the elastic stresses even more [23]. All this make the flow increasingly turbulent, until a kind of a saturated dynamical state is finally reached. This state implies some mutually consistent fields of average stresses and velocities and their fluctuations. Those fields are connected with each other by the equation of motion and the constitutive relation.

3.4.2. Scaling with set-up size and fluid viscosity. We explored variation of different characteristics of the elasticity-induced turbulent flow with viscosity of the polymer solution and

with size of the set-up. It was in sharp contradiction with the general principles of Newtonian fluid mechanics, but quite in line with the concept of viscoelastic similarity suggested for purely elastic flow transitions [13]. So, velocity required for excitation of inertial turbulence in a Newtonian fluid is proportional to viscosity of the fluid. This is connected with the fact that characteristic inertial time (viscous diffusion time), $t_{vd} = d^2/v$, is inversely proportional to viscosity. In contrast, polymer relaxation time, λ , usually grows proportionally to viscosity of the solvent [3]. When the system gap ratio and the viscosity ratio are both kept constant, purely elastic transitions should occur at a constant Weissenberg number, $Wi = V\lambda/L$. Therefore, increase in viscosity of the solvent should lead to transition to elastic turbulence at lower velocity. To verify whether the PAAm solutions were prepared with the same η_p/η and a relaxation time larger than 3.4 s, a few other sugar syrups with higher viscosities were used as solvents. The transition to elastic turbulence in these solutions occurred at the same Wi, indeed, while the angular velocity of the upper plate and Re at the transition threshold were lower, scaling as $\Omega \sim 1/\eta_s$ and $Re \sim 1/\eta_s^2$, respectively. (In fact the situation was very similar to the case of purely elastic instabilities in CT flow [13].) The dependence of τ_w/τ_w^{lam} on Wi for those solutions were close to those in curve 2 in figure 3. (In order to obtain a similar dependence, though, it was important to vary the shear rate slowly, in proportion with the increase in λ .) Using a very viscous sugar syrup with $\eta_s \simeq 7.2 \,\text{Pa}$ s, we prepared a polymer solution with a relaxation time of about 30 s, and observed a transition to elastic turbulence at $\Omega \approx 0.05 \, \text{s}^{-1}$ and $Re \approx 3 \times 10^{-3}$ [23].

Furthermore, the viscous diffusion time, t_{vd} , is proportional to the square of the size of the system. Therefore, when the size of the set-up is proportionally reduced, the linear and angular velocities required for excitation of inertial turbulence should increase as $V \sim 1/d$ and $\Omega \sim 1/d^2$, respectively. In contrast, the polymer relaxation time, λ , does not depend on system size. Therefore, Ω (and $\dot{\gamma}$) at transition to the elastic turbulence should not depend on the size of the set-up, as long as its proportions are preserved. A natural scale of stress in the purely elastic flow regime is given by η_p/λ , and it also does not depend on the size of the system. (It may depend on η_p/η_s , however.) Therefore transition to the elastic turbulence is expected to occur at the same characteristic stress, and dependence of τ_w/τ_w^{lam} on Wi is supposed to be the same for all system sizes.

Figure 14 shows dependence of τ_w/τ_w^{lam} on Wi in a set-up with all the dimensions proportionally reduced by a factor of four compared with the standard configuration. The plot looks quite similar to curve 1 in figure 3. One can see that the transition to the elastic turbulence indeed occurred at the same $\dot{\gamma}$ as in the standard set-up, although *Re* was 16 times lower. To compensate for lower sensitivity of the rheometer to shear stress due to smaller total torque, the rate of variation of $\dot{\gamma}$ was reduced. This may be the reason for a smaller region of hysteresis in figure 14 compared with figure 3. The plot in figure 14 also implies that the transition to elastic turbulence occurred at the same τ_w and Wi, and dependence of τ_w on $\dot{\gamma}$ was also almost the same as in the standard configuration.

The parameters of the system relevant to the elastic turbulence are d, R, η_p , η_s , λ and Ω . (Density of the fluid should be of no importance, since inertial effects are negligible.) There seems to be no apparent way to construct, out of these parameters, a scale of length other than the size of the system (with a possible pre-factor depending on η_p/η_s and $\Omega\lambda$). It implies that, when the dimensions of the system, d and R, are proportionally changed, while η_p , η_s , λ and Ω are preserved, the turbulent velocity field should remain the same up to linear rescaling with size of the system.



Figure 14. Dependence of the upper plate stress ratio, τ_w/τ_w^{lam} (actual polymer solution flow and a laminar flow with the same boundary conditions, cf figure 3) on the shear rate for a set-up with dimensions reduced four-fold compared with the standard configuration of the flow between two plates. Black and grey curves correspond to increasing and decreasing shear rate respectively.

This suggestion is qualitatively confirmed by the mixing patterns presented in figure 15. The snapshots in the left column are the same as in figure 9. They show consecutive stages of mixing of a droplet of ink in a polymer solution upon sudden inception of shear flow. The dimensions of the system were twice as small as those of the standard configuration. The snapshots in the right column show mixing in the same polymer solution in a system with dimensions reduced four-fold compared with the standard configuration. In both cases the area of the snapshots corresponds to the area of the upper plates. So, the snapshots in the right column are magnified by a factor of two compared with the snapshots in the left column. (The volume of the ink droplet added to the solution in the right column was four times smaller than for the left column in order to match the amount of ink per unit area.) The angular velocities of the upper plates were the same, $\Omega = 1.5 \text{ s}^{-1}$, and so were the shear rates, $\dot{\gamma} = 5.6 \text{ s}^{-1}$, and the Weissenberg numbers. However, *Re* for the right column was four times smaller, *Re* = 0.125 compared with *Re* = 0.5 for the left column.

The mixing patterns in figure 15 are irregular in both columns, but the two sequences look rather similar and make an impression of two realizations of the same stochastic process. So the transition to the elastic turbulence seems to go similarly in the two systems, and the mixing seems to progress with about the same characteristic time. This implies that characteristic velocities of the irregular flow scale linearly with the size of the system and with velocity of the primary flow. Patterns, which appear in the snapshots taken at equal elapsed times, seem to have similar sizes and similar contrast levels. This suggests that, in elastic turbulence, characteristic spatial scales of velocity field are indeed proportional to the size of the system.

Quantitative data on the dependence of the turbulent velocity field on the system size are obtained from velocity spectra measured at a point with a large mean flow velocity. As it



Figure 15. Mixing patterns in polymer solution flow in set-ups with flow between two plates and with dimensions reduced two-fold (left column) and four-fold (right column) compared with the standard configuration. Photographs in consecutive rows were taken at equal times elapsed from sudden inception of the upper plate rotation at $\Omega = 1.47 \text{ s}^{-1}$ ($\dot{\gamma} = 5.6 \text{ s}^{-1}$). Images in the right column are magnified by a factor of two to match the sizes. The area of the photographs corresponds to the area of the white upper plate.

was argued above, those measurements provide information about both the power of the velocity fluctuations and spatial structure of the velocity field. Spectra of fluctuations of the azimuthal and radial velocity components measured at r = 2d, z = d/4 in the standard and in the half-size set-ups are shown in figure 16. The shear rate was 4 s^{-1} (corresponding to Wi = 13.5) in both the systems, the same as for the spectra in figure 8 and for PDFs in figures 12 and 13.

From the above similarity arguments, fluctuating velocity was expected to scale as d/λ . Rms of the fluctuations of V_{ϕ} and V_r were 1.55 and 0.97 mm s⁻¹, respectively, for the standard configuration, and 0.78 and 0.48 mm s⁻¹, respectively, for the half-size system. So the



Figure 16. Power, *P*, of velocity fluctuations in set-ups with flow between two plates as a function of frequency, *f*. Curves 1 and 3 are for the radial velocity component, V_r , and curves 2 and 4 are for the azimuthal velocity, V_{ϕ} . Black curves (1 and 2) are for the standard set-up, and grey curves (3 and 4) are for a set-up with two-fold reduced dimensions.

corresponding fluctuation amplitudes in the half-size system were indeed halved. Therefore, to match spectra in the two systems, the velocity measured in the half-size system was multiplied by a factor of 2. One can see that, after this velocity rescaling, the spectra of both V_{ϕ} and V_r become very close. The apparent difference at high frequency is due to lower signal-to-noise ratio in the half-size system. Indeed, one of the main sources of noise in the LDV measurements is finite size of the crossing region of the laser beams, over which the measured velocity is averaged. Characteristic velocity gradients (scaling with Ω and $1/\lambda$) are expected to be equal in the two systems, and dimensions of the laser beam crossing region are the same as well. Therefore, the level of the gradient noise should not change when the size of the system is reduced, whereas the level of the signal (characteristic fluctuating velocity) decreases.

The average azimuthal velocities were 6.34 and 3.37 mm s⁻¹ for the standard and the halfsize systems, respectively, with their ratio being rather close to 2 again. Therefore, applying the Taylor hypothesis to the power spectra in figure 16, we conclude that spatial structure of the velocity fluctuations in the two systems is statistically identical up to rescaling by a factor of 2 in the *k*-space. Thus, the experimental results in figure 16 provide further, more quantitative support to the suggestion of simple linear rescaling of the turbulent velocity field with size of the system at constant Ω (and η_p , η_s and λ).

3.4.3. Flow at small scales and near walls. The structure of flow at small scales and near boundaries gives another example of substantial differences between the elastic and inertial turbulence. It is instructive to summarize some relevant basic facts about the latter first. In high-*Re* inertial turbulence, the range of excited spatial scales is restricted from below by the Kolmogorov dissipation length, l_0 . It can be estimated from the condition $Re_{l_0} \simeq 1$ [1, 27], with

the local Reynolds number, Re_l , defined as $V_l l/\nu$, where V_l is a characteristic turbulent velocity difference at the scale *l*. Since V_l decreases as *l* becomes smaller (as $V_l \sim l^{1/3}$ in Kolmogorov's picture of turbulence [1, 27]), Re_l decreases faster than linearly with decreasing *l*. So, the dissipation length l_0 necessarily exists in any inertial turbulent flow.

An issue related to the problem of the smallest excited length scale is the stability of boundary layers near solid walls. The latter can be most simply approached by introduction of a local Reynolds number, $Re(z) = \overline{V}(z)z/\nu$, which is a growing function of distance, z, from the wall. (The usual convention is to use y for distance from a wall (transverse direction), and z as a co-ordinate in the spanwise direction, but we would like to stay consistent with the notation used in figure 11 and cylindrical geometry.) Near a wall the mean flow velocity, $\overline{V}(z)$, increases linearly with z, giving $Re(z) \sim z^2$ at small z. Therefore, Re(z) remains low in some vicinity of a wall, and the flow is a laminar shear. This laminar shear layer near a wall (usually called the viscous sublayer) ends at a distance z (decreasing with growth of global Re), where Re(z) becomes sufficiently large, and the shear flow becomes unstable and turbulent.

Analogously to the inertial turbulence, we can introduce a local Weissenberg number, $Wi_l = \lambda V_l/l$. It does not decrease as *l* becomes smaller, though. Indeed, the ratio V_l/l is a characteristic gradient of the turbulent velocity smoothed over the length *l*, and it can only grow as *l* decreases. Since Wi_l does not become small at small *l*, there appears to be no straightforward way to construct an analogue of the dissipation length, l_0 , for the elastic turbulence. So, there may be no macroscopic length setting a lower limit to the range of scales of the turbulent motion. (Something may change, though, at a scale comparable with size of the polymer molecules or with their characteristic diffusion length.)

This suggestion is consistent with the plots shown in figure 16. Indeed, the spectra of both V_{ϕ} and V_r in figure 16 have broad frequency ranges where the power of fluctuations decays as $P \sim f^{-\alpha}$. The value of α is close to the values found for the other velocity spectra (figures 5 and 8). The highest frequency in a power-law-decay region corresponds to the smallest spatial (or temporal) scale, at which the turbulent flow is excited. The power-law-decay regions of the spectra in figure 16 are cut off at high frequencies by plateaux rather than by abrupt drops, however. As we argued before, those plateaux appear solely due to the instrumental noise. The power-law-decay range and the apparent range of excited spatial scales deduced from it become broader, when the signal-to-noise ratio becomes higher (figure 16). However, this lack of a cut-off frequency from above does not lead to any kind of high frequency divergence (ultraviolet catastrophe). Indeed, since $\alpha > 3$, the total power of the velocity gradients with their power scaling as $P \sim k^{-(\alpha-2)}$ in the spatial domain. (We assume again that the Taylor hypothesis is applicable.)

Similarly to Re(z), one can introduce a local Weissenberg number near a wall, $Wi(z) = \lambda \overline{V}(z)/z$ ($\overline{V}(z)$ here is the difference between the local mean flow velocity and the velocity at the wall at z = 0), and easily see that it does not become small as $z \to 0$. In fact, we can learn from the average velocity profiles in the elastic turbulence (figure 11) that the region near the upper plate corresponds to the highest average shear rate and to the highest Wi(z). On the other hand, the velocity fluctuations clearly decline near the upper plate (figure 11) and existence of such a stable high-shear-rate boundary layer in the elastic turbulence may appear rather puzzling.

A possible mechanism providing stability of the flow near the upper plate is significant shear thinning of the relaxation time, λ . As was argued above, in the elastic regime, flow stability depends on the parameter $K = (\eta_p/\eta)(d/R)Wi^2$. Adopting it to the boundary layer, we can suggest that stability should be defined by a local parameter $K(z) = (\eta_p/\eta)(z/R)Wi^2(z)$, where $Wi(z) = \lambda(\dot{\gamma})\bar{V}(z)/z$, $\lambda(\dot{\gamma})$ is the shear-rate-dependent relaxation time, and *R* is the radial position of the region of interest. We can consider the case of a simple sigmoidal profile, where the whole growth of the mean flow velocity from zero to \bar{V} occurs across thin layers of widths z_b near the boundaries. If \bar{V} , η_p/η and *R* are all kept constant, we obtain $K(z_b) \sim \lambda^2(\dot{\gamma})/z_b$, where $\dot{\gamma} = \bar{V}/z_b$. Therefore, if the relaxation time decreases with the shear rate, faster than $\lambda \sim \dot{\gamma}^{-1/2}$, $K(z_b)$ is a decreasing function, and the boundary layer remains stable as it becomes thinner and the slope, \bar{V}/z_b , increases.

Such fast decays of λ with $\dot{\gamma}$ would also suggest that a more global (although radiusdependent) parameter for the flow between two plates $K = (\eta_p/\eta)(d/R)[\lambda(\Omega R/d)\Omega R/d]^2$ would be decreasing with *R*, and the elastic instability would be limited to a region of sufficiently small *R*. This kind of behaviour was indeed reported for a different polymer solution and a system with small d/R [20]. In a more general case of shear thinning, with the relaxation time scaling as $\lambda = \lambda_0 \dot{\gamma}^{-\alpha}$ (and $\alpha < 1/2$), the condition $K < K_c$ (where K_c is a critical value for instability) is satisfied for $\bar{V}(z) < [(K_c \eta R)/(\eta_p \lambda_0^2)]^{1/(2-2\alpha)} z^{(1-2\alpha)/(2-2\alpha)}$, which is reminiscent of the $\bar{V}(z)$ profiles in figure 11.

Dependence of λ on $\dot{\gamma}$ in a shear flow can be inferred from measurements of normal stress difference, N_1 . At $\lambda \dot{\gamma} \ll 1$ it is usually found that $N_1 = 2\lambda \eta_p \dot{\gamma}^2$ with constant λ and η_p . Therefore, $\lambda(\dot{\gamma}) = N_1/(2\eta_p \dot{\gamma}^2)$ can be used as an estimate of shear thinned relaxation time at higher $\dot{\gamma}$. Measuring this way $\lambda(\dot{\gamma})$ for solutions of a few lower molecular weight PAAm samples we usually found $\lambda \sim \dot{\gamma}^{-\alpha}$, with α around 0.3 [12]. (Resolution of our rheometer was not sufficient to measure normal stresses in the solution used in the experiments on the elastic turbulence at *Wi* in the range of interest [12].) Therefore the shear thinning may indeed play an important role in the stabilization of the boundary layer.

Finally, it is worth noting that the whole concept of boundary layers is less straightforward in the case of elastic turbulence, and the condition of their stability may be less restrictive than for inertial turbulence. Indeed, in a high *Re* turbulent flow, characteristic fluctuating velocity, V'(z), generated by a flow instability originating at a distance z from a wall scales as $V'(z) \sim v/z$, and the boundary layer must become stable at some z, since $V'(z) \rightarrow \infty$ as $z \rightarrow 0$. In contrast, characteristic fluctuating velocity generated by an elastic instability should be proportional to z/λ and approach zero at $z \rightarrow 0$.

4. Couette-Taylor flow

4.1. Experimental set-up and results

Experiments were performed to study the Couette–Taylor flow in a small set-up mounted vertically on the AR-1000 rheometer. The stationary outer cylinder was made of glass, had a radius $R_2 = 13.5$ mm, was open from above and closed from below by a stainless steel plug with a flat top, figure 17. The rotating inner cylinder had a radius $R_1 = 9$ mm and was conical at the bottom with an angle of $\tan^{-1}(0.5)$, so that the gap ratio in the cone-and-plate flow at the bottom was the same as in the annular gap, $d/R_1 \equiv (R_2 - R_1)/R_1 = 0.5$. The cylindrical surface of the inner cylinder was 72 mm long. Both the cylinders were coaxial with the shaft of the rheometer and the inner cylinder was attached to it. The working liquid was filled

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Figure 17. Schematic drawing of the Couette–Taylor set-up, mounted on top of the AR-1000 rheometer. A circular cap with a hole for the rotating shaft, which covered the glass cylinder from above, is not shown.

above the upper rim of the inner cylinder. In order to reduce evaporation of the fluid from its open surface, a special cover (vapour trap) was used. The outer cylinder was enclosed in a thermal jacket with a square cross-section that was made out of plexiglass. The temperature of the system was set by a fast stream of water pumped through the jacket from a refrigerated circulating bath. The polymer solution was the same as in the experiments on the flow between two plates (80 ppm of PAAm in 65% sucrose, 1% NaCl in water), and the experiments were carried out at the same temperature of 12 °C. The characteristic shear rate in the flow was calculated using $\dot{\gamma} = \Omega R_1/d$, and the Weissenberg number was taken to be $Wi = \lambda \Omega R_1/d$ with $\lambda = 3.4$ s.

First, we measured flow resistance in the polymer solution as a function of $\dot{\gamma}$ and Wi. The protocol was very similar to that described above for the flow between two plates. At the beginning, pure solvent (with a viscosity η_s) was filled into the system, and the torque, T_s , applied to the inner cylinder was measured as a function of angular velocity, Ω . The dependence was linear over the whole range of Ω explored. The suggested dependence of torque for an (imaginary) laminar flow of the polymer solution was calculated from it as $T_{lam} = T_s \eta(\dot{\gamma})/\eta_s$. Then the system was filled with the polymer solution, Ω was gradually increased starting from a low point corresponding to Wi = 0.68, and the torque, T, required to drive the actual polymer solution flow was measured. The ratio of the two torques, T/T_{lam} , is equal to the ratio of the average stresses, τ/τ_{lam} , in the actual and the imaginary laminar flow of the polymer solution. It is a quantitative expression of growth of the flow resistance compared with the laminar flow. The dependence of τ/τ_{lam} on Wi is shown in figure 18 (thin black line) for Wi varying over almost two orders of magnitude. One can see that the resistance ratio τ/τ_{lam} is equal to unity at low values of Wi corresponding to laminar flow in the system. Then at Wi of about 4 an abrupt transition occurs, and the flow resistance starts growing quickly and non-linearly. Re at the transition point is quite small, $Re = \Omega R_1 d\rho/\eta = 0.07$. Therefore, the transition is of a purely elastic nature. The non-linear growth of the flow resistance is quite significant; at the



Figure 18. The ratio of average shear stress in the actual flow of the polymer solution and the stress corresponding to a laminar flow with the same $\dot{\gamma}$, τ/τ_{lam} , as a function of the Weissenberg number, $Wi = \lambda R_1 \Omega/d$. Thin black line: increasing rotation rate, starting from Wi = 0.7; thick grey line: decreasing rotation rate, starting from Wi = 7.

highest tested value of Wi, the ratio τ/τ_{lam} is more than 4. Undulations of the curve in figure 18 reflect the fluctuating nature of the flow resistance above the transition and are due to limited integration time during the measurements.

To explore the behaviour of the flow resistance at decreasing shear rate, we carried out another experiment, where Wi was first set to 7, well above the transition point, and gradually reduced afterwards. Dependence of τ/τ_{lam} on Wi in this run is shown by a thick grey curve in figure 18. The two curves in figure 18 overlap at Wi above 4.7 and below 1.8, whereas in the region in between, a well-pronounced hysteresis occurs. This broad region of hysteresis agrees quite well with previously reported results for Couette–Taylor flow in a purely elastic regime, which were obtained in a system with a smaller gap ratio and a different PAAm solution [12, 13]. The plot in figure 18 is quite similar to the dependence of τ/τ_{lam} on Wi for the flow between two plates shown in figures 3 and 14.

The non-linear velocity transition was accompanied by the emergence of a disordered and randomly fluctuating pattern of vortices on top of the basic shear flow. When visualized with the light reflecting Kalliroscope flakes, the fluctuating pattern looked similar to the Disordered oscillations reported before at similar flow conditions in a larger CT system with $d/R_1 = 0.2$ [12]. To measure the velocity associated with the fluctuating flow we used the LDV set-up again. Two horizontal laser beams were crossing inside the polymer solution seeded with polystyrene beads in the middle of the annular gap at $r = (R_1 + R_2)/2$ and at about half height from the bottom. The beams were adjusted in such a way that it was the radial component of the flow velocity, V_r , perpendicular to the basic shear flow that was measured. Power spectra of fluctuations of V_r at a few different Wi above the non-linear transition are shown in figure 19. As Wi is raised, the total power of the velocity fluctuations increases and fluctuations at higher frequencies are excited. Nevertheless, all four curves share the same



Figure 19. Power of fluctuations of the radial component of the flow velocity, V_r , measured at $r = (R_1 + R_2)/2$, as a function of frequency, f. Curves 1–4 correspond to Wi = 8.5, 13.6, 21.8 and 35, respectively. Two power-law fits for curve 4 in different regions are shown to guide the eye.

general appearance: there are two plateau regions at low and high f (the latter is probably be due to the instrumental noise) and two contiguous regions of a power-law decay in between, with two different exponents and a distinct inflection point separating them. Although characteristic frequencies increase with Wi, those exponents are rather close for all four curves, around -1.1at lower frequencies and around -2.2 in the high-frequency power-law-decay regions. Apart from the inflection points, none of the curves exhibits any distinct peak. Amplitude of the velocity fluctuations in all four cases was quite small compared with average flow velocity. The latter was directed along the circumference (azimuthally) and is equal to about $\Omega R_1/2$. (See also figure 20 and discussion thereof.) Therefore, the spectra in figure 19 should actually reflect spatial variation of V_r in the azimuthal direction.

In order to obtain additional information about the fluctuating flow velocity, we used the same time series to calculate velocity statistics. The PDF of V_r at different Wi are shown in figure 20. One immediately observes that, although the average velocity is always zero, the shapes of the distributions are essentially non-Gaussian. Moreover, unlike the velocity distributions for the flow between two plates in figure 12, PDFs in figure 20 cannot be satisfactory fitted by skewed Gaussians (functions of the form $A \exp[-(V - \bar{V}_r)^2/(2a^2) + b(V - \bar{V}_r)^3/a^3]$). One of the most notable features of the PDFs in figure 20 is the relatively high probability for large negative values of V_r ('long tails' of the distributions at negative V_r), which implies asymmetry for radial flows inwards and outwards. Although most striking in the curve for Wi = 5.4, this feature is still quite well expressed at Wi = 35. In fact, the curve at Wi = 5.4 is also quite similar to a PDF of V_r reported before in the CT flow with $d/R_1 = 0.2$ (see figure 12 in [12]), at comparable Wi in the regime of disordered oscillations. In the latter case, the asymmetry in the radial velocity distribution was due to an abundance of coherent structures having a shape of couples of vortices ('diwhirls') with a narrow region of intensive radial flow



Figure 20. Probability distribution functions of the radial component of the flow velocity, V_r , measured at $r = (R_1 + R_2)/2$, at different *Wi*. Curves 1–4 correspond to Wi = 5.4, 13.6, 21.8 and 35, respectively.

inwards in the middle and slow radial flow directed outwards on the sides of a vortex couple. Such an asymmetric shape of the diwhirls is closely related to the mechanism of generation of the elastic instability in CT flow, and it allows efficient pumping of energy from the primary shear flow into the secondary fluctuating velocity field [12]. Therefore, the asymmetric shapes of PDFs in figure 20 suggest that the coherent structures of that diwhirl type are present in the CT flow even at high *Wi* and may have significant influence on its organization and statistical properties.

The distribution functions in figure 20 also allow us to find characteristic rates of extension in the flow in the radial direction, $\partial V_r/\partial r$, and Weissenberg numbers, $Wi' = \lambda \partial V_r/\partial r$, associated with them. Indeed, we have got boundary conditions of $V_r = 0$ and $\partial V_r/\partial r = 0$ at $r = R_1$ and $r = R_2$. Therefore, $\partial V_r/\partial r$ in the bulk can be estimated as $4V_r^{rms}/d$, where V_r^{rms} is the rms of the radial velocity fluctuations. The resulting values of $\partial V_r/\partial r$ calculated from PDFs in figure 20 are then 0.059, 0.071, 0.097 and 0.15 s⁻¹ at *Wi* of 5.4, 13.6, 21.8 and 35, respectively. Those values of $\partial V_r/\partial r$ correspond to *Wi'* of 0.2, 0.24, 0.33 and 0.52, respectively. These estimated numbers are rather high and are generally consistent with the picture of strong elongation of the polymer molecules by elements of extensional flow in the fluctuating velocity field [28], [31]–[33].

4.2. Discussion and comparison with the flow between two plates

Summarizing the results shown in figures 18–20, we conclude that, in the purely elastic regime, the CT flow behaves similarly to the flow between two plates described in section 3. The purely elastic non-linear flow transition occurs at Wi = O(1) and (arbitrarily) small *Re* in both cases; the transition is rather sharp, has pronounced hysteresis and it is accompanied by a

significant growth of flow resistance. The fluctuating velocity field above the transition is quite random for both the flows and its spectra do not exhibit any distinct peaks, cf figures 8 and 19. Therefore, although characterized less extensively, the CT flow above the non-linear transition fits quite well the definition of elastic turbulence introduced above.

Nevertheless, the elasticity-induced turbulent flow in the CT system differed from the turbulent flow between two plates quantitatively, and the turbulent flow effects in the CT system were less dramatic. Indeed, the non-linear transition in the CT system occurred at $Wi \approx 4$ (figure 18), which is higher than critical Wi in the flow between two plates, 3.2 and 2, for d/R = 0.26 and 0.52, respectively (figure 3). The growth of flow resistance in the CT flow above the transition was notably smaller than in the flow between two plates—a factor of 4.2 (figure 18) versus factors of 11 and 19 (figure 3) at the highest Wi explored. (However, the ratio τ/τ_{lam} in the CT flow kept growing with Wi, and it was the increasingly fast mechanical degradation of the polymers that prevented us from raising Wi higher, figure 18.) Characteristic amplitudes of fluctuating velocity in the CT flow (figure 19) were significantly smaller than in the flow between two plates (figures 11 and 12), both in terms of d/λ (0.06 versus 0.5 at $Wi \simeq 13$) and of the maximal azimuthal flow velocity (0.005 versus 0.075 at $Wi \simeq 13$). This comparison is certainly not quite fair, since V_r in the CT flow is a transversal velocity component. Hence, it should be best compared with V_z in the flow between two plates, which was not measured. However, the general tendency appears to be rather clear.

Characteristic estimated rate of extension due to the fluctuating velocity field in the CT flow was also considerably smaller than in the flow between two plates, $0.24/\lambda$ versus $0.83/\lambda$ at $Wi \simeq 13$. (We note here again, however, that these estimates were obtained from different types of data sets, figure 20 versus figure 13.) Nevertheless, even the relatively low value of the rate of extension in the CT flow is reasonably consistent with the general condition for it to be of the order of $1/\lambda$ to cause significant elongation of the polymer molecules, growth of the elastic stresses and notable flow resistance increase.

As was already suggested above, more favourable conditions for the development of the elastic turbulence and stronger turbulent effects in the flow between two plates may be due to the large toroidal vortex. This vortex, which is driven by the distribution of normal stresses in the basic shear flow, may provide a major channel for pumping of energy into the turbulent velocity field. CT flow certainly has a more limiting geometry, without any coherent three-dimensional large-scale flow of comparable magnitude and with fluid motion confined to the annular gap. The latter confinement limits possible curvature of flow trajectories and may also make the general requirement of axial symmetry for excited vortex structures (as dictated by the system geometry) more severe and restrictive, somewhat impeding development of the turbulence. In fact, axial symmetry of the vortex patterns is a well-pronounced feature in inertial turbulence in CT flow even at very high Re [34].

Among all the discrepancies between the elastic turbulence in the CT system and in the flow between two plates, the principal one appears to be the difference in the decay exponents in the power spectra of velocity fluctuations (cf figures 5, 8 and 16 versus figure 19). In the case of the flow between two plates the decay exponents were around -3.5 in both spatial and temporal domains, while the power spectra in the CT flow had two distinct power-law-decay regions with significantly lower exponents of -1.1 and -2.2. As shown in section 5 below (figure 25), velocity spectra in an elastic turbulent flow in a curvilinear channel have regions of power-law decay as well. The decay exponents are about -3.3, suggesting that it is the CT flow that behaves 'abnormally'.

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Figure 21. Schematic drawing of the curvilinear channel showing the inlet, a region of observation and the outlet.

A clue to the behaviour of the CT flow may be the inflection points of the spectra in figure 19, which appear at frequencies $f_c = (0.45 \pm 0.05)\Omega/(2\pi)$ at all four *Wi*. These frequencies are close to average rates of rotation (in revolutions per second) of the fluid in the measurement point. Together with the non-Gaussian and asymmetric shapes of PDFs in figure 20, that brings us to a suggestion that the power spectra in figure 19 are significantly influenced by the axisymmetric coherent structures (possibly of the diwhirl type) discussed above. Those coherent structures may dominate the low frequency part of the spectra, $f < f_c$, and also feed into the high frequency part, $f > f_c$ (where the 'original' power of fluctuations should be rather low to begin with; see figures 8 and 16), reducing the apparent absolute values of the decay exponents for $f > f_c$.

5. Flow in a curvilinear channel

5.1. Experimental set-up and procedure

The main motivation for the experiments on flow in a curvilinear channel (Dean flow) was to carry out detailed quantitative study of mixing in elastic turbulence [35]. It is an open flow that allows extended continuous experimental runs with reproducible and well-controlled initial conditions, and easy gathering of extensive data at different stages of mixing. The channel is schematically shown in figure 21. It had a uniform depth d = 3 mm, was machined in a transparent bar of perspex, and it was sealed from above by a transparent window. The channel consisted of a sequence of smoothly connected half-rings with the inner and outer radii $R_1 = 3$ mm and $R_2 = 6$ mm, respectively; it was square in its cross-section, and had quite a high gap ratio, $d/R_1 = 1$. The high gap ratio was intended to facilitate development of an elastic instability at low *Wi* and of intensive irregular flow above the instability threshold. The channel had 30 repeating units, 18 mm in length each, see figure 21.

The liquids to be mixed were fed into the channel by two identical syringe pumps through two separate tubing lines, always at equal discharge rates. Chemical composition of the two liquids was always identical as well, with the only difference of a small concentration, $c_0 = 2$ ppm, of a fluorescent dye (fluorescein) added to one of them. They were prepared from the same stock of a carefully filtered liquid, which was divided into two equal parts. A small amount of a concentrated solution of the dye was added to one part, while the other part was diluted by an equal amount of pure water. (Each of the liquids was carefully mixed afterwards.) This method of preparation provided very good matching of densities and refraction indices of the liquids.





Figure 22. Photographs of the flow taken with the laser sheet visualization (figure 21) at different N. The field of view is $3.07 \text{ mm} \times 2.06 \text{ mm}$, and corresponds to the region shown in figure 21 (rotated 90° counterclockwise). Bright regions correspond to high concentration of the fluorescent dye. (a) Flow of the pure solvent at N = 29; (b–d) flow of the polymer solution at Wi = 6.7 and at N = 8, 29, 54, respectively.

The channel was illuminated from a side by an argon-ion laser beam converted by two cylindrical lenses to a broad sheet of light with a thickness of about 40 μ m in the region of observation. It produced a thin cut in the three-dimensional mixing pattern, parallel to the top and bottom of the channel at a half of the channel depth.

Fluorescent light emitted by the liquid in the direction perpendicular to the beam and to the channel plane was projected on to a CCD array by a camera lens and digitized by an 8-bit, 512×512 frame grabber. Using homogeneous solutions with different amounts of the dye, we found intensity of the fluorescent light captured by the camera to be linearly proportional to the dye concentration. Therefore, concentration of the dye was evaluated from the fluorescent light intensity. (The liquid without the dye appeared completely dark (figure 22) and signal from it was below the noise level of the camera.)

The flow was always observed near the middle of a half-ring on the side from which the laser beam was coming. So, the number, N, of a unit (figure 21) counted from the inlet was a natural linear co-ordinate along the channel. Since the total rate of liquid discharge, Q, was constant, N was also proportional to the average time of mixing. In order to observe further stages of mixing (corresponding to N > 30), we carried out a series of experiments, where the liquids were pre-mixed before they entered the channel. A shorter channel of the same shape was designed for this purpose and put before the entrance to the original one. In multiple calibration experiments we found good matching between states of mixing at N = 2 with the pre-mixer and at N = 27 without it [35]. Therefore, for the experiments with the pre-mixing, a number of 25 was added to the actual co-ordinate along the channel to calculate the effective N.

Flow velocity was measured directly with the aid of the LDV set-up. Because of the small width of the channel, a special effort was made to obtain high spatial resolution by reduction of the region of space, where the two laser beams crossed, and reduction of distance between the interference fringes. Focusing lenses with a small focal length (about 25 mm) were used, and the angle between the beams was raised to about 90° in the air (and about 60° in the liquid). As a result, the region of the beam crossing was brought down to $15 \times 15 \times 40 \,\mu$ m, and the distance between the fringes was 0.44 μ m.

We used the same polymer solution as in the experiments on the flow between two plates and on the CT flow. (It was 80 ppm of PAAm, 65% sugar, 1% NaCl in water.) This time, however, the experiments were made at room temperature, 22.5 ± 0.5 °C. So, the solvent viscosity was $\eta_s = 0.153$ Pa s, and viscosity of the solution was $\eta = 0.198$ Pa s at a shear rate $\dot{\gamma} = 4 \text{ s}^{-1}$. The relaxation time, λ , estimated from phase shift between the stress and the shear rate in oscillatory tests with a shear rate amplitude of 3 s^{-1} , was 1.4 s. An estimate for the diffusion coefficient of the dye was given by that for the saccharose molecules, $D = 8.5 \times 10^{-7} \text{ cm}^2 \text{ s}^{-1}$. The characteristic shear rate and the Weissenberg number in the flow were estimated as $\dot{\gamma} = (2Q/d^2)/(d/2) = 4Q/d^3$ and $Wi = \lambda(4Q/d^3)$, respectively. *Re* was calculated to be $Re = 2Q\rho/(d\eta)$.

5.2. Results

Re in the flow was always quite small, reaching only 0.6 for the highest Q that we explored. Therefore, flow of the pure solvent always remained laminar and no mixing occurred (figure 22(a)). The boundary separating the liquid with and without the dye was smooth and parallel to the direction of the flow and only became somewhat smeared due to molecular diffusion as the liquid advanced downstream. Flow of the polymer solution was laminar and stationary up to a value of Q corresponding to $Wi_c = 3.2$ (and Re = 0.06), at which an elastic instability occurred. This instability led to irregular flow and fast mixing of the liquids.

A few typical mixing patterns at different N in the polymer solution above the instability threshold are shown in photographs in figures 22(b)-(d). More insight about the structure and evolution of the mixing patterns can be obtained from space-time diagrams. Representative diagrams taken at Wi = 6.7 at four different N are shown in figure 23. Brightness profiles along a single line going perpendicularly to the channel near the middle of a half-ring (a horizontal line going through the middle of a snapshot in figure 22) were captured even at time intervals of 80 ms and plotted from top to bottom. The diagrams in figure 23 share the same chaotic appearance and show features at comparable scales, but they lose contrast as the liquid advances downstream and gets progressively mixed.

As is illustrated by the space-time diagrams in figure 23, mixing in the polymer solution flow above the instability threshold was a random process calling for statistical analysis [35]. A simple parameter characterizing homogeneity of the mixture is an rms of deviations of the dye concentration from its average value, $\bar{c} = c_0/2$, divided by the average value itself,

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Figure 23. Representative space–time diagrams of the polymer solution flow at Wi = 6.7 taken at different positions, N, along the channel.

 $c_{rms} = \langle (c - \bar{c})^2 \rangle^{1/2} / \bar{c}$. A small value of c_{rms} means high homogeneity and good mixing of the liquids. At the channel entrance, where the two injected liquids are perfectly separated, c_{rms} is unity, and it should become zero for a perfectly mixed liquid.

Dependence of c_{rms} on Wi near the exit of the channel, at N = 29, is presented in figure 24. Statistics of the dye concentration was evaluated from space-time diagrams similar to those in figure 23. However, regions near the walls of the channel with widths of 0.1*d* were excluded from the statistics, because of possible image aberrations. In a stationary flow regime (*Wi* below 3.2), when the concentration profile did not change with time, the brightness profiles were measured over short time intervals (about 100 s). In the regime of an irregular flow, however, the profile of concentration was strongly fluctuating. So, in order to obtain representative statistics of *c*, the measurements of the brightness profile were taken during quite long intervals of time (about 20–30 min), which typically corresponded to the total liquid discharge of about $10^3 d^3$.

The plot in figure 24 is somewhat analogous to those in figures 3 and 18, which show the dependence of the flow resistance on Wi. Indeed, the decrease in c_{rms} is an integral result of mass transfer produced by the irregular flow in the channel, just as growth of the flow resistance is an integral characteristic of increase of momentum transfer in the elastic turbulence. At Wi below 2, dye distribution patterns in the polymer solution were similar to those in the solvent

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Figure 24. Dependence of c_{rms} (normalized rms of concentration deviations from the average) on the Weissenberg number, Wi, measured near the channel exit at N = 29 (semi-logarithmic co-ordinates).

(figure 22(a)) with c_{rms} close to unity. Some decrease in c_{rms} at low Wi is due to the large residence time and diffusional smearing of the boundary between the liquids.

At Wi of about 2, reduction of c_{rms} from 1 to about 0.9 occurred, which was the result of a transition in the flow. Although this transition produced rather complex spatial distribution of the dye (data not shown), the patterns were stationary and mixing was rather minor. This transition probably related to the onset of stationary vortices with vorticity directed along the mean flow, as it was predicted in [36]. Recently, those vortices have been observed directly using particle image velocimetry [37] (see also below). The most striking feature of the plot in figure 24 is certainly an abrupt drop in c_{rms} at $Wi_c = 3.2$, where the irregular motion of the liquid set in. It is worth noting here that since the mixing and reduction in c_{rms} is a cumulative effect of stirring and diffusion in the flow starting from the inlet, the actual value of c_{rms} above the transition is defined by the distance from the inlet. On the other hand, the onset of the fluctuating flow occurred rather consistently along the whole channel, and we expect Wi_c to be virtually independent of the channel length, when the latter is sufficiently large.

We studied the dependence of c_{rms} on N at Wi = 6.7, corresponding to highest homogeneity of the mixture near the channel exit (figure 24), and found c_{rms} to decay exponentially with the distance from the inlet with a characteristic decay length ΔN of about 15 segments [35]. One can learn from figure 24 that, if Wi is raised above 6.7, c_{rms} starts to increase again. The most plausible explanation for this is saturation of growth of the velocity fluctuations together with reduction of the residence time in the flow at growing Wi (and average flow velocity, $\bar{V} = Q/d^2$). If the ratio between fluctuating and average flow velocities remains constant, while they both increase, the stirring in the flow remains the same, but there is less time available for molecular diffusion, and homogeneity is reduced as a result of it. This situation can be qualitatively described by growth of the Peclet number, $Pe = \bar{V}d/D$. It was recently found for the flow of a polymer solution in a channel of the same shape and at similar Wi, that the characteristic length ΔN increases as $Pe^{0.25}$ [38]. This suggests that c_{rms} should start increasing with Wi once the growth of the velocity fluctuations is slowed down.



Figure 25. Power, *P*, of fluctuations of velocity in the middle of the channel at N = 12 as a function of frequency, *f*. The spectra in the polymer solution flow at Wi = 6.7 for the velocity components along and across the mean flow are shown by curves 1 and 2, respectively. Curve 3 shows velocity spectrum across the mean flow for the pure solvent at the same *Q*.

Typical time of mixing in the channel at Wi = 6.7 was found to be 3–4 orders of magnitude shorter than the diffusion time, d^2/D , for the small molecules of fluorescein [35]. Dependence of the efficiency of mixing at the optimal flow conditions (for the 80 ppm solution it was Wi = 6.7; figure 24) on the concentration of the polymers was surprisingly weak (although Wi_c grew fast, as the polymer concentration was decreasing). So, for a solution with the polymer concentration of 10 ppm ($\eta/\eta_s = 1.03$), c_{rms} of as low as 0.29 could be reached at N = 29. (It was measured at Re = 0.065, where inertial effects were still negligible.) Excitation of irregular flow and active mixing was observed down to a polymer concentration of 7 ppm.

Figure 25 shows power spectra of fluctuations of longitudinal and transversal components of the velocity in the polymer solution at Wi = 6.7. The measurements were done at N = 12, near the middle of the half-ring in the middle of the channel. A spectrum of the velocity fluctuations in the flow of the pure solvent at the same Q, giving just instrumental noise, is shown for comparison. The mean velocity was $\bar{V} = 6.6 \text{ mm s}^{-1}$; the rms velocity of fluctuations, V_{rms} , was $0.09\bar{V}$ and $0.04\bar{V}$ in the longitudinal and transversal directions, respectively. Measuring the transversal velocity component in a few off-centre positions, we found non-zero averages, which typically persisted for a few minutes and changed their sign rather randomly in time. This situation can be explained by the presence of persistent longitudinal vortical structures in the flow. These vortices should be filling the whole channel cross-section, with their vorticity direction randomly jumping between parallel and anti-parallel to the mean flow. Those vortices may first appear at Wi = 2 but remain stationary below $Wi_c = 3.2$.

To measure resistance as a function of the flow rate, we used a micro-fabricated channel with the same proportions but with a 30 times smaller segment size [39]. It had $d = 100 \,\mu\text{m}$, $R_1 = 100 \,\mu\text{m}$, $R_2 = 200 \,\mu\text{m}$ and consisted of 46 segments. The polymer solution had the



Figure 26. Ratio of a pressure drop across the channel in the polymer solution flow, ΔP , to a pressure drop for a laminar flow, ΔP_{lam} , as a function of the rate of liquid discharge, Q (in semi-logarithmic co-ordinates).

same concentration of 80 ppm by weight of the same PAAm sample. However, the Newtonian solvent was significantly thinner, containing 35% of sucrose and 1% NaCl, and having a viscosity of 4.2×10^{-3} Pa s at the room temperature of 22 °C. The polymer solution viscosity was $\eta = 5.6 \times 10^{-3}$ Pa s at $\dot{\gamma} = 50$ s⁻¹.

The flow was driven by a difference of hydrostatic pressures, ΔP , applied between the inlet and outlet. The dependence of the flow rate, Q, on ΔP was measured using an *in situ* compensation technique [40] (see details in [39]). Figure 26 shows the resistance ratio $\Delta P/\Delta P_{lam}$ as a function of Q, where ΔP_{lam} is the pressure difference expected in a laminar flow for a liquid with a viscosity, $\eta(\dot{\gamma})$, at the same Q. (Average $\dot{\gamma}$ was calculated as $4Q/d^3$ again.) One can see a distinct transition and an onset of non-linear growth of the flow resistance at $Q \approx 8.5 \text{ nl s}^{-1}$, corresponding to $\dot{\gamma} = 34 \text{ s}^{-1}$ and $Re \approx 0.017$. The low value of Re suggests that the transition is of a purely elastic nature. Relaxation time, λ , of the polymer solution was low and could not be measured directly. For the purpose of estimation we can assume $\lambda \sim \eta_s$ and plug into this relation the values for the more viscous solution used in the table-top channel, $\lambda = 1.4 \text{ s}$ at $\eta_s = 0.153 \text{ Pa} \text{ s}$. Then we get $\lambda = 0.04 \text{ s}$ and Wi = 1.3 at the instability threshold for the current solution, which is in reasonable agreement with the results in figure 24. As Q increases to 60 nl s⁻¹ (corresponding to $Wi \simeq 9$), the flow resistance growth reaches a factor of 2.8, which is comparable with the CT flow, although significantly lower than in the flow between two plates.

5.3. Discussion

The experimental results in figures 24 and 26 show that, above the elastic instability threshold, the flow of the polymer solution in the channel exhibits two major features of turbulent flows: a significant increase in the rate of mixing and in the flow resistance. Furthermore, solid evidence

for the turbulent character of the flow is given by the power spectra of fluctuating velocity in figure 25. The spectra of both longitudinal and transversal velocity components do not exhibit any distinct peaks and have broad regions of a power-law decay with a power of about -3.3. Since the power spectra in figure 25 were measured at a point with a high mean flow velocity (10 times higher than the characteristic fluctuating velocity), we can use the Taylor hypothesis and argue that the spectra in figure 25 actually reflect the spatial structure of the flow. Then the power-law-decay region can be transferred to the spatial domain, with the power of the velocity fluctuations scaling as $P \sim k^{-3.3}$ with the wave number, k. The absence of peaks in the spectra in figure 25 and significant extension of the frequency region, where P follows the power law, are strong evidence of fluid motion in a broad range of spatial scales and of the turbulent character of the flow. We also notice here that an exponent of -3.3 in figure 25 is very close to those measured in the flow between two plates (figures 5, 8 and 16), which varied from -3.3 to -3.6 depending on the position. So, one can suggest that decay of the power of the velocity fluctuations with an exponent around -3.5 is a rather general feature of the elasticityinduced turbulent flows. It, however, does not appear in the radial velocity spectrum of the CT flow (figure 19), probably because of abundance of the coherent axisymmetric structures (see section 4.2).

The functional form of the velocity power spectra, $P \sim k^{-3.3}$, deduced from figure 25, suggests that the fluctuations of velocity gradients scale as $k^{-1.3}$. An integral of $k^{-1.3}$ diverges at $k \to 0$ and converges at $k \to \infty$. It means that the main contribution to the fluctuations of the velocity gradients and the velocity differences at all scales comes from the biggest eddies, having dimensions of the whole system (diameter of the channel). This conclusion has an immediate implication for mixing in the flow: it should result in the same type of patterns and in functionally the same statistics as in the case of a completely homogeneous flow, $\vec{V}(\vec{r}, t) = \vec{V}_0(t) + (\partial V_i(t)/\partial r_j)(\vec{r} - \vec{r}_0)$, randomly varying in time.

Such a flow is a realization of the so-called Batchelor regime of mixing [41], and the problem of statistics of a tracer (dye) distribution in it has been solved analytically recently [42]–[44]. The Batchelor regime occurs at small scales (below the Kolmogorov dissipation scale [1, 27]) in the usual, high *Re* turbulence, and it is rather difficult to realize in a laboratory otherwise. Therefore, the elastic turbulent flow in the channel provided a very convenient experimental system for a quantitative study of mixing in this regime [35]. The experimental results on correlation functions and PDF of dye concentration, and on their dependence on time of mixing agreed very well with the theoretical predictions [35]. A practical message of the experiments is that very viscous liquids can be efficiently mixed in curvilinear channels at very low flow rates by adding high-molecular-weight polymers at very low concentrations. This method of mixing, we believe, can find some industrial and laboratory applications [45].

6. Concluding remarks

We studied flows of dilute solutions of a flexible high-molecular-weight polymer in three different flow set-ups, which shared the same feature of high curvature of the flow lines, quantitatively expressed in large gap ratios, d/R. High viscosity of the solvent and long polymer relaxation times ensured an elasticity-dominated-flow regime, where effects of the non-linear constitutive relation between polymer stress and rate of deformation prevailed over the inertia-related non-linearity of the Navier–Stokes equation. As a result, all flow transitions were induced by the non-linear elasticity, for *Wi* of the order of unity and at very low *Re*. In all three systems, a flow above the instability threshold was randomly fluctuating in time and space, and exhibited some or all of the main features of turbulence: fluid motion excited in a broad range of spatial and temporal scales, and significant increase in the rates of momentum and mass transfer (flow resistance and mixing). The extent of the flow resistance growth varied rather substantially between the systems (factors of 20, 4.2 and 2.8 for the flow between two plates, the CT flow, and the flow in the curvilinear channel, respectively) and so did the exponents in the power-law-decay regions of the velocity spectra (from -3.3 to -3.6 for the flow between two plates, however, that all the experimental findings are fitting rather well to the same general framework of a phenomenon which we call elastic turbulence.

Observation of the elastic turbulence and investigation of its properties have been greatly facilitated by the choice of the systems with large d/R, where the elastic instability occurs at low Wi, and by the choice of the high molecular weight polymer, which is only moderately extended (compared with its full contour length) at the instability threshold. Nevertheless, we believe that elements of the elastic turbulence should generally appear in three-dimensional flows of polymer solutions at high Wi and low Re. Possible realizations of this regime may range from flows of polymer melts in industrial reactors to small scale flows in drag reducing aqueous polymer solutions. Dependence of the elastic turbulence-onset conditions on the parameters of the system and on the properties of the polymer solution (discussed in section 3.4.2) implies that by using polymer solutions with sufficiently high elasticity, one can excite turbulent motion at arbitrary low velocities and in arbitrary small tanks. (In some cases it may be the only way to produce a chaotic flow, which may be needed for fluid mixing.) Irregularly fluctuating flows showing multiple features of the elastic turbulence have been recently demonstrated in a rotating flow between two plates with a 300 μ m gap [28] and in a 100 μ m thick curvilinear channel [39]. An apparent restriction is, however, that the size of the tank still has to be large compared with the size of the polymer coils.

Recently, there have been some serious advances in the theory of random flows of polymer solutions in both high and low *Re* regimes [25, 26, 32, 33]. In particular, the following explanation was proposed for the shapes of velocity spectra seen in figures 5, 8, 16 and 25 [26]. Fast decay of the fluctuation power with k implies a velocity field, where the main contribution to deformation and stirring (stretching and folding) at all scales comes from a randomly fluctuating velocity field at the largest scale of the system. Therefore, it is suggested that the leading mechanism for generation of small scale (high k) fluctuations in the elastic stress is advection of the fluid (which carries the stress) in this fluctuating large-scale velocity field. Hence, the fluctuating velocity field and stress tensor can both be decomposed into large- and small-scale components, and the leading mechanism for the generation of the small-scale (high k) portions is advection by the fluctuating large-scale flow. (The theory considers the elastic stress tensor to be passively advected in a random velocity field, which is analogous to the concept of a passively advected vector in the magnetic dynamo theory [26].) The small-scale velocity fluctuations are a product of the small-scale stress fields. The elastic stresses are mainly pumped by the large-scale flow field, and the smaller scale stress fields created by the advection are permanently decaying because of the polymer relaxation. These simultaneously occurring advection and relaxational decay of τ_p result in a quick decay of τ_p fluctuations at large k, which should produce $P \sim k^{-\alpha}$ velocity spectra with $\alpha > 3$. The current understanding of the elastic turbulence still remains

quite incomplete, however, and this phenomenon certainly awaits further detailed experiments and numerical simulations.

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