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Andriy Pogrebnyak, Volodymyr Pogrebnyak

National Technical University of Oil and Gas, Ivano-Frankivsk, Ukraine

HYDROCUTTING OF FROZEN FOOD PRODUCTS

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Abstract. The article aims to determine the peculiarities of macromolecule deformation behavior under conditions of a jet-shaping head that would allow to solve the issue related to the mechanism of increasing water-jet cutting power with polymer additions. The roles of longitudinal and transverse velocity gradients in the manifestation of distinctive features of polymer solution flows are analysed.

In converging polyethyleneoxide solution flow macromolecules are forced by a hydrodynamic field to rather strong stretching that causes the dynamic structure formation in solutions. There have been studied experimentally velocity fields and their gradients as well as the degree of macromolecule unrolling under pattern conditions of a jet-shaping head in polyethyleneoxide solutions flow. In converging polymer solution flow macromolecules are forced by a hydrodynamic field to rather strong (~ 60 % and more) stretching that causes the field restructuring. The determined regularities of macromolecules behavior in the flow under conditions of a jet-shaping head and manifested in this case effects of elastic deformations have paramount importance in understanding the mechanism of "anomalously" high cutting power of water-polymer jet. The work for the first time makes it possible to explain the nature of increased water-jet cutting power with polymer additions when cutting food products. Understanding the nature of increased cutting power of water-polymer jet will make it possible to develop recommendations on choosing regimes for water-polymer jet processing of food products by cutting.

Keywords: jet-shaping head, water-jet cutting, polymer solution flow, hydrodynamic field, elastic deformation, food products, water-polymer jet processing.

Introduction

In the works [1–3] the complex studying of the process of hydro-cutting frozen food products is carried out. It has given the chance to offer the most expedient ways of intensification of the process of hydrocutting frozen food products. It has been experimentally proven that using polyethyleneoxide (PEO) water solutions as a working liquid while cutting frozen food products substantially increases efficiency of hydro-cutting process and quality of the cut surface.

The works [3–6] deal with the regularities of the frozen food products hydrocutting to increase its efficiency and to improve the quality of cut surface by working liquid modification. The influence of PEO concentration on cutting depth and rate for food frozen at -25 °C by pressure water-polymer jet of 100 MPa

flow pressure and $0.37 \cdot 10^{-3} m$ nozzle diameter is investigated. It is experimentally proved that when PEO water solutions as a working liquid are used the optimum distance between nozzle edge and food surface increases 15 times, cutting depth at cutting speed of 0.100 m/s – of 4 times, and the quality of the cut surfaces is also improved. In connection with this, it is important to know physical mechanism of the observed effect.

Among the attempts to explain the nature of the effect of water-polymer jet anomalously high cutting power a special place is held by an approach based on deformation impact of hydrodynamic field on macromolecules. To substantiate this approach it is necessary to prove experimentally the presence of strong deformation impact of hydrodynamic field under conditions of a jet-shaping head. The research of

converging currents has shown that it is possible to generate flow with predominantly longitudinal velocity gradient, i.e. to simulate conditions that appear in a jet-shaping head, with the help of a short capillary tube [4, 5].

Objective of the article

The objective of the article is to determine peculiarities of macromolecule deformation behavior under conditions of a jet-shaping head that would allow to solve the issue related to the mechanism of increasing water-jet cutting power with polymer additions, and the quality of the frozen food products cut surfaces.

Methods and Materials

The hydrodynamic activity of polymers in high-speed flows experiment was studied out with the use of an industrially fabricated pulsed water jet. The description of its hydrodynamic unit, parameters, and method of determination of the mean velocity, momentum, and energy of the ultrajet are given elsewhere [5]. To determine the parameters of the ultrajet, shooting was performed with a physical pendulum, whereas for the qualitative determination of energy possibilities of the jet the interaction of the jet with a target installed at the pendulum was studied experimentally. The target was fabricated from steel (St3) plates with the dimensions $(220 \times 220 \times 3) \ 10^{-3}$ m with mounting holes $16 \cdot 10^{-3}$ m at the corners. A gap of width 5 10^{-2} m was left between the plane of the target and the body of the physical pendulum.

In this study, velocity and velocity gradient fields as well as the degree of the coil-stretch transition at the entrance of the capillary were investigated for various flow regimes. A flow viscometer with an entrance angle of 180° was used. The instrument contained a cell having a rectangular cross-section $(10 \times 17) \cdot 10^{-3} m$ and height of $8 \cdot 10^{-2} m$ and two short removable capillaries having the following diameter and length, respectively: $0.5 \cdot 10^{-3}$ and $0.21 \cdot 10^{-3} m$ (capillary I) and $0.37 \cdot 10^{-3} m$ and $1.1 \cdot 10^{-3} m$ (capillary II). The velocity field at the entrance of the capillaries was measured using a laser Doppler anemometer according to the method [5]. The average flow rate \bar{u} was measured volumetrically using a photo-electronic system; the flow downstream the capillary inlet was submerged. PEO having the viscosity-average molecular weight of $M_{PEO} = 3 \cdot 10^6$, $4 \cdot 10^6$, $6 \cdot 10^6$ The methods of Δn_{∞} , calculations and experimental procedure of Δn measurements are given elsewhere [6, 7]. The solutions were prepared in the following manner. A previously (one week before) prepared 0.1 % solution of PEO was diluted with distilled water. Additives of 0.05 % potassium iodide were introduced to exclude degradation of the polymer solutions upon storing.

Results and Discussion

Hydrodynamic activity of polymers in high-speed flows. Results of the experiments are presented in Table 1, whence it is seen that:

1) the maximum distance from the water jet to the target when punching is still observed comprises 1.5 m for water, whereas for 0.0008 % polymer solution it comprises 2.5 m;

2) the efficiency of polymer additions was observable starting from a concentration of 0.0004 %;

3) polymer additions lower the mean jet velocity.

The experimental data obtained in the experiments can be explained rather well from the viewpoint of the strong deformational action of the hydrodynamic field on the molecular coils [1, 3]. It was proved that deformation of a molecular coil in a longitudinal field is substantially greater than in a shear one, since in the case of a longitudinal flow, positive feedback arises between molecular dimensions increasing as a result of deformation and the velocity gradient, and the unfolding process starts at relatively low critical velocity gradients, building up in an avalanchelike manner. In the case of a shear hydrodynamic field, negative feedback arises instead of the positive one, as in the case of the longitudinal field, between the dimensions of the molecules and the velocity gradient and, naturally, considerable deformations of coils are hindered.

Table 1

No. of experiment	Distance from the pendulum, m	Polyethyleneoxide concentration, %	Mean jet velocity, m/sec	Target punching character
1	0.5	_	255	Hole diameter (18–20)·10 ⁻³ m, punched holes
2	1.5	_	255	Hole diameter (15–18)·10 ⁻³ m, punched holes
3	1.75	_	250	No punching
4	2.5	_	250	No punching
5	1.5	0.0004	240	Hole diameter (15–18)·10 ⁻³ m, punched holes
6	1.75	0.0004	240	Depression with rupture $(3x10) \cdot 10^{-3}$ m
7	2	0.0004	235	No punching
8	2.5	0.0008	230	Hole diameter $(8-10) \cdot 10^{-3}$ m

The effect of PEO concentration in water and distance from the pendulum on character interaction of hydrojet with target

If one calculates the longitudinal velocity gradient realized in the barrel (in the case of Newtonian liquid flow) it appears that it did not exceed (in our experiments) $1.8 \cdot 10^{-3} \text{ sec}^{-1}$, i.e., despite such high velocities (U = 250 m/sec) and consequently high transverse velocity gradients the deformation conditions for macromolecules in the cone barrel of the water jet compare well with those at the entrance to a capillary with a rectangular entrance at a mean flow velocity of ~ 1 m/sec [3–5]. This circumstance bears witness to the fact that in movement of a polymer solution along a water jet barrel the molecules as in the case of a short capillary [4,5] are, first of all, subjected to the strong deformational action of the longitudinal hydrodynamic field since the velocity gradients \mathcal{E}_{cr} realized under these conditions are higher than certain critical values (connected with the relaxation time θ_c of the macromolecules by the relationship $\theta_c \cdot \mathcal{E}_{cr} = 0.5$ [8]) sufficient to ensure unfolding of the macromolecules. Therefore the outgoing water jet appears to be armed with macromolecules unfolded to a great extent. A portion of its energy goes into structural reorganization of the flow [1], which causes an increase in the compactness of the jet with polymer additives and decreases its mean velocity. The latter circumstance is favorable for an increase in the threshold distance from which the target is still broken through.

The vanishing of differences in the behavior of jets of water and polymer solution observed in the experiments at such velocities when the longitudinal velocity gradient becomes less than \mathscr{E}_{cr} , whereas at the same time the transverse gradient can exceed substantially (by 1–2 orders) the value of \mathscr{E}_{cr} , also bears witness that the extent of the hydrodynamic action on the molecular coils is practically independent of the transverse velocity gradient.

The experiments considered confirm the above conclusion that the extent of the hydrodynamic action on polymer solutions is determined primarily by the value of the longitudinal velocity gradient achieved [4] and the duration of the action of the stretching hydrodynamic field [6, 7].

Dynamic structure formation in solutions polymers. Let's examine the tests allowing to reveal the ability of solutions polymers to dynamic structure formation effected by hydrodynamic field with stretching. To create hydrodynamic field with stretching there has been used a flow of Newtonian liquid converging to a small outlet $0.3 \cdot 10^{-3} m$ in diameter. At quite a distance from the outlet there have been

injected in this flow some jets of PEO solutions. Polymer solution velocity in points of injection agreed with velocity of the main liquid flow. Visualization of the flow in the inlet area was done with the help of dye additives injected into polymer solution. Under small outflow velocities dyed jets of polymer solution visualize stream-lines of the main flow (Fig. 1). In his case behavior of PEO solution jets doesn't differ at all from the behavior of water jets injected into converging liquid flow. Reaching some critical flow rate of liquid through the orifice the character of jet flow of polymer solution drastically changes. Rather thick polymer jets transform into thin threads that change their length with time flow (Fig. 2).

When observing the dynamics of forming and destructing separate threads (here lies the moving pulsation character of the flooded polymer jets) the following regularity comes to life. At the beginning when polymer jets approach the orifice there can be traced their gradual bend towards the orifice. Here their velocity growth along these curve trajectories becomes more noticeable as the jet thickness gets reduced. Near the orifice discontinuous reduction of jets thickness takes place as they are transformed into thin threads. Transformation area of a thick jet into a thin thread starts to shift up along the jets resulting in the increase of the thread length. As jets approach the orifice not concurrently the length of threads spun by the flow from separate polymer jets is different at each given moment. When reaching some critical length a thread cuts off in close proximity to the orifice. After that the remaining at the top thread loses its elasticity and sags marking one of the stream lines of the main flow. When a polymer jet approaches the orifice again the whole thing is repeated anew. This process is resumed with time interval from fractions of a second to several seconds depending upon the outflow mode of the main flow, polymer molecular mass, polymer type, solvent quality and temperature, as well as polymer concentration in solution.





 $M_{PEO} = 6 \cdot 10^6$, $C_{PEO} = 0.03\%$

Fig. 1. Picture of flow in the inlet area of a capillary when injecting PEO solution into water during precritical mode of outflow

Fig. 2. Picture of flow at the point of wire probe effecting a polymer thread spun by hydrodynamic field

Outflow velocity growth of converging water flow as well as increase of molecular mass and polymer concentration result in the increase of thread length and reduction of their length-changing rate. Temperature growth leads to the opposite effect. Tests with acetone and dioxan (solvents with poorer thermodynamic qualities than water) showed that under the same velocities as in tests with water pulsation rate of thread length is bigger, but the amplitude of these changes is less than in case of injecting polymer solution into water.

So, the results obtained by us show that dynamic structure formation and periodic processes subjected to Prigozhin's principles of self-organization may occur in polymer solutions in flow with stretching, i.e. under jet-shaping head in model conditions.

Distortions of the Molecular Shape of Polymers under jet-shaping head conditions. Data describing the influence of discharge velocity on effective viscosity of water PEO solutions with different concentrations for molecular weights of $4 \cdot 10^6$ and $6 \cdot 10^6$ at 25 °C are given on Fig. 3.



Fig. 3. Dependence of effective viscosity of water PEO solutions on average discharge velocity: $a - d_k = 0.34 \cdot 10^{-3} m$, $l_k = 10^{-3} m$, $M_h = 4 \cdot 10^6$, C_{PEO} : 1 - 0.01 %,

 $2 - 0.02 \%; 3 - 0.05 \%; 4 - 0.1 \%; 5 - water; b - d_k = 0.12 \cdot 10^{-3} m; l_k = 0.2 \cdot 10^{-3} m; M_h = 6 \cdot 10^6; C_{PEO}: 1 - 0.0005 \%; 2 - 0.001 \%; 3 - 0.002 \%; 4 - 0.003 \%; 5 - 0.004 \%; 6 - water$

It can be seen, that the phenomena, unusual for purely viscous mediums are characteristic of such currents. At certain critical (threshold) values of average exhaust velocity \overline{u} the relative pressure differential begins sharply to increase, and it is the sharper the more is the concentration of polymer in a solution. The marked character of dependence $x = f(\overline{u})$ testifies about high dissipation (sometimes, than is on 2 orders of magnitude more) of energy during the course of solutions of polymers through an slot i.e. the increased hydrodynamic resistance on supercritical flow rates is observed.

Transition to a mode of current with an increased dissipation of energy is accompanied by formation of the source flooded jet as "cord" enclosed by secondary currents in the shape of a ring-shaped vortex. In case of supercritical mode of current for area of the concentration lying between very diluted and moderately concentrated solutions of polymers, there happens rather strong deformation effect of a hydrodynamic field on molecular chains.

To interpret the data, the structure of the hydrodynamic field and degree of the molecular shape distortions induced by the field should be evaluated. Distribution of the flow rate along the flow axis for 0.05 % PEO solution in dimensionless coordinates is depicted in Fig. 4. It can be seen that, before the critical flow regime is attained, the increase of the effective viscosity is not exhibited and the axial distributions of velocity for the polymeric solution and pure water are almost the same (curve 3) and filled circles on curve 3, respectively. After passing through the critical flow regime, the curves exhibit a considerable deformation and development of the axial velocity profile (curves 1 and 2 in Fig. 4).

The latter curves have at least two linear regions. Using the experimental velocity distributions along the flow axis, the respective rate gradient distributions were calculated (Fig. 5). It can be seen that the maximal rate gradient ε max, occurs not at the entrance of the capillary, but at some distance from that (Fig. 5, *a*, curve 2). The \mathscr{C}_{max} at the entrance of the capillary for polymeric solution is considerably lower that for water. Hence, the hydrodynamic field results in perturbation of the macromolecules, which, in turn, affects the velocity field in such a way that the longitudinal velocity gradients are decreased.

Thus, the longitudinal velocity gradient at the flow axis does not exceed $30 s^{-1}$. An increase in molecular weight of the polymer and its concentration also results in decrease of the longitudinal gradient at the flow axis. Thus, the respective value for 0.05 % PEO solution and capillary I at the average flow rate

 $2 m \cdot s^{-1}$ is equal to $10 s^{-1}$. Fig. 5, *b* illustrates that the longitudinal rate gradient at the axis of the flooded jet as "cord" (curve 3) and maximal value of the rate gradient at the entrance of capillary (curve 2) are only slightly dependent on the average flow rate through the capillary.



Fig. 4. Velocity distribution for the polymer solution along the flow axis: $M_h = 4 \cdot 10^6$, flow through the capillary I at the average flow rate $\overline{u} = 2.5 \text{ m} \cdot \text{s}^{-1}(1)$, flow through the capillary II at $\overline{u} = 0.8 \text{ m} \cdot \text{s}^{-1}(2)$ and through the capillary II at $\overline{u} = 0.5 \text{ m} \cdot \text{s}^{-1}(3)$, filled circles represent the data for water when $\overline{u} = 0.8 \text{ m} \cdot \text{s}^{-1}$

The observed changes in the structure of the hydrodynamic field can be associated with large distortions of the macromolecular coils induced by hydrodynamic field, leading to non-linear elasticity effects. The degree of the coil-stretch transition may be estimated from the value of the deformational factor $(\Delta n/\Delta n_{\infty})_{max}$, where Δn is the experimental flow birefringence value, while the Δn_{∞} , is the limiting value of the flow birefringence calculated at the given concentration of polymer [8].



Fig. 5. *a* – velocity gradient distribution along the flow axis at $\overline{u} = 0.8 \text{ m} \cdot \text{s}^{-1}$ for water (1) and 0.05 % PEO solution (2). Measurements were performed with the capillary I; *b* – dependence of the maximal longitudinal velocity gradient on the average flow rate for water (1) and 0.05 % PEO solution (2) and in the vortex for 0.05 % PEO solution (3)

Results of the studies of the influence of hydrodynamic field on the polymeric solution are depicted in Fig. 6. The value of deformational factor increases when approaching the entrance of the capillary for the average flow rate equal to the critical value (curve 1) and reaches its maximum at rather high \bar{u} values (curve 2). The maximum position of the deformational factor $(\Delta n/\Delta n_{\infty})_{max}$, at the entrance of the capillary corresponds to the domain with maximal longitudinal velocity gradient (Fig. 5, *a*, curve 2).



Fig. 6. The distribution of the coil-stretch transition degree over distance from the entrance of the capillary for aqueous PEO solution: $C_{PEO} = 0.03 \% (1), \ \overline{u} = 1 \ m \cdot s^{-1}; \ C_{PEO} = 0.03 \% (2), \ \overline{u} = 2.5 \ m \cdot s^{-1};$ $C_{PEO} = 0.1 \% (3), \ \overline{u} = 1 \ m \cdot s^{-1}$

The obtained distribution of the deformational factor over the flow axis at the entrance of the capillary (Fig. 4) envisages the possibility of a high degree of coil-stretch transition under the free-converging flow conditions. The flow birefringence ratio attains the value of 0.34, which corresponds to ~60 % and more coil-stretch transition degree. Increasing the polymer concentration results in a decrease of the deformational factor (curve 3).

Fig. 7 envisages deformational behavior of macromolecules at the different flow rates. It can be seen that the $(\Delta n/\Delta n_{\infty})_{max}$, versus average flow rate dependence may be divided onto three domains. Domains 1 and 3 are characterized by a monotonic increase of the coil size with growth of the flow rate, while in the intermediate domain 2 the transition is rather sharp. From a comparison of Fig. 3 and 5, it can be concluded that an increase of the \bar{u} in the domain I is accompanied by growth of the longitudinal velocity gradient, which results in an increase of the size of the coils. Otherwise, the reverse stretch-coil transition occurs [9,10] and macromolecules decrease their influence on the velocity field, which, in turn, results in a steep increase of the velocity gradient. This will lead to the coil-stretch transition and all the above processes will happen once again.

Hence, the stable state is characterized by the minimal value of the velocity gradient which is sufficient for a sharp coil-stretch transition. Increase in the flow rate results in some additional growth of the deformational factor due to nonlinear effects (Fig. 7) sufficient for stability of the rate gradient field of the chosen polymeric system (Fig. 5, b, curve 2).

Conclusions

The determined regularities of macromolecules behavior in the flow with longitudinal velocity gradient and manifested in this case effects of elastic deformations have paramount importance in

understanding the mechanism of anomalously high cutting power of hydro-jet water-polymer processing of foodstuffs by cutting. The nature of the anomalously high efficiency of hydro-jet water-polymer processing of food products by cutting is explained quite well from the point of view of the strong deformation effect of the hydrodynamic field on molecular coils. The outflow of water appears to be armed macromolecules and dynamic structures stretching to a large extent. Part of its energy goes into a structural reorganization of the flow, which leads to an increase in the compactness of the jet with polymer additives. The latter circumstance is favorable for increasing the threshold distance from which the material is still cut, and the quality of the cut frozen food products surfaces is also improved.



Fig. 7. Dependence of the coil-stretch transition degree on average flow rate: $d_k = 0.5 \cdot 10^{-3} m$; $l_k = 0.21 \cdot 10^{-3} m$; $C_{PEO} = 0.03 \%$; $M_{PEO} = 4 \cdot 10^6$

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