The response of nano-ceramic doped fluids in heat convection models: A characteristic-based numerical approach

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**KEYWORDS**

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**Abstract.** In this paper, forced, free, and mixed convections in incompressible flow were studied numerically. Nano-sized Al\textsubscript{2}O\textsubscript{3}, TiO\textsubscript{2}, MgO, and ZnO ceramics with water were considered as nano-fluids. This study simulated a cavity flow with different boundary conditions and aspect ratios as well as flow over stationary and rotating cylinders. The mean Nusselt number ($\overline{Nu}$) and friction factor for cavity flow and ($\overline{Nu}$) for flow over a cylinder were compared for different nano-fluids. A new code was developed in FORTRAN 95 for numerical simulations. A fifth-order Runge-Kutta method for time discretization and a characteristic-based scheme for convective terms were used in this code. The averaging scheme on the secondary cells was employed to obtain viscous fluxes. Primary results were validated in comparison to other research outputs. Results demonstrated that MgO-water and ZnO-water had maximum and minimum heat transfer rates, respectively. Moreover, maximum and minimum shear stresses were recorded for the Al\textsubscript{2}O\textsubscript{3}-water and TiO\textsubscript{2}-water, respectively. Using nano-fluid increased the heat transfer rate by 15-37\%, depending on the Richardson number and selected nano-particles.

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1. Introduction

Optimization of thermal systems and use of renewable energy sources are the inevitable ways of attenuating the threat and intensity of air pollution, global warming, and environmental concerns as well as decreasing other related costs. Enhancement of heat transfer in thermal systems is the basic part of system optimization. There are two major categories of heat transfer enhancement: active and passive methods \cite{1}. In active methods, some types of external energies including electric field, magnetic field, and acoustic force are employed to perturb the fluid flow and increase the heat transfer rate, whereas, in passive methods, the enhancement is achieved by making changes to the geometrical configuration of the system or manipulating the working fluid \cite{2}. Nano-fluids, as newborn working fluids, belong to the group of passive methods. The term nano-fluids refer to stable
dispersion of nano-sized particles in a base fluid. Solid materials generally possess high thermal conductivities than conventional fluids; therefore, the addition of solid particles improves their thermal properties [3]. The interesting properties of nano-fluids are the reasons behind the researchers’ and engineers’ motivation to investigate their application in various fields of engineering including Micro Electro-Mechanical Systems (MEMS), fuel cells, and heat treatment of metals [4]. Still, a wide range of experimental works are needed to provide a reliable database to establish a comprehensive theory about nano-fluids. Conducting accurate experimental researches is costly and time-consuming. Moreover, working with nanoparticles is subject to its particular dangers about the safety of the experimenter. On the other hand, extracting accurate data is difficult and sometimes impossible, e.g., determining the temperature of each point in micro-channels. Uncertainties related to measuring instruments and sources of error including instrumental, environmental, procedural, and human are the other drawbacks of using experimental methods. Numerical methods, however, represent a powerful tool for analyzing almost all physical phenomena and they can provide detailed information concerning each desired location.

Maiga et al. [5] managed to study the effect of Al_2O_3 dispersion in two different media, water and ethylene glycol. They investigated the forced convection of the mentioned nano-fluid numerically and reported that using alumina nanoparticles enhanced the heat transfer of both fluids. They found that the ethylene glycol-Al_2O_3 combination achieved better heat transfer enhancement than water-Al_2O_3. However, wall shear stress was higher at the same time. Kefayat [6] studied natural convection flow with Cu-water nano-fluid by the Lattice-Boltzmann scheme in a cavity at different Aspect Ratios (ARs). The magnetic field was applied in the flow domain. The results demonstrated that the effect of the nanoparticles increased at large Hartmann numbers as the AR increased. Mahmoodi [7] studied the effect of particle type on natural convection of the cavity flow with different heaters in it. They employed the water as a base fluid and performed a series of numerical simulations using the SIMPLER algorithm. Their results demonstrated that a horizontally located heater would have a higher Nu value at small Ra numbers than a perpendicularly placed heater, but the location of the radiator did not affect Nu at large Ra numbers. Rahmati and Tahery [8] simulated the laminar natural convection in a cavity using TiO_2-water nano-fluid. They proposed some obstacles in the base geometry and used the Lattice-Boltzmann method to predict the velocity field and thermal behavior of the nano-fluid. They found that the (Nu) value increased upon an increase in Rayleigh number and the volume fraction of nanoparticles. They also reported that the obstacle dimension had a direct effect on the thermal behavior of the cavity such that the (Nu) improved despite the obstacle dimensions up to 0.5 L; however, the Nusselt number deteriorated as the obstacle dimensions increased to 0.7 L.

Akbarinia and Behzadmehr [9] studied numerically the mixed convection of Al_2O_3-water nano-fluid within a tube. Their results illustrated that the concentration of nanoparticles had no direct effect on the secondary flow, axial velocity, and friction coefficient. They claimed that the rising volume fraction of nanoparticles increased Nu and reduced friction factor (f). Heydari et al. [10] simulated the three-dimensional nano-fluid flow in a heat exchanger. They used SiO_2, CuO, Au, Cu, Fe, Al_2O_3, and Fe_2O_3 in ethylene glycol and water-based fluids. It was reported that the ethylene glycol-based nano-fluids had higher efficiencies. Uysal et al. [11] performed a three-dimensional simulation in a circular tube using ZnO-ethylene glycol nano-fluid. Their results showed 10% enhancement of Nusselt number in a concentration of 1%. Sunil and Kumar [12] used the Lattice-Boltzmann scheme to simulate the Al_2O_3-water nano-fluid. They found that increase in the heat transfer rate was not dependent on Reynolds number variation. Al-Kouz et al. [13] two-dimensionally simulated the cavity using Al_2O_3 – Air nano-fluid and employed two fins on the hot wall to increase the effect of nano-fluid on the rising Nusselt number. Natural laminar convection flow was considered in this work and Knudsen number ranged between 0 and 0.1, while the Rayleigh number ranged between 10^7 and 10^9. They found that Nu was a function of Rayleigh number and inverse of Knudsen number. They proposed a new relation for the Nusselt number as Nu = 0.219Ra_{0.082}Kn_{-0.514}d_{10.14}.

Zhang et al. [14] conducted a numerical study to simulate nano-fluid flow and employed an adaptive grid method to ensure accuracy. Using industrial nano-fluid to improve heat transfer rate has increased in recent years. The numerical method costs less than experimental methods. Hence, numerical methods are excellent for gaining an initial understanding of nano-fluid flow behavior. As a result, different numerical studies of nano-fluid flow have been done so far [15–18]. The literature review points to the valuable scope of numerical simulations performed to predict the behavior of nano-fluids. However, the unknown nature of the involved mechanisms is still a big challenge to completely cover the thermal behavior of these fluids such that in some cases, the numerical simulations are considerably distant from, or inconsistent with, the experimental data. Therefore, it is required to examine different schemes to reduce the related numerical errors. The present work aims to solve the governing
equations of the nano-fluids using the characteristic-based scheme proposed previously by our group [19-21]. To investigate the potential of the present scheme, four types of nanoparticles including Al₂O₃, TiO₂, MgO, and ZnO ceramics in base water are considered. Different types of convections are simulated numerically for nano-fluids and a pure fluid. The present scheme is applied to two different geometries: cavity flow and flow over a cylinder. The cavity flow is simulated at different ARs, boundary conditions, and Richardson numbers. The flow over the cylinder is simulated in two cases of stationary and rotating cylinders.

2. The governing equations and the numerical procedure

Nano-fluids can be considered as single-phase fluids in numerical studies in which their physical characteristics depend on the characteristics of the base fluid and the nanoparticles. The following are Navier-Stokes equations for a two-dimensional nano-fluid with heat transfer:

\[ \frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} + v \frac{\partial u}{\partial y} = -\frac{1}{\rho} \frac{\partial p}{\partial x} + \frac{\partial}{\partial y} \left( \frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} \right), \]

\[ \frac{\partial v}{\partial t} + u \frac{\partial v}{\partial x} + v \frac{\partial v}{\partial y} = -\frac{1}{\rho} \frac{\partial p}{\partial y} + \frac{\partial}{\partial y} \left( \frac{\partial v}{\partial x} + \frac{\partial v}{\partial y} \right), \]

\[ \frac{\partial T}{\partial t} + u \frac{\partial T}{\partial x} + v \frac{\partial T}{\partial y} = \frac{1}{\rho C_p} \left( \frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} \right) + \frac{k}{\rho C_p} \left( \frac{\partial^2 T}{\partial x^2} + \frac{\partial^2 T}{\partial y^2} \right) + \frac{\mu}{\rho C_p} \Phi, \]

\[ \Phi = \left( \frac{\partial u}{\partial y} + \frac{\partial v}{\partial x} \right)^2 + 2 \left( \left( \frac{\partial u}{\partial x} \right)^2 + \left( \frac{\partial v}{\partial y} \right)^2 \right). \]  

(1)

The artificial compressibility of Chorin [22] is applied and the gradient of pressure is added to the continuity equation. As a result, the pressure can be obtained by numerical schemes. Also, the nature of governing equations varies and the characteristic-based scheme proposed by ourselves can be used as a numerical scheme. This scheme solves the problems of the averaging scheme [19-21]. The instability of the averaging scheme is a serious problem solved by using our proposed characteristic-based scheme.

\[ \frac{1}{\varepsilon} \frac{\partial p}{\partial t} + u \frac{\partial p}{\partial x} + v \frac{\partial p}{\partial y} = 0, \]  

where \( \varepsilon \) is the artificial compressibility factor [22]. The governing equations can be transformed into dimensionless forms and shown in the matrix shape as follows:

\[ \begin{align*}
\frac{\partial Q}{\partial t} + \frac{\partial F}{\partial x} + \frac{\partial G}{\partial y} &= \frac{\partial R}{\partial x} + \frac{\partial S}{\partial y} + H, \\
Z &= \begin{bmatrix} p \\ u \\ v \\ T \end{bmatrix}, \\
F &= \begin{bmatrix} \beta u \\ p + u^2 \\ w v \end{bmatrix}, \\
G &= \begin{bmatrix} \beta v \\ u w \\ T \end{bmatrix}, \\
R &= \frac{1}{Re} \begin{bmatrix} 0 \\ \frac{\partial u}{\partial x} \\ \frac{\partial v}{\partial y} \end{bmatrix}, \\
S &= \frac{1}{Re} \begin{bmatrix} 0 \\ \frac{\partial u}{\partial y} \\ \frac{\partial v}{\partial x} \frac{\partial^2 u}{\partial y^2} \end{bmatrix}, \\
H &= \begin{bmatrix} 0 \\ 0 \\ \frac{\partial \Phi}{\partial x} \frac{T}{Re} \end{bmatrix}. 
\end{align*} \]  

(3)

The reference length for a cavity flow is equal to the width of the cavity and for the flow over a cylinder is the diameter of the cylinder. The reference velocity and density are the far-field velocity and density, respectively. The Mach number of the flow is less than 0.3, hence, the flow is considered incompressible flow, but the Boussinesq assumption is applied to ensure density change in the y-direction because of the temperature gradient in the flow field. Reference temperatures include the temperatures of warm and cold walls for cavity as well as the far-field and wall of the cylinder for flow over the cylinder. The properties of water, nanoparticles, and nano-fluids are extracted from Ref. [23]. The following equations are used in the case of nano-fluids in order to determine their thermodynamic characteristics [24]:

\[ \frac{\rho_{nf}}{\rho_{bf}} = 1 - C + \frac{\rho_p}{\rho_{bf}}, \quad \frac{\mu_{nf}}{\mu_{bf}} = 1 + C N v, \]

\[ \frac{(\rho C_p)_{nf}}{(\rho C_p)_{bf}} = 1 - C + \frac{(\rho C_p)_{p}}{(\rho C_p)_{bf}}, \quad \frac{k_{nf}}{k_{bf}} = 1 + C N c. \]  

(4)
\[
\frac{\beta_{nf}}{\beta_{hf}} = \frac{\rho_{nf}}{\rho_{hf}} \left( 1 - C \right) + C \left( \frac{\rho_{hf} \beta_{nf}}{\rho_{nf} \beta_{hf}} \right) .
\]  
\]  
(5)

The governing equations are discretized by the FVM, where the fifth-order Runge-Kutta is used for time discretization as follows:

\[
Z_{ijk}^{(q)} = Z_{ijk}^{(n)} - F(Z_{ijk}^{(n-1)}).
\]  
\]  
(6)

The time step is considered \(10^{-4}\) for a stable solution. The usual upwind scheme can be used for time discretization, but the Runge-Kutta method is much more stable \([25,26]\). The convective fluxes are calculated through our proposed characteristic-based scheme \([20,21]\). The first-order derivatives (convective fluxes) in cell centers are changed into normal parameters on cell boundaries based on Green’s theorem. The averaging method is a simple method for calculating the convective fluxes. In this respect, the convective fluxes are obtained through simple averaging of cell center data. However, this method is not stable and this is the reason why it is replaced by the characteristic-based method introduced by the authors. The introduced scheme is stable. The pseudo wave propagation is considered in this method and convective fluxes are calculated by the pseudo waves. This scheme was elaborated in detail in \([20,21]\). The viscous fluxes (the second-order derivatives) in the cell center are replaced by the first-order derivatives on the cell boundaries. Obtained first-order derivatives are replaced by normal parameters on the boundaries of secondary cells based on Green’s theorem (Eq. (7)).

\[
\frac{\partial \zeta}{\partial x} \bigg|_{AB} = \frac{1}{A} \int_{A} \frac{\partial \zeta}{\partial x} \, ds = \frac{1}{A} \int_{A} \zeta dy = \frac{1}{A} \sum_{k=1}^{4} \zeta \Delta y_k ,
\]  
\]  
\( \zeta = u, v, T, \) 
\]  
(7)

where \( f \) and \( Nu \) are calculated as follows:

\[
f = 2 \frac{\mu_{nf}}{\Re \mu_{hf}} \frac{\partial u}{\partial y} = 2 \frac{\mu_{nf}}{\Re \mu_{hf}} \frac{u_2 - u_1}{y_2 - y_1},
\]  
\]  
\]  
\[ Nu = 2 \frac{k_{nf}}{\Re \kappa_{hf}} \frac{\partial T}{\partial y} = 2 \frac{k_{nf}}{\Re \kappa_{hf}} \frac{T_2 - T_1}{y_2 - y_1} .
\]  
\]  
(8)

Also, \( \overline{f} \) and \( \overline{(Nu)} \) for cavity flow are obtained through the following equations:

\[
\overline{f} = \int_{0}^{1} f dx = \frac{1}{N} \sum_{j=1}^{N} f_j ,
\]  
\]  
\]  
\[ \overline{Nu} = \int_{0}^{1} Nu dx = \frac{1}{N} \sum_{j=1}^{N} Nu_j .
\]  
\]  
(9)

\( \overline{Nu} \) for flow over a cylinder is obtained using Eq. (6):

\[ U = 1, v = 0, T = \sin(x) \]

\[ U = 0, v = 0, dT/dx = 0 \]

\[ U = 1, v = 0, T = \cos(x) \]

\[ \omega = C, T = 1, u = 0, v = \ell \]

\( T = 0, U = 1, v = 0 \)

**Figure 1.** The boundary conditions for cavity flow and flow over a circular cylinder.

\[ \overline{Nu} = \int_{0}^{2\pi} Nudd\theta = \frac{1}{2\pi} \sum_{j=1}^{N} Nu_j \Delta \theta . \]  
\]  
(10)

Boundary conditions for flow over the circular cylinder and cavity flows are shown in Figure 1.

3. Results and discussion

3.1. Grid independence and convergence history

The quadrilateral grids were used in this work. Different grid sizes were selected and simulation was conducted to ensure grid independence. The grid independence check for flow over a cylinder and cavity flow is shown in Figure 2, in which horizontal velocity variations at the vertical centerline of the cavity for different grids are compared. In addition, Nusselt number variations on cylinder wall for different grids are compared in terms of grid independence. The error is calculated below:

\[ Error = \frac{\sum_{j=1}^{N} \sum_{i=1}^{M} (u_{i,j}^{k+1} - u_{i,j}^{k})}{NM} . \]  
\]  
(11)

The convergence history is obtained and displayed in Figure 3.
3.2. Validation

For validation, the results of the proposed numerical method for cavity flow are compared with those of MuthamilSelvan et al. [27] in Table 1. The applied boundary conditions are similar to the conditions set by MuthamilSelvan et al. [27] at Ri = 1. Table 1 shows that the applied novel numerical approach can predict the Nu number quite accurately in different nano-fluid concentrations. To ensure greater certainty, validation is done through comparisons with the results of Ghia et al. [28]. To this end, the velocities in the $x$ and $y$ directions are obtained on the centerlines of the cavity and are compared with those of Ghia et al. [28] (Figure 4.).

Good agreement is observed, which proves that our new characteristic-based scheme can solve the governing equation numerically with acceptable accuracy.

3.3. Results of simulation of cavity flow

The $\text{Al}_2\text{O}_3$-water nano-fluid flow in a cavity was simulated by the proposed novel numerical approach, and the streamlines at different Ri numbers are shown in Figure 5. The $\text{Al}_2\text{O}_3$ concentration in the fluid was assumed to be 5%. At larger Ri numbers, the dominant heat transfer mechanism is natural convection, while at small Ri numbers, the forced convection
Table 1. Comparison of average Nu numbers for cavity flow obtained by the present numerical approach and that of Mutthamiliselvan et al. [27].

| Nanoparticles concentration % | $C = 2\%$ | $C = 4\%$ | $C = 6\%$ |
|-------------------------------|-----------|-----------|-----------|
| Mutthamiliselvan et al. [27]  | 2.40      | 2.56      | 2.73      |
| Present work                  | 2.41      | 2.54      | 2.76      |

Figure 5. Streamlines at different Richardson numbers for Al$_2$O$_3$-water nano-fluid at $C = 5\%$.

Figure 6. Isotherms at different Richardson numbers for Al$_2$O$_3$-water nano-fluid at $C = 5\%$ and Re = 20.

is the dominant one. At low Richardson numbers, two vortices are produced because of hydrodynamic boundary conditions. Upper and lower walls move to the right and force fluids on the boundary layer to move to the right. Therefore, two almost symmetrical vortices are made in the upper and lower parts. With increase in Ri number, the natural convection becomes more important, but it is not dominant. Although there are two vortices at medium Richardson numbers the same as small ones, they are not symmetrical anymore. At Ri = 10, the natural convection affects the hydrodynamics of the flow and the number of vortices increases. In addition, the formation of four vortices is shown in Figure 5. Due to the effect of the moving upper and lower walls, there are two vortices suppressed to the right wall. Two other ones are the result of the buoyancy effect, which is stronger on the right side because of the higher temperature difference between lower and upper walls. The isotherms at different Ri numbers for 5\% Al$_2$O$_3$-water nano-fluid are given in Figure 6, indicating that forced convection remains dominant at small Richardson numbers. The effect of the thermal boundary layer is significant in these cases. The isotherms show that heat penetration from/into both upper and lower walls is high and the value of the isotherms is greater at Ri = 0.1 than that at Ri = 10. Another important parameter for the cavity flow is AR. The novel developed code was also employed to simulate the cavity flow with AR = 2. In this part, the nano-fluid was assumed to be ZnO-water with $C = 5\%$. The streamlines are shown in Figure 7 at different Ri numbers. Figure 7 shows that at Ri =
two almost symmetrical vortices are formed the same as the ones in Figure 5 since natural convection is negligible. At $\text{Ri} = 10$, due to the effect of gravity and buoyancy effect, two more vortices are formed, being one more than AR = 1. Due to the thermal boundary conditions at the upper and lower walls, the buoyancy effect is more powerful on the right-hand side of the cavity. Therefore, on the top right side of the cavity, the temperature of the upper wall is lower than that of the fluid and the buoyancy effect reinforces the formation of larger vortices in this part. Nu values for the upper and lower walls of the cavity are compared at different Ri numbers in Figure 8.

MgO-water nano-fluid with $C = 5\%$ is considered as our case-study nano-fluid. The effect of Ri is negligible on Nu of the upper wall since the upper wall is warm and the buoyancy cannot influence the flow regime considerably. However, at the lower wall, the temperature gradient results in higher buoyancy effect and Nu increases when the Richardson number decreases. There are some extremums in the Nu-X chart since hydrodynamic and thermal boundary conditions are very complicated and periodic. Effects of different nano-fluids and Ri number on the average friction factor ($\overline{f}$) at the upper and lower walls are shown in Figures 9 and 10, respectively. $\text{Al}_2\text{O}_3$-water and TiO$_2$-water nano-fluids have maximum and minimum $\overline{f}$ values for upper and lower walls, respectively, because the relative viscosities of the $\text{Al}_2\text{O}_3$-water and the TiO$_2$-water are maximum and minimum, respectively. $\overline{f}$ is minus for the lower wall because it moves to the right. As a result, shear stress and $\overline{f}$ are minus in the lower wall. Of note, the value of the average friction factor at the lower wall increases when the Richardson number decreases. However, at the upper wall, increase in Ri numbers elevates the average friction factor. It

Figure 7. Streamlines for free, mixed, and forced convections (ZnO-water nano-fluid at $C = 5\%$, Re = 20, and AR = 2).

Figure 8. Nu at upper and lower walls at different Ri numbers (MgO-water, $C = 5\%$, and Re = 20).

Figure 9. $\overline{f}$ at the upper wall for different nano-fluids at Re = 20 and $C = 5\%$. 
the result of the temperature boundary condition on the lower and upper walls. As mentioned previously, the hot upper wall cannot considerably influence the hydrodynamics of flow, while for the lower wall, it is completely an effective parameter in the boundary layer formation. The buoyancy tends to separate the fluid from the lower wall; therefore, at larger Ri numbers, when the natural convection is dominant, the average friction factor decreases. Average Nu values for different nano-fluids at the upper and lower walls versus Ri are demonstrated in Figures 11 and 12, respectively. The MgO-water and ZnO-water nano-fluids have maximum and minimum Nu values for the upper and lower walls, respectively, which can be attributed to the higher thermal conductivity of these nano-fluids. Nu decreases when the Richardson number increases in all cases, since natural convection indicates lower heat transfer than forced convection. Since the effect of natural convection on the upper wall is not as much as that on the lower wall, for mixed and forced convections (Ri = 0.1 and 1), the Nu at the upper wall remains almost unchanged. The effect of AR on the average Nu and friction factor of the cavity flow is shown in Table 2. The nano-fluid was assumed MgO-water with $C = 5\%$ and the Ri was 0.1. In this case, forced convection is dominant and the moving walls cannot move the fluid at the center of the cavity easily when the AR increases. Therefore, both Nu and $\bar{f}$ decrease when the AR increases.

3.4. Results of simulation of flow over a cylinder

In this section, flow over stationary and rotating cylinders is simulated. Al$_2$O$_3$, TiO$_2$, MgO, and ZnO are used as ceramic dopants in water. Streamlines and isotherms point to the flows over stationary and rotating cylinders in Figures 13 and 14, respectively. MgO-water with $C = 5\%$ is considered as a nano-fluid in this case. Streamlines and isotherms are symmetrical in terms of the stationary cylinder. No vortex is observed for forced convection flow over the

| W/L ratio | $\bar{Nu}$ | Upper wall | Lower wall | $\bar{f}$ | Upper wall | Lower wall |
|-----------|-------------|------------|------------|-----------|------------|------------|
| 0.5       | 5.423       | 12.668     | 4.398      | -4.404    |            |            |
| 1.0       | 4.270       | 12.061     | 3.035      | -3.061    |            |            |
| 2.0       | 3.582       | 10.666     | 2.258      | -2.282    |            |            |
| 4.0       | 2.659       | 8.128      | 1.589      | -1.606    |            |            |

Table 2. The $\bar{f}$ and the $\bar{Nu}$ for MgO-water nano-fluid at $C = 5\%$, $Re = 20$, and $Ri = 0.1$. 
rotating cylinder. Angular velocity of cylinder forces fluid near the cylinder to rotate around the center of the cylinder. Therefore, the vortices disappear around the rotating cylinder, compared to the stationary one. However, it should be noted that despite the stationary cylinder, the lift force acts on the rotating one due to the asymmetric pressure distribution. Streamlines and isotherms are compared for different kinds of convections in flow over the stationary cylinder in Figures 15 and 16, respectively. ZnO-water nano-
Figure 16. Comparison of isotherms at different Richardson numbers in the stationary cylinder (ZnO-water nano-fluid, Ri = 0, and C = 5%).

Figure 17. Comparison between shear stress at different Richardson numbers in the stationary cylinder (TiO$_2$-water nano-fluid and C = 5%).

Figure 18. Comparison of pressure coefficient at different Richardson numbers in the stationary cylinder (TiO$_2$-water nano-fluid and C = 5%).

Figure 19. Comparison of Nu at different Ri values in the rotating cylinder (Al$_2$O$_3$-water nano-fluid, C = 5%, and w = 10).
Combination between the mixed convection and the effect of cylinder rotation results in complex Nusselt number variation. Nusselt number is maximum at the lower part of the cylinder at smaller and medium Richardson numbers and is minimum at the upper part of the cylinder at larger Richardson numbers. The $\overline{N_u}$, calculated for four selected nano-fluids and pure water, is compared at different Richardson numbers in Figure 20. MgO-water has maximum $\overline{N_u}$ between selected nano-fluids at all Richardson numbers. Al₂O₃-water has minimum $\overline{N_u}$ at small Richardson numbers, and ZnO-water has minimum $\overline{N_u}$ at medium and large Richardson numbers.

4. Conclusions

Different models of heat convections were simulated in this work. The case studies were cavity flow and flow over stationary and rotating cylinders. Results demonstrated that when Richardson number increased, the number of vortices increased in cavity flow, but vortices vanished for flow over a stationary cylinder. Also, when Richardson number increased, $\overline{f}$ increased and $\overline{N_u}$ decreased in cavity flow. $\overline{f}$ and $\overline{N_u}$ decreased when the aspect ratio of cavity increased. Comparisons between different nano-fluids demonstrated maximum and minimum $\overline{N_u}$ values for MgO-water and ZnO-water, respectively, for flow over the rotating cylinder. When rotational speed increased, $\overline{N_u}$ and Nusselt number variations decreased on the cylinder wall.

Nomenclature

| Symbol | Description |
|--------|-------------|
| $A$    | Area        |
| $AR$   | Aspect Ratio|
| $C$    | The volume fraction of nanoparticle |
| $C_p$  | Specific heat in constant pressure (J/kg/K) |
| $D$    | Diameter (m) |
| $F$    | Friction factor |
| $\overline{f}$ | Mean friction factor |
| $Gr$   | Grashof number |
| $K$    | Thermal conductivity (W/m/K) |
| $M$    | Number of cells in x-direction |
| $N$    | Number of cells in y-direction |
| $NC$   | Thermal conductivity parameter |
| $\overline{N_u}$ | Local Nusselt number |
| $Nu$   | Mean Nusselt number |
| $Ne$   | Dynamic viscous parameter |
| $P$    | Pressure |
| $Re$   | Reynolds number |
| $Ri$   | Richardson |
| $T$    | Temperature |
| $Pr$   | Prandtl number |
| $u, v$ | $x, y$ velocity components |
| $t$    | Time |
| $U, v$ | Velocities |
| $x, y, r$ | Coordinates |

Greek symbols

| Symbol | Description |
|--------|-------------|
| $\beta$ | Thermal expansion coefficient (K⁻¹) |
| $\mu$  | Coefficient of viscosity (kg/m/s) |
| $\omega$ | Rotational speed |
| $\nu$  | Kinematic viscosity (m²/s) |
| $\rho$ | Density (kg/m³) |
| $\tau$ | Shear stress |
| $\varepsilon$ | Artificial compressibility |

Subscripts

| Symbol | Description |
|--------|-------------|
| $ref$ | Reference |
| $bf$  | Body fluid |
| $nf$  | Nano-fluid |
| $p$   | Particle |
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