INTRODUCTION

Reverse flow reactor (RFR) can regenerate the heat through periodically reciprocating flow (PRF), and it can be recognized as one of most effective ways to increase the thermal efficiency of chemical reactor. Since the concept of RFR was first introduced and utilized to purify by Cottrell in 1935, afterward, many efforts have been devoted to its theoretical and practical development. Although RFR is not new in concept, it has still been widely applied in various industrial field in recent years, such as the oxidation of volatile organic compounds (VOC), oxidation of sulfur dioxide (SO2), the synthesis of methanol, and others.
It is recognized that early scientists at Matros Technologies Inc, and Alternative Fuel Systems, Inc, applied the concept of RFR to remove pollutants from natural gas diesel engine and dual-fuel engine (DFE). Their research work demonstrated that PRF could compensate for insufficient catalyst activity with the higher temperature achieved through regenerative heat exchange inside the catalyst monolith. And PRF could completely destruct the combustible components in the exhaust gas from DFE. Afterward, they developed and tested a PRF aftertreatment device of DFE. The conversion rates of reactive HC and CO are above 95%, and the conversion rates of CH₄ are around 90% when using the PRF aftertreatment device. In the meantime, the RFR technique was also used to facilitate the regeneration of diesel particulate filter (DPF) by Konstandopoulos and Kostoglou. Their simulation results showed that PRF mechanism that could capture a high-temperature region in the DPF substrate could be capable of reducing the supplemental energy during regeneration process and thus to improve the overall thermal efficiency of the engine.

Later on, Liu et al, Zheng and Reader, and Zheng et al carries out a series of theoretical analysis and experimental study on the performance of PRF aftertreatment device. From investigation of PRF regeneration of DPF, Zheng et al concluded that the temperature gradients could defer regeneration process and result in additional thermal stress. Therefore, they proposed a PRF aftertreatment structure that two inert monolith blocks were installed at the both ends of a DPF and analyzed the effect of two inert parts on device performance. Their results indicated that the inert monolith blocks could (a) enhance the heat recuperation, (b) reduce the temperature gradient, (c) decrease the substrate temperature, (d) increase the conversion efficiency of soot oxidation, and (e) decrease dependence on the thermal properties of the substrate materials and the operating conditions. Furthermore, Zheng et al performed an experimental study on an aftertreatment system of diesel oxidation catalysts (DOCs) plus noncatalytic DPFs under PRF operations. Their results pointed out that the “light-off” temperature region could be maintained at certain locations. The critical energy efficiency could be improved with nonuniform supplemental energy distribution. Thus, it can be seen that the excellent performance of PRF aftertreatment mainly depends on the structure optimized of the device and the reasonable temperature distribution formed inside the device.

In order to further improve the performance of PRF aftertreatment, Deng et al proposed a PRF aftertreatment system consisting of a catalyzed DPF and two DOCs. In their previous study on the operation performance of the aftertreatment system, they have validated its advantages that the mixed passive-active regeneration of DPF could be realized using a small amount of external fuel, the HC and CO escaped from upstream can also be oxidized in downstream DOC, and the ash accumulated in the filter could be blown away by reverse flow. These also further demonstrated that the PRF aftertreatment system was an ideal diesel aftertreatment system and a development goal of diesel aftertreatment technology. However, the PRF regeneration modes of DPF can also lead to low filtration efficiency, new or secondary particle escaped, and the nucleation of semivolatile particle because of the influence of reverse flow disturbance on particle deposited, the rapid oxidation of particles deposited, lower temperature at inlet and outlet, and the higher temperature in chemical reaction zone.

The significant increase in PM emission levels from DPF during regeneration compared with that of non-regeneration has caused many scholars' attention. A number of studies have been conducted to investigate the influence of regeneration conditions on the PM emission performance of DPF. The various mechanisms obtained from these investigations...
could reasonably explain the increase phenomenon of PM emission from DPF during regeneration. But these past investigations main focused on the regeneration performance of traditional unidirectional flow DPF. In our current work, we will carry out study on the characteristics of the pollution emissions, the concentration and size distribution of PM emissions, the temperature distribution, and the energy efficiency of the PRF aftertreatment applied in high power diesel engine.

2 EXPERIMENTAL APPARATUS AND METHODS

In the current study, a test rig for diesel aftertreatment system with PRF is set up. The test rig mainly consists of a PRF aftertreatment system, a diesel engine, exhaust pipelines, and measurement system, as shown in Figure 1.

2.1 PRF aftertreatment system, diesel engine, and exhaust pipelines

PRF aftertreatment system is mainly composed of two DOCs, a DPF, flow pipelines, and a mixer. Two DOCs are arranged on both sides of the DPF, and a symmetrical structure of DOC+DPF+DOC is formed. The characteristics of DOCs and DPF have been described in detail in literature. In the current study, for the purpose of completeness, they are listed in Table 1. Two pairs of synchronously opening and closing valves in the flow pipelines are used to control the PRF of the aftertreatment system. The exhaust gas flows from right to left through the aftertreatment system is called the forward flow, whereas the flow from left to right is defined as the backward flow. A 30-L container is used as the mixer before the outlet. In the PRF experiment, the reciprocating flow cycle \( T_c \) is separately set at 50, 80, and 100 s. The P1-P8 measuring holes are arranged in the system and used to extract samples.

The diesel engine is a six-cylinder, turbo-charging, intercooling, and 110kW heavy-duty diesel engine. The diesel engine burns 0 # diesel oil (National V emission standard) and is operated under idle conditions. The diesel engine runs in the ambient temperature of \(-10°C\). During the experiment, the volume flow of exhaust gas \( q_v \) is, respectively, set at 260, 300, and 340 m\(^3\)/h through adjusting the throttle valve of the diesel engine.

The exhaust pipelines are set a bypass line with a 30 KW electrical heater. At the startup of the system, the bypass line is opened while the main line is closed. The electrical heater is used to heat the exhaust gas to the temperature starting the system. Meanwhile, a certain amount of propane gas is introduced at the inlet in order to initiate a thermal regeneration of DPF. As a reasonable temperature distribution in the system is established, the main line is opened and the bypass line is closed. During the experiment, 0.16 m\(^3\)/h of the propane gas is continually introduced at the inlet. When the temperatures periodically fluctuate and the fluctuating amplitude is nearly equal at a measuring position, the system is considered to be in quasisteady states.

2.2 Measurement system

In Figure 2, the K-type thermocouples T2-T14 are separately located in two DOCs at intervals of about 30 mm and the DPF at intervals of about 42 mm to measure the temperature of the substrate along the centerline. Thermocouples T1 and T15 are placed at the outsides of two DOCs in order to measure the temperature of the exhaust gas. In the following analysis of temperature distribution, we have ignored the space between DOCs and DPF.

The flow rate of the exhaust gas will be measured using a vortex flow meter, and the flow rate of the propane gas will be measured using a glass rotameter. The pressures at P3 and P6 are measured with piezoresistive pressure transmitters. The concentrations of PM emission in the exhaust gas are measured using a SMG 100 Dust Measuring Device. The measuring range of particle diameters is from 0.1 to 10 μm using Dust Measuring Device. The components of gaseous pollutants in the exhaust gas are measured using a Testo 350 flue gas analyzer. In order to measure the average value of CO, PM, NO, and NO\(_2\) components in exhaust gas under the PRF operation, the exhaust gas firstly flows through the mixer before the outlet. For different reciprocating flow cycles \( T_c \), the average values of total mass concentration of particulates and gas components are calculated from the

| TABLE 1 Characteristics of DOCs and DPF |
|----------------------------------------|
| **DOC** | **DPF** |
| Pore density/cpsi | 300 | 200 |
| Main material | Cordierite | Cordierite |
| Porosity/% | 35-40 | 50-60 |
| Wall thickness of channel/mm | 0.25 | 0.3 |
| Carrier average pore diameter/μm | 3-7 | 8-20 |
| Noble metal catalysts | Pt, Pd | Pt, Pd |
| Catalyst load/(g m\(^{-3}\)) | 706.2 | 353 |
| Catalyst carriers | Al\(_2\)O\(_3\) + Ce(Zr)O\(_2\) | Al\(_2\)O\(_3\) + Ce(Zr)O\(_2\) |
| Length/mm | 152.4 | 254 |
| Diameter/mm | 240 | 240 |
parameters continuously measured in 30 $T_c$ as the system is under quasisteady-state conditions.

The size distributions of PM are measured using sequential mobility particle sizer and counter (SMPS+C) system. The exhaust gas will be firstly extracted into a dilution device from measuring hole and is diluted with ambient air to a dilution ratio of 80 before introduced into SMPS+C system. According to literatures 31 and 32, if the total dilution ratio is higher than 60, particle agglomeration should be remarkably reduced and the measuring results will be close to the practical situation. The sampling pipe of SMPS+C system connects to the dilution device, the sampling aerosol flow is at 0.3 lpm, the sheath airflow is at 3 lpm, and the measuring range of particle diameters is from 11 to 1100 nm. The particle number and mass size distributions are calculated from the data recorded and multiplied by the dilution ratio of the exhaust gas. The mean value of particle number and mass size distribution will be the average of three consecutively measured numbers.

As the exhaust gas flows through the exhaust pipe and the aftertreatment system from the diesel engine, there are the nucleation of particle inception, the surface growth of very small particles, and the coalescence and agglomeration of small particles.33 These formation processes of particulates not only depend on the working conditions of the diesel engine but also are determined by the temperature, the pressure, and the gas component of the exhaust gas in the exhaust pipe. Therefore, in the current study, the number size distribution, mass size distribution and total mass concentration of particulates, and gas components at from P1 to P8 measuring points are measured and they are used to analyze the evolution features of PM emission under different operation conditions.

### 2.3 Uncertainty analysis of experimental results

The uncertainties of the experiment measurements are mainly dependent on the experimental conditions and the measurement instruments. The specifications of the measurement instruments used in the experiment have been described in detail in literature 22. In the current study, for the purpose of completeness, they are listed in Table S1.

An uncertainty analysis could be performed for the experimental data using the propagation of error method described by Moffat and Giechaskie et al.34,35 The uncertainty analysis processes are described in Table S2.

### 2.4 Energy efficiency analysis

In order to make an energy efficiency comparison under different operating conditions, the specific energy consumption $\beta$ is defined the Equation (1) according to the idea of reference.36

$$\beta = \frac{m_g c_{p,g} (T_s - T_g)}{H_f} \times 100\%$$

where $m_g$ is the mass flow rate [g/s], $c_{p,g}$ is the specific heat of gas at constant pressure [J/kg K], $T_s$ is the average temperature of substrate in DPF [K], $T_g$ is the temperature of exhaust gas at inlet [K], and $H_f$ is the external supplemental energy [W].

The critical energy efficiency $\eta_c$ is defined the Equation (2) according to reference 16.

$$\eta_c = \frac{\Delta E_a}{\Delta E_a + \Delta E_{sup}} \times 100\%$$

where $\Delta E_a$ is the energy consumed by the diesel engine under the critical threshold gas condition, and $\Delta E_{sup}$ is the external supplemental energy required to achieve critical threshold gas property.

In the long-running operations under PRF operating conditions, there are not the accumulation of ashes in the filter as we observe the section of DPF, as shown in Figure S1. This further validated the ash accumulated in the filter could be blown away by reverse flow under PRF operating conditions. Therefore, the PM filter efficiency $\alpha_{PM}$ is defined the Equation (3).

$$\alpha_{PM} = \frac{c_{8PM}}{c_{2PM}}$$

where $c_{2PM}$ and $c_{8PM}$ are, respectively, the mean concentrations of PM at P2 and P8 measuring points.
The CO conversion rate $a_{CO}$ is defined the Equation (4).

$$a_{CO} = \frac{c_{8CO}}{c_{2CO}}$$  \hspace{1cm} (4)

where $c_{2CO}$ and $c_{8CO}$ are, respectively, the mean concentrations of CO at P2 and P8 measuring points.

3 | RESULTS AND DISCUSSIONS

3.1 | Characteristics of unidirectional flow

Figure 3 shows the particle size distribution at P1, P2, and P7 measuring points as the exhaust gas flows the main line and $q_v$ is 260 m$^3$/h under unidirectional operation condition. Left longitudinal coordinate is particle number concentration while right longitudinal coordinate is particle mass concentration in Figure 3.

The idealized diesel exhaust particle size distributions are that most of the particle mass exists in the accumulation mode in the 0.1-0.3 μm diameter range while the nuclei mode in the 0.005-0.05 μm diameter range contains 1%-20% of the particle mass and more than 90% of the particle number. In the current experiment, the exhaust gas flows from exhaust port to P1 through an exhaust pipe, the temperature of the exhaust gas at P1 drops to 120°C, and most of nuclei mode particles have been accumulated into mode particles through coalescence and agglomeration at P1. Therefore, the particle number size distribution is a bi-modal curve which higher peak locates at a particle size of 190 nm and lower peak locates at a particle size of 349 nm. The particle number size is mainly distributed within 75-398 nm. The particle mass size distribution is also a bi-modal curve which lower peak locates at a particle size of 243 nm and higher peak locates at a particle size of 349 nm. The particle mass size is also mainly distributed within 75-398 nm. Both number and mass sizes of particulates in

![Size distribution of PM at P1 and P2](image)

![Size distribution of PM at P2 and P7](image)
the exhaust gas lie in the accumulation size range that is 50 nm < Dp < 1000 nm.

The temperature of the exhaust gas transitions from 120 to 80°C as the exhaust airflows directly from P1 to P2. Because the exhaust gas cools down, some of small size particles are converted into larger size particles through collision, coalescence, and agglomeration. It can be seen clearly from Figure 3A that both number and mass distribution ranges of particle size slightly become narrow and shift toward larger size particles, the position of two peaks is almost unchanged and the peak values increase at P2 compared with that at P1.

As the exhaust gas flows the main line, the mean components of O₂, CO, NO, NO₂, and PM in the exhaust gas at different measuring points under unidirectional operation condition are listed in Table 2. Total mass concentration of particle increases up from 140 to 197 mg/m³ when the exhaust gas flows from P1 to P2 in Table 2.

When the exhaust gas is kept at about 80°C at P2, the temperature is lower than the light-off temperatures of PM, hydrocarbons, and CO. Only a small number of the hydrocarbons and CO are oxidized into H₂O and CO₂ under the action of bimetallic Pt/Pd catalytic agent and strong oxidant NO₂ in DOCs and DPF. Therefore, the mean component of CO and NO₂ decreases at P7 as listed in Table 2. As the exhaust gas flows through DPF, most of large PM are directly removed by filter. Therefore, total mass concentration of particle is reduced to 45.8 mg/m³ from 197.1 mg/m³ as exhaust gas flow from P2 to P7 in Table 2. But some of ultrafine particles can seep through the DPF and most of aromatic hydrocarbons as particulate precursor in exhaust gas can also be carried through the DPF. As a result, new particles are formed from the aromatic hydrocarbons through nucleation while the ultrafine particles are agglomerated into larger particles in second DOC under the cooling condition. Furthermore, the particle size distribution shifts to large size compared with that at P2 and the range of particle number and mass size distribution is mainly within 349-1100 nm at P7, as illustrated in Figure 3B. These further illustrate that the PMs detected are formed from the agglomeration of ultrafine particles and is belong to second particles. Thus, it can be seen, as the temperature is lower than the light-off threshold of combustible gas in the exhaust gas, the aftertreatment system can only remove the PM existed in the exhaust gas and cannot restrain the formation of second particles in atmospheric environment, under the low-temperature condition of unidirectional flow.

### Table 2: Components of exhaust gas at various measuring point

| Measuring Point | O₂/% | CO/ppm | NO/ppm | NO₂/ppm | PM/mg/m³ |
|-----------------|------|--------|--------|---------|----------|
| P1              | 18.0 | 226.9  | 162.0  | 124.0   | 140.4    |
| P2              | 17.9 | 226.7  | 169.0  | 119.0   | 197.1    |
| P3              | 17.9 | 221.8  | 173.7  | 114.3   | 265.3    |
| P7              | 17.8 | 171.92 | 178    | 105.0   | 45.8     |

3.2 Influence of temperature distribution

Figure 4 shows the symmetrical temperature distribution curves during a reciprocating flow cycle as q_v is 260 m³/h, fuel supplement is 0.16 m³/h of C₃H₈ at the inlet, the temperature at P2 is 80°C, and T_s is 50 s. Figure 5 shows the asymmetrical temperature distribution curves during a reciprocating flow cycle under the same conditions as that in Figure 4. The mean components of O₂, CO, NO, NO₂, and PM in the exhaust gas at P2 in the two cases of symmetrical and asymmetrical temperature distribution are same as that in Table 2. For the two cases, utilizing a small amount of external fuel, the combined effects of the reversal flow, exothermic reaction, and solid heat retention result in a temperature distribution which exists a high-temperature region in the middle region and has high-temperature gradients on both sides in the aftertreatment system as shown in Figures 4 and 5.

As the initial high temperature is located in DPF, the symmetrical temperature distribution will be formed in the aftertreatment system under quasisteady states with PRF operation. The symmetrical temperature distribution is a parabola profile that the DPF is always kept at higher temperature and each DOC has a temperature gradient as is shown in Figure 4. The processes of alternately storing and releasing heat result in the periodic fluctuation of the temperature in the same location with PRF operation as is shown in Figure S2. The dividing point between endothermic regions and exothermic regions during the forward and backward flow is located in the same position, that is about X = 0 mm. From another perspective, during a forward flow operation the thermal wave propagates downstream and conversely for a backward flow operation, both following the directions of the exhaust gas flow. Thus, temperature distribution profile shifts toward the rear of the system during every half-cycle as shown in Figure 4A,B.

As the initial high temperature is located in left DOC or right DOC, the heat transfer between DOC and DPF is not achieved by heat conduction due to the existence of the space between them. The exothermic reaction as well as storing heat and releasing heat of the substrate could only been maintained in near initial high-temperature region during PRF operation. Therefore, an asymmetrical temperature distribution will be formed in the aftertreatment system under quasisteady states during PRF operation. An asymmetrical temperature distribution is formed because the initial high temperature is located in left DOC, as is shown in Figure 5. The asymmetrical temperature distribution is a trapezoidal profile which exists
a high-temperature region within $-218 \text{ mm} \leq X \leq 45 \text{ mm}$ and has high-temperature gradients on both sides. The range of the high-temperature region in the case of asymmetrical temperature distribution is obviously larger than that in the case of symmetrical temperature distribution. The processes of alternately storing and releasing heat separately exist within four regions and the dividing points of the endothermic and exothermic regions separately locate at $X = -220 \text{ mm}$, $X = -120 \text{ mm}$, and $X = 10 \text{ mm}$.

For the two cases, the CO conversion rate $\alpha_{\text{CO}}$, the PM filter efficiency $\alpha_{\text{PM}}$, and the specific energy consumption $\beta$ are listed in Table 3. Based on above analysis, the symmetrical temperature distribution could form better mechanism of heat recuperation, and it has higher specific energy consumption $\beta$, as shown in Table 3.

Figure 6 shows the fluctuating curves of the total mass concentration of PM in the two cases of symmetrical and asymmetrical temperature distributions as $T_c = 50 \text{ s}$. For symmetrical temperature distribution, the DPF is always kept at higher temperature than the lighting off temperature of soot oxidation. And each DOC has a temperature range from 80 to 450°C, most of the time the temperature is lower
than 400°C in the every half-cycle. Therefore, the chemical reaction equilibrium of NO₂ and NO is on the NO₂ side in the every half-cycle. In upstream DOC, the hydrocarbons and CO in exhaust gas are oxidized into H₂O and CO₂ during every half-cycle. Most of the particles attached on the walls of DPF passages are completely oxidized with the NO₂ catalytic action. The little incombustible residue as ash deposits on the walls. The HC and CO from incomplete oxidation conversion of soot are further oxidized into H₂O and CO₂ in downstream DOC. As switching to next a half-cycle from a half-cycle, the flow resided in the aftertreatment system carries PM suspended and flows reversely out. This includes the flow pipelines, the DOC channels, and the inlet passages of DPF at the end of the previous half-cycle before switching to next a half-cycle. The ashes deposited on the walls are also blown away the walls and carried out the aftertreatment system by the reverse flow. Therefore, as switching to next a half-cycle, the PM content rapidly increases to a peak value and then rapidly decreases. The content of PM periodically fluctuates, and the fluctuating amplitude is nearly equal for symmetrical temperature distribution, as shown in Figure 6A.

For asymmetrical temperature distribution, the DPF is also always kept at higher temperature than the lighting off temperature of soot oxidation. But temperature is in a range from 80 to 845°C, the temperature in the part region is higher than the 550°C in left DOC. Therefore, the chemical reaction equilibrium of NO₂ and NO is on the NO side in left DOC. The temperature is in a range from 80 to 300°C, most of the time the temperature is lower than 200°C in the every half-cycle in right DOC. Most of the time the conversion from NO to NO₂ could not been achieved at the temperature lower than 200°C in right DOC. Most of soot particles attached on the internal walls of the DPF are oxidized with O₂ alone and without the NO₂ catalytic action during every half-cycle. A small amount of particles incompletely oxidized deposit and reside on the internal walls of the DPF during every half-cycle. Every time switching to next a half-cycle, the flow resided in the aftertreatment system reversely flows out the

FIGURE 6 Total mass concentration of PM at P7 for Tc = 50 s. A, Under symmetrical temperature condition. B, Under asymmetrical temperature condition

FIGURE 7 Size distribution of PM at P8 for symmetrical and asymmetrical temperature distribution
system. Subsequently, the residual particles deposited on the walls of the DPF during the previous half-cycle are stirred and gradually carried out by the reverse flow during the half-cycle. Therefore, for asymmetrical temperature distribution, as following the flow direction switch the PM content periodically fluctuates is same as that of symmetrical temperature distribution, but the fluctuating amplitude is different between forward flow cycle and backward flow cycle and every fluctuating amplitude is higher than that of symmetrical temperature distribution, as shown in Figure 6B.

Figure 7 shows the distribution curves of the particle size at P8 in the two cases of symmetrical and asymmetrical temperature distributions as $T_c = 50$ s. The mean components of O$_2$, CO, NO, NO$_2$, and PM in the exhaust gas at P8 in the two cases of symmetrical and asymmetrical temperature distribution are listed in Table 4.

For symmetrical temperature distribution, most of soot particles attached are oxidized and the continuous regeneration is realized in the DPF. Most of aromatic hydrocarbons as particulate precursor in the exhaust gas is also catalytically oxidized in the DOCs. The formation of second particles from nucleation, coalescence, and agglomeration can be effectively restrained. The particles escaped are mainly from the reverse flow resided in the aftertreatment system as switching to next a half-cycle from a half-cycle. Therefore, compared with the size distribution at P2 in Figure 3, the peak value of particle number is reduced by three orders of magnitude, while the peak value of particle mass is reduced by two orders of magnitude, as shown in Figure 7. The ranges of particle size distribution obviously narrow and shift toward smaller size particles. Total mass concentration of particle changes from 197.1 to 38 mg/m$^3$ as the exhaust gas flows from P2 to P8 across the aftertreatment system as listed in Tables 2 and 4.

According to the characteristics of asymmetrical temperature distribution, the particles deposited on the internal walls could not be completely oxidized in the system. The particles escaped are not only from the reverse flow resided in the aftertreatment system but also from the particles deposited on the internal walls. Therefore, compared with the symmetrical temperature distribution in Figure 7, the peak values of particle number and mass obviously increase, the ranges of particle size distribution widen and shift toward larger size particles. The PM filter efficiency $\alpha_{PM}$ decreases compared with the symmetrical temperature distribution, as shown in Table 4. Total mass concentration of particle changes from 197.1 to 38 mg/m$^3$ as the exhaust gas flows from P2 to P8 through the aftertreatment system as listed in Tables 2 and 4.

For PRF operation, the downstream DOC can be used to oxidize HC and CO produced from incomplete oxidation conversion of soot during every half-cycle. Therefore, CO conversion rate is high and CO emission is also kept at low level in both cases, as listed in Tables 3 and 4.

It can be seen that the symmetrical temperature distribution conditions can not only sustain above “light-off” temperature region in DPF, but also maintain high-temperature gradients in two DOCs. Thus, the symmetrical temperature distribution not only can ensure the catalyst coating having the best performance but also can reduce thermal stress in DPF. Compared with the asymmetrical temperature distribution under the same conditions, the aftertreatment system formed the symmetrical temperature distribution has better performance of trapping particles and regeneration.

### 3.3 Influence of reciprocating flow cycle

Figure 8 shows the symmetrical temperature distribution profiles in the aftertreatment system during forward half-cycles as $q_i$ is 260 m$^3$/h, fuel supplement is 0.16 m$^3$/h of C$_3$H$_8$ at the inlet, the temperature at P2 is 80°C, and $T_c$ is 50 s, 80 s and 100 s, respectively. As $T_c$ is 50 s, 80 s and 100 s, respectively, the CO conversion rate $\alpha_{CO}$, the PM filter efficiency $\alpha_{PM}$ and the specific energy consumption $\beta$ are listed in Table 5.

For the case of every reciprocating flow cycle, the temperature distribution curve at the end of previous backward half-cycle is almost same as that at the beginning of the next forward half-cycle, as shown in Figure 4, Figure S3, and Figure S5. For the case of every reciprocating flow cycle, the heat released heat solid substrate and forms thermal wave in DOCs and DPF after the chemical reactions are lighted-off at the beginning of every half-cycle. Subsequently, the thermal wave propagates downstream during the half-cycle. The thermal wave propagates

| TABLE 4 | Components of exhaust gas at P8 for symmetrical and asymmetrical temperature distribution |
| --- | --- | --- | --- | --- |
| State | O$_2$/% | CO/ ppm | NO/ ppm | NO$_2$/ ppm | PM/ mg/m$^3$ |
| Symmetry | 17.4 | 13.7 | 245.9 | 20.0 | 15.0 |
| Asymmetric | 17.7 | 19.1 | 207.7 | 5.0 | 37.9 |

![FIGURE 8](image)
downstream nearer to the outlet of downstream DOC, the temperature at the outlet of downstream DOC becomes higher, and the heat loss is more during the half-cycle. Therefore, the temperature in the high-temperature region gradually decreases along with temperature distribution profile shifting toward downstream during the half-cycle, as shown in Figure 4, Figure 8, Figure S3, and Figure S5.

With $T_c$ increasing, because the time of thermal wave propagation downstream increases during a half-cycle, the moving distance toward downstream of the temperature distribution profile increases during the half-cycle and the high-temperature zone in the profile slightly becomes wider. As $T_c$ increases up to 80 s from 50 s, the temperature of each DOC outlet at the end of every half-cycle slightly increases corresponding to the time increases in thermal wave propagation downstream as shown in Figure 8. Therefore, the heat loss increases slightly comparing with that at $T_c = 50$ s, and the specific energy consumption $\beta$ changes little comparing with that at $T_c = 50$ s, as shown in Table 5. As $T_c$ increases up to 100 s from 80 s, the temperature of left DOC outlet at the end of backward half-cycle significantly increases corresponding to the time increases in thermal wave propagation downstream, and the fluctuating amplitudes of temperature increase in the same location, as shown in Figure S4, Figure S5, Figure S6, and Figure S7. Therefore, the heat loss increases and the temperature in high-temperature region of the system at $T_c = 100$ s decreases comparing with that at $T_c = 80$ s, and as shown in Figure 8. The specific energy consumption $\beta$ decreases significantly, as shown in Table 5.

Figure 9 shows the distribution curves of the particle size at P8 as $T_c$ is 50 s, 80 s, and 100 s, respectively. Left longitudinal coordinate is particle number concentration while right longitudinal coordinate is particle mass concentration in Figure 9. As $T_c$ is 50 s, 80 s, and 100 s, respectively, the mean components of $O_2$, $CO$, $NO$, $NO_2$, and PM in the exhaust gas at P8 are listed in Table 6.

Temperature in central zone of DPF is always kept at higher temperature than the lighting off temperature of soot oxidation in all three cases of $T_c = 50$ s, 80 s, and 100 s, even if the temperature fluctuates during every half-cycle. Therefore, the oxidation conversation rate of soot particles captured in DPF almost is not affected by $T_c$ increasing. Nevertheless, because the time that downstream DOC is kept at low temperature increases during a half-cycle, and thus, the oxidation conversation rates of aromatic hydrocarbons and CO decrease in the downstream DOC with $T_c$ increase. Therefore, the amount of aromatic hydrocarbons escaped from the upstream of the system increase, and new relatively small PMs formed through nucleation increase in the downstream DOC under the cooling condition. This results into that the number of relatively small size particle increases with $T_c$ increase as $T_c$ increases up to 100 s from 80 s, the temperature of left DOC outlet at the end of backward half-cycle significantly increases corresponding to the time increases in thermal wave propagation downstream, and the fluctuating amplitudes of temperature increase in the same location, as shown in Figure S4, Figure S5, Figure S6, and Figure S7. Therefore, the heat loss increases and the temperature in high-temperature region of the system at $T_c = 100$ s decreases comparing with that at $T_c = 80$ s, and as shown in Figure 8. The specific energy consumption $\beta$ decreases significantly, as shown in Table 5.

**TABLE 5** $α_{CO}, α_{PM}$, and $\beta$ for $T_c = 50, 80,$ and 100 s

| Symmetric cycle | $α_{CO}$ | $α_{PM}$ | $\beta$ |
|-----------------|----------|----------|---------|
| 50 s            | 93.96%   | 92.38%   | 124.22% |
| 80 s            | 88.53%   | 88.93%   | 124.19% |
| 100 s           | 83.41%   | 88.64%   | 117.69% |

**FIGURE 9** Size distribution of PM at P8 for $T_c = 50, 80,$ and 100 s

**TABLE 6** Components of exhaust gas at P8 for $T_c = 50, 80,$ and 100 s

| Symmetric cycle | $O_2$/% | $CO$/ppm | $NO$/ppm | $NO_2}$/ppm | PM$/mg/m^3$ |
|-----------------|---------|----------|----------|-------------|-------------|
| 50 s            | 17.4    | 13.7     | 245.9    | 20.0        | 15.0        |
| 80 s            | 17.5    | 26.0     | 254.2    | 18.1        | 21.8        |
| 100 s           | 17.5    | 37.6     | 251.1    | 17.6        | 22.4        |
shown in Figure 9. Conversely, with $T_c$ increase the frequency of flow direction switching decreases, the mean number of the particles carried in the reverse flow has a certain amount of decrease. This results into that the number of relatively big size particle decreases with $T_c$ increase as shown in Figure 9. The mass fraction of the relatively big size particle carried in the reverse flow is higher than the relatively small size particle formed through nucleation in the exhaust gas. Therefore, the mass of relatively big size particle decreases with $T_c$ increase and particle mass size distribution region widens as shown in Figure 9. Based on the above reasons, the number peak value of particle size increases, the mass peak value of particle size decreases, and particle size distribution shifts toward less size particles at P8 with increasing $T_c$, as listed in Table 6.

### 3.4 Influence of volume flow of exhaust gas

Figure 10 shows the symmetrical temperature profiles in the aftertreatment system during forward flow cycles as $T_c$ is 50 s for $q_v = 260, 300,$ and $340 \text{ m}^3/\text{h}$

![Temperature profiles as $T_c = 50$ s for $q_v = 260, 300,$ and $340 \text{ m}^3/\text{h}$](image)

**TABLE 7** $\alpha_{CO}, \alpha_{PM}$ and $\beta$ for different volume flow

| Volume flow/m³/h | $\alpha_{CO}$ | $\alpha_{PM}$ | $\beta$ |
|------------------|---------------|---------------|--------|
| 260              | 93.96%        | 92.38%        | 124.22%|
| 300              | 86.83%        | 87.82%        | 137.91%|

With $q_v$ increasing from 260 to 300 $\text{ m}^3/\text{h}$, the high-temperature zone in the system slightly narrows and temperature peak value slightly decreases. Therefore, both CO conversion rate $\alpha_{CO}$ and the PM filter efficiency $\alpha_{PM}$ decrease with increasing $T_c$ as listed in Table 5. Total mass concentration of particle emissions slightly increases, and the mean concentration of CO emissions increases at P8 with increasing $T_c$, as listed in Table 6.

**CONCLUSIONS**

An experimental study is carried out to investigate the characteristics of the pollution emissions, the concentration and size distribution of PM emissions, and the temperature distribution of a PRF aftertreatment system applied in high power diesel engine. The effect of $T_c$ and $q_v$ of exhaust gas on the performance of the aftertreatment system is discussed in a systematic manner. The energy efficiency the aftertreatment

**TABLE 8** Components of exhaust gas at P2 for different volume flow

| Volume flow/m³/h | $O_2$/% | CO/ppm | NO/ppm | NO₂/ppm | PM/mg/m³ |
|------------------|---------|--------|--------|---------|----------|
| 260              | 17.9    | 226.7  | 169.0  | 119.0   | 197.1    |
| 300              | 16.7    | 142.0  | 208.1  | 20.3    | 317.0    |
TABLE 9 Components of exhaust gas at P8 for different volume flow

| Volume flow/m³/h | O₂%/ppm | CO/ ppm | NO/ ppm | NO₂/ ppm | PM/ mg/m³ |
|------------------|----------|---------|---------|----------|-----------|
| 260              | 17.4     | 13.7    | 245.9   | 20.0     | 15.0      |
| 300              | 17.1     | 16.1    | 245.6   | 17.6     | 38.6      |

system is also analyzed. The main findings from the current work could be summarized as follows:

1. As the temperature of substrate is lower than the light-off threshold of combustible gas in the exhaust gas, the aftertreatment system could only remove the PM existed in the exhaust gas and could not restrain the formation of second particles in atmospheric environment, under the low-temperature condition of unidirectional flow.

2. The aftertreatment system has excellent performances of trapping particles and regeneration as the symmetrical temperature distribution is formed. The PM filter efficiency $\eta_{PM}$ is 92%, and the specific energy consumption $\beta$ is 124% for symmetrical temperature distribution.

3. The increase in $T_c$ leads to the shifting of the temperature profile toward downstream, which would affect the particle number and mass size distribution as well as the characteristic of PM and CO emissions.

4. A certain increase in volume flow of exhaust gas could not alter the excellent temperature distribution characteristics.

5. The critical energy efficiency $\eta_c$ of the system could reach 96.61%.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.

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