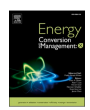
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Direct numerical simulation of packed and monolith syngas catalytic combustors for micro electrical mechanical systems

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ABSTRACT

In this work, a catalytic combustor for micro electrical mechanical system for syngas was designed and analysed using Direct Numerical Simulation (DNS) in conjunction with finite rate chemistry. The effect of catalyst (platinum (Pt), palladium (Pd), palladium oxide (PdO), and rhodium (Rh)), bed type (packed with twelve catalyst shapes and four catalyst monolith), shapes (packed: cylinder, hollow cylinder, four cylinder, single cylinder, single cylinder, cross-webb, grooved, pall-ring, hexagonal, berl-saddle, cube, intalox-saddle, and sphere, monolith: triangular, rectangular, hexagonal, and circular), and operating conditions (inlet temperature and velocity, fuel/air ratio, different concentrations $\mathrm{CH_4-H_2-CO}$) on combustion efficiency and pressure drop were studied using different parameters (combustion efficiency (η), pressure drop, effectiveness factor (Ψ), and fuel conversions ($\mathrm{H_2}$ and $\mathrm{CH_4}$ conversions)). Analysis under different operating conditions reveals that the designed combustor can operate effectively with syngas of varying compositions with a high combustion efficiency of over 85%. Combustion mainly takes place on the surface of the catalyst without gas phase reaction with pressure drops between 18 Pa and 155 Pa. The intalox saddle shape catalysts resulted in the bed effectiveness factor 0.93.\(^1\) The Damköhler for hydroxyl radicals (OH) over the entire length of the reactor is uniformly distributed and well below 3, suggesting uniform combustion.

Introduction

The worldwide demand for generating clean energy necessitates the application of renewable fuels and novel combustion systems [1]. This has spawned a tremendous research effort for microburners as they are compact and efficient for occurrence of combustion under the desired conditions [2]. An effective design of microburners is needed to ensure the autothermal operation of these systems with clean alternative fuels. The catalytic combustors emerge as one of the promising technologies for the occurrence of the combustions in microchannels [3]. They do provide a higher surface area per reactor volume for the chemical reactions, thereby broadening the stability limits [4], facilitating the ignition at low temperatures [5], and suppressing the flame instabilities [6]. Indeed, catalytic combustors enable clean combustion of both low and high calorific values on the surface of the catalyst. The small size of these systems improves the conjugate heat transfer from the reactor body to the preheat zone, stablising the flame [7]. The combustion in

catalyst combustors is shared between the gas phase reactions and those under the control on the surface of catalyst. This technology can alter the combustion chemistry by selective adsorption of materials, oxidising the radical on solid surfaces, and desorbing the formed stable gases, thereby limiting the homogeneous reactions [8]. The microburner can also be utilised in chemical reactors, hydrogen generators [9], thermoelectric [10] and thermophotovoltaic [11]. They could provide a reliable source of heat for endothermic processes such as steam reforming and ammonia decomposition when integrated with other microchannel systems to produce fuel cell applications [12].

Conventionally, the occurrence of homogeneous combustion at large scale was used to generate energy [13]. The homogeneous reactions of renewable fuels at the industrial scale are usually challenging as the clean alternative fuel characteristics are usually different from the conventional ones with some like biogas having lower calorific values [14] and the others like hydrogen having wide flammability range and high reactive propensity [15]. These different extreme characteristics of the alternative fuels are limiting factors in the occurrence of

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¹ An ideal reactor effectiveness factor is 1 which 0.5 is equivalent to 100% combustion efficiency and 0.5 is from 0% pressure drop at the combustor outlet.

Nome	nclature	ν	Kinematic viscosity [m ² .s ⁻¹]
		ω	Bed effectiveness components
Symbol	ls	Ψ	Bed effectiveness factor
Α	surface area [m ²]	Γ	Diffusivity coefficient [m ² .s ⁻¹]
Da	Damköhler number	C. I	
D	Diameter [m]	Subscrip	
E	Equivalent emission	1	inlet
E_f	Fluid energy [J. kg^{-1}]	i, j, k	Species indices
H	Enthalpy [J. mol^{-1}]	eff	effective
h_o	Heat transfer coefficient [W/m ² K]	ave	average
I	Unit matrix	w .	wall
J	Molecular diffusion flux	w, i	wall inlet
\overrightarrow{J}	Diffusion flux vector	w, o	wall outlet
K	thermal conductivity [W/m K]	Superscr	int
k_n	Knudsen number	h	enthalpy [J]
L	length [m]	T	transpose
l*	characteristic length [m]	-	a anopose
P	pressure [pa]	Abbrevia	ution
Q	Thermal flux [W/m ²]	CVODE	C-language Variable-coefficients
R	Production rate of chemical reactions [mol. m^{-3} . s^{-1}]	DEM	Discrete element method
R	Coefficient of determination	DNS	Direct numerical simulation
S	catalytic molar production [mol. m ⁻² . s ⁻¹]	EDC	Eddy Dissipation Concept
S	source term [mol. s ⁻¹]	ER	Equivalence Ratio
Re	Reynolds number	FSI	Fluid Solid Interactions
Sc	Schmidt number	HDPE	High density of polyethylene
T	Temperature [K]	LDPE	Low density polyethylene
\overrightarrow{u}	Velocity vector [m.s ⁻¹]	MEMS	Micro-Electro-Mechanic-Systems
X	Mole fraction	NO_X	Nitrogen oxides
Y	Mass fraction	ODE	Ordinary differential equation
1	Mass fraction	OH	hydroxyl radicals
Greek l	letters	Palladiu	
α	Inertial resistance	Palladiu	ım oxide Pd-O
ρ	Density [kg. m ⁻³]	Platinun	n Pt
λ	Mean free path [m]	PET	Polyethylene terephthalate
η	Combustion efficiency	Rhodiun	
M	Fluid viscosity [kg. m ⁻¹ .s ⁻¹]	PS	Polystyrene
θ	coverage	UHC	Unburned hydrocarbons
σ	Site occupancy	VOC	Volatile organic compounds

homogeneous combustion in the majority of practical systems (i.e. boiler, gas turbine) [16], making the conventional combustor designs inept for green fuels. The problems associated with clean alternative fuels are severe in the small-scale micro combustors, purposefully designed to save energy and minimise combustion heat loss. If not carefully designed, the small size of the microcombustors hampers the occurrence of ignition by instigating the radical recombination, extinguishes the flame appearance by dissipating the combustion heat [11]. It can also lead to the blow-off if the outlet velocities transcend the fuel burning velocity [17]. They may also lead to pollutant formation if hydrocarbon chain branching reactions occur without external control, as in conventional combustion systems [18]. Overall, combustion is more under control on the solid surface than in a homogeneous mixture where the chain branching reactions spontaneously take place without any hindrance.

Previous experimentation confirmed that the flame propagation at the microscale is feasible [19] by the interplay among the kinetic and transport of species [20]. Different reactors such as: porous, fixed and fluidised beds have been designed to enhance combustion and diminish the technical problems associated with the micro-scale microburners. Swiss roll burners [21] and porous media are widely used for the catalytic combustion of fuels. The application of renewable fuels, due to their different physical characteristic, for maintaining NOx emissions, effective flow distribution, the combustor's ability to light, tolerate

sudden changes in power requirement, efficient combustion, and the achievement of the required life is possible at microscale MEMS combustion systems by a design of catalytic microcombustor. In this work, a new catalytic packed bed reactor for combustion of syngas fuel is designed and analysed in terms of combustion performance and effectiveness under various loads and variables. The operation of the designed reactor is also investigated when loaded with different packings and catalyst monoliths.

Today, the application for carbon capture technologies and renewable energy are at the same order of importance for smooth transition from hydrocarbon based resources [22]. Syngas can be synthesized from variety of feedstocks through different processes [23] and can work as a future fuel and an important hydrogen career for transition from non to renewable energy sources. This work will facilitate the application of hydrogen careers and syngas in small combustors by presenting a novel procedure for design of a catalytic combustor. Sometime, fuel purification could be challenging and production of 100% hydrogen from the gasification process won't turn out techno-economically feasible. In these cases, the utilisation and direct combustion of syngas can be a wise choice for the future design and application of renewable fuels.

Catalytic combustor

A purified syngas gas mixture with 92 wt% methane and 8 wt%

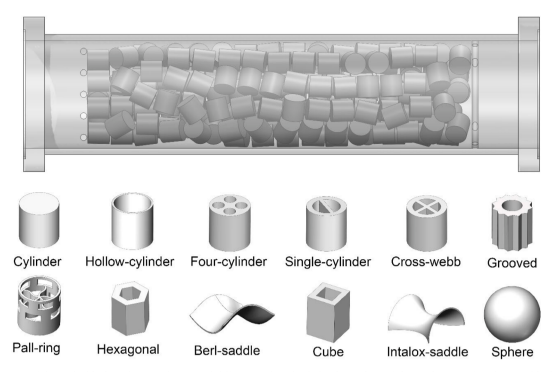


Fig 1. Schematic of the packed bed catalytic combustor and different packing loads for the analysis of the combustion of methane and hydrogen fuel.

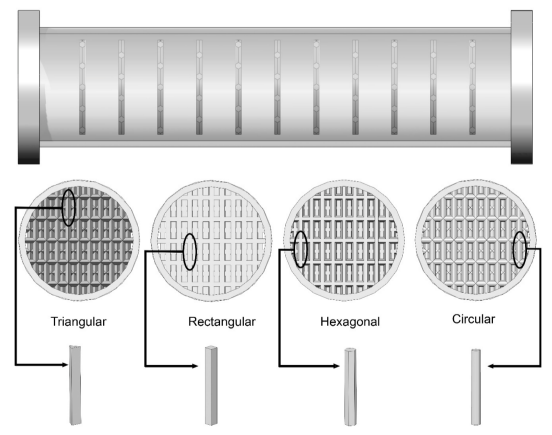


Fig 2. The schematic of catalyst monolith loaded with different catalytic monolith used to load the reactor for catalytic combustion of methane and hydrogen fuel.

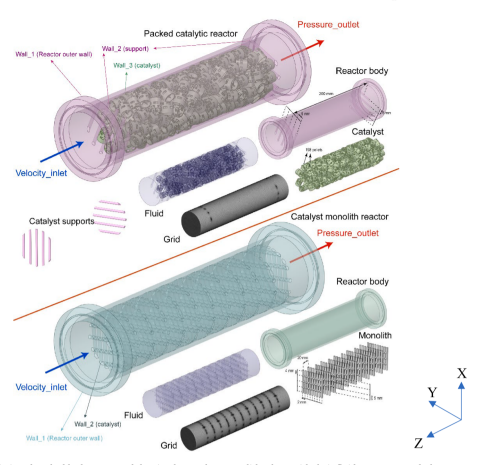


Fig 3. Schematic of the designed packed bed reactor and the circular catalyst monolith, along with their fluid parts extracted, the computational domain made from the reactors fluid, and the boundary conditions in the numerical simulation.

hydrogen was first as a baseline fuel to design the reactor, although syngas with CO and different compositions of species is considered in the sensitivity analysis for show the ability of the combustor to operate with different grades of syngas. This (CH₄: 92 wt% and H₂: 8 wt%) is a baseline which is obtained from the gasification of waste plastic and polymers. The waste material is considered a mixture of polystyrene (PS), high density of polyethylene (HDPE), low density polyethylene (LDPE) and Polyethylene terephthalate (PET) with identical mass composition (each 25%). It is assumed that the CO is fully converted to CH₄ using water–gas shift reaction before considered as a fuel for the catalytic combustor.

The dimensions of the reactor were obtained based on the complete combustion of the fuel/air mixture under equivalence ratio of 0.99 and inlet velocity of 1 ${\rm ms}^{-1}$ for the premixed mixture syngas/air. This design characteristic will the give the input Re number of 2500. It was envisaged that complete combustion is achieved at the first 1/3 entire length of the combustor with the least pressure drop possible and heat loss. The surface/volume ratios in microburners are usually high [24] making them susceptible to heat loss and low thermal efficiency [25]. The design objective was to control the gas phase reactions so the main contribution in the combustion was from the surface catalyst reactions.

Reactor dimensions

A 220 mm long cylindrical container with an internal diameter of 50 mm was considered for fuel combustion. The thickness of the reactor wall was 3 mm. Two beds support were considered at distances of 20 $\,$

mm from the reactor inlet and outlet planes. The bed supports are five rows of cylinders, 3 mm in diameter each located at ZX plane near the reactor inlet plane and ZY plane near the reactor outlet plane, respectively. The center-to-center distance of the bed support cylinder was 10 mm. They are well able to limit the movement and transverse of pellets, keeping them at the very middle parts of the combustor. Around 200 pellets (exactly 195) can be placed in the designed reactors. The length and diameter of pellets are both 10 mm. Different shapes for the pellets are assumed to fill the reactor within two bed supports. Fig. 1 gives the schematic of the designed catalytic reactor filled with cylinder types of pellets and other shapes considered for the investigations.

For catalytic monolith reactors, the same cylinder for catalytic beds is exploited to form the body of the combustor. Four different monoliths, namely rectangular, triangular, hexagonal and circular, were employed for the analysis of the combustion, Fig. 2. The circular monolith had a diameter of 3 mm. The other three catalyst monolith has been designed the same way. The triangular, rectangular, and hexagonal catalysts monolith rows were circumscribed within the circle diameter of 3 mm. Two sets of catalysts at two perpendicular planes YZ and XZ, in 11 different columns, placed 18.2 mm from each other, were considered. In the YZ plane, the catalytic part of the reactor is with a column including five rows with a distance of 10 mm. Another column is created at a distance of 18.2 mm from the first one with four rows of catalysts. The latter four rows were placed to cover the space between the two nearby rows of catalysts in the first column. These two sets of columns have been repeated five times along the reactor, with the last column like the first column. For the flow to see the catalyst during the passage of the reactor, more rows of monoliths in every column in XZ plane were added. In the first column, 9 rows of monolith were embedded into the reactor with a center-to-center distance of 5 mm. In the second column,

 $^{^{2}\,}$ Grade here connotes different composition of CH₄, H₂ and CO in the syngas.

Table 1 CH₄/H₂/O₂/Pt combustion mechanism, $k = AT^n \exp(-E/RT)$.

No	Reaction	A (cm, mol,	n	E (kcal
		s)		mol ⁻¹)
(R1)	$H_2 + 2PT(S) \Rightarrow 2H(S)$	4.46×10^{10}	0.5	0.0
(R2)	$2H(S) \Rightarrow H_2 + 2PT(S)$	3.70×10^{21}	0.0	67400.0
(R3)	$O_2 + 2PT(S) \Rightarrow 2O(S)$	1.80×10^{21}	-0.5	0.00
(R4)	$O_2 + 2PT(S) \Rightarrow 2O(S)$	$2.30 \times$	0.00	0.00
		10^{-02}		
(R5)	$2O(S) \Rightarrow O_2 + 2PT(S)$	3.7×10^{21}	0.00	213200.0
(R6)	$H_2O + PT(S) \Rightarrow H_2O(S)$	7.50 ×	0.00	0.00
		10^{-01}		
(R7)	$H_2O(S) \Rightarrow H_2O + PT(S)$	1.00×10^{13}	0.00	40300.00
(R8)	$OH + PT(S) \Rightarrow OH(S)$	1.00×10^{00}	0.00	0.00
(R9)	$OH(S) \Rightarrow OH + PT(S)$	1.00×10^{13}	0.00	192800.0
(R10)	$H(S) + O(S) \Leftrightarrow OH(S) + PT(S)$	3.70×10^{21}	0.00	11500.0
(R11)	$H(S) + OH(S) \Leftrightarrow H_2O(S) +$	3.70×10^{21}	0.00	17400.0
	PT(S)			
(R12)	$OH(S) + OH(S) \Leftrightarrow H_2O(S) +$	3.70×10^{21}	0.00	48200.00
	O(S)			
(R13)	$CO + PT(S) \Rightarrow CO(S)$	1.62×10^{20}	0.5	0.00
(R14)	$CO(S) \Rightarrow CO + PT(S)$	1.00×10^{13}	0.00	125500.0
(R15)	$CO_2(S) \Rightarrow CO_2 + PT(S)$	1.00×10^{13}	0.00	20500.0
(R16)	$CO(S) + O(S) \Rightarrow CO_2(S) +$	3.70×10^{21}	0.00	105000.0
	PT(S)			
(R17)	$CH_4 + 2PT(S) \Rightarrow CH_3(S) +$	4.63×10^{20}	0.50	0.0
(D10)	H(S)	21	0.00	20000
(R18)	$CH_3(S) + PT(S) \Rightarrow CH_2(S)s +$	3.70×10^{21}	0.00	20000.0
(R19)	H(S) $CH_2(S)s + PT(S) \Rightarrow CH(S) +$	0.70 1021	0.00	20000.0
(K19)	$H(S) \Rightarrow CH(S) + H(S)$	3.70×10^{21}	0.00	20000.0
(R20)	$CH(S) + PT(S) \Rightarrow C(S) + H(S)$	3.70×10^{21}	0.00	20000.0
(R21)	$C(S) + O(S) \Rightarrow CO(S) + PT(S)$	3.70×10^{21} 3.70×10^{21}	0.00	62800.0
(R21)	$CO(S) + O(S) \Rightarrow CO(S) + PT(S)$ $CO(S) + PT(S) \Rightarrow C(S) + O(S)$	1.00×10^{18}	0.00	184000.0
(1144)	$CO(3) + FI(3) \rightarrow C(3) + O(3)$	1.00 × 10 ¹⁰	0.00	104000.0

^a Averaged for fuel species and hydroxyl at 20Efficiency factor for $\epsilon_{H_2O}=12$ and $\epsilon_{H_2}=12$.

10 rows were considered, so the centre of each monolith in the second column is exactly in the front of the middle point of an imaginary line connecting two centres of the monolith located in the first column. This configuration was to cover all the spaces of the reactor cylinder, so the flow from the inlet sees at least a row of catalyst monolith while passing the reactor. Again, 11 rows of monolith have been created inside the

reactor in XZ plane with each 18.2 mm.

Geometry

3-D geometries for both the catalytic bed and catalyst monolith were prepared using SolidWorks 2022 and are demonstrated in Figs. 1 and 2. This forms the solid parts of the computational domain. The volume parts of the model were extracted and forms the computational domain for the combustion modelling and analysis. The solid walls are made of stainless steel 310. The catalyst pellets with coatings: platinum, rhodium, palladium, and palladium and nickel are considered for packing of the microcombustor. Fig. 3 gives the schematic of the solid and fluid parts for pall ring packed bed and circular catalyst monolith. The boundary conditions for solving the governing equations are also presented in Fig. 3. The velocity inlet was used for the inlet boundary condition, and for the outlet, the pressure outlet was used. Three wall types are considered for the packed bed- 1. catalyst wall providing the surface for mass and heat transfer of methane and hydrogen, 2. bed wall for keeping the catalyst in the reactive zone of the microburner, and 3. reactor outer wall for transferring some part of combustion heat to the

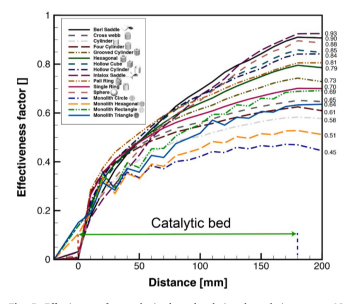


Fig. 5. Effectiveness factor obtained at the designed catalytic reactors 19 transverse plane for different packings and catalytic monolith.

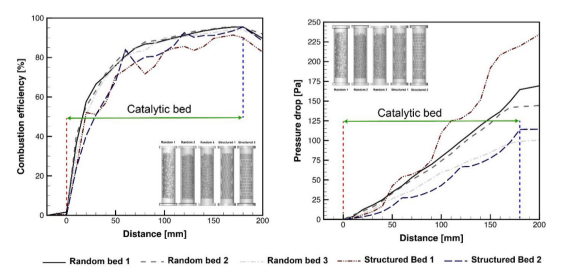


Fig. 4. The combustion efficiency and pressure drop along 19 transverse planes for three random packed beds and two structured beds.

^b Troe parameter is $F_c=0.8$. Efficiency factor for $\varepsilon_{H_2O}=12$. Efficiency factor for $\varepsilon_{H_2O}=11$, $\varepsilon_{H_2}=2$ and $\varepsilon_{O_2}=0.78$.

^c (R14) and (R19) are expressed as the sum of the two rate expressions.

^d Troe parameter is $F_c = 0.5$. Efficiency factor for $\varepsilon_{H_2O} = 12$ and $\varepsilon_{H_2} = 2.5$.

Table 2Operating conditions for hydrogen combustion in micro combustor.

Shape	Packed bed performance parameter					
	η [\%]]	ΔP [Pa]	Ψ	α	C_2	
Berl saddle	87	25.5	0.90	0.33555	0.1255	
Cross Webb	93	105.36	0.65	0.00082	3.6375	
Cylinder	94	125.33	0.58	0.00020	11.239	
Four Cylinder	89	109.55	0.61	0.00096	3.2617	
Grooved Cylinder	91	80.13	0.73	0.00122	2.8669	
Hexagonal	92	66.33	0.79	0.00151	2.4810	
Hollow Cube	91	47.40	0.85	0.00334	1.5119	
Hollow Cylinder	96	55.49	0.84	0.01985	0.4861	
Intalox Saddle	85.03	18.55	0.93	0.41933	0.1147	
Pall ring	92.91	61.91	0.81	0.00169	0.5165	
Single ring	94.42	92.69	0.70	0.00118	2.7952	
Sphere	89.47	35.07	0.88	0.00090	3.8651	
Monolith circle	89.09	154.78	0.45	0.01404	0.6249	
Monolith hexagonal	89.66	137.52	0.51	0.02045	0.5285	
Monolith rectangle	91.31	90.80	0.69	0.3551	0.3902	
Monolith triangle	92	107.05	0.64	0.0851	0.2420	

Table 3Operating conditions for hydrogen combustion in catalytic combustor.

No.	Operation p	oarameter				
	u [m/s]	T [K]	$Y_{CH_{4}} \\$	Y_{H_2}	$Y_{\rm O_2}$	N
1 ^a	1	300	0.05	0.0045	0.23	200
2		350				
3		400				
4		500				
5	0.5					
6	1.5					
7	2					
8			0.0545	0	0.23	
9			0.0327	0.0218	0.23	
10			0.0218	0.0327	0.23	
11			0.0109	0.0436	0.23	
12			0	0.0545	0.23	
13						110
14						50
15						10
16						5

All compositions are mass fractions.

Empty means that the operating point is the same as the base case.

environment. In catalyst monolith, just the first and third walls existed in the model, and the support walls are not considered in the simulation. For the catalyst wall, the surface reactions have been defined to represent the absorption, reactions, and desorption of species. Support walls are assumed to be adiabatic. The reactor's outer wall was assumed to have natural convection with air at temperature 283 K and convective heat transfer coefficient (h) $10~{\rm W/m^2.K.}$

Model formulation

The model used for modelling of the combustion is a typical CFD-DEM model. However, the solution of each part (CFD and DEM) was not obtained simultaneously. It was assumed that the packings are stagnant, and the fluid motion does not influence the solids that much, nor the solid particles random movement influence the fluid. The fluid movement was considered turbulent, although the upstream Reynolds number is around 2500 which is at the transition state. The DEM models were used for preCFD modelling and generation of the computational domain. The reactor monolith catalytic structures are considered stagnant as they are fixed over particular shapes entrenched and embedded into the body of the reactor. For this work, the pellets are also located carefully between two sets of supports, one at the beginning of the and the other at the very end of the reactor. The DEM was used to provide the

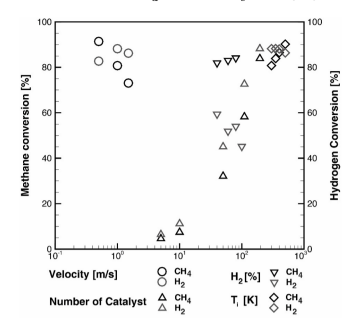


Fig. 6. The operating graph of the catalytic reactor in terms of methane and hydrogen conversion in relation to premixed reactant velocity, temperature, percentage of hydrogen and number of catalysts in the bed.

 Table 4

 The surface combustion mechanisms on different catalyst pellets.

Fuel	Catalyst	No of reac.	Ref
CH ₄ /H ₂	Platinum [Pt]	22	[26]
H_2	Rhodium [Rh]	12	[42]
H_2	Palladium [Pd]	13	[43]
CO	Rhodium [Rh]	10	[44]
CH_4/H_2	Palladium [Pd]	54	[45]
CH ₄ /H ₂	Palladium oxide [PdO]	77	[46]

randomness in the structure of packings when they were loaded by different spatial orientation via Newton's laws of motion. The combusting gas flow will impact and may cause some vibrational and some slight rotational but not really apparent translational movement. This might lead to a solid-fluid interaction (FSI) that should be considered in the modelling. However, in this work, for a matter of simplicity the FSI are not considered in the modelling as we aim to offer a design that in the packed solids are fixed and stagnant.

Governing equations

The conservation equations for mass, momentum, total internal energy (E), and species mass fractions, Y_j for (N-1) chemical species in a low Mach incompressible format are defined for the gas phase. The equations with standard notations are:

The mass balance:

$$\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \, \overrightarrow{u}) = 0 \tag{1}$$

The momentum balance:

$$\frac{\partial \rho \overrightarrow{u}}{\partial t} + \rho(\overrightarrow{u}.\nabla \overrightarrow{u}) = -\nabla p + \nabla \cdot \left(\mu \left[\nabla \overrightarrow{u} + (\nabla \overrightarrow{u})^T - \frac{2}{3}\nabla \cdot \overrightarrow{u}I\right]\right)$$
(2)

The energy balance:

^a Base case.

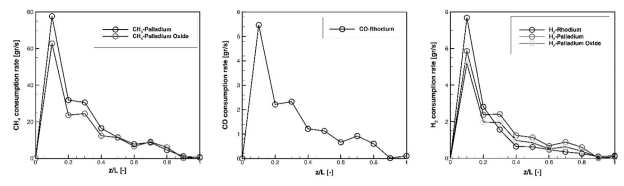


Fig. 7. The time averaged rate of fuel consumption along the reactor dimensionless length for catalytic combustion of h₂-Rhodium, H₂-Palladium, and CO-Rhodium for intalox packing.

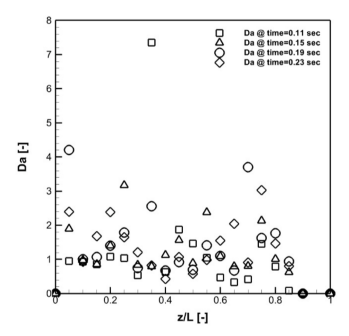


Fig. 8. Temporal evolution of da for ch_4 , H_2 and OH at physical times 0.03, 0.04, 0.07, 0.11, 0.15, 0.23 averaged at 20 transverse planes along the reactor length.

$$\frac{\partial \rho E}{\partial t} + \nabla \cdot \overrightarrow{u} \left(\rho E_f + p \right) = \nabla \cdot \left[k_{eff} \nabla T - \left(\sum_j h_j \overrightarrow{J_j} \right) + \left(\mu \left[\nabla \overrightarrow{u} + (\nabla \overrightarrow{u})^T - \frac{2}{3} \nabla \cdot \overrightarrow{u} I \right] \cdot \overrightarrow{u} \right) \right] + S_f^h$$

The gaseous species transport:

$$\frac{\partial \rho Y_j}{\partial t} + \nabla \cdot (\rho \overrightarrow{u} Y_j) = -\nabla \overrightarrow{J_j} + R_j, \alpha = 1, ..., N-1$$
(4)

The surface species:

$$\frac{\partial \theta_k}{\partial t} = \sigma_k \frac{\dot{s}_k}{\Gamma} \ k = 1, ..., N_s \tag{5}$$

These sets of equations are closed using the equation of state (EOS), which in this study is assumed as the ideal gas, considering the temperature is high enough for the gas to have negligible intermolecular forces

$$p_0 = \frac{\rho RT}{\overline{W}} \overline{W} = \frac{1}{\sum_{i=1}^{N_g} \frac{Y_i}{W_i}}$$
 (6)

To model the catalytic reaction along with the gas phase flow, The diffusion terms in the species equation were modelled using:

$$-\nabla \overrightarrow{J_j} = -\nabla \left(\rho Y_j V_j\right) \tag{7}$$

In this equation, the diffusion term was modelled based on the gas phase diffusion velocity, V_i :

$$V_j = \widetilde{V}_j + \underline{V}_c \tag{8}$$

 $\underline{V_c} = \sum_{i=1}^N Y_j \underline{\widetilde{V_j}}$ indicates the correction velocity which is added to ensure the conservation of mass. $\underline{\widetilde{V_j}}$ is obtained from the mixture-averaged diffusion model:

$$\widetilde{V}_i = -\left(D_i/X_i\right)\underline{\nabla}X_i \tag{9}$$

where D_j indicates jth species mixture-averaged diffusivity and X_j denotes the jth mole fraction.

The following boundary condition between the gas-phase and gaswall interface are considered:

$$J_{j,w} = \underline{n} \cdot \rho Y_i \left(\underline{u} + \underline{V_j} \right) \Big|_{W} = \dot{s}_i w_i, \ j = 1, 2, ..., N$$
 (10)

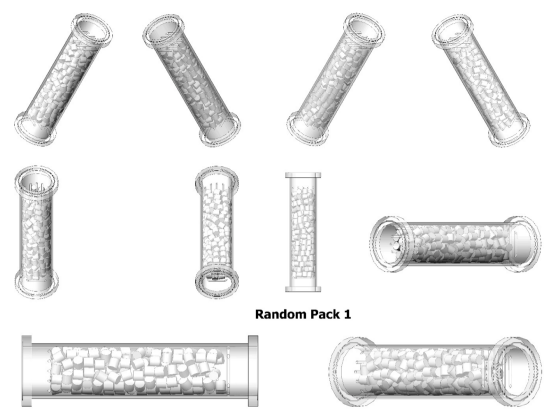
 $\dot{s_i}$ represents the molar catalytic production rate, $J_{j,w}$ indicates the mass flux normal to the wall for the jth gas-phase species, \underline{n} denotes the outward normal vector at the catalyst surface, and $\underline{u}\Big|_{W}$ is the Stefan velocity, given by:

$$\underline{n}.\underline{u}\Big|_{W} = \frac{1}{\rho} \sum_{i=1}^{N} \dot{s}_{i} W_{i}, \ i = 1, ..., N_{g}.$$
 (11)

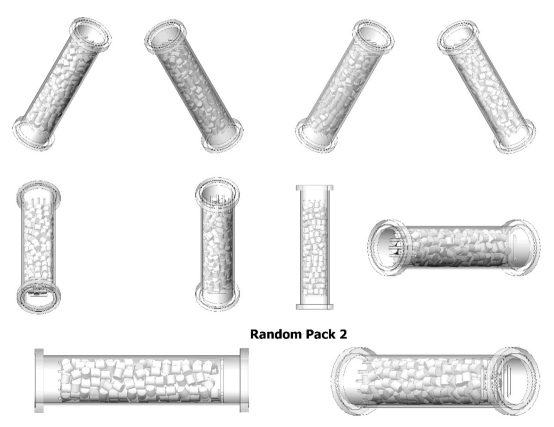
Combustion chemistry

For the steady simulation, wall temperature was observed on the catalyst surface around $T_{\rm w}=1290~{\rm K}$. This temperature is chosen in design optimisation phase testing different surface catalyst temperature to find one with maximum effectiveness factor while it leads to the expected combustion state. This gives the average temperature of the $T_{\rm average_volumetric}=800~{\rm K}$ for the mixture in the reactor which is well below the minimum temperature required for lean combustion of methane and hydrogen. The numerical simulation with and without the gas-phase reactions leads to insignificant difference based on preliminary calculations in the species concentrations in the reactor. Hence, only the surface reactions were considered for design and modelling of the reactor. For this study, the detailed catalytic chemistry reactions by Deutschmannet al. [26] has been used to simulate the catalytic combustion and design of the reactor. The reaction, given in Table 1, includes 8 volumetric, 11 sites and 1 solid species.

(3)



 $\textbf{Fig. A1.} \ \ \textbf{The designed combustor random pack 1 different views.}$



 $\textbf{Fig. A2.} \ \ \textbf{The designed combustor random pack 2 different views.}$

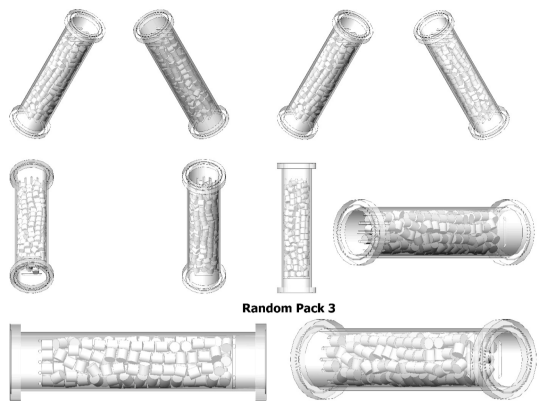


Fig. A3. The designed combustor random pack 3 different views.

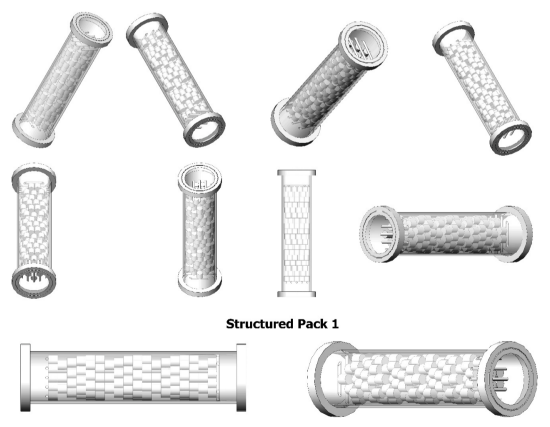


Fig. B1. The designed combustor structured pack 1 different views.

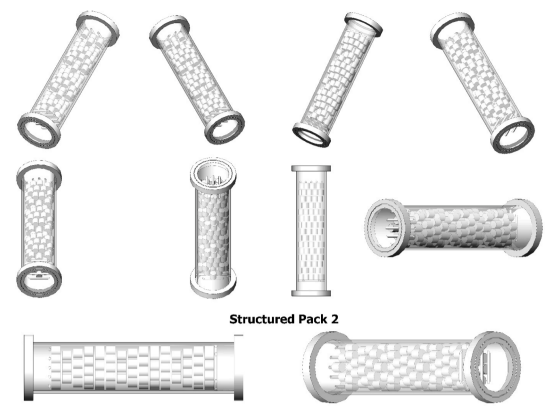


Fig. B2. The designed combustor structured pack 2 different views.

Numerical method

Solution algorithm

The computational domain including the volumetric space for the transverse motion of the syngas reactive mixture is segregated into the hexahedral elements with identical surfaces and dimensions. The conservation equations are separated using a high-order splitting scheme [27] and are discretised using Nth-order Legendre-Lagrangian polynomial [28] to give the solution in each spatial dimensions. The thermochemical state of the microcombustor is resolved using the implicit stiff ODE integrator CVODE [29]. For this work, the reaction source terms in the gas and surface are directly solved although they are massive sets of stiffs ODEs.

The analysis of this work includes the findings of 44 independent different simulations. Five distinct numerical analysis is associated with the different strategies for packing the bed (randomness or structured. Sixteen simulations belong to the structural analysis of the reactor for different catalyst types (monolith and packed beds). Another sixteen sets are for the optimisation of the reactor and analysis at different operating points which gives the operating line of the reactor at different variable loads, different fuel grades, different number of catalytic pellets, and different syngas grades.

Computational domain

Since the flow is considered turbulent, the domain with a high resolution of elements is considered for the simulations. The reactor fluid part, i.e., computational domain, is split into several conforming hexahedral elements. In span-wise direction x-direction, the domain is discretized into semi-uniform 30 elements. Around 35 elements are clustered close to the wall in spanwise-y direction (r-direction) near the combustor walls and catalyst surface. Around 500 uniform elements in the streamwise-z direction z are allocated to the domain. The

discretization was done using the seventh order interpolating for each element in spatial directions. This gives a total number of 180,000,000 grid points. The resolution of the domain is most exhaustive in the proximity of catalyst pellets and monolith where adsorption-reactions-desorption are taking place. The parameters, values and variables are normalized and non-dimentionalised using associated boundary and initial conditions before solving the governing equations and presentation of results.

Results and discussion

Effect of catalyst packing

The operation of the designed reactor is evaluated for different ways for loading the bed with catalyst. Three random structures and two structured versions of the microcombustor filled with only cylinder catalyst type have been prepared and analysed in terms of pressure drop and combustion efficiency. The minimum pressure drops, and a maximum combustion efficiency are desired for both stationary and mobile combustion systems. Fig. 4 gives the trend of pressure drop and combustion efficiency along the reactor from inlet to outlet. The dp and CE were averaged at 19 planes perpendicular of the flow direction and inlets and outlet planes. The schematic of the reactors has also been shown in a small subgraph within the trendline box.

The random structures are the same in number 195 and different in the orientation and arrangement of catalysts in the reactor. The two structured packed reactors are different in number of catalysts and spaces between catalysts in each row of the reactors. In structured reactor number 1, the catalysts were placed at the top of those below without any space considering for each row of catalysts. This reactor includes 200 catalysts. In structured reactor number 2, the catalysts were placed in series of sieves each with distance 2 mm from the bottom catalysts, therefore there is a gap between every series of catalysts in the reactor. The number of catalysts in this reactor is 171.

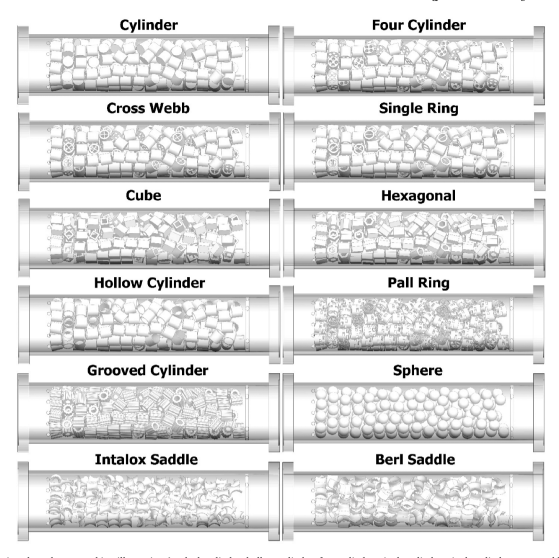


Fig. C1. The designed combustor packing illustration (packed: cylinder, hollow cylinder, four-cylinder, single cylinder, single cylinder, cross-webb, grooved, pall-ring, hexagonal, berl-saddle, cube, intalox-saddle, and sphere).

The results demonstrate that the combustion of $CH_4 + H_2$ completes along the 180 mm length of both the random and structured packed reactor. The combustion efficiency of the random beds 1, 2, and 3 reaches the values 90%, 92%, and 89% at the combustor outlet plane. The combustion efficiency of the structured beds 1 and 2 go up to the value of 83% and 88% at the reactor outlet. The pressure drops, however, for the studied reactors are quite different. In terms of pressure drop, maximum pressure drop is for the structured bed 1 although the catalysts are perfectly arranged within the reactor body. Indeed, the reactor volume is playing the main role in the pressure drop. The reactor with the most space available for the reacting mixture has the least pressure drop and vice versa. The restriction and constriction of flow increases the inter-catalytic fluid velocity, thereby decreasing the pressure of the fluid. The free volume inside the structured packed reactor 1 and 2 is 312 and 335 mm³, respectively. For random packed reactors, the free volume of the reactor is 316 mm³.

Note that the combustion efficiency within [83 95]% is comparable to the combustion efficiency of industrial scale gensets [30–33] and other micro combustor units (microturbine combustors: [34]). The net combustion efficiency of diesel fuel and biodiesels is obtained more than 90% [33]. In microgas turbines the combustion efficiency of nearly 100% [35] could be obtained as the air staging leaves no room for poor mixing and insufficient combustion [36]. In conventional combustor, an air swirler is implemented to improve the mixing and reduction of

nitrogen oxides [37,38]. Here, the mixing is achieved by bifurcation and microchanneling of the fluid over the catalyst surface. This improves the rate of adsorption and catalytic reactions.

Effects of catalyst shape

For mechanical devices, the pressure drop along with combustion efficiency is important for the efficient operation of MEMS. The effectiveness factor has been defined to give both the contribution of the pressure drop and combustion efficiency. The ideal reactor is with effectiveness factor 1 which both satisfies the 0.5 for combustion efficiency and pressure drop. The definition of the effectiveness factor (Ψ) is given:

$$\Psi = \omega_{Combustionefficiency} + \omega_{Pressuredrop} \tag{12}$$

The contribution of the combustion efficiency ($\omega_{Combustion\ efficiency}$) is universal and represent the heat loss by unburned materials. It is simply half of the combustion efficiency value. The contribution from the pressure drops ($\omega_{Pressure\ drop}$) is specific to this study which gives the value 0 for the reactor with maximum pressure drop and the value 1 for reactor with minimum pressure drop. It is also distributed within [0 0.5] similar to $\omega_{Combustion\ efficiency}$ from inlet to outlet. Such a definition gives the effect of packing on both the pressure drop and combustion efficiency.

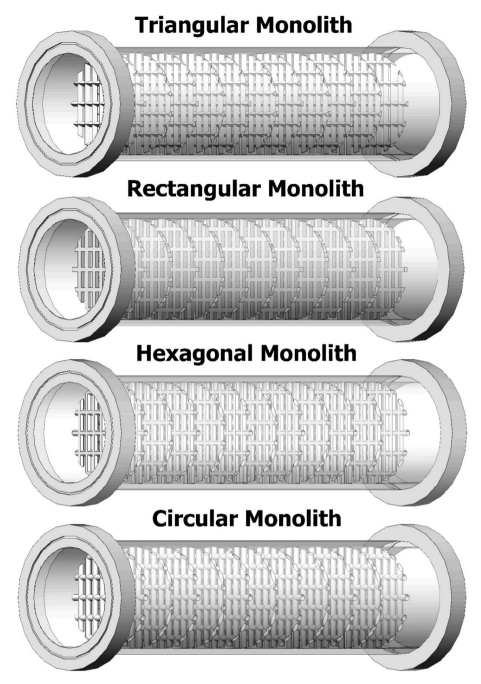


Fig. D1. The designed combustor packing illustration (monolith: triangular, rectangular, hexagonal, and circular).

$$\Psi = \left(\frac{0.169008[OH] + 0.21[H_2O]}{[OH] + [H_2O] + [H_2]} + \frac{0.18705[CO] + 0.29[CO_2]}{[CO_2] + [CO] + [CH_4]}\right) + \frac{\Delta P_{Reactor,local}(\Delta P_{\text{maximum}} - \Delta P_{Reactor,outlet})}{2\Delta P_{Reactor,outlet}(\Delta P_{\text{maximum}} - \Delta P_{Minimum})}$$

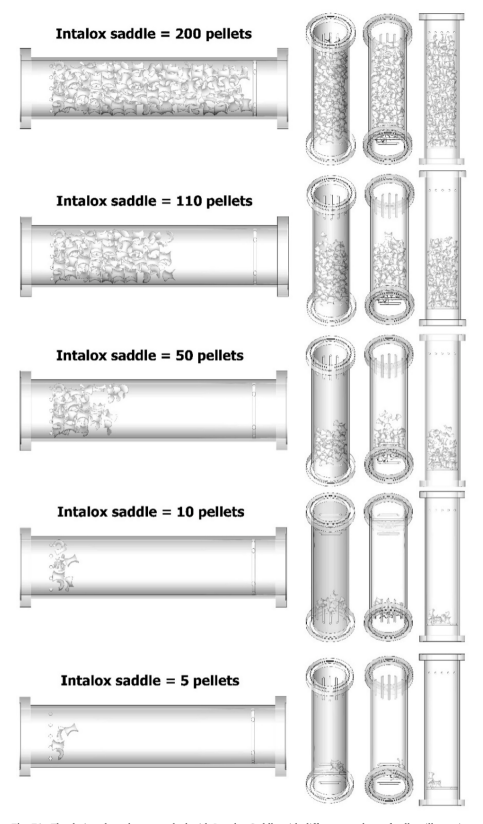
$$\tag{13}$$

where [X] denotes either the mole fraction or mole concentration of the specie X, $\Delta P_{\text{max}imum}$ and $\Delta P_{\text{Minimum}}$ are constant and represent the maximum and minimum pressure drop for the reactor with all investigated packings. In this study, $\Delta P_{\text{max}imum}$ is for the monolith circle with 155 Pa and $\Delta P_{\text{Minimum}}$ belongs to the intalox saddle packing with 18 Pa. $\Delta P_{\text{Reactor,local}}$ and $\Delta P_{\text{Reactor,outlet}}$ specify the local, i.e., spatial, pressure drop at pressure drop at the reactor outlet.

The spatial effectiveness factor for the catalytic and monolith reactor

is given Fig. 5. The intalox and berl saddles provide the effectiveness factor 0.93 and 0.90, respectively which is higher than other packings. The analysis has shown that the combustion efficiency for all the packings and monolith reaches almost the value 1 at the combustor outlet plane. Thus, the effectiveness factor for this reactor is more under the influence of pressure drop along the catalytic beds. The catalytic monoliths have the minimum effectiveness factor although their combustion efficiency is slightly higher than any other packings.

The inertial resistance $(\alpha = \frac{D_p^2}{150} \frac{\varepsilon^3}{(1-\varepsilon)^2})$ and viscous resistance $(C_2 = \frac{1.75\rho}{D_p} \frac{(1-\varepsilon)}{\varepsilon^3})$ are two parameters that can be used to evaluate the operation of packed beds, catalyst monolith or other porous medium. For this reactor, the effectiveness factor is a direct function of the inertial resistance and inverse function of the viscous resistance. Indeed, the



 $\textbf{Fig. E1.} \ \ \textbf{The designed combustor packed with Intralox Saddle with different numbers of pellets illustration.}$

presence of more pore volume in the case of saddle packing gives rise to their highest effectiveness factor and their better performance for catalytic combustion. Table 2 gives the combustion efficiency, pressure drop, effectiveness factor, inertial and viscous resistance for different

packings loaded into the designed reactor at the outlet plane. The combustion efficiency for hollow cylinder, single ring and cylinder is noticeably higher than any other packings, although they are in the middle hierarchy of packings in terms of effectiveness factor.

Table F1 $H_2/O_2/Pd$ combustion mechanism, $k = AT^n \exp(-E/RT)$.

No	Reaction	A (cm, mol, s)	n	E (kJ mol ⁻¹)
	$H_2 + 2Pd => 2H(s)$	0.70	0.0	0.0
	$2H(s) => H_2 + 2Pd(s)$	$4.800 \times 10^{+21}$	0.0	84.0
	COV/H(s)	0.0	0.0	-15.0/
	H + Pd(s) => H(s)	1.00	0.0	0.0
	$O_2 + 2Pd(s) => 2O(s)$	0.400×10^{-00}	0.0	0.0
	$2O(s) => O_2 + 2Pd(s)$	$7.100 \times 10^{+21}$	0.0	230.0
	O + Pd(s) => O(s)	1.00	0.0	0.0
	$H_2O + Pd(s) => H_2O(s)$	0.75	0.0	0.0
	$H_2O(s) => H_2O + Pd(s)$	$1.300{ imes}10^{+13}$	0.0	44.0
	OH + Pd(s) => OH(s)	1.00	0.0	0.0
	OH(s) => OH + Pd(s)	$1.300 \times 10^{+13}$	0.0	213.0
	H(s) + O(s) = OH(s) + Pd(s)	$3.70 \times 10^{+21}$	0.00	11.5
	$H(s) + OH(s) = H_2O(s) + Pd(s)$	$3.70 \times 10^{+21}$	0.00	17.4
	$OH(s) + OH(s) = H_2O(s) + O(s)$	$3.70 \times 10^{+21}$	0.00	48.2

Table F2 $H_2/O_2/Rh$ combustion mechanism, $k = AT^n \exp(-E/RT)$.

No	Reaction	A (cm, mol, s)	N	E (kJ mol ⁻¹)
(R1)	$H_2 + Rh(s) + Rh(s) => H(s)$	3.000×10^{-02}	0.000	0.000
	+ H(s)			
(R2)	$O_2 + Rh(s) + Rh(s) => O(s)$	1.000×10^{-02}	0.000	0.000
	+ O(s)			
(R3)	$H_2O + Rh(s) => H_2O(s)$	1.000×10^{-01}	0.000	0.000
(R4)	H(s) + H(s) => Rh(s) + Rh(s)	$5.574 \times 10^{+19}$	0.239	59.691
	$+ H_2$			
(R5)	O(s) + O(s) => Rh(s) + Rh(s)	$5.329 \times 10^{+22}$	-0.137	386.995
	$+ O_2$			
(R6)	$H_2O(s) => Rh(s) + H_2O$	$6.858 \times 10^{+14}$	-0.280	44.993
(R7)	O(s) + H(s) => OH(s) + Rh(s)	$8.826 \times 10^{+21}$	-0.048	73.365
(R8)	OH(s) + Rh(s) => O(s) + H(s)	$1.000 \times 10^{+21}$	0.045	48.041
(R9)	$OH(s) + H(s) => H_2O(s) + Rh$	$1.743 \times 10^{+22}$	-0.127	41.731
	(s)			
(R10)	$H_2O(s) + Rh(s) => OH(s) + H$	$5.408 \times 10^{+22}$	0.129	98.220
	(s)			
(R11)	OH(s) + OH(s) => O(s) +	$5.736 \times 10^{+20}$	-0.081	121.594
	$H_2O(s)$			
(R12)	$O(s) + H_2O(s) => OH(s) +$	$1.570 \times 10^{+22}$	0.081	203.407
	OH(s)			

Effect of inlet velocity, H_2 concentration, temperature, and number of pellets

One important criterion which shows the operation of the reactor under possible variable loads, catalyst inactivity, temperature oscillation and fuel type is the operating graph of the catalytic reactor. The operating graph is obtained for this designed reactor from the simulations of Table 3 to show how the conversation of methane and hydrogen is influenced by the inlet jet velocity, inlet temperature, reactor number of pellets and percentage of hydrogen in the fuel stream.

Fig. 6 demonstrates the operating graph in terms of CH_4 and hydrogen conversations in relation to the velocity, temperature, number of catalyst in the bed and H_2 volumetric percentage in the fuel stream. The graph has two markers-1. Black showing the conversion of methane and 2- Gray showing the hydrogen conversion. The logarithmic X-axis could be velocity, number of catalysts in the reactor, inlet temperature, and hydrogen percentage in the reactant.

The operating graph exhibits the conversion trendlines over number of catalysts and hydrogen percentage to be more deterministic of the methane and hydrogen conversion. When the number of catalysts goes below 100, the conversion of methane and hydrogen significantly diminishes. For feed hydrogen percentage, the situation is different. When the volumetric percentage of hydrogen in the fuel is going from 30% to 100%, the $\rm H_2$ conversion is fixed within [50 60], although the conversion of methane regardless of feed hydrogen percentage is as high as

Table F3 CO/O₂/Rd combustion mechanism, $k = AT^n \exp(-E/RT)$.

No	Reaction	A (cm, mol, s)	n	E (kJ mol ⁻¹)
(R1)	$O_2 + Rh(s) + Rh(s) => O(s)$ + $O(s)$	1.000×10^{-02}	0.000	0.000
(R2)	$CO_2 + Rh(s) => CO_2(s)$	4.800×10^{-02}	0.000	0.000
(R3)	CO + Rh(s) => CO(s)	4.971×10^{-01}	0.000	0.000
(R4)	$O(s) + O(s) => Rh(s) + Rh(s) + O_2$	$5.329 \times 10^{+22}$	-0.137	386.995
(R5)	CO(s) => Rh(s) + CO	$1.300 \times 10^{+13}$	0.295	134.070
	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-47.000/
(R6)	$CO_2(s) => Rh(s) + CO_2$	$3.920 \times 10^{+11}$	0.315	20.505
(R7)	O(s) + C(s) => CO(s) + Rh(s)	$1.173 \times 10^{+22}$	0.000	92.142
(R8)	CO(s) + Rh(s) => O(s) + C(s)	$6.390 \times 10^{+21}$	0.000	174.758
	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-47.000/
(R9)	O(s) + CO(s) => CO2(s) + Rh (s)	$6.183 \times 10^{+21}$	0.034	129.982
	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-47.000/
(R10)	$CO_2(s) + Rh(s) => O(s) + CO$ (s)	5.752×10 ⁺²²	-0.175	106.492

85%. The reactor response to the feed stream temperature, velocity oscillation is in an efficient way that can tolerate 50% overload and drop load (50% increase and decrease in the mass flow rate). In such cases, the conversion for both methane and hydrogen still remains almost 80%.

The spatial temperature in the designed combustor is obtained below 1500 K. This temperature is considered low for the formation of NO and prompt NO [39–41]. The negligible contribution of gas phase reaction in the operation of the catalyst leaves no room for dissociation of nitrogen and attack of the hydrocarbon radicals to nitrogen.

Effect of catalyst

The influence of different catalysts including Platinum, Rhodium, Palladium and Palladium oxide- are considered to determine the operation of designed catalytic reactor in combustion of hydrogen, methane, and carbon monoxide for intalox pellets. The characteristics of the employed surface chemical reactions for the analysis of different catalyst are given Table 4. The chemistry set and list of chemical reactions are given in the appendix at the end of the manuscript.

The spanwise time-averaged rate of fuel consumption along the

Table F4 CO/O₂/Pt combustion mechanism, $k = AT^n \exp(-E/RT)$.

	· · · · · · · · · · · · · · · · · · ·	,	/	
No	Reaction	A (cm, mol, s)	n	E (kJ mol ⁻¹)
(R1)	$O_2 + (Pt) => O_2(Pt)$	5.000×10^{-02}	0.000	0.000
(R2)	$O_2(Pt) => (Pt) + O_2$	$5.243 \times 10^{+11}$	-0.069	19.573
(R3)	$O_2(Pt) + (Pt) => O(Pt) + O$ (Pt)	$8.325 \times 10^{+13}$	-0.000	39.933
(R4)	$O(Pt) + O(Pt) => O_2(Pt) + (Pt)$	$4.444 \times 10^{+21}$	0.000	264.067
	COV/O(Pt)	$0.000 \times 10^{+00}$	0.000	-88.200/
(R5)	CO + (Pt) => CO(Pt)	8.400×10^{-01}	0.000	0.000
(R6)	CO(Pt) => (Pt) + CO	$7.635 \times 10^{+12}$	-0.139	143.145
	COV/CO(Pt)	$0.000 \times 10^{+00}$	0.000	-29.300/
(R7)	$CO_2 + (Pt) => CO_2(Pt)$	3.193×10^{-03}	-0.035	2.686
(R8)	$CO_2(Pt) => (Pt) + CO_2$	$1.894 \times 10^{+10}$	0.139	21.855
(R9)	$CO(Pt) + O_2(Pt) => CO_2(Pt) + O(Pt)$	$4.124 \times 10^{+13}$	0.069	9.494
	COV/O(Pt)	$0.000 \times 10^{+00}$	0.000	44.100/
(R10)	$CO_2(Pt) + O(Pt) => CO(Pt) + O2(Pt)$	$2.910 \times 10^{+23}$	-0.069	272.506
	COV/CO(Pt)	$0.000 \times 10^{+00}$	0.000	29.300/
(R11)	CO(Pt) + O(Pt) => CO2(Pt) + (Pt)	4.764×10 ⁺¹³	0.069	101.361
	COV/CO(Pt)	$0.000 \times 10^{+00}$	0.000	-29.300/
(R12)	$CO_2(Pt) + (Pt) => CO(Pt) + O(Pt)$	$6.297 \times 10^{+20}$	-0.069	140.239
	COV/O(Pt)	$0.000 \times 10^{+00}$	0.000	44.100/

Table F5 CH₄/H₂/CO/O₂/Ni combustion mechanism, $k = AT^n \exp(-E/RT)$.

No	Reaction	A (cm, mol, s)	N	E (kJ mol ⁻¹)
	$H_2 + 2NI(s) => 2H(s)$	1.000×10^{-002}	0.00	0.0
	$2H(s) => H_2 + 2NI(s)$	$2.545 \times 10^{+019}$	0.00	81.2
	$O_2 + 2NI(s) => 2O(s)$	1.000×10^{-002}	0.00	0.0
	$2O(s) => O_2 + 2NI(s)$	$4.283\times10^{+023}$	0.00	474.9
	$CH_4 + NI(s) => CH_4(s)$	8.000×10^{-003}	0.00	0.0
	$CH_4(s) => CH_4 + NI(s)$	$8.705\times10^{+015}$	0.00	37.5
	$H_2O + NI(s) => H_2O(s)$	1.000×10^{-001}	0.00	0.0
	$H_2O(s) => H_2O + NI(s)$	$3.732\times10^{+012}$	0.00	60.8
	$CO_2 + NI(s) => CO_2(s)$	1.000×10^{-005}	0.00	0.0
	$CO_2(s) = > CO_2 + NI(s)$	$6.447 \times 10^{+007}$	0.00	26.0
	CO + NI(s) => CO(s)	5.000×10^{-001}	0.00	0.0
	CO(s) => CO + NI(s)	$3.563 \times 10^{+011}$	0.00	111.3
	COV/CO(s)	$0.000 \times 10^{+000}$	0.00	-50.0/
	H(s) + O(s) => NI(s) + OH(s)	$5.000 \times 10^{+022}$	0.00	97.9
	NI(s) + OH(s) => H(s) + O(s)	$1.781 \times 10^{+021}$	0.00	36.1
	H(s) + OH(s) => NI(s) + H2O(s)	$3.000 \times 10^{+020}$	0.00	42.7
	$NI(s) + H_2O(s) => H(s) + H(s)$	$2.271{\times}10^{+021}$	0.00	91.8
	$2OH(s) => H_2O(s) + O(s)$	$3.000 \times 10^{+021}$	0.00	100.0
	$H_2O(s) + O(s) => 2OH(s)$	$6.373 \times 10^{+023}$	0.00	210.9
	C(s) + O(s) => NI(s) + CO(s)	$5.200 \times 10^{+023}$	0.00	148.1
	NI(s) + CO(s) => C(s) + O(s)	$1.354 \times 10^{+022}$	-3.00	116.1
	COV/CO(s)	$0.000 \times 10^{+000}$	0.00	-50.0/
	CO(s) + O(s) => NI(s) + CO2(s)	$2.000 \times 10^{+019}$	0.00	123.6
	COV/CO(s)	$0.000 \times 10^{+000}$	0.00	-50.0/
	NI(s) + CO2(s) => CO(s) + O(s)	$4.653 \times 10^{+023}$	-1.00	89.3
	NI(s) + HCO(s) => H(s) + CO(s)	$3.700 \times 10^{+021}$	0.00	0.0
	COV/CO(s)	$0.000 \times 10^{+000}$	0.00	50.0/
	H(s) + CO(s) => NI(s) + HCO(s)	$4.019 \times 10^{+020}$	-1.00	132.2
	NI(s) + HCO(s) = CH(s) + O(s) $NI(s) + HCO(s) = CH(s) + O(s)$	$3.700 \times 10^{+024}$	-3.00	95.8
	CH(s) + O(s) => NI(s) + HCO(s) $CH(s) + O(s) => NI(s) + HCO(s)$	4.604×10 ⁺⁰²⁰	0.00	110.0
	$NI(s) + CH_4(s) = H(s) + CH_3(s)$	$3.700 \times 10^{+021}$	0.00	57.7
	H(s) + CH3(s) => H(s) + CH3(s) H(s) + CH3(s) => H(s) + CH4(s)	$6.034 \times 10^{+021}$	0.00	61.6
	$NI(s) + CH_3(s) => H(s) + CH_2(s)$	$3.700 \times 10^{+024}$	0.00	100.0
	H(s) + GH2(s) => H(s) + GH2(s) H(s) + CH2(s) => NI(s) + CH3(s)	1.293×10 ⁺⁰²³	0.00	55.3
	$NI(s) + CH_2(s) = > H(s) + CH(s)$ $NI(s) + CH_2(s) = > H(s) + CH(s)$	$3.700 \times 10^{+024}$	0.00	97.1
		$4.089 \times 10^{+024}$		79.2
	H(s) + CH(s) => NI(s) + CH2(s)	$3.700 \times 10^{+021}$	0.00	
	NI(s) + CH(s) => H(s) + C(s)		0.00	18.8
	H(s) + C(s) => NI(s) + CH(s)	$4.562 \times 10^{+022}$ $1.700 \times 10^{+024}$	0.00	161.1
	$CH_4(s) + O(s) => OH(s) +$	1./00×10	0.00	88.3
	$CH_3(s)$ $OH(s) + CH_3(s) => CH_4(s) + O$	$9.876 \times 10^{+022}$	0.00	30.4
	(s)	, 0,,10	0.00	
	$CH_3(s) + O(s) => OH(s) +$	$3.700 \times 10^{+024}$	0.00	130.1
	CH ₂ (s)			
	$OH(s) + CH_2(s) => CH_3(s) + O$ (s)	$4.607 \times 10^{+021}$	0.00	23.6
	$CH_2(s) + O(s) => OH(s) + CH(s)$	$3.700\times10^{+024}$	0.00	126.8
	$OH(s) + CH(s) => CH_2(s) + O(s)$	1.457×10 ⁺⁰²³	0.00	47.1
	$CH(s) + CH(s) = > CH_2(s) + C(s)$ CH(s) + O(s) = > OH(s) + C(s)	$3.700 \times 10^{+021}$	0.00	48.1
	Cir(s) + C(s) = > Cir(s) + C(s)	$1.625 \times 10^{+021}$	0.00	128.6

reactor is given in Fig. 7 for different combustion systems- H_2 rhodium, H_2 palladium and CO rhodium. The designed reactor can burn hydrogen and carbon monoxide on the surface of precious metals such as rhodium and palladium. The findings, not shown here, confirmed that the combustion efficiency of these systems is up to 90% at the combustor outlet plane. As the trendlines in Fig. 7 exhibits, the converter most reactive zone is from 0 to 20 mm where the surface reactions are at the maximum rate of occurrence. The combustion in all of the cases is taking place at near 1300 K with surface catalytic reaction prevailing the gas phase reaction. The gas phase reactions, if any are volumetric as the temperature within the convertor does not go beyond the ignition flame temperature of the fuels.

Transient analysis of the bed

How well the designed microcombustor will behave in time can be addressed by transient analysis of the bed in progressing the combustion and to burn the fuels. The premix mixture of methane, hydrogen and oxygen will be injected to the reactor where they will be adsorbed to the

Table F6 CH₄/H₂/CO/O₂/Pd combustion mechanism, $k = AT^n \exp(-E/RT)$.

lo	Reaction	A (cm, mol, s)	n	E (kcal mol ⁻¹)
R1)	$H_2 + Pd(s) + Pd(s) => H(s)$ $+ H(s)$	2.445×10 ⁻⁰¹	-0.076	0.000
R2)	$O_2 + Pd(s) + Pd(s) => O(s)$ $+ O(s)$	5.712×10^{-02}	0.012	1.945
R3)	$CH_4 + Pd(s) => CH_4(s)$	8.557×10^{-03}	0.017	0.000
R4)	$H_2O + Pd(s) => H_2O(s)$	1.402×10^{-01}	0.033	0.000
R5)	$CO_2 + Pd(s) => CO_2(s)$	8.186×10^{-03}	-0.037	0.000
R6)	CO + Pd(s) => CO(s)	6.716×10^{-01}	0.015	0.935
R7)	$CH_4 + O(s) + O(s) => H_3CO$ (s) + OH(s)	8.928×10^{-03}	0.004	159.978
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-22.500/
R8)	H(s) + H(s) => Pd(s) + Pd (s) + H ₂	8.393×10 ⁺¹⁹	0.303	85.036
R9)	O(s) + O(s) = Pd(s) + Pd $(s) + O_2$	$2.932 \times 10^{+22}$	-0.049	221.818
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-22.500/
R10)	$H_2O(s) => Pd(s) + H_2O$	$1.244 \times 10^{+14}$	-0.134	46.593
R11)	CO(s) = Pd(s) + CO	$8.565 \times 10^{+13}$	-0.062	138.558
.(11)	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-31.000/
R12)	$CO_2(s) => Pd(s) + CO_2$	$1.392 \times 10^{+12}$	0.000	23.156
R13)	$CH_4(s) = Pd(s) + CH_4$	$1.865 \times 10^{+13}$	-0.068	29.499
R14)	$H_3CO(s) + OH(s) => O(s) + O(s) + CH_4$	2.046×10 ⁺¹⁸	-0.017	248.686
R15)	O(s) + H(s) => OH(s) + Pd (s)	3.598×10 ⁺²²	-0.060	85.044
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-11.250/
R16)	OH(s) + Pd(s) => O(s) + H (s)	4.169×10 ⁺²⁰	0.060	66.456
R17)	OH(s) + H(s) => H2O(s) + Pd(s)	$3.017 \times 10^{+20}$	-0.059	-0.823
R18)	$H_2O(s) + Pd(s) => OH(s) + H(s)$	$4.972 \times 10^{+22}$	0.059	109.623
R19)	OH(s) + OH(s) => O(s) + H2O(s)	$2.516 \times 10^{+19}$	0.001	54.533
R20)	O(s) + H2O(s) => OH(s) + OH(s)	$3.578 \times 10^{+23}$	-0.001	183.567
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-11.250/
R21)	O(s) + C(s) => CO(s) + Pd(s)	$5.434 \times 10^{+23}$	0.004	21.097
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-11.250/
R22)	CO(s) + Pd(s) => O(s) + C(s)	$2.392 \times 10^{+21}$	-0.004	214.903
	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-31.000/
R23)	O(s) + CO(s) => CO2(s) + Pd (s)	6.874×10 ⁺¹⁸	0.052	71.247
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-11.250/
	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-31.000/
R24)	$CO_2(s) + Pd(s) => O(s) + CO$ (s)	$2.400 \times 10^{+21}$	-0.052	132.753
R25)	$CH_4(s) + Pd(s) => CH_3(s) +$	$1.301{\times}10^{+22}$	0.064	66.978
R26)	H(s) $CH_3(s) + H(s) => CH_4(s) +$ Pd(s)	$1.052{\times}10^{+21}$	-0.064	38.422
R27)	$CH_3(s) + Pd(s) => CH_2(s) + H(s)$	$2.811{\times}10^{+24}$	0.064	104.253
R28)	$CH_2(s) + H(s) => CH_3(s) + Pd(s)$	$4.871{\times}10^{+21}$	-0.064	37.147
R29)	$CH_2(s) + Pd(s) => CH(s) + H(s)$	$8.888 \times 10^{+23}$	0.064	100.278
R30)	CH(s) + H(s) => CH2(s) + Pd(s)	$1.540{\times}10^{+25}$	-0.064	63.722
R31)	CH(s) + Pd(s) => C(s) + H(s)	$2.811 \times 10^{+23}$	0.064	20.753
R32)		4.871×10 ⁺²⁴		
-	C(s) + H(s) => CH(s) + Pd(s)		-0.064	165.847
R33)	O(s) + CH4(s) => CH3(s) + OH(s) $O(s) + CH4(s) => CH3(s) + CH3(s) + CH4(s) + CH3(s) + CH3(s$	2.591×10 ⁺²⁴	0.004	96.622
R34)	COV/O(s) $CH_3(s) + OH(s) => O(s) +$	$0.000 \times 10^{+00}$ $2.427 \times 10^{+21}$	-0.000 -0.004	-11.250/ 49.478
R35)	$CH_4(s)$ $O(s) + CH_3(s) => CH_2(s) +$	$2.611{\times}10^{+25}$	0.004	97.347
	OH(s)	0.000, 10+00	0.000	11.0507
R36)	COV/O(s) $CH_2(s) + OH(s) => O(s) +$	$0.000 \times 10^{+00}$ $5.243 \times 10^{+20}$	-0.000 -0.004	-11.250/ 11.653
R37)	CH3(s) O(s) + CH2(s) => CH(s) +	8.257×10 ⁺²¹	0.004	88.322

(continued on next page)

Table F6 (continued)

No	Reaction	A (cm, mol, s)	n	E (kcal mol^{-1})
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-11.250/
(R38)	CH(s) + OH(s) => O(s) +	$1.658 \times 10^{+21}$	-0.004	33.178
	CH ₂ (s)			
(R39)	O(s) + CH(s) => C(s) + OH	$2.611 \times 10^{+21}$	0.004	36.847
	(s)			
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-11.250/
(R40)	C(s) + OH(s) => O(s) + CH	$5.243 \times 10^{+20}$	-0.004	163.353
	(s)			
(R41)	C(s) + OH(s) => CO(s) + H	$1.898 \times 10^{+22}$	0.064	0.503
	(s)			
(R42)	CO(s) + H(s) => C(s) + OH	$7.213 \times 10^{+21}$	-0.064	212.897
	(s)	0.000 10±00		
(D. 10)	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-31.000/
(R43)	$H_3CO(s) + O(s) => H_2CO(s)$ + OH(s)	$2.351 \times 10^{+24}$	0.017	65.714
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-11.250/
(R44)	$H_2CO(s) + OH(s) => H_3CO$	$5.824 \times 10^{+21}$	-0.017	56.886
	(s) + O(s)			
(R45)	H2CO(s) + O(s) => HCO(s)	$2.351 \times 10^{+24}$	0.017	31.014
	+ OH(s)	$0.000 \times 10^{+00}$	0.000	11.0507
(R46)	COV/O(s)	$5.824 \times 10^{+21}$	0.000 -0.017	-11.250/ 14.986
(N40)	$HCO(s) + OH(s) => H_2CO$ (s) + O(s)	3.624×10	-0.017	14.960
(R47)	HCO(s) + O(s) => CO(s) +	$2.351 \times 10^{+24}$	0.017	12.114
(1(47)	OH(s)	2.551 \ 10	0.017	12.117
	COV/O(s)	$0.000 \times 10^{+00}$	0.000	-11.250/
(R48)	CO(s) + OH(s) => HCO(s) +	5.824×10 ⁺²¹	-0.017	111.186
,	O(s)			
	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-31.000/
(R49)	CO(s) + OH(s) => COOH(s)	$4.167 \times 10^{+19}$	0.018	46.493
	+ Pd(s)			
	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-31.000/
(R50)	COOH(s) + Pd(s) => CO(s)	$3.251 \times 10^{+21}$	-0.018	27.807
	+ OH(s)			
(R51)	$COOH(s) + Pd(s) => CO_2(s)$	$1.871 \times 10^{+19}$	0.094	27.009
	+ H(s)			
(R52)	$CO_2(s) + H(s) => COOH(s)$	$7.228 \times 10^{+21}$	-0.094	125.791
	+ Pd(s)	.10		
(R53)	COOH(s) + H(s) => CO(s) +	$2.532 \times 10^{+18}$	-0.076	61.084
	H ₂ O(s)			
(R54)	$CO(s) + H_2O(s) => COOH$	$5.349 \times 10^{+18}$	0.076	190.216
	(s) + H(s)	0.000 10+00	0.000	01.000 /
	COV/CO(s)	$0.000 \times 10^{+00}$	0.000	-31.000/

surface of catalyst for the combustion with concomitant desorption of water vapor and carbon dioxide which will exit from hot end of the reactor. The Damköhler number (Da) is used here to show the capability of the reactor to simultaneously burn the hydrogen and methane. It is the ratio of the turbulent timescale to the chemical reaction timescale. It has been estimated for methane, hydrogen and OH as the following.

$$Da \sim \frac{\text{mixing time scale}}{\text{chemical time scale}}$$
 (13)

The mixing rate is estimated from the turbulent mixing of the components in the gas phase and surface chemical reactions associated with the those considered species in the turbulent mixing are for the chemical time scales. The hydrogen is destructed through the surface reaction R1 $(H_2 + 2PT(S) \Rightarrow 2H(S))$ and a possibility for surface formation and desorption exists from R2 $(2H(S) \Rightarrow H_2 + 2PT(S))$. The OH will be produced and consumed R8 $(OH + PT(S) \Rightarrow OH(S))$ and R9 $(OH(S) \Rightarrow OH + PT(S))$, respectively. The methane is also consumed only from the reaction R17 $(CH_4 + 2PT(S) \Rightarrow CH_3(S) + H(S))$. In laminar flow through the catalytic converter, Da can be estimated from the ratio of advection to reaction source terms for the species. The results of the Da analysis at different times when the feedstock is injected to the reactor are presented in Fig. 8.

The Da number is averaged for fuel species and hydroxyl at 20 transverse planes along the reactor starting from the part of reactor loaded with catalysts. The distance between the planes was 20 mm. In

Table F7 CH₄/H₂/CO/O₂/PdO combustion mechanism, $k = AT^n \exp(-E/RT)$.

No	Reaction	A (cm, mol, s)	n	E (kcal mol ⁻¹)
R1)	$CH_4 + PdO + PdO => CH_3(s)$	4.582E-02	-0.001	33.155
R2)	+ H(s) $CH_3(a) + H(b) => PdO + PdO$ $+$ CH_4	1.034E + 21	0.003	106.180
R3)	$CH_4 + (s) + OH(s) => CH_3(s) + H_2O(s)$	1.510E-02	0.001	28.860
R4)	$CH_3(s) + H_2O(s) => OH(s) + PdO + CH_4$	7.071E + 19	-0.004	104.259
R5)	$CH_4 + PdO + O(s) => CH_3(s) + OH(s)$	3.082E-02	0.007	26.234
R6)	$CH_3(s) + OH(s) => O(s) + PdO + CH_4$	2.695E + 20	-0.029	192.865
R7)	$O_2 + PdO => O_2(s)$	5.710E-02	0.000	0.000
R8)	$O_2(a) => (a) + O_2$	6.702E + 15	-0.031	63.541
R9) R10)	$O_2 + PdO => O(b)$ $O(b) => PdO + O_2$	5.710E-02 7.024E +	$0.000 \\ -0.027$	0.000 159.975
R11)	$H_2O + PdO + O(s) => OH(s)$ + $OH(s)$	15 1.400E-01	0.000	0.000
R12)	OH(s) + OH(s) => O(s) + PdO + H_2O	3.307E + 21	0.013	191.178
R13)	$+ H_2O + PdO => H_2O(s)$	1.400E-01	0.000	0.000
R14)	$H_2O(s) => PdO + H_2O$	6.293E + 12	0.045	99.946
R15)	$CO_2 + PdO => CO_2(s)$	4.910E-02	0.000	0.000
R16)	$CO_2(s) => PdO + CO_2$	4.087E + 14	0.029	65.097
R17)	$CH_3(s) + PdO => PdO + CH_3(s)$	1.494E + 22	0.008	131.139
R18)	$PdO + CH_3(s) => CH_3(s) + PdO$	2.441E + 23	-0.008	181.201
R19)	$O(s) + CH_3(s) => OH(s) + CH_2(s)$	1.250E + 22	0.012	33.366
R20)	OH(s) + CH2(s) => O(s) + CH3(s)	1.855E + 22	-0.012	303.074
R21)	OH(s) + CH3(s) => H2O(s) + CH2(s) $CH2(s) = CH3(s) + CH3(s) +$	1.398E + 22	-0.004	33.193
R22)	$H_2O(s) + CH_2(s) => OH(s) + CH_3(s)$	1.111E + 22	0.004	211.667
R23)	$CH_3(s) + O(s) => CH_2OH(s) + PdO$	3.534E + 21	0.016	30.953
R24)	$CH_2OH(s) + PdO => CH_3(s) + O(s)$	2.466E + 23	-0.016	333.447
R25)	$CH_3(s) + OH(s) => CH_2(s) + H_2O(s)$	2.014E + 21	0.004 -0.004	73.229 116.851
R26) R27)	$CH_2(s) + H_2O(s) => CH_3(s) + OH(s)$ $CH_2(s) + OH(s) => CH_2OH(s)$	1.175E + 22 3.420E +	-0.004 -0.004	12.581
R28)	+ PdO $CH_2OH(s) + PdO => CH_2(s) +$	2.191E +	0.004	180.219
R29)	$CH_2OH(s) + PdO => CH_2(s) + OH(s)$ $CH_2OH(a) + PdO => CH_2O(s)$	2.191E + 22 1.917E +	0.004	12.654
R30)	+ H(s) $- CH_2OH(a) + PHO = > CH_2O(s)$ $+ H(s) = > CH_2OH(s)$	22 3.348E +	-0.012	57.646
R31)	+ PdO $CH_2O(s) + OH(s) => CHO(s)$	20 5.608E +	0.012	51.570
R32)	$+ H_2O(s)$ $+ H_2O(s)$ $- CHO(s) + H_2O(s) => CH_2O(s)$	21 2.119E +	-0.012	192.230
R33)	+ OH(s) $+ OH(s) => CO(s) +$ $+ OH(s) => OH(s) +$	22 5.921E +	0.008	65.968
R34)	$H_2O(s)$ CO(s) + H2O(s) => CHO(s) +	22 8.990E +	-0.008	267.532
R35)	$OH(s)$ $CH_3(s) + PdO => CH_2(s) + H$	21 1.827E +	0.000	123.976
R36)	(s) $CH_2(s) + H(s) => CH_3(s) +$	21 5.137E +	-0.000	165.224
R37)	PdO $CH_2(s) + PdO => PdO + CH2$	22 1.257E +	0.000	13.103
	(s)	22		

(continued on next page)

Table F7 (continued)

Table F/	(continued)			
No	Reaction	A (cm, mol, s)	n	E (kcal mol ⁻¹)
(R39)	$CH_2(s) + PdO => CH(s) + H(s)$	3.793E + 22	0.008	112.541
(R40)	$CH(a) + H(s) => CH_2(s) + PdO$	2.987E + 22	-0.008	163.159
(R41)	CH(s) + PdO => CHO(s) + PdO	2.103E + 21	0.008	24.821
(R42)	CHO(s) + PdO => CH(s) + PdO	1.218E + 21	-0.008	215.219
(R43)	PdO + CH2(s) => CH2O(s) + PdO	2.155E + 22	0.008	91.222
(R44)	$CH_2O(s) + PdO => PdO + CH_2(s)$	2.425E + 20	-0.008	9.038
(R45)	$CH_2O(s) + PdO => CHO(s) + H(s)$	3.274E + 21	0.008	69.676
(R46)	CHO(s) + H(s) => CH2O(s) + PdO	5.962E + 22	-0.008	207.964
(R47)	CHO(s) + PdO => CO(s) + H (s)	1.088E + 23	0.004	55.610
(R48)	CO(s) + H(s) => CHO(s) + PdO	7.956E + 22	-0.004	254.800
(R49)	CHO(s) + PdO => PdO + CHO (s)	2.746E + 22	0.018	76.549
(R50)	PdO + CHO(s) => CHO(s) + PdO	6.754E + 21	-0.018	106.611
(R51)	PdO + CHO(s) => H(s) + CO (s)	2.732E + 21	-0.002	141.937
(R52)	H(s) + CO(s) => PdO + CHO (s)	2.123E + 21	0.002	263.113
(R53)	CO(s) + PdO => PdO + CO(s)	1.313E + 21	0.011	33.868
(R54)	PdO + CO(s) => CO(s) + PdO	2.212E + 21	-0.011	16.262
(R55)	$PdO + CO(s) => CO_2(s) + PdO$	1.410E + 21	0.010	42.023
(R56)	$CO_2(s) + PdO => PdO + CO(s)$	9.554E + 20	-0.010	78.477
(R57)	PdO + H(s) => OH(s) + PdO	1.886E + 22	0.000	137.800
(R58)	OH(s) + PdO => PdO + H(s)	4.222E + 21	0.000	27.900
(R59)	$O_2(s) + PdO => O(s) + O(s)$	3.815E + 22	0.005	174.961
(R60)	$O(s) + O(s) => O_2(s) + PdO$	1.744E + 21	-0.005	59.139
(R61)	PdO + O(s) => O(s) + PdO	2.380E + 22	0.027	116.625
(R62)	O(s) + PdO => PdO + O(s)	1.797E + 21	-0.027	107.875
(R63)	O(s) + PdO => PdO + PdO	8.012E + 21	0.020	48.597
(R64)	PdO + PdO => O(s) + PdO	1.387E + 22	-0.020	252.103
(R65)	$O_2(s) + PdO => O(s) + PdO$	1.196E + 24	0.025	81.408
(R66)	$O(s) + PdO => O_2(s) + PdO$	9.469E + 22	-0.025	169.092
(R67)	PdO + H(s) => H(s) + PdO	6.133E + 22	0.002	148.173
(R68)	H(s) + PdO => PdO + H(s)	9.505E + 21	-0.002	117.827
(R69)	OH(s) + CHO(s) => H2O(s) + CO(s)	2.992E + 22	0.001	2.168
(R70)	$H_2O(s) + CO(s) => OH(s) + CHO(s)$	3.113E + 22	-0.001	156.062
(R71)	PdO + CHO(s) => HCOO(s) + PdO	3.658E + 21	0.000	11.580
(R72)	HCOO(s) + PdO => PdO + CHO(s)	1.210E + 21	0.000	8.680
(R73)	$HCOO(s) + OH(s) => CO_2(s)$ + $H_2O(s)$	2.305E + 23	0.011	183.191
(R74)	$CO_2(s) + H2O(s) => HCOO(s)$ + $OH(s)$	4.912E + 23	-0.011	376.439
(R75)	$HCOO(s) + PdO => CO_2(s) + H(s)$	1.681E + 23	0.007	177.623

Table F7 (continued)

_	No	Reaction	A (cm, mol, s)	n	E (kcal mol ⁻¹)
	(R76)	$CO_2(s) + H(s) => HCOO(s) + PdO$	1.726E + 24	-0.007	368.497
	(R77)	$H_2O(s) + PdO <=> OH(s) + H$ (s)	3.659E + 21	0.000	12.500

fuel species (H2 vs CH4) Da for hydrogen is slightly more than the methane which is likely due to more propensity of reactor catalyst to absorb and combust hydrogen. Da for methane is almost constant along the reactor while it shows an increasing trend towards hot end. For flame marker OH, similar to fuel species, Da also remains within a limited bound and does not go beyond 10. Indeed, the affinity of the surface catalyst for OH is higher to remove the main flame marker from the gas phase, possibly resulting in more intense surface than gaseous phase reactions. The spatial Da also demonstrates that the evolution of the combustion is from the surface of catalysts locating everywhere in the chamber. The length is sufficient to achieve around 90% combustion efficiency and 93% effectiveness factor in the case of intalox saddle. The Da for other case studies (catalyst pellets and monolith), not shown here, demonstrates that the designed reaction is with appropriate dimensions to maintain overall combustion over the entire length of the combustor at temperature near the ignition temperatures. At such a temperature, the gaseous reaction phase, if any, evolves throughout the entire combustor volume and combustion is nearly flameless. The reactor temperature was below stoichiometric mixture temperature.

Conclusions

This work delivers a new proof of concept for the invention of catalytic microcombustors to burn syngas. The DNS and finite chemistry were applied to model the reactive flow within the designed combustor and to analyse the combustion performance under different operating conditions. Under the first phase of the design procedure, the premix combustion within the micocombustor is evaluated to maintain low NO_X emission, the combustor ability to light, toleration of sudden changes in power loads, efficient combustion, and achievement of the required life span by using various design parameters. Within twelve different shapes of catalyst pellet and four monolithic structures, the microcombustor demonstrates the best combustion performance with the catalyst in the shape of intalox saddle with a combined least possible pressure drop and maximum combustion efficiency. The designed reactor is also well able to respond to the variation in fuel composition and catalyst material, temperature and mass flow rate fluctuations by maintaining the combustion efficiency of more than 90%. The microcombustor can control the combustion on the surface of the catalyst and limit the gaseous phase reactions with proper equivalence ratio.

The evolution of surface averaged Da demonstrates that the entire 200 mm reactor is well able to participate in the dissociation reaction without leaving any part ineffective and appearance of both cold and hot spots. The low spatial gaseous concentrations of OH gives rise to comparatively higher contribution of the surface reactions in the combustion. Here, the combustion is uniform and no significant rise in combustion temperature in the gas phase is observed throughout the reactor.

The fluid structure inside of the packed bed reactors is very complex, especially when as fluid channeling, fluid-solid interactions are intermingled with the stochastic nature of the fluid motion. To provide a thorough and comprehensive analysis, the reactor should be built and experimented using proper laser diagnostic techniques, the models should be validated, and employed to delve into the physics and thermochemical state of the combustion within the designed micro combustor.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Data for this aricle can be accessed through the Brunel University London repository at: $\frac{https://doi.org/10.17633/rd.brunel.23706051.}{v1}.$

Appendix A. Random pack illustration³

Random Pack 1

Fig. A1.

Random Pack 2

Fig. A2.

Random Pack 3

Fig. A3.

Appendix B. Structured bed illustration

Structured Pack 1

Fig. B1.

Structured Pack 2

Fig. B2.

Appendix C. Packing illustration

Fig. C1

Appendix D Monolith illustration

Fig. D1.

Appendix E Surface chemistry

Fig. E1.

Appendix F. Surface chemistry

Hydrogen (H_2) surface dissociation chemistry set on Pd/Rh. Tables F1 and F2.

Carbon monoxide (CO) surface dissociation chemistry set on Pd/Rh. Tables F3 and F4.

Syngas (CH₄/H₂/CO) surface dissociation chemistry set on Pd/Rh. Tables F5 - F7.

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