Article

Turbulence Enhancement and Mixing Analysis for Multi-Inlet Vortex Photoreactor for CO$_2$ Reduction

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1. Introduction

Fossil fuels are the predominant energy source in today’s society. Although most of the current research in the field of renewable energies focuses on the study and design of solar panels and wind turbines [1–3], these technologies only represent the 5% of the worldwide energy generation, while oil, coal, and natural gas have more than 85% [4]. The use of fossil fuels leads to the generation of greenhouse gases, where CO$_2$ is the most abundant anthropogenic gas [5]. One of the possible flanks of attack on this problem is the synthesis of useful chemical products using CO$_2$ as raw material. It is in this way that CO$_2$ emission could be reduced, starting a new stage of energy supply-based in circular economy.

Photocatalysis is one of the ways that allow CO$_2$ conversion to chemical products. This type of technology provides, in addition to chemical products, a way to long-term energy storage [6,7]. In the photocatalytic process, photoactive materials absorb sunlight, which promotes the electronic excitation of the material, generating the hollow-electron pair, a process that is important to carry out reduction or oxidation reactions [8–14]. Although the study of new photoactive materials is a fundamental part of CO$_2$ transformation, the systematic study of transport phenomena and reactor engineering is fundamental to achieve high performance systems [15–20]. There is a consensus concerning the geometry and arrangement of a photoreactor since it affects both the yield and selectivity of the chemical process that is carried out [21,22]. In addition, recent literature [23] has reported
the relevance of combining light and heat to enhance the catalytic reaction. In particular, the CO\(_2\) reduction using water as a reactant molecule (that is called the wet process CO\(_2\) reduction) aims to obtain products such as methanol, formaldehyde, etc. The vision of chemical transformations within experimental set-ups where sunlight is used has strong relevance and is the aim of green technology.

Two reactor configurations are used extensively for applications in CO\(_2\) reduction, with photocatalytic methods, the continuous flow system and the batch system. The batch system is one of the most studied. However, its photocatalytic efficiency is low, and given its condition, it becomes imprecise compared to other methods [24]. The key limitation of the batch reactor system is the products accumulation inside the reactor, which can lead to changes in the concentration of the reactants and reabsorption at the surface of the photocatalyst. On the other hand, although, continuous flow reactors have better efficiency, the production of compounds is inadequate due to the short residence time of the reactants inside the reactor chamber, reducing the contact time with the photocatalytic material [25,26].

The fluid dynamics dependence on the geometric configuration of the reactor, makes the optimization process a complex problem to solve, since an analytical representation of the problem cannot be proposed. One of the alternatives to solve such type of optimization problems, in addition to numerical methods, are stochastic algorithms. In this work, Genetic Algorithms, one of the most popular stochastic methods [27], are used to search for the geometric configuration that increases the residence time. Once the best design is obtained, a fine-tuning process is performed to determine the inlets flow that maintains a prolonged residence time, high turbulence intensity near the photocatalytic bed, and an adequate molar fraction of gases. The mixing capability and turbulence enhancement of the optimized photoreactor are evaluated. Likewise, the effect of the cone-shaped geometry of the reactor is studied.

2. Methodology
2.1. Optimization Process

Genetic Algorithm

The implemented GA consists of six main steps; four of them correspond to the evolutionary loop, where the selection of parents, crossing, mutations, creation of offspring and the choice of individuals for the next generation takes place; the other two steps correspond to the initialization and termination of the evolutionary loop, where the maximum number of iteration (MaxIt) is reached, see Figure 1. The algorithm can find a local minimum of a real and real-valued objective function \( f(x) \). The objective function corresponds to the volume rate (\( \Phi_{V f} \) mL/min) that passes through the photocatalytic bed. In all the runs for the GA the three inlets have a volume flow rate of 83.3 mL/min with CO\(_2\) at 298 K, and the outlet has a pressure opening at 100 kPa, as boundary conditions for the CFD.

There are seven geometric design variables that define each member of the population for the GA. Three design variables correspond to the angle around the main chamber of the three inlets, namely \( \theta_A \), \( \theta_B \), and \( \theta_C \). These inlets are always tangent to the circle formed by the cut of the cone of the main chamber at their specific height, and have a diameter of 1.59 mm. Another three design variables are the height of the inlets, namely \( H_A \), \( H_B \) and \( H_C \), all the heights are measured from the surface of the photocatalytic bed. Finally, the wall angle in the main chamber defines the cone shape of the main chamber, which is labeled as \( \theta_D \).
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Only two dimensions are kept constant, the height of the main chamber, with a value of 14.30 mm, and the diameter of the photocatalytic bed, with a value of 10 mm. The gene of a specific individual in the GA will be, 

$$y_m = (\theta_A, \theta_B, \theta_C, \theta_D, H_A, H_B, H_C).$$

(1)

The design limits for each variable are shown in Table 1. For $\theta_A$, $\theta_B$, and $\theta_C$ the values were selected to guarantee that they do not overlap, as well as $H_A$, $H_B$ and $H_C$.

Table 1. Limit values for the reactor design variables.

| Name  | Maximum Value | Minimum Value |
|-------|---------------|---------------|
| $\theta_A$ | 360.0° | 310.0° |
| $\theta_B$ | 168.0° | 70.0° |
| $\theta_C$ | 288.0° | 190.0° |
| $\theta_D$ | 150.0° | 90.0° |
| $H_A$ | 13.5 mm | 2.0 mm |
| $H_B$ | 13.5 mm | 2.0 mm |
| $H_C$ | 13.5 mm | 2.0 mm |

The evolutionary loop was executed several times before the selection of the optimized geometric configuration, the design variables obtained by GA are presented in Section 2.2.1.

2.2. Tunning Process for the Optimized Photoreactor

2.2.1. Reactor Configuration

The study photoreactor, called optimized photoreactor, is characterized by having a quartz window at the top, which allows light to enter the main chamber. It has three inlets that surround the cone-shaped body of the photoreactor (Inlet A, Inlet B and Inlet C) Figure 2a. At the bottom, before the outlet, the photocatalytic bed is placed, Figure 2b. For the CFD the photocatalytic bed is simulated as an isotropic porous support with 95% of porosity and 1.00 mm thickness. The height from the catalytic bed to the quartz window...
is 14.30 mm. The mean feature of the optimized photoreactor is the position of the three inlets around the photoreactor. The inlets are at different heights with respect to the photocatalytic bed, namely $H_A$, $H_B$ and $H_C$, and at different angles, namely $\theta_A$, $\theta_B$, and $\theta_C$. The rise of the cone of the main chamber, namely $\theta_D$ is 23.0° Figure 2c. All the values described are in Table 2.

Table 2. The optimized set of parameters for the reactor configuration, obtained by Genetic Algorithms.

| Name   | Value  |
|--------|--------|
| $\theta_A$ | 313.4° |
| $\theta_B$ | 74.0° |
| $\theta_C$ | 240.9° |
| $\theta_D$ | 23.0° |
| $H_A$     | 11.0 mm |
| $H_B$     | 7.2 mm  |
| $H_C$     | 6.3 mm  |

The reference reactor used for comparative purposes to the prototype reactor is depicted in Figure 3b. It is important, especially for explaining the fluid behavior, to emphasize that the gas inlet in both the prototype and the reference reactor are formed by a hole with a chamfer arrangement that makes it relevant to define properly which inlet is considered (A, B or C).

The criteria to evaluate the performance of both optimized configuration of the prototype reactor and the reference are the value of the turbulence and the composition (in mole fraction) at the region along the reactor vessel and, especially, at the zone of the catalytic bed. Because of this, the optimized prototype reactor was divided into five different regions, whereas the reference reactor was divided into four related to the planes depicted in Figures 4 and 5. The reference reactor was proposed based on the characteristics commonly found in multi-inlet vortex reactors [28,29]. The height of the reference and the optimized reactors are equal but the wall in the main chamber of the former is 90° since it is related to a cylindrical vessel. In addition, the gas inlets are equidistant from each other and located at the same height of the vessel. Nevertheless, it is essential to remind that inlet A, B, and C can affect the turbulence intensity since gas flow through inlet A hits directly to C, gas flow through inlet B hits directly to A and gas flow through inlet C hits directly to B, see Figure 5b. The effect of this configuration in the turbulence intensity within the reference reactor is one topic of interest of the CFD simulations.
Figure 3. Reactors for study: (a) shows the optimized photoreactor by genetic algorithm, and in (b) there is the Reference photoreactor.

Figure 4. Planes of the imaginary surfaces (a) of the optimized photoreactor vessel and that related to the vertical cut (b) of reactor body.

Figure 5. Planes of the imaginary surfaces (a) of the reference reactor vessel and that related to the vertical cut (b) of reactor body.
2.2.2. Inlets Gas Flow Considerations

After obtaining the best reactor design, through GA, the optimal model was studied varying the substance type at the inlets as well as their volume flow rate, to determine the turbulent behavior near the surface of the photocatalyst including the mole fraction. In all the studies, the volume flow rate of two inlets (either A and B fixed and C varying, or B and C fixed and A varying or, finally, A and C fixed and B varying) were kept at 60 mL/min. The configuration for the substance inlet for the case of the prototype reactor is shown in Table 3. There are always two inlets with H$_2$O at 374 K and one inlet with CO$_2$ at 298 K. The reason for this configuration, is to have a molar ratio CO$_2$:H$_2$O equal to 1:2, as previously mentioned, to foresee the formation of molecules such as methanol (CH$_3$OH), formaldehyde (CH$_2$O) and molecular oxygen (O$_2$) as expected reaction products. It is obvious that other heavier molecules might be formed and that might imply a different reactant molar ratio, but as molecular oxygen is expected to be formed, this would balance the chemical reaction. Briefly, the presence of two moles of water provides enough hydrogen to obtain the reduced products. The difference in temperature of the reactant gas flows leads to evaluate the changes in thermal energy within the reactor that will be analyzed and discussed later.

Table 3. Gas configuration for the turbulence and Mole Fraction Tuning.

| Factor          | Level Name | Level Value |
|-----------------|------------|-------------|
| Gas configuration | Inlet A    | CO$_2$      |
|                 | Inlet B    | H$_2$O      |
|                 | Inlet C    | H$_2$O      |
| 2               | Inlet A    | H$_2$O      |
|                 | Inlet B    | CO$_2$      |
|                 | Inlet C    | H$_2$O      |
| 3               | Inlet A    | H$_2$O      |
|                 | Inlet B    | H$_2$O      |
|                 | Inlet C    | CO$_2$      |

2.3. Computational Fluid Dynamics Simulation

SolidWorks Flow Simulation™ is a solver for Computational Fluid Dynamics (CFD) that implements the finite volume methodology using a Reynolds-averaged Navier–Stokes approach, alongside the modified k-ε turbulence model with damping functions proposed by Lam and Bremhorts, using the SmartCell® approach for mesh generation [30]. The solver describes laminar, turbulent and transitional flows of homogeneous fluids, and employs two transport equations, one for the turbulent kinetic energy ($k$), Equation (2), and the second for the turbulent dissipation ($\varepsilon$), Equation (3) [31].

\[
\frac{\partial \rho k}{\partial t} + \frac{\partial \rho k u_i}{\partial x_i} = \frac{\partial}{\partial x_j} \left( \left( \mu + \frac{\mu_t}{\sigma_k} \right) \frac{\partial k}{\partial x_j} \right) + \tau_{ij}^k \frac{\partial u_i}{\partial x_j} - \rho \varepsilon + \mu_t P_B \tag{2}
\]

\[
\frac{\partial \rho \varepsilon}{\partial t} + \frac{\partial \rho \varepsilon u_i}{\partial x_i} = \frac{\partial}{\partial x_j} \left( \left( \mu + \frac{\mu_t}{\sigma_\varepsilon} \right) \frac{\partial \varepsilon}{\partial x_j} \right) + C_{\varepsilon 1} \frac{\varepsilon}{k} \left( f_1 \tau_{ij}^R \frac{\partial u_i}{\partial x_j} + C_{\varepsilon 2} \mu_t P_B \right) - f_2 C_{\varepsilon 2} \frac{\rho \varepsilon^2}{k} \tag{3}
\]

with

\[
\tau_{ij} = \mu s_{ij}, \tag{4}
\]

\[
\tau_{ij}^R = \mu_t s_{ij} - \frac{2}{3} \rho k \delta_{ij}, \tag{5}
\]

\[
s_{ij} = \frac{\partial u_i}{\partial x_j} + \frac{\partial u_j}{\partial x_i} - \frac{2}{3} \delta_{ij} \frac{\partial u_k}{\partial x_k}, \tag{6}
\]

\[
P_B = -\frac{g_i}{\rho_0} \frac{1}{\rho} \frac{\partial \rho}{\partial x_i}. \tag{7}
\]
In which $C_{\mu} = 0.09, C_{\epsilon_1} = 1.44, C_{\epsilon_2} = 1.92, \sigma_{\epsilon} = 1.3, \sigma_B = 0.9, \sigma_B = 1$ if $P_B > 0$, $\sigma_B = 0$ if $P_B < 0$.

The turbulent viscosity is determined by:

$$\mu_t = f_\mu \frac{C_{\mu} \rho k^2}{\epsilon},$$  \hspace{1cm} (8)

The Lam and Bremhorst’s damping function $f_\mu$ is determined by:

$$f_\mu = \left(1 - e^{-0.025 R_y}\right)^2 \left(1 + \frac{20.5}{R_t}\right),$$  \hspace{1cm} (9)

where

$$R_y = \frac{\rho \sqrt{k_y}}{\mu},$$  \hspace{1cm} (10)

$$R_t = \frac{\rho k^2}{\mu \epsilon},$$  \hspace{1cm} (11)

in this case, $y$ is the distance from a point to the wall and Lam and Bremhorst’s damping functions $f_1$ and $f_2$ are determined by:

$$f_1 = 1 + \left(\frac{0.05}{f_\mu}\right)^3,$$  \hspace{1cm} (12)

$$f_2 = 1 - e^{R_t^2}.$$  \hspace{1cm} (13)

The heat flux is defined by:

$$q_i = \left(\frac{\mu}{Pr} + \frac{\mu_t}{\sigma_{\epsilon}}\right) \frac{\partial h}{\partial x_i}, i = \{1, 2, 3\},$$  \hspace{1cm} (14)

where, $\sigma_{\epsilon} = 0.9$, $Pr$ is the Prandtl Number, and $h$ is the thermal enthalpy.

Another important quantity for analysis is the turbulence intensity, which is defined as,

$$I \equiv \frac{u'}{U},$$  \hspace{1cm} (15)

where $u'$ is the root-mean-square of the turbulent velocity fluctuations and $U$ is the mean velocity (Reynolds-averaged), and it can be computed as

$$I \equiv \sqrt{\frac{1}{3} \left(u_x'^2 + u_y'^2 + u_z'^2\right)} = \sqrt{\frac{1}{3} k},$$  \hspace{1cm} (16)

The mean velocity can be obtained from the three mean velocity components as

$$U \equiv \sqrt{\frac{1}{3} \left(u_x^2 + u_y^2 + u_z^2\right)},$$  \hspace{1cm} (17)

All the thermodynamic properties of the gases used in this work were obtained with the engineering data base of SolidWorks Flow Simulation™. The mesh selection for the reactors could be reviewed in the S1 Supplementary Material.

3. Results and Discussion
3.1. CFD Simulations Results
3.1.1. Turbulence Intensity and Mole Fraction Profile

Two additional aspects of the performance of the reactor were considered by calculating the turbulence intensity and the composition (mole fraction) over the whole system and in the region close to the catalyst bed (plane 1). As mentioned in Section 2.2.2, the
total gas flow rate was split into two flows of water and one for CO₂. In this case, each volumetric flow rate was 83.3 mL/min. The set of calculations amounted to 4 in which the turbulence intensity was evaluated, as depicted in Figure 6. It can be observed that the turbulence intensity within the optimized reactor vessel in bulk and on the surface of the catalyst bed changes drastically depending on the gas inlet configuration as compared to the reference reactor. It must be important to mention that the number following the optimized photoreactor (Opt) code refers to design detailed in Table 3. The CO₂ inlet flow configuration affects the turbulence intensity depending on which injection port is used. Nevertheless, the tendency indicates that the turbulence intensity increases if CO₂ is fed from top to bottom keeping the volumetric flow rate constant. The case of the reference reactor shows that the turbulence is higher in the bulk of the system but, at the zone of the catalyst zone, it becomes lower than the Opt-3 (Opt reactor with CO₂ fed at the lower inlet port). In this case, since the flow rate was kept constant and equal in all the inlets, it is irrelevant to detail which inlet (A, B or C) is used to fed CO₂ in the reference reactor. This means that in the case of no variation of the volumetric inlet flow rate is set, the net effect of changing the CO₂ inlet port in the reference reactor has no impact on the turbulence intensity or mass fraction distribution.

![Average turbulence intensity using Table 2 of gas inlet configuration. Data from the main chamber and at the photocatalytic bed surface.](image)

Figure 6. Average turbulence intensity using Table 2 of gas inlet configuration. Data from the main chamber and at the photocatalytic bed surface.

A different behavior is observed when the volumetric inlet gas flow rate is changed. Figure 7a,b show that the turbulence intensity increases as the gas flow changes keeping in mind that the assigned inlet port for CO₂ is C in both reference and Opt-3. However, to enhance and discuss these results properly, it should be mentioned that the gas flow was varied for both Opt-3 and the reference reactor in all the inlet ports (A, B or C) and that logically, changes in gas flow through A and B means changes in the volumetric flow rate of water. Those volumetric flow rates kept constant were set at 60 mL/min in all cases. In addition, the turbulence intensity values plotted in this figure are referred to the plane close to the catalytic bed (plane 5 in Opt-3 and plane 4 in the reference reactor). This indicates that the catalyst will find those values of turbulence intensity, which is implicitly is related to the energy content for the chemical reaction to occur. Turbulence is linked to both homogeneity of the system (stoichiometric requirement) and kinetic energy associated with the movement of the reactant molecules. The higher the value of the turbulence intensity the better the energy requirement will be fulfilled for the chemical reaction. This behavior was expected because, as the volumetric flow rate increases, more mass is entering for which its velocity is higher so that the turbulence intensity also increases. The higher values
of turbulence intensity were found in the reference reactor case although the observed difference is relatively low as compared to the Opt-3 case.

When varying the water inlet flow rate (inlet A and B), the turbulence intensity increases in a smooth shape. On the contrary, a steep increase in turbulence is observed as the gas flow increases through inlet C. This result is important since CO\textsubscript{2} is fed through inlet C, which indicates that varying its flow rate affects the turbulence within the reactor. The values of the turbulence intensity are close between both reference and Opt-3 reactors. Hence, it can be concluded that the effect of changing the volumetric gas flow rate affect proportionally to the turbulence intensity with slightly better performance in the reference reactor.

Together with turbulence, there is the mass fraction concern while varying the volumetric gas flow rate and the inlet port in the reactor. Again, inlet C is used for CO\textsubscript{2} and A and B for water. In the Opt-3 reactor, inlet C is located at the lower part of the reactor vessel. Figure 8 shows the behavior of mole fraction as a function of the volumetric gas flow rate in both reference and Opt-3 reactors. The calculated plotted mole fraction values were estimated in the proximity of the catalytic bed surface (plane 5 in Opt-3 and plane 4 in the reference reactors). The emphasis of this result lies in the molar ratio obtained at the catalytic site that corresponds to the stoichiometric requirement for the chemical reaction (CO\textsubscript{2}:H\textsubscript{2}O 1:2). No significant difference between reference and Opt-3 reactor can be observed. This is also related to the good turbulence achieved in the whole system and at the closeness of the catalyst. With this result, it has been established that the operating conditions in which the optimal dynamics of the system in the Opt-3 reactor were reached are those summarized in Table 4.

Table 4. Optimal operating conditions based on turbulence intensity and stoichiometric CO\textsubscript{2}:H\textsubscript{2}O molar ratio in Opt-3 reactor.

| Factor          | Level Name | Level Value | Volume Flow Rate (mL/min) |
|-----------------|------------|-------------|----------------------------|
| Gas configuration | Inlet A    | H\textsubscript{2}O | 60                          |
|                 | Inlet B    | H\textsubscript{2}O | 90                          |
|                 | Inlet C    | CO\textsubscript{2} | 60                          |
Table 4. Optimal operating conditions based on turbulence intensity and stoichiometric CO2 sub-

Figure 7. Turbulence intensity profile as a function of the volumetric gas flow rate for both the reference (dotted lines) and Opt-3 (solid lines) reactors. Test case varying volumetric water (a,b) CO2 flow rate. The constant volumetric flow rate was set at 60 mL/min. For further details, refer to Figures 3 and 4.

Figure 8. Mole fraction profile of CO2 in both reference (dotted lines) and Opt-3 (solid lines) reactors. Test case varying volumetric water (a,b) CO2 flow rate. The constant volumetric flow rate was set at 60 mL/min. For further details, refer to Figures 3 and 4.

Those settings of Table 4 led to the residence time distribution depicted in Figure 9. The residence time was obtained from a particle study, based on the Lagrangian particle tracking approach [32], where 30,000 representative particles of the corresponding substance were injected at each inlet, after obtaining the fluid dynamics, for each particle the time elapsed inside the computer domain was tracked. The residence time plays an important role whenever a chemical reaction is about to occur. The case of the reference reactor indicates that the distribution of the particles is divided into two big lumps with relatively close residence time values (that it has been called bimodal). On the other hand, the residence time distribution of the named Opt-3 tuned reactor (based on the aforementioned operating conditions) shows a multimodal distribution with slightly different time values among them. Considering the average values, the residence time of the reference reactor yields 0.35 s, while that of the Opt-3-tuned reactor yields 0.85 s. This means that the time used for the reactant molecules within the prototype reactor is almost three times the value obtained for the reference. Again, this might be priceless at the moment in which the chemical reaction takes place.

Figure 9. Residence time distribution of the Opt-3 tuned (red) and reference (blue) reactors. The operating conditions used for calculation are summarized in Table 4.
3.1.2. Graphical Profile of Reference and Opt-3-Tuned Reactor

A better look at the effluent behavior within the reference and Opt-3-tuned reactors is depicted by the images in Figures 10–15. Again, the different coloring regions help visualize how the reactant mixture behaves depending on the studied variable. For instance, Figure 10 shows that under conditions of Table 4, the turbulence at the interior of each reactor vessel behaves differently depending on the location. However, for the sake of clarity, the turbulence found at the catalyst region is somewhat similar in both systems, see Figure 10c,d.

Notably, the turbulence at the center and on the upper region of the Opt-3-tuned reactor has low turbulence (dark blue spots) since there is the vortex formation (whirl) due to the gas entrance, the arrangement of this inlet and the shape of the hole (chamfer shape). As expected, in both reactors, once the reactant mixture has passed through the catalyst support, the gas leaves the reactor in a laminar regime.

![Figure 10. Turbulence intensity profile in bulk (plane 4 for reference (a) and plane 5 for Opt-3-tuned (b) reactors) and at the region of the catalytic bed (plane 1 in both reference (c) and Opt-3-tuned (d) reactors).](image)

Concerning the composition within the system for the Opt-3-tuned reactor, Figure 11 shows the calculated mass fraction profile at different layers of the current reactor vessel (see the horizontal planes 1–4 in Figure 4). For the sake of knowledge, the mass fraction depicted in this figure is what the program yields, and that is the reason the mole fraction is not currently plotted. Nevertheless, the correspondence between one and another is that for the stoichiometric molar requirement of the reaction (CO$_2$:H$_2$O 1:2) the CO$_2$ mole fraction should be 0.33, equivalent to the CO$_2$ mass fraction equal to 0.55. Therefore, the CO$_2$ mass fraction near 0.55 refers to the molar ratio needed for the chemical reaction of
current interest. Therefore, according to the mass composition in the Opt-3-tuned reactor, on the surface of the catalyst, there will be the amount of reactant needed to perform the chemical reaction. This, together with the turbulence requirement previously discussed, can be set as the conditions to perform the chemical process without having mass transport limitations and an apparent energy requirement fulfilled.

Another perspective of the CO₂ mass fraction profile along both the reference and the Opt-3-tuned reactor is shown in Figure 12. Once CO₂ and water enter the reactor vessel, the mixture becomes more homogeneous in prototype rather than in reference reactor, which is reflected in the yellow to greenish color that reaches the catalyst area in the reactor. It seems that although turbulence is slightly higher in the reference reactor, there is still a gradient of reactants concentration that reaches the catalytic bed, which is much less pronounced in the Opt-3-tuned reactor. For the latter case, the conical shape enhances the homogeneous distribution of reactants even though turbulence is, under these conditions, one value lower than the reference case.
Figure 11. CO$_2$ mass fraction profile as a function of the position within the Opt-3-tuned reactor. The mass fraction at the photocatalyst surface (a); red color (b) corresponds to CO$_2$ gas inlet. Blue color (c, d) refers to H$_2$O gas inlet.

Figure 12. CO$_2$ mass fraction profile in the bulk of the reference (a) and Opt-3-tuned (b) reactors.

Figure 13. Calculated particle trajectory velocity vector for reference and Opt-3-tuned reactor at the vertical plane (a,b) and above the catalytic bed surface (c,d), respectively.

Two additional aspects of interest, while dynamic simulations are performed are the vectorial performance of the particle trajectory and the streamlines related to the eddy
formation during the gas mixing. The calculated speed vector of the trajectory of the particle, Figure 13, shows that once the volumetric gas flow rate enters the reactor vessel, the velocity of the particles becomes almost zero at any point in the bulk of both types of reactors. This means that inertial and gravitational forces rather than the turbulence effect might govern the movement of the particle. In other words, the direction and speed of the particles seem to depend on the inlet flow rate, and its configuration in the reactor vessel is also influenced by its geometrical shape (cylindrical and conical). It seems a logical result to find that the velocity vector in Opt-3-tuned is better distributed than in the reference reactor because, besides the conical shape, the upper inlet port exerts this inertial force enhanced by the other two inlet ports as the gas effluent reaches the exit. A situation in reference reactor cannot be observed because all the inlet ports are located at the same height from the reactor exit.

The qualitative eddies formation and their size and distribution are better observed in the image of Figure 14. The streamlines depicted in this figure seem to be larger and rounded in the Opt-3-tuned reactor, whereas elongated streamlines are observed in the reference reactor. Only to emphasize the importance of this picture, large eddies are related to the higher kinetic energy of the particles in the gas effluent. In contrast, small eddies promote an increase in thermal energy because more friction forces are involved. It can be concluded that the behavior of the dynamics of the gas reactant mixture for both types of reactors are rather similar because eddies of comparative size, but different shapes are formed in both reactors.

Figure 14. Streamlines profile in both reference and Opt-3-tuned reactor view along the reactor vessel (a,b) and on the closeness of the catalyst reactor (c,d).
Finally, but no less important for the current work is the temperature profile of the reactant gas mixture for both types of reactors. A marked temperature gradient in both reactors that is related to the temperature of the water (as vapor) inlet flow rate and that of the CO$_2$ (gas) inlet flow rate. To feed water in the vapor phase, this must be evaporated, which means to raise temperature to its boiling point (~373.15 K). On the other hand, CO$_2$ is a gas at room temperature, and there is no need to be heated up. A rough calculation of the equilibrium temperature, based on specific heat and mass of both reactants accordingly to the volumetric standard flow rate, yields that the temperature at thermal equilibrium is 346.3 K which corresponds to the yellow color in Figure 15. The slightly more uniform temperature profile observed within the Opt-3-tuned reactor might be relevant when the chemical reaction occurs since the thermal energy is better distributed in this reactor than in the reference one.

![Figure 15](image1.png)

**Figure 15.** The temperature profile in the reference and Opt-3-tuned reactors at the bulk (a,b) and on the closeness of the catalytic bed (c,d).

### 3.1.3. Main Chamber Cone Influence

To verify the influence of the main chamber cone of the prototype photoreactor, which was generated from the variable $\theta_D$ (see Figure 2); a new design was established, namely Opt-3-Tuned-No-Cone Figure 16. This new reactor design has the inlet locations with the same values as those shown in Table 2 in Section 2.2.1. This new design considers $\theta_D = 90^\circ$, which is the value used in the reference reactor. For comparative purposes, the inlet fluids were taken from Section 3.1.1 Table 4, where the optimal operation values were found.
Figure 16. Planes of the imaginary surfaces of the Opt-3-Tuned-No-Cone (a) of the prototype reactor vessel and that related to the vertical cut (b) of reactor body. (c) Isometric view of the prototype reactor vessel in which the gas inlet configuration is highlighted.

Figure 17 shows the calculated distribution of the residence time for Opt-3-Tuned-No-Cone reactor, reference reactor and the Opt-3-tuned reactor and the one-inlet reactor. The reference reactor has two main regions in its distribution, while the Opt-3-Tuned-No-Cone reactor has a single area distributed in time. As mentioned in Section 3.1.1, the longest residence time was obtained for the optimized photoreactor and the shortest time for that of the one-inlet reactor design. The later configuration seems to be a completely non-efficient system since the reactant gas mixture passes through the reactor almost without interacting due to the short contact time (see Table 5). On average, the residence time of the Opt-3-Tuned-No-Cone and that of the reference reactor are the same besides their distribution profile is entirely different.

Figure 17. Residence time distribution of the Opt-3 tuned (red), reference (blue) and Opt-3-Tuned-No-Cone (pink) reactors. The operating conditions used for calculation are summarized in Table 4.
Table 5. Reactors mean residence time using operation conditions described in Table 4.

| Reactor                  | Mean Residence Time (s) |
|--------------------------|-------------------------|
| Opt-3-Tuned              | 0.85                    |
| Opt-3-Tuned-No-Cone      | 0.35                    |
| One Inlet                | 0.20                    |
| Reference                | 0.35                    |

The average turbulence intensity near the surface of the photocatalytic bed of all the reactor systems is shown in Table 6. The best result was obtained for the Opt-3-Tuned-No-Cone photoreactor followed by the Opt-3-tuned with a slightly different values for the reference and one-inlet reactors. This result might be due to the position of the inlets that enhances the eddies formation that appears to be stable and remains without changing along the body of the No-cone reactor vessel, a situation that is not assured in the cone reactor vessel, logically, because of the reactor geometry. Although the position of the inlets controls the degree of turbulence (that remains in the cylindrical reactor vessel), the cone-shaped geometry controls the residence time, which is also an essential parameter in catalytic processes.

Table 6. Reactors average Turbulence Intensity at the Photocatalyst surface.

| Reactor                  | Photocatalyst Surface Turbulence Intensity (%) |
|--------------------------|------------------------------------------------|
| Opt-3-Tuned              | 13.07                                           |
| Opt-3-Tuned-No-Cone      | 17.97                                           |
| One Inlet                | 11.10                                           |
| Reference                | 12.17                                           |

For the sake of comparison of the mass fraction, velocity vectors, turbulence and temperature profile with those of the reference and optimized photoreactor, Figures 18–22 show the results obtained with the reactor vessel of the Opt-No-cone system. Rather similar velocity vectors are found for both the conical and No-cone reactor vessels (see Figure 13). However, there is a slight mass fraction distribution difference between these reactors which is due to the geometrical shape. It seems that a more uniform distribution of reactants is obtained using the conical reactor vessel that the non-conical one (see Figures 12 and 19). As mentioned, the turbulence profile is better in the non-conical reactor vessel than that in the conical one (see Figures 10 and 20) which also can be qualitatively verified by the eddy formation depicted in Figures 14 and 22. Concerning the temperature profile, there is a more notorious temperature gradient in the non-conical reactor vessel than in the conical. Again, the geometry of the vessels plays the role of reaching a better distribution of heat in the conical configuration than in the cylindrical one (see Figures 15 and 21).

The configuration of cone-shaped geometry of the reactor with the use of broadband radiation is compatible, using natural or artificial solar light. We consider light-assisted CO$_2$ conversion over heterogeneous catalyst supported by the catalytic bed surface inside the optimized reactor chamber. The fused quartz window at the top of the photoreactor, which allows light to enter the main chamber, is very efficient for transmitting UV-Visible and Infrared radiation. The bed surface area can be irradiated by focusing an external solar simulated light from a commercial lamp, selected wavelength lasers, or optically filtered source. Light-assisted is attributed to electron-hole pairs generated in the catalysts by the photon energies presented in the solar spectral range [22,23]. Thus, we can study multiphoton capture, light penetration depth, and uniform photon access of bed surfaces (foams, films, packed bed, catalysts prepared by sputtering, among others) [33]. Even the light that is potentially absorbed and overlapped, not only in the catalytic bed surface with active species for the photoreduction, but also an increment in the vibration and certain polarization of the molecules of CO$_2$ and H$_2$O is expected, promoting an increment in their reactivity. Additionally, catalyst temperature with high average power can be optically
increased [21]. Therefore, efficiently absorbed with respect to available irradiated photons is typically considered. On the other hand, the product molecules respect for such absorbed photons is also required to study the optimal thickness and optical properties of the photocatalyst. The combination control of gas mixing and the selection of solar radiation allows the selective product molecules with the practical option of residence time. The residence time is an excellent tool for improving light-matter interaction, studying thermodynamics, kinetics, and photocatalytic activity properties [34,35], and the concentration control of the reactants and reabsorption at the surface of such photocatalyst is possible.

**Figure 18.** Calculated flow trajectories for the Opt-3-Tuned-No-Cone reactor on the closeness of the catalytic bed (a) and at the bulk (b).

**Figure 19.** Mass fraction of Carbon dioxide for the Opt-3-Tuned-No-Cone reactor on the closeness of the catalytic bed (a) and at the bulk (b).
From a fundamental point of view, the mechanistic (synergistic) steps involved, while a photocatalytic process is performed, are related to alternative pathways compared to those involved in thermal (dark) processes. Although photonics is always complicated and many speculations emerge when it is used for chemical transformations, greater and greater insight into catalyst characterization has been gained that allows the establishment of more convincing reaction mechanisms based on both the molecular electronic behavior and their interactions [21–23]. Prototype reactors, hence, become important because it sets the path for scaling-up. Assuring the latter, and with the long residence time that is currently evaluated, it is likely to obtain those aforementioned eco-fuels from the wet process CO$_2$ reduction.

**Figure 20.** Turbulence Intensity for the Opt-3-Tuned-No-Cone reactor on the closeness of the catalytic bed (a) and at the bulk (b).

**Figure 21.** Temperature distribution for the Opt-3-Tuned-No-Cone reactor on the closeness of the catalytic bed (a) and at the bulk (b).
temperature with high average power can be optically tuned to study the optimal thickness and optical properties of natural or artificial solar light. We consider light processes that allow the selective product molecules with the practical option of residence time. The following are available online at https://www.mdpi.com/article/10.3390/pr9122237/s1. Figure S1. Mesh independence test: (a) Reference photoreactor; (b) Optimized photoreactor; (c) Opt-Non-Cone; (d) One-Inlet. Figure S2. Mesh cross-section: (a) Reference photoreactor; (b) Optimized photoreactor; (c) Opt-Non-Cone; (d) One-Inlet. Figure S3. Mesh cut refinement: (a) For the reference photoreactor; (b) For the optimized photoreactor. Table S1. Number of cells for the studied reactors.

**Supplementary Materials:** The following are available online at https://www.mdpi.com/article/10.3390/pr9122237/s1. Figure S1. Mesh independence test: (a) Reference photoreactor; (b) Optimized photoreactor; (c) Opt-Non-Cone; (d) One-Inlet. Figure S2. Mesh cross-section: (a) Reference photoreactor; (b) Optimized photoreactor; (c) Opt-Non-Cone; (d) One-Inlet. Figure S3. Mesh cut refinement: (a) For the reference photoreactor; (b) For the optimized photoreactor. Table S1. Number of cells for the studied reactors.

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