CFD SIMULATION OF TWO-PHASE FLOW IN A VENTURI SCRUBBER: VALIDATION AND COMPARISON OF SECONDARY ATOMIZATION MODELS

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Abstract. In order to study the two-phase flow dynamics inside an industrial Venturi scrubber, the present work compares numerical simulations employing the standard k-ε turbulence model and the Eulerian-Lagrangian method with different droplet breakup models, as implemented in ANSYS CFX 15.0, with experimental data. It is also compared the influence of liquid injection in the droplet dispersion inside the Venturi scrubber, which was achieved by comparing injecting droplets with the same diameter and injecting droplets with a size-distribution (Rosin-Rammler). A better liquid dispersion was achieved using CAB model.

Keywords: computational fluid dynamics (CFD), model validation, secondary atomization, multiphase flow modeling

1. INTRODUCTION

Venturi scrubbers are widely used in industry as gas cleaning devices to control pollution emissions, due to their high collection efficiency, simple structure and implementation, according to Ahmadvand and Talaei (2010). This type of scrubber consists generally of three segments: convergent section, throat and divergent section. In the convergent section, gas flow accelerates due to a reduction in cross-section area, hence reducing static pressure, occurring the opposite in the divergent section, in which gas velocity is reduced due to an increase in cross-section area and pressure regain occurs.

In those equipment, the injection of liquid occurs in convergent or throat sections and production of droplets of liquid are achieved by a jet atomization or spray injection. The high relative velocity between the gas and liquid phases causes the jet breakup, whilst a pressure difference in a spray nozzle causes the liquid dispersion. Liquid droplets produced collect contaminants in the gas flow as they travel through the equipment’s length by three main mechanisms, known as, inertial impaction, interception and diffusion, as explained in Heumann (1997). This contaminant collection efficiency is largely affected by droplet size, relative velocity between phases, liquid atomization, liquid injection method, droplet dispersion in cross-section area and others, as noted by Ali et al. (2012).

Since interaction between liquid droplets and gas phase is a complex problem, the prediction of multiphase dynamic behavior in industrial equipment with computational fluid dynamics (CFD) has been gaining highlights, as noted by Guerra et al. (2012). However, these numerical simulations need to be checked against experimental data in order to verify their feasibility.

In order to study the two-phase flow dynamics inside an industrial Venturi scrubber, the present compares CFD simulations employing standard k-ε turbulence model and Eulerian-Lagrangian method, as implemented in ANSYS CFX 15.0 (Ansys, 2013), with experimental data obtained by Silva (2008) inside a plant pilot size Venturi scrubber and empirical correlations (Hsiang and Faeth, 1992; Wert, 1995). It is also compared the influence of liquid injection in the droplet dispersion inside the Venturi scrubber, which was achieved by comparing injecting droplets with the same diameter and injecting droplets with a size-distribution (Rosin-Rammler), with parameters given by Gonçalves et al. (2003).

2. LITERATURE REVIEW

Venturi scrubbers have been extensively studied in literature in order to model and understand their multiphase flow, according to Silva (2008). One important design parameter is pressure loss through the equipment and so Boll (1973) developed a one-dimensional momentum balance equation in order to predict pressure drop in a Venturi scrubber. The proposed model considered pressure loss due to wall friction and momentum exchange due to acceleration of liquid droplets and gas acceleration. Viswanathan et al. (1985) predicted pressure drop inside a Venturi scrubber as an annular-flow, with momentum transfer similar to approach used by Boll (1973). For more information
about pressure loss equations for Venturi scrubbers, it is recommended the work of Gonçalves et al. (2002), in which important models and correlations available in literature for estimating pressure loss were compared with experimental data.

Another important design parameter is pollutant collection efficiency which is affected by liquid dispersion inside the scrubber. Usually, higher collection efficiency is achieved by uniform liquid dispersion, since augments collision probability between collector droplet and contaminant, according to Gonçalves et al. (2004). Fathikalajahi et al. (1995) developed a three-dimensional droplet dispersion model, which assumed a coefficient of eddy diffusion for droplets transport, in order to obtain liquid dispersion. The model was compared with experimental data obtained by Viswanathan et al. (1984) and a good degree of agreement was achieved for droplet concentration in end of throat section. Gonçalves et al. (2004) developed a mathematical model for droplet dispersion, which added a jet dynamic based on Gonçalves et al. (2003) droplet transport was achieved by assuming fluctuations in droplet concentration and velocity, agreeing well with experimental data. Talai et al. (2008) modified droplet dispersion model proposed by Fathikalajahi et al. (1995) in order to account for droplet size distribution and to estimate collection efficiency. The authors obtained a better agreement for droplet dispersion than the one previously done, as well as for collection efficiency estimation.

None of the previously mentioned models considered the influence of droplets in gas flow and so, CFD simulations becomes a powerful tool for predicting liquid dispersion in scrubbers, as noted by results of Ahmadvand and Talaei (2010). In order to simulate a multiphase flow using a CFD method, two commonly approaches, known as Euler-Euler and Euler-Lagrangian, have been used. The former models the fluids as an interpenetrating continuum medium, solving for each phase, momentum balance equations, whilst the latter solves the continuum phase as an Euler approach and particle trajectory is computed for each droplet by a balance force equation in a Lagrangian approach.

Pak and Chang (2006) solved the gas flow field in a Venturi scrubber using an Eulerian approach with a standard k-ε turbulence model and trajectory of liquid droplets and dust particles were obtained by a force balance done for each representative particle. The authors also took in consideration primary and secondary atomization and achieved good agreement for pressure loss and collection efficiency. Goniva et al. (2009) solved the multiphase flow with a Euler-Lagrangian method using an open-source CFD package (OpenFOAM). Droplet breakup was considered using a Taylor analogy breakup (TAB) model, as well as liquid entrainment in a film flowing in the scrubbers wall. Majid et al. (2013) employed Euler-Lagrange method implemented in ANSYS CFX in order to predict particle removal efficiency in a Venturi scrubber. The authors modeled turbulence with a RNG k-ε model and droplet breakup were achieved using a Cascade atomization and breakup (CAB) model. The removal efficiencies predicted by simulation with CFD were validated with experimental data.

Guerra et al. (2012) employed a Volume of fluid (VOF) with a standard k-ε turbulence model in order to obtain pressure drop and liquid dispersion inside a Venturi scrubber, obtaining visually good agreement with experimental images. Sharifi and Mohebbi (2014) compared the influence of population balance equations to predict pressure drop in a Venturi scrubber. Flow field was solved with an Eulerian-Eulerian method and droplet breakup was considered. As a result, a good agreement for pressure loss was achieved with population balance equations.

3. MODEL DESCRIPTION

3.1 Eulerian-Lagrangian method

For calculating gas flow field a Eulerian approach was used and a set of time-averaged mass and momentum conservation equations was solved, with turbulence modeled with a standard k-ε model. Mass and momentum conservation equations are given, respectively, as Eq. (1) and Eq. (2).

\[
\frac{\partial (\rho u)}{\partial t} + \nabla (\rho u u) = 0 \tag{1}
\]

\[
\frac{\partial (\rho u^2)}{\partial t} + \nabla (\rho u^2 u) = -\nabla P + \rho g + \nabla \tau_c + S_m \tag{2}
\]

With \( \rho_c \) being continuum phase density, \( u_c \) is continuum phase velocity, \( P \) is pressure, \( g \) is gravitational force, \( \tau_c \) is stress tensor and \( S_m \) is a source term.

Particle trajectories were obtained by a force balance for each representative particle, known as parcel, in order to reduce computational time. Each computational parcel represents a number of droplets in real flow and as an assumption all droplets within a parcel have the same velocity, diameter and position. Since ratio between disperse phase and continuum phase density is high, forces such as Basset-history term, added mass and fluid inertia are negligible, according to Lain and Sommerfeld (2007). For force balance as shown in Eq. (3), only drag force and buoyancy were considered affecting particle trajectory.

\[
m_d \frac{d u_d}{d t} = F_D + F_B = \frac{1}{2} C_d \rho_d A_d \left( |U_c - U_d| (U_c - U_d) + m_d g \right) \left( 1 - \frac{\rho_c}{\rho_d} \right) \tag{3}
\]
With \( m_d \) being droplet mass, \( \mathbf{U}_d \) is droplet velocity, \( \mathbf{F}_d \) and \( \mathbf{F}_B \) are, respectively, drag force and buoyancy force, \( C_d \) is drag coefficient for spheres, \( \rho_d \) is droplet phase density, \( \mathbf{U}_c \) is the instantaneous gas velocity and \( A_d \) is droplet cross-section area.

Drag coefficient was obtained by Schiller-Naumann correlation as implemented in ANSYS CFX 15.0 with a correction proposed by Liu et al. (1993) due to droplet deformation. A two-way coupling between phases was considered, in which liquid droplets affect gas flow by a source term in Eq. (2). Source term due to particles was added for each finite volume where particles passed in a transient simulation.

Lagrangian time step for particles were estimated as being smaller than two relevant particle time scales, being the time required for a particle to cross a finite volume and the particle response time scale (Lain et al., 1999; Lain et al., 2002; Lain and Sommerfeld, 2003). Since particle response time depends upon particle diameter and velocity, it was assumed liquid droplets diameter inside the Venturi scrubber ranging from 10 µm to 2000 µm, with estimated relative velocities between phases ranging from 0 to 70 m/s based on Silva (2008). Finite cell dimensions near liquid injection were obtained as having a height of 3 mm (parallel to gas flow) and a length of 7.5 mm (direction of gas flow), thus obtaining the time required for a particle to cross a cell volume. It was obtained a value of 1.07*10^4 s for the smallest particle time scale. Since a value of 25% of the smallest particle time scales were used in the simulations and three integration time steps for each particle were utilized, a value of 5*10^3 s for Euler time step were obtained.

Transient simulations were conducted until 0.3 s (3000 Euler time steps) and through monitoring pressure and velocity for probe points along the Venturi scrubber, it was determined that after 0.15 s, a stable condition was achieved. For convergence criteria a value of 10^-3 for residuals (RMS) was assumed.

### 3.2 Droplet breakup

As shown by Hsiang and Faeth (1992), different modes for droplet breakup in secondary atomization can be classified according to two nondimensional parameters: Weber (\( We \)) and Ohnesorge (\( Oh \)). The former represents the relation between disruptive aerodynamic forces with restorative surface tension force, as shown in Eq. (4), whilst the latter compares the effect of liquid viscosity in hindering droplet deformation (dissipating energy supplied by aerodynamic forces), which reduces droplet fragmentation probability, with restorative surface tension force, as shown in Eq. (5), according to Guildenbecher et al. (2011).

\[
We = \frac{\rho_d U_c^2 - \rho_d U_d^2}{\sigma} \\
Oh = \frac{\mu_d}{(\rho_d D_d \sigma)^{\frac{1}{2}}} 
\]

With \( |\mathbf{U}_c - \mathbf{U}_d| \) representing slip velocity between phases, \( D_d \) is droplet diameter, \( \sigma \) is surface tension and \( \mu_d \) is liquid viscosity.

According to Pilch and Erdman (1987) and Hsiang and Faeth (1992), different modes for droplet breakup, resulting in different droplet child diameters, are achieved depending upon Weber number. However, for values of \( Oh < 0.1 \), a constant value for transition Weber between modes is obtained, as shown in Tab. 1.

**Table 1. Droplet breakup modes for values of \( Oh < 0.1 \), according to classification of Pilch and Erdman (1987) and Hsiang and Faeth (1992)**

<table>
<thead>
<tr>
<th>Breakup mechanism</th>
<th>Weber number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vibrational breakup</td>
<td>( We &lt; \sim 12 )</td>
</tr>
<tr>
<td>Bag breakup</td>
<td>( \sim 12 &lt; We &lt; \sim 35 )</td>
</tr>
<tr>
<td>Multimode breakup</td>
<td>( \sim 35 &lt; We &lt; \sim 80 )</td>
</tr>
<tr>
<td>Sheet-thinning breakup</td>
<td>( \sim 80 &lt; We &lt; \sim 350 )</td>
</tr>
<tr>
<td>Catastrophic breakup</td>
<td>( We &gt; \sim 350 )</td>
</tr>
</tbody>
</table>

In order to predict droplet breakup using CFD techniques, models for droplet breakup were developed. For this work only Taylor analogy breakup (TAB) and Cascade atomization and breakup (CAB) models were considered, being recommended the work of Chryssakis et al. (2011) for other breakup models.

O’Rourke and Amsden (1987) developed a droplet breakup model, in which droplet oscillation and deformation was modeled as a forced, damped, harmonic oscillator, being an analogy to a mass-spring system, as shown in Eq. (6)

\[
m_d \frac{d^2x}{dt^2} = F_a - kx - \eta \frac{dx}{dt} 
\]
With \( x \) being displacement from equilibrium position (droplet deformation), \( F_a \) is external force applied in the system (aerodynamic force), \( k \) is spring stiffness (related to surface tension) and \( \eta \) is damping coefficient (related to liquid viscosity).

Rewriting Eq. (6) in order a dimensionless deformation equation and considering the analogy explained before, one can write Eq. (7), as implemented in ANSYS CFX (ANSYS, 2013) and explained in Marek (2013).

\[
\frac{d^2 \hat{x}}{dt^2} = \frac{c_{f} \rho v_{c} - u_d^2}{c_{f} \rho \hat{x}} - \frac{c_{b} \sigma}{\rho_d \hat{x}} \frac{d}{dt} - \frac{c_{k} \sigma}{\rho_d \hat{x}^3} \frac{d^2}{dt^2} \quad (7)
\]

With \( \hat{x} \) being dimensionless deformation, \( r_d \) is droplet radius and \( C_f, C_b, C_k \) and \( C_\eta \) are coefficients with values, respectively, 0.33, 0.5, 8 and 5.

Droplet breakup occur only if \( \hat{x} \) exceeds unity and child droplet radius is obtained according to Eq. (7) based on energy conservation balance.

\[
\frac{r_d}{r_{\text{child}}} = 1 + 0.4K + \frac{\rho_{r_d} \hat{x} \frac{d}{dt} (6K - 5)}{\sigma} \quad (8)
\]

With \( r_{\text{child}} \) being child droplet radius after breakup and \( K \) have a value of 3.33.

Tanner (2004) developed a CAB model using the same criteria for droplet deformation given by O’Rourke and Amsden (1987), however, 3 breakup modes were used for determining child droplet diameter, as shown in Eq. (9) and Eq. (10).

\[
\frac{r_d}{r_{\text{child}}} = e^{-K_{bu} t_{bu}} \quad (9)
\]

\[
K_{bu} = \begin{cases} 
  k_1 \omega & 5 < We < 80 \\
  k_2 \omega We^{0.5} & 80 < We < 350 \\
  k_3 \omega We^{0.75} & We > 350
\end{cases} \quad (10)
\]

With \( t_{bu} \) being the time necessary for \( \hat{x} \) to exceed unity, \( \omega \) is droplet oscillation frequency and \( k_1, k_2 \) and \( k_3 \) are constants given in ANSYS CFX (ANSYS, 2013).

### 4. RESULTS AND DISCUSSION

Pressure drop and Sauter mean diameter obtained by Silva (2008) throughout a large-scale Venturi scrubber were compared with numerical simulations carried out in ANSYS CFX 15.0. A computational mesh was created around a geometry similar to the one used experimentally by Silva (2008) and is outlined in Fig. 1, with dimensions given in Tab. 2.

![Figure 1. Schematic of a Venturi scrubber used by Silva (2008) with four liquid displacement positions](image)

Table 2. Dimensions used for mesh creation in CFD software

<table>
<thead>
<tr>
<th>Definition</th>
<th>Dimensions (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diameter (D)</td>
<td>250</td>
</tr>
<tr>
<td>Throat diameter (Dt)</td>
<td>122.5</td>
</tr>
<tr>
<td>Inlet length (l_{in})</td>
<td>500</td>
</tr>
<tr>
<td>Convergence section length (l_{c})</td>
<td>230</td>
</tr>
<tr>
<td>Throat section length (l_{t})</td>
<td>300</td>
</tr>
</tbody>
</table>
Divergent section length ($l_d$) & 740  \\
Outlet length ($l_{out}$) & 1500  \\

Computational mesh was achieved with a grid generation tool in the CFD software (Meshing), by first creating a meshed region in the inlet face and afterwards, using this mesh as a base for creation of other element faces throughout equipment, with a method called Sweep. The generated mesh consisted of 1,117 thousand hexahedral elements and element cells are shown in Fig. 2 for different plane cuts. In bottom left a transversal cut in the beginning from convergent section shows cell faces, while in bottom right, cell faces are show in the beginning from throat section.

![Computational mesh for a circular Venturi scrubber](image)

Figure 2. Computational mesh for a circular Venturi scrubber, in which is shown for different plane cuts, convergent and throat sections (upper part); transversal cut in convergent section (bottom left); and transversal cut in throat section (bottom right).

According to Silva (2008), water was injected in convergent section through four different nozzles, as shown in Fig. 1, with liquid flow rates varying from 0.0134 to 0.0583 kg/s and gas flow rates from 0.483 to 0.987 kg/s. In order to compare with experimental data, simulations were conducted for a set of three gas flow rates (0.483, 0.861 and 0.987 kg/s) with a value of 0.0134 kg/s for water flow rate.

In order to compare droplet breakup models, a size distribution for liquid injection was not initially adopted and therefore, computational parcels were injected near the wall with diameter of 1 mm. For gas flow rate of 0.861 kg/s, computational particles positions were obtained with their current diameter in last time step (0.5 s) for TAB and CAB models, as shown in Fig. 3.
Comparing the two droplet breakup models, one obtains a similar droplet diameter reduction for all particles with TAB model, whereas with CAB model, a variation in diameter reduction was achieved, resulting in droplets with different diameters, along with a better liquid dispersion transversal to gas flow after droplet breakup, if compared with TAB model. According to Fig. 3, one perceives that particles followed the same trajectory paths, with breakup occurring in the beginning of throat section for TAB model, whilst breakup simulated by CAB model occurred along throat section. For a higher gas flow rate, the same patterns were observed for the two models, not being reported in this work.

In order to compare the numerical simulation results with the experimental data obtained by Silva (2008) and literature correlations, as shown in Tab. 3, the Sauter mean diameter ($D_{32}$) was analyzed. Venturi scrubber length was divided in a series of control volumes with width of 2 cm and computational particles between simulation times of 0.15 s to 0.3 s were analyzed with Eq. (11). As a result, Sauter mean diameter through the equipment was obtained, as shown in Fig. 4, in which coordinate axis is given as distance to liquid injection.

$$D_{32} = \frac{\sum_{i=1}^{m} n_{d_i} d_{d_i}^3}{\sum_{i=1}^{m} n_{d_i} d_{d_i}^2}$$  \hspace{1cm} (11)

With $D_{32}$ being Sauter mean diameter, $n_{d_i}$ is particle number rate, in which each computational particle relates to a real number of droplets, $d_{d_i}$ is computational particle diameter and $m$ represents total number of particles inside a determined control volume.
For the lowest gas flow rate, no breakup of the droplets for both models was achieved, which resulted in a bigger difference between the experimental data and the simulated one, as seen by Fig. 4 and Tab. 3. However for higher gas flow rates, the results obtained are in accordance with experimental data, with a Sauter mean diameter of 515 µm obtained with numerical simulation, while a value of 211.5 µm for the experimental data (Silva, 2008) and a value of 193 µm with correlation of Wert (1995) was achieved.

For the present reason, along with a better droplet dispersion achieved, CAB model in conjunction with a droplet size distribution injection was studied.

According to Alonso et al. (2001) droplet size distribution through Venturi scrubbers can be described by Rosin-Rammler equation, as given in Eq. (12), with parameter $n_1$ varying from 1.7 to 2.2. A value of 2.15, as in Gonçalves et al. (2004), along with a value of 1 mm for Sauter mean diameter.

$$1 - \varphi = \exp\left(-\left(\frac{D_d}{X_1}\right)^{n_1}\right)$$

With $\varphi$ being cumulative volume fraction of droplets with diameter lower than $D_d$ and $X_1$ is a Sauter mean diameter corrected by a gamma function.
Figure 5 compares droplet Sauter mean diameter through Venturi scrubber obtained with a Rosin-Rammler droplet size distribution using CAB model with a 1 mm diameter particle injection, as previously done. One perceives that using a size distribution, droplet breakup was achieved in a shorter distance to liquid injection if compared to a monodisperse injection, since a reduction in Sauter mean diameter occurred a shorter distance before throat section. For a gas flow rate of 0.483 kg/s droplet breakup with CAB model was achieved, resulting in a lower value for Sauter mean diameter for a droplet size distribution injection, since it is present droplets with bigger diameters, possessing bigger Weber numbers and having higher probabilities of reducing their diameter. For a gas flow rate of 0.861 kg/s, a lower value for Sauter mean diameter was obtained if compared to the previously simulation, whereas for highest flow rate, a similar value was achieved. It was noted that using a droplet size distribution to model liquid injection resulted in droplet breakup for all three cases.

In order to analyze droplet size evolution through the Venturi scrubber, droplet size distribution was evaluated in four control volumes with 0.02 m width located from liquid injection: 0 m (liquid injection); 0.23 m (beginning from throat section); 0.53 m (beginning from divergent section); 1.27 m (end of divergent section). Droplet size distribution was obtained by separating each particle inside a control volume based in their diameter and adding their mass contribution in their size interval. Figure 6 shows droplet size distribution for 3 gas flow rates distant 0.23 m from liquid injection.

According to Fig. 6 one can obtain that droplet breakup was achieved for the three gas flow rates, since droplet size distribution was dislocated to smaller droplet sizes. It was seen that increasing mass gas flow rate, smaller droplet size distributions were achieved, since higher values for droplet Weber number were achieved. However, for the other two control volumes, a quasi-uniform droplet size distribution was obtained for gas flow rates of 0.861 kg/s and 0.987 kg/s. Due to the quasi-uniform droplet size distribution, rather than using a constant value of 1 mm for droplet diameter in Rosin-Rammler function for droplet injection, droplet diameter for Rosin-Rammler function was chosen as being the Sauter mean diameter after breakup have occurred for different gas flow rates.

Figure 7 shows droplet size distribution along the Venturi scrubber length for a gas flow rate of 0.861 kg/s. One perceives that droplet breakup still occurred, even though droplets with smaller diameter were present, resulting in lower Weber numbers. Using the mentioned criteria for determining droplet diameter in liquid injection resulted in a droplet size distribution with a broader size distribution, if compared to the previous simulation results. For the highest gas flow rate (0.987 kg/s) the same pattern were observed.
5. CONCLUSIONS

The flow field inside a wet scrubber was simulated with a RANS model using Computational Fluid Dynamics (CFD) in which the multiphase flow was solved by an Euler-Lagrange approach. TAB and CAB models were evaluated for droplet breakup inside a Venturi scrubber. Broader size distributions were achieved using a Rosin-Rammler function with droplet diameter obtained after droplet breakup. With the simulated data it can be seen that droplet breakup due to secondary atomization plays an important role in varying droplet size distribution along the Venturi scrubber.

6. ACKNOWLEDGEMENTS

CAPES is gratefully acknowledged.

7. REFERENCES

8. RESPONSIBILITY NOTICE

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