CFD simulation of dense gas dispersion in neutral atmospheric boundary layer with OpenFOAM

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Abstract In this study, MOST (Monin-Obukhov Similarity Theory) is used to specify the profiles of velocity, turbulent kinetic energy ($k$) and eddy dissipation rate ($\epsilon$) in ABL (Atmospheric Boundary Layer) flow. The OpenFOAM standard solver $buoyantSimpleFoam$ is modified to simulate neutrally stratified ABL. The solver is able to obtain equilibrium ABL. For gas dispersion simulation, $buoyantNonReactingFoam$ is developed to take into accounts fluid properties change due to temperature, buoyancy effect and variable turbulent Schmidt number. The solver is validated for dense gas dispersion in wind tunnel test and field test of LNG (Liquefied Natural Gas) vapour dispersion in neutrally stratified ABL.

Keywords LNG dispersion · CFD · ABL · OpenFOAM
1 Introduction

Many human activities are affected by the atmospheric boundary layer (ABL). This is also where most air pollution phenomena occur. Understanding of the processes taking place in the ABL has attracted various research studies. Some typical applications of ABL related research topics are wind engineering, urban flows, weather forecast, air pollution and risk assessment of hazardous material spills in industrial sites.

One hazardous dense gas is liquefied natural gas (LNG), which is an effective solution for long-distance natural gas transfer. LNG has become the preferred option for international trading of natural gas. However, LNG storage, handling, transportation are exposed to serious risks for humans, equipment and the environment due to thermal hazards associated with combustion events such as pool fire, vapour cloud fire, explosion or rapid phase transition. Safety assessment and hazard mitigation methods should be applied to lower the possibilities of catastrophic disaster relating to the LNG industry. The scope of this study is constrained to the discussion of dense gas dispersion when released into the ABL.

Computational Fluid Dynamics (CFD) is increasingly being used in simulation of ABL flows. Open source CFD tool is a more powerful research tool in comparison to proprietary software because of its flexibility to incorporate new implementation of fields calculation and post-processing. OpenFOAM is an open source CFD software package that attracts users from both industry and academia. Using a general CFD code such as OpenFOAM for simulating ABL flow and gas dispersion also encourages research sharing and reusing code in this specific field where in-house code is usually adopted.
An important task before modelling gas dispersion in the ABL is obtaining the correct ABL flow prior to the release of gas source. One approach to achieve this is using equilibrium ABL, i.e. zero stream-wise gradients of all variables, as a steady state ABL flow. For neutral ABL, Richards and Hoxey (1993) proposed appropriate boundary conditions of mean wind speed and turbulence quantities for the standard $k - \epsilon$ model based on Monin-Obukhov similarity theory (MOST). These profiles were derived assuming constant shear stress with height and were used to model ABL as horizontally homogeneous turbulent surface layer (HHTSL). However, HHTSL was hard to achieve mostly due to the ground boundary conditions (Yang et al., 2009), which manifested in a decay of velocity profile due to a spike in the turbulent kinetic energy close to the ground. However, consistency between wall boundary conditions, turbulence model with associated constants and numerical schemes was shown to achieve HHTSL (Jonathon and Christian, 2012; Parente et al., 2011; Yan et al., 2016). These authors adopted proprietary CFD software for their simulation. Applying these implementations in open-source CFD code also require extensive modifications of the source code to successfully simulate equilibrium ABL. OpenFOAM was previously used for atmospheric buoyant (Flores et al., 2013) and dense gas dispersions (Mack and Spruijt, 2013; Fiates et al., 2016; Fiates and Vianna, 2016). However, the validation of these solvers in simulation of equilibrium ABL was not reported. Therefore, the atmospheric turbulence might not be correctly solved throughout the computational domain.

In this study, MOST is used to model the profiles of velocity, turbulent kinetic energy ($k$) and eddy dissipation rate ($\epsilon$) of ABL according to an approach proposed by Richards and Hoxey (1993). These profiles are used as the boundary conditions at the inlet of ABL flow simulation. OpenFOAM application **buoyantSimpleFoam**
is modified to simulate neutrally stratified ABL turbulence. For gas dispersion simulation, `buoyantNonReactingPimpleFoam` is developed to take into account the buoyancy effect and variable turbulent Schmidt number. The solver is validated for dense gas dispersion cases from wind tunnel and field tests of LNG vapour dispersion in neutrally stratified ABL.

2 Methodology

2.1 Models

The $k-\epsilon$ model is used for turbulence modelling. It is based on expression of turbulent dynamic viscosity $\mu_t$ by Jones and Launder (1972):

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

Two additional transport equations for turbulence kinetic energy $k$ and turbulence dissipation rate $\epsilon$ are required. To include the effect of buoyancy, the transport equations for $k$ and $\epsilon$ are:

$$\frac{D}{Dt}(\rho k) = \frac{\partial}{\partial x_i} \left[ \left( \mu + \mu_t \right) \frac{\partial k}{\partial x_i} \right] + G_k + G_b - \rho \epsilon$$ \hspace{1cm} (2)

$$\frac{D}{Dt}(\rho \epsilon) = \frac{\partial}{\partial x_i} \left[ \left( \mu + \mu_t \right) \frac{\partial \epsilon}{\partial x_i} \right] + C_{1\epsilon} \frac{\epsilon}{k} G_k + C_{1\epsilon} \frac{\sigma_k}{k} G_b - C_{2\epsilon} \rho \frac{\epsilon^2}{k}$$ \hspace{1cm} (3)

where $\rho$ is the fluid density; $C_\mu = 0.09$, $\sigma_k = 1$, $\sigma_\epsilon = 1.3$, $C_{1\epsilon} = 1.44$ and $C_{2\epsilon} = 1.92$ are model constants as proposed in original paper (Launder and Spalding, 1974). The value of $C_{3\epsilon}$ is calculated using:
\[ C_{3k} = \tanh \left( \frac{v}{u} \right) \quad (4) \]

where \( v, u \) are vertical and horizontal velocity accordingly.

\( G_k \) is production of turbulence kinetic energy due to the mean velocity gradients. \( G_b \) is the buoyancy source term:

\[ G_b = -\frac{\mu_t}{\rho Pr_t} \left( g \cdot \nabla \rho \right) = -C_g \frac{\mu_t}{\rho} \left( g \cdot \nabla \rho \right) \quad (5) \]

where \( C_g = 1/Pr_t \) is used as a model constant to take into account the user-defined value of turbulent Prandtl number \( Pr_t \). \( g \) is gravitational vector.

Energy, heat and transport properties are determined by a set of thermophysical models (Greenshields, 2017) in OpenFOAM. This set defines mixture type, transport and thermodynamic properties models, choice of energy equation variable and equation of states.

The fluid in a simulation is defined as a mixture of fixed compositions. Enthalpy is chosen as energy equation variable. Transport and thermodynamic properties are determined using models based on the density \( \rho \), which are calculated from pressure and temperature fields. Polynomial functions of order \( N \) are used to relate transport property \( \mu \), the specific heat \( c_p \) and density \( \rho \) with temperature field \( T \):

\[ \mu = \sum_{i=1}^{N-1} a_{\mu i} T^i \]
\[ c_p = \sum_{i=1}^{N-1} a_{c_p i} T^i \]
\[ \rho = \sum_{i=1}^{N-1} a_{\rho i} T^i \quad (6) \]

where \( a_{\mu i}, a_{c_p i} \) and \( a_{\rho i} \) are the polynomials coefficients.
2.2 Boundary conditions

2.2.1 ABL air inlet

Most has been validated for the surface layer of ABL by many empirical studies (Foken, 2006). It assumes horizontally homogeneous and quasi-stationary flow field, i.e., profiles of flow variables are only varying in the vertical direction and their vertical fluxes are assumed constant. The inlet boundary conditions proposed by Richards and Hoxey (1993) based on MOST are widely used in CFD study of atmospheric flow. The velocity, turbulent production rate $k$ and dissipation rate $\epsilon$ profiles in vertical direction $z$ are written as:

$$u(z) = \frac{u^*}{\kappa} \ln \frac{z + z_0}{z_0}$$

$$k = \frac{u^2_z}{\sqrt{C_\mu}}$$

$$\epsilon(z) = \frac{u^3_\kappa}{\kappa(z + z_0)}$$

where $z_0$ is aerodynamic roughness length, $u^*$ is friction velocity, $C_\mu$ is $k - \epsilon$ model constant.

These profiles are implemented in OpenFOAM as `atmBoundaryLayer` class and its subclasses. Required parameters are flow and vertical direction, reference velocity, reference height and aerodynamic roughness length. The friction velocity is calculated as:

$$u^* = \frac{\kappa \cdot u_{ref}}{\ln((z_{ref} + z_0)/z_0)}$$

Parente et al. (2011) presented an elaborate procedure to ensure the consistency for arbitrary inlet profile of turbulent kinetic energy $k$. Instead of altering model
constants as Yang et al. (2009), the effect of non-constant \( k \) on momentum and \( \epsilon \) equation can be characterised by deriving an equation for \( C_\mu \):

\[
C_\mu(z) = \frac{u_\tau^4}{k(z)^2}
\]  

(9)

Source terms are added to \( k \) and \( \epsilon \) transport equations to ensure equilibrium condition:

\[
S_k = \frac{\rho u_\tau \kappa}{\sigma_k} \frac{\partial}{\partial z} \left( (z + z_0) \frac{\partial k}{\partial z} \right)
\]

\[
S_\epsilon = \frac{\rho u_\tau^4}{(z + z_0)^2} \left( \frac{(C_{c2} - C_{c1}) \sqrt{C_\mu}}{\kappa^2} - \frac{1}{\sigma_\epsilon} \right)
\]

(10)

Richards and Norris (2011) revisited the problem of modelling the HHTSL by deriving the inlet profiles directly from the conservation and equilibrium equations. This allows various inlet profiles to be specified by varying the turbulence model constants. For standard \( k - \epsilon \) models, the inlet profiles of velocity and turbulence properties are identical to Equation (7). However they suggested to change the von-Karman constant \( \kappa \) according to model constants as:

\[
\kappa = \sqrt{(C_{c2} - C_{c1}) \sigma_\epsilon \sqrt{C_\mu}}
\]

(11)

Using the standard \( k - \epsilon \) model constants, we obtain \( \kappa_{k-\epsilon} = 0.433 \).

2.2.2 Wall boundary conditions

In CFD, the below approximation is used to calculate wall shear stress:

\[
\tau_w = \nu_\tau \frac{\partial u}{\partial n} \bigg|_w \approx \nu_\tau \frac{(u_P - u_w)}{y_P}
\]

(12)
where $y_P$ is distance to wall of the wall adjacent cell. Subscript $w$ and $P$ denote field value evaluated at wall and wall adjacent point respectively.

However, this approximation is inaccurate when wall velocity gradient is significantly larger than velocity difference between the adjacent cell and the wall. This is the case for most ABL flows. Turbulent kinematic viscosity $\nu_t$ wall function is used to calculate the wall shear stress $\tau_w$ from the wall velocity difference. To take into account the aerodynamic roughness length $z_0$, the calculation of turbulent kinematic viscosity at wall adjacent cell is:

$$\nu_t = \frac{k u_* y_P}{\ln \left( \frac{y_P + z_0}{z_0} \right)}$$

where friction velocity $u_*$ can be calculated from a simple relation derived by Launder and Spalding (1974), assuming that generation and dissipation of energy are in balance:

$$u_* = C_1^{1/4} k^{1/2}$$

$\epsilon$ wall function is used to calculate value of $\epsilon$ at wall adjacent cell $\epsilon_P$ as:

$$\epsilon_P = \frac{C_0^{0.75} k^{1.5}}{k y_P + z_0}$$

The wall is usually defined as non-slip condition where velocity is zero. However, to account for the effect of aerodynamic roughness length, a new boundary condition for velocity is implemented in OpenFOAM as:

$$u_P = \frac{u_*}{k} \ln \left( \frac{y_P + z_0}{z_0} \right)$$
2.2.3 Top, side and outlet boundaries

At the outlet boundary, the flow is assumed fully developed and unidirectional. All flow variables are supposed to be constant at this boundary.

The top and side of the computational domain are external boundaries representing the far fields of flow. If a constant pressure is applied in these boundaries, this may alter the inlet wind profile in case the prescribed pressure is not matched with the boundary velocity (Luketa-Hanlin et al., 2007). The zero gradient boundary condition, which set normal velocity to zero and all others variables are set equal to the inner values, or symmetry condition can be used at the top and side boundaries to reserve the wind profile and eliminate the effect of changing the inlet profiles.

Hargreaves and Wright (2007) showed that zero gradient velocity at the top boundary resulted in a decay of velocity downstream, due to the extraction of energy at the wall with respect to the wall shear stress. A driving shear stress, zero flux of turbulent kinetic energy and a flux of dissipation rate $\epsilon$ are imposed at the upper boundary:

\[
\begin{align*}
\frac{du}{dz} &= \frac{u_*}{Kz} \\
\frac{\mu_t}{\sigma_\epsilon} \frac{d\epsilon}{dz} &= -\frac{\rho u_*^4}{\sigma_\epsilon z}
\end{align*}
\]

2.3 Numerical tool and data sets

OpenFOAM is an open source CFD software package based on finite volume method, co-located variables and unstructured polyhedral meshes. In this study, buoyantSimpleFoam is used to simulate ABL turbulence. The application buoyantNonReactingFoam
is developed based on rhoReactingBuoyantFoam solver, previously used for dense
gas dispersion by Fiates et al. (2016), to simulate atmospheric turbulence un-
der neutral stability for dispersion of dense gas continuous source in flat terrain.

buoyantNonReactingFoam uses polynomial thermophysical models to account for
the change of fluid properties due to temperature. The solver takes into account
the buoyancy effect and the variable turbulent Schmidt number. Algorithms used
in these two solvers are presented in Algorithm 1 and 2.

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**Algorithm 1 buoyantSimpleFoam solver algorithm**

1: Initializing variables such as: time variables, mesh, solution control, fields and continuity errors

2: while $t < t_{end}$ do

3:   Solving momentum equation

4:   Solving energy equation for enthalpy and correcting thermal properties

5:   Solving pressure correction equation for $p_{rgb}$ and calculating pressure field

6:   Correcting turbulent properties

7:   Writing fields

8: end while
Algorithm 2 buoyantNonReactingFoam solver algorithm

1: Initializing variables such as: time variables, mesh, solution control, fields and continuity errors

2: \textbf{while } $t < t_{end}$ \textbf{do}

3: \hspace{1em} $t = t + \Delta t$

4: \hspace{1em} Solving continuity equation for density

5: \hspace{1em} \textbf{while} PIMPLE outer correctors \textbf{do}

6: \hspace{2em} Solving momentum equation

7: \hspace{2em} Solving species transport equation

8: \hspace{2em} Solving energy equation for enthalpy and correcting thermal properties

9: \hspace{2em} \textbf{while} PIMPLE inner correctors \textbf{do}

10: \hspace{3em} Solving pressure correction equation for $p_{rgh}$ and calculating pressure field

11: \hspace{3em} Solving continuity equation for density

12: \hspace{3em} Calculating time step continuity errors

13: \hspace{2em} \textbf{end while}

14: \hspace{2em} Correcting turbulent properties

15: \hspace{2em} \textbf{end while}

16: Writing fields

17: \textbf{end while}

Boundary conditions used for ABL flows are developed as new libraries in OpenFOAM. These include velocity inlet, turbulent kinetic energy, dissipation rate inlet and wall boundary conditions.

A set of full scale field tests and experimental wind tunnel tests for LNG dispersion model validation was reported in (Ivings et al., 2013). Most data of these tests were available in REDIPHEM database (Nielsen and Ott, 1996). The data contains physical comparison parameters of each test. These are maximum arc-wise concentration, i.e. the maximum concentration across an arc at the specified
distance from the source and point-wise concentration, i.e. the concentration at specific sensor locations. Two wind tunnel data DA0120 and DAT223 are used to validate OpenFOAM solver in prediction of dense gas dispersion over a flat, unobstructed terrain in simulated neutral ABL. In these tests, continuous source of SF$_6$ gas was released in flat terrain without obstructions. For field test, we select Burro9, which is continuous LNG spills under neutral ABL.

3 Neutral ABL simulation

3.1 Domain and mesh

A 2D domain of 5000 m $\times$ 500 m with the resolution of 500 $\times$ 50 cells is used for the simulation of neutral ABL over flat terrain. The mesh is uniform in stream-wise direction and stretched in vertical direction with the expansion ratio of 1.075.

3.2 Numerical setting

The boundary conditions of the cases are represented in Table 1.

| Table 1 | Boundary conditions for neutral ABL simulation |
|---------|-----------------------------------------------|
| ABL inlet | profiles of $k$, $u$, $\epsilon$ | Eq. (7) |
| ABL outlet | zeroGradient for all variables | |
| | fixedValue for static pressure | |
| ABL side | zeroGradient for all variables | |
| ABL top | zeroGradient for all variables | |
| | fixedFlux/zeroGradient for $u$ and $\epsilon$ | Eq. (17) |
| ABL bottom | noSlip for $u$ | |
ABL parameters used to define inlet variable profiles are listed in Table 2 according to the reference case of Hargreaves and Wright (2007):

| $u_*$ (m/s) | $z_0$ (m) | $u_{ref}$ (m/s) | $z_{ref}$ (m) |
|------------|-----------|----------------|--------------|
| 0.625      | 0.01      | 10             | 6            |

Steady state simulation is employed using buoyantNonReactingSimpleFoam described in previous section. OpenFOAM discretisation schemes, velocity-pressure coupling algorithm as well as linear solvers are listed below:

- Time schemes: steadyState
- Gradient schemes: Gauss linear
- Divergence schemes: Gauss limitedLinear 1
- Surface normal gradient schemes: corrected
- Laplacian schemes: Gauss linear corrected
- Interpolation schemes: linear
- Solving algorithm: SIMPLE
- Linear solver for p: GAMG with DICGaussSeidel preconditioner
- Linear solver for U, h, k, epsilon: PBiCGStab with DILU preconditioner

Residual control is set at three order of magnitude for pressure and four order of magnitude for other variables such as $U$, $k$, $\epsilon$ and $h$. Modification of $k - \epsilon$ (Equation (11)) are used to simulate neutral ABL and comparing with standard models. These three cases are summarised in Table 3.

Different levels of inlet kinetic energy are obtained by altering $C_\mu$ according to Equation (9). The source term by Pontiggia et al. (2009) is implemented us-
Table 3 Turbulence models setting for neutral ABL simulation

| Turbulence model       | standard $k - \epsilon$ | modified $k - \epsilon$ | Eq. (10) |
|------------------------|---------------------------|---------------------------|----------|
| Wall functions         | nutkWallFunction for $\nu_t$ | epsilonWallFunction for $\epsilon$ | kqRWallFunction for $k$ |

Modification of $k - \epsilon$ models achieve the matched results as shown in Figure 1. Including source terms as in Equation (10) is sufficient to compensate terms deflection from calculation of von-Karman constant $\kappa_{k-\epsilon} = 4.3$ from model constants (Equation (11)) and $\kappa = 4.1$ used in Monin-Obukhov theory.

Results from modelling different turbulence kinetic energy by varying $C_\mu$ are presented in Figure 2. The profiles of velocity and dissipation rate are perfectly matched with the Monin-Obukhov profiles. In the $C_\mu = 0.017$ simulation, the value of $k$ near ground is smaller than the theoretical value, however, the kinetic energy level is matched with the theory at greater height. The smaller value of $k$ at the wall adjacent cell is due to the wall function, where wall treatment used with the default $C_\mu = 0.09$ is implemented. However, the overall results are acceptable for verifying the proposed model in simulating different levels of kinetic energy.
Fig. 1 Comparing velocity, turbulent kinetic energy and turbulent dissipation rate profiles at the outlet boundary from simulation of neutral ABL using standard $k-\epsilon$ (kEps), modified $k-\epsilon$ (kEpsMod) turbulence model and MOST inlet profiles (MOST).

Fig. 2 Comparing velocity, turbulent kinetic energy and turbulent dissipation rate profiles from simulations of different kinetic energy levels by varying $C_\mu = 0.09$ (KEpsCmu09) $C_\mu = 0.017$ (KEpsCmu017) and MOST inlet profiles (MOST).
4 Dense gas dispersion in wind tunnel tests

4.1 Numerical setting

The effect of the turbulent Schmidt number $S_{Sc}$ is investigated in dense gas dispersion. Three test cases are summarised in Table 4. The effect of turbulent models is examined by applying the modified $k - \epsilon$ which is already validated in simulating ABL over flat terrain in Section 3.

| Case  | Case 2 | Case 3 |
|-------|-------|-------|
| $S_{Sc}$ | 1     | 0.7   | 0.3   |
| Label (Fig. 4, 10) | FOAM_ORIG | FOAM_Sc07 | FOAM_Sc03 |

Firstly, the steady simulation using `bouyantSimpleFoam` is performed to establish the steady ABL flow prior to the dense gas release. The solver which includes buoyancy effects `bouyantSimpleFoam` is used to account for density stratification in dense gas flow. The atmospheric inlet profiles are specified by MOST with parameters in Table 5. Standard $k - \epsilon$ with modifications is used to study the ability to simulate the ABL with each model. Secondly, the transient simulation is performed using steady simulation solutions as initial fields. A modified version of `rhoReactingBouyantFoam` is studied to model multi-species flow where mixture considered are air and dense gas SF$_6$. The wind tunnel tests were conducted in isothermal condition, therefore constant thermal and transport properties are used for both gases.
Table 5  Hamburg flat, unobstructed test case parameters (Nielsen and Ott, 1996)

| Substance | DA0120 | DAT223 |
|-----------|--------|--------|
| Density   | kg/m³  | 6.27   | 6.27   |
| z₀        | m      | 0.0001 | 0.0001 |
| Wind speed| m/s    | 0.54   | 0.74   |
| Reference height | m   | 0.00718 | 0.01367 |
| Ambient temperature | °C | 20    | 20     |
| Source diameter | m | 0.07   | 0.07   |
| Spill rate  | kg/s   | 0.0001743 | 0.000872 |

In simulations of DA0120 and DAT223 tests, the discretisation schemes and linear solver setting are identical to the simulation of neutral ABL (Section 3).

4.2 Results and discussion of gas dispersion in wind tunnel tests

4.2.1 Peak concentration prediction

The steady state plumes at ground level of DAT0120 and DAT223 tests are plotted in Figure 3. Under higher release volume flow rate and higher wind speed, DAT223 plume is wider and is spreading further downstream than the DAT0120 plume.

The predicted and measured peak gas concentration are compared at several distances from the spill in Figure 4. Turbulent Schmidt number $S_{Ct}$ has significant effect in predicting dense gas dispersion. The original rhoReactingBouyantFoam code, with assumption of species diffusivity equals to viscosity, is shown to over-predict concentration with a factor of three. The modified code takes into account the variable species diffusivity $S_{Ct}$ by reading this parameter from user input. The
value of $Sc_t = 0.3$ is shown to yield a perfect match with the experimental data. However, there is a slightly acceptable over-predicted species concentration at a point near the source release.

Results from the DAT223 simulation are presented in Figure 5. Satisfactory over predicted peak concentration is similar to DA0120 case.

4.2.2 Point-wise concentration

Figure 6 presents gas concentration at the downwind distance $X = 1.84$ of the DA0120 test. The simulation can reproduce the averaged gas concentration. The first incidence time of gas concentration is earlier than observed in experiments. However, the time for reaching averaged maximum concentration is well predicted.

4.2.3 Statistical model evaluation

Statistical Performance Measures (SPMs) are means to compare prediction parameters and the measured ones for model evaluation. The SPM chosen should reflect
Fig. 4  Peak concentration for DA0120 test

Fig. 5  Peak concentration for DAT223 test
the bias of these predictions. In the context of LNG vapour dispersion model evaluation, Ivings et al. (2013) proposed five SPMs including mean relative bias (MRB), mean relative square error (MRSE), the fraction of predictions within the factor of two of measurements (FAC2), geometric mean bias (MG) and geometric variance (VG). Definition and acceptability criteria for each SPMs are presented in tabular form as Table 6 where $C_m$, $C_p$ are the measured and simulated concentration, respectively, and $\bar{A}$ denotes the mean operation of variable $A$.

Statistical performance of OpenFOAM results are compared with the specialised commercial code for gas dispersion FLACS in Table 6. FLACS results are extracted from (Hansen et al., 2010). The performance of current OpenFOAM code is considerably better than FLACS. In fact, FLACS is based on the porosity distributed resistance (PDR) approach. Therefore, this modelling of the boundary layer close to solid surfaces might contribute to the outperformance of the Open-

Fig. 6 Concentration at $X = 1.84$ of DA0120 test
FOAM model in the comparison. In conclusion, even though larger tests were validated in FLACS, the proposed model in OpenFOAM is a promising tool for further investigation of atmospheric dense gas dispersion.

| SPM  | MRB | RMSE | FAC2 | MG   | VG   |
|------|-----|------|------|------|------|
| Definition | \( \frac{C_m - C_p}{0.5(C_m - C_p)} \) | \( \frac{(C_m - C_p)^2}{0.25(C_m + C_p)^2} \) | \( \frac{C_m}{C_p} \) | \( \exp \left( \ln \frac{C_m}{C_p} \right) \) | \( \exp \left( \ln \frac{C_m}{C_p} \right)^2 \) |
| Acceptable range | [-0.4,0.4] | < 2.3 | [0.5, 2] | [0.67, 1.5] | < 3.3 |
| Perfect value | 0 | 0 | 1 | 1 | 1 |
| FLACS (Hansen et al., 2010) | 0.25 | 0.29 | 0.89 | 1.34 | 1.61 |
| FOAM | -0.06 | 0.02 | 1.07 | 1.06 | 1.02 |

5 LNG vapour dispersion in field tests

5.1 Numerical setting

The steady simulation uses the atmospheric inlet specified by MOST. Standard \( k-\epsilon \) with modifications is used to study the ability to simulate the ABL with each model. All required meteorological parameters are tabulated in Table 7, where \( u_{ref} \) and \( T_{ref} \) are air velocity and temperature at the height of 2 m respectively.

The transient simulation is divided into two steps. The first step is during the spill duration, i.e. from the time of zero to when the spill ends. The second step is after the spill stops to the end time of simulation. The gas inlet is treated as a ground boundary in this later step.

The gas inlet condition is usually obtained from separate source term modelling. There is not much information about the vaporisation of LNG from the ex-
perimental data. Therefore, uncertainty arises at the setting of this condition. Mass flux of LNG or the LNG vaporization rate is used to derive source term of LNG spilling. Luketa-Hanlin et al. (2007) reviewed a number of experiments conducted to estimate the LNG vaporization rate of the spill on water, the range of this value varied between approximately 0.029 to 0.195 kg m$^{-2}$ s$^{-1}$. In the case of Burro test, the simulated vaporisation rate is assumed to be $m'_{LNG} = 0.167$ kg m$^{-2}$ s$^{-1}$. Density of LNG vapour is similar to that of CH$_4$ at boiling point $\rho_{LNG} = 1.76$ kg m$^{-3}$ (Luketa-Hanlin et al., 2007). The spill diameter is derived from the vaporization rate, reported spill mass $m_{spill}$ and duration $t_{spill}$:

$$D_{spill} = \sqrt{\frac{4m_{spill}}{\pi m'_{LNG} t_{spill}}}$$  \hspace{1cm} (18)

The volume spill rate is used as gas inlet condition:

$$\dot{V}_{spill} = \frac{m_{spill}}{\rho_{LNG} t_{spill}}$$  \hspace{1cm} (19)

The LNG spill variables used in simulation are also tabulated in Table 7.

| Parameters | $u_{ref}$ | $u_*$ | $z_0$ | $T_{ref}$ | $t_{spill}$ | $D_{spill}$ | $\dot{V}_{spill}$ |
|-----------|-----------|-------|-------|-----------|------------|-------------|-----------------|
| Unit      | m/s       | m     | m/s   | K         | s          | m           | m$^3$/s         |
|           | 5.7       | 0.252 | 2E-4  | 308.55    | 79         | 32.2        | 77.17           |
5.2 Results and discussion of LNG vapour dispersion

5.2.1 The steady simulation

Profiles of velocity and turbulence quantities are sampled at the outlet boundary and compared with Monin-Obukhov theory profiles which are used as inlet boundary conditions. The steady state simulation of ABL with $k-\epsilon$ reveals that wind velocity and turbulence profiles are accurately reproduced as presented in Figure 7. The success of the modified $k-\epsilon$ model proves that the proposed model can adequately reproduce the Monin-Obukhov ABL profiles in the full scale simulation.

![Fig. 7 Comparing ABL profiles at outlet boundary of Burro9 simulation using modified $k-\epsilon$ model (kEpsMod) and MOST inlet profiles (MOST)](image)

5.2.2 Mesh sensitivity study

Maximum concentration at the arcs of 57 m, 140 m, 400 m and 800 m downwind are used as performance parameters for the mesh sensitivity study. Three meshes
with refined factors as summarised in Table 8 are used to simulate LNG gas dispersion under adiabatic thermal wall condition. Results from four peak arc-wise concentrations are plotted in Figure 8.

### Table 8 Burro test computational domain and mesh parameters

|                  | Mesh 1 | Mesh 2 | Mesh 3 |
|------------------|--------|--------|--------|
| Domain region    | [(-150, 0, 50), (850, 300, 50)] |
| Refined region   | [(-100, 0, 5), (400, 100, 5)] |
| Mesh size (m)    | 10     | 5      | 2      |
| Mesh refined size (m) | 5      | 2      | 1      |

![Fig. 8 Result of Burro mesh sensitivity study](image)

Increasingly mesh refinements help to resolve maximum concentration more accurately. The difference of gas concentrations between meshes are significantly
reduced with refinement. Due to computational restriction, no further mesh is
used for mesh sensitivity study and Mesh 3 parameters (Table 8) is chosen for the
following study.

5.2.3 Ground heat transfer sensitivity study

Three different models of heat transfer from the ground are used to study their
effect on the numerical results, which are summarised in Table 9. For constant
heat flux case, the value of 200 W/m$^2$ is used.

| Table 9 | Wall thermal boundary conditions in Burro tests |
|---------|------------------------------------------------|
|         | Case 1       | Case 2        | Case 3       |
| Heat transfer model | Adiabatic wall | Constant Heat Flux | Wall temperature |
| Label (Fig. 9)      | Adiabatic    | fixedFlux     | fixedTem     |

The effect of ground heat in predicting peak gas concentration is plotted in
Figure 9.

The adiabatic case results in a better prediction of experimental data than
the fixed flux and fixed temperature cases. However, all simulations yield under-
predicted results. This may be due to that the buoyancy effect is over-predicted
and consequently the gas concentration is zero in the fixed flux case at downwind
arcs (at 400 and 800 m).
5.2.4 Turbulence Schmidt number, $Sc_t$, sensitivity study

Two values of $Sc_t = 1$ and $Sc_t = 0.3$ are used for studying the sensitivity of the proposed model in predicting the maximum gas concentration. Results are compared in Figure 10.

$Sc_t = 0.3$, which was used previously in wind tunnel dense gas dispersion is shown to be appropriate for accurate prediction of maximum gas concentration at the 57 m array and 140 m array. Further downwind, at 400 m array and 800 m array, there is no significant difference between the two values.
Fig. 10 Result of Burro9 turbulent Schmidt number Sc study

5.2.5 Isosurface contour

The vertical isosurface contours at X = 140 are illustrated in Figure 11. Under-predicted cloud height are revealed in all tests indicating that the cloud buoyancy is not correctly solved.

Horizontal isosurface contours at height Z = 1 are shown in Figure 12. The gas concentration contour is plotted side by side with the contour from experimental data, where the left is the result of interpolating concentration at some concentration data points (presented in plots by black dot points), the right is from experimental data. Overall, the cloud height is considerably well predicted but the cloud width is over-predicted. Furthermore, it can be seen that the gas moves downwind slower than experimental data, which under-estimates the downwind spreading of the gas cloud.
Fig. 11 Vertical isosurface at X = 140, Top: Simulation, Bottom: Experimental data of Burro9 test.
Fig. 12 Horizontal isosurface at Z = 1; Top: Simulation, Bottom: Experimental data of Burro9 test
5.2.6 Concentration predictions

FDS (Fire Dynamics Simulator) (McGrattan et al., 2013) is a low Mach number code using the LES turbulence model. The computational domain is discretised into a connected rectilinear mesh. The governing equations are discretised using finite-difference method. A second-order scheme is used for space discretisation and an explicit second-order Runge-Kutta scheme for time discretisation. OpenFOAM concentration results are compared with FDS data extracted from (Mouilleau and Champassith, 2009).

The comparison of OpenFOAM, FDS, and experimental results for Burro9 test is shown in Figure 13. FDS is over-predicted, while OpenFOAM is under-predicted. However, OpenFOAM is accurate in prediction at 800 m arc.

**Fig. 13** Maximum arc-wise gas concentrations Burro9 test.
Figure 14 is the plot of gas concentration at 1 m elevation at 140 m downwind of Burro9 experiment (EXP) and simulations using the developed solver (FOAM) and FDS (FDS). For the developed solver result, the peak concentration is under-estimated while the temporal trend of changing concentration generally shows good agreement with validation data. The concentration magnitude is fairly matched except during local maximum/minimum durations. Also shown in Figure 14 is the result from FDS simulation which is generally over-predicted. However, the developed solver cannot capture the fluctuation, while FDS yields fluctuating gas concentration over the time period. This is an advantage of LES over RANS turbulence model. The over-prediction of FDS may be due to that a constant coefficient Smagorinsky model was adopted in the simulation. However, the dynamic Smagorinsky model was shown to improve the gas dispersion prediction (Ferreira Jr. and Vianna, 2016). This indicates that it would be a promising approach to use LES in order to enhance the performance of the developed solver.

5.2.7 Statistical model evaluation

Overall statistical performance of OpenFOAM results are compared versus FLACS with data extracted from (Hansen et al., 2010) in Table 10. The predictions do not match all SPMs. However, some important SPMs are within the acceptable range. All gas concentrations are within a factor of two (FAC2=1) and better than FLACS (FAC2 = 0.94).
Fig. 14  Point concentration at 140 m of Burro9 test

Table 10  Statistical performance measures of Burro tests

|                | MRB | RMSE | FAC2 | MG  | VG  |
|----------------|-----|------|------|-----|-----|
| FLACS (Hansen et al., 2010) | 0.16| 0.12 | 0.94 | 1.18| 1.14|
| FOAM Burro9    | 0.44| 0.23 | 1    | 0.63| 1.28|

6 Conclusions

A solver is developed to reproduce horizontal homogeneous atmospheric surface layer in neutrally stratified ABL using OpenFOAM. The empirical atmospheric boundary layer model MOST is used to specify the inlet boundary conditions for velocity, turbulent kinetic energy and dissipation rate. Flow variable profiles at outlet boundary are successfully maintained and consistent with their profiles at the inlet boundary. This demonstrates the effectiveness of the solver in simulating
the horizontal homogeneous atmospheric surface layer. It can also predict different levels of ABL turbulence kinetic energy.

A solver for ABL gas dispersion simulation taking into account buoyancy effect, variable turbulence Schmidt number and ground heat transfer is developed using the OpenFOAM platform. In the study of dense gas dispersion in neutral simulated ABL, the model is successfully validated by reproducing maximum gas concentration. SPMs from simulation results are better than those from the specialized commercial software for gas dispersion, FLACS.

In the study of LNG accidental release, a dense cold gas vapour dispersion in ABL with three ground heat transfer assumptions are simulated and compared with the full scale field measurements. The gas peak concentration is used as validation parameters. Adiabatic wall assumes zero heat flux from ground to the gas cloud, whereas, the fixed temperature model assumes isothermal ground where the ground temperature remains unchanged when in contact with the cold gas cloud. The real heat flux to the gas cloud would be in between these two cases. The other model assuming a fixed flux of heat to the gas cloud is also included. Of the three ground heat transfer models, adiabatic wall gives the closest prediction of gas peak concentration. The model is shown to accurately predict vertical buoyancy while the cloud spreading downwind is under-predicted. SPMs from the simulation results are compared with the LES code FDS and specialized dispersion code FLACS, showing that the solver is more accurate in predicting gas concentration in neutrally stratified ABL. Further investigation is required to validate the OpenFOAM solver in ABL with thermal stratification.
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CFD simulation of dense gas dispersion in neutral atmospheric boundary layer with OpenFOAM

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