CONTRIBUTION TO THE DEVELOPMENT OF IMPLICIT LARGE EDDY SIMULATIONS METHODS FOR COMPRESSIBLE TURBULENT FLOWS

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iii
ABSTRACT

CONTRIBUTION TO THE DEVELOPMENT OF IMPLICIT LARGE EDDY SIMULATIONS METHODS FOR COMPRESSIBLE TURBULENT FLOWS

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This work is intended to compare Large Eddy Simulation and Implicit Large Eddy Simulation (LES and ILES) for a turbulent, non-reacting or reacting high speed H2 jet in co-flowing air, typical of scramjet engines. Numerical simulations are performed at resolutions ranging from 32×32×128 to 256×256×1024, using a 5th order WENO scheme. Physical LES are carried out with the Smagorinsky and the Selective Structure Function models associated to molecular diffusion. Implicit LES are performed with and without molecular diffusion, by solving either the Navier-Stokes or the Euler equations. In the nonreacting case, the Smagorinsky model is too dissipative. The Selective Structure Function leads to better results, but does not show any superiority compared to ILES, whatever the grid resolution. In the reacting case, a molecular viscous cut-off in the simulation is mandatory to set a physical width for the reaction zone in the ILES approach, hence to achieve grid-convergence. It is also found that ILES/LES are less sensitive to the inlet conditions than the RANS approach. The first chapter is an introduction to the context of this study. In the second chapter, the governing equations for multispecies reacting flows are presented, with emphasis on the thermodynamic and transport models. In the third chapter, physical LES equations and explicit sub-grid modeling strategies
are detailed. Some properties of the numerical scheme are also investigated. In chapter four, the numerical scheme and some aspects of the solver are explained. Finally, non-reacting and reacting numerical experiments are presented and the results are discussed.

Keywords: turbulent, combustion, LES, implicit, compressible
ÖZ

SIKİŞTİRİLӘBLİR VE TEPKİMELİ TÜRBÜLANSLI AKIMLAR İÇİN ZIMNI BÜYÜK GİRÐAP BENZETİMİ YÖNTEMLERİ GELİŞ TİRİLMESİNİ KATKI

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Bu çalışmada scramjet motorlarına özgü yüksek hızda, türbülanslı, tepkimesiz veya tepkimeli, eş-hava akışında H2 jeti için uygulanan Büyük Girdap Benzetimi ve Zimni Büyük Girdap Benzetimi (BGB ve ZBGB) metotlarının karşılaştırılması amaçlanmaktadır. Sayısal simülasyonlar, 5. seviye WENO şemasını kullanarak, 32 × 32 × 128, 256 × 256 × 1024 arasında değişen çözünürlüklerde yapılmıştır. Fiziksel BGB, Smagorinsky ve Seçici Yapısal Fonksiyon modelleriyle bilikte moleküler difüzyon uygulanılarak yansıtılmaktadır. Örtülü BGB, moleküler difüzyon eklenip çıkarılarak elde edilen Navier-Stokes veya Euler denklemleri çözüllerek yapılmıştır. Tepkimesiz durumda, Smagorinsky modeli çok tüketimlidir. Seçici Yapısal Fonksiyonu daha iyi sonuçlar vermektedir, ancak ZBGB ile karşılaştırıldığında, ag çözünürlük ne olursa olsun herhangi bir üstünlük göstermez. Tepkimeli durumda, ZBGB yaklaşımı için reaksiyon bölgesinde fiziksel bir et kalınılığı ve ag yakınsaması sağlayan moleküler viskozite zorunludur. Ayrıca, BGB ve ZBGB metotlarının giriş şartlarına RANS yaklaşımına göre daha az duyarlı olduğu görülmüştür. İlk bölümde, çalışma bağlamında bir giriş ve çözümleme yaklaşımlar temsil eden denklemler, termodinamik ve taşıma modelleri üzerinde durulurak sunulmuştur. Üçüncü bölümde, fizik-
sel BGB denklemler ve ag-altı modelleme stratejileri ayrıntılı olarak verilmektedir, sayısal şemanın bazı özellikleri de incelenmiştir. Dördüncü bölümde, sayısal şema ve çözümünün bazı yönleri açıklanmıştır. Son olarak, tepkimesiz ve tepkimi sayısal deneylerin sonuçları tartışılmaktadır.

Anahtar Kelimeler: türbülanslı, yanma, ZBGB, Zımnı, sıkıştırılabilir
dedicated to my family.
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TABLE OF CONTENTS

ABSTRACT .......................................................... iv
ÖZ ................................................................. vi
ACKNOWLEDGMENTS ........................................ ix
TABLE OF CONTENTS .......................................... x
LIST OF TABLES ................................................ xv
LIST OF FIGURES ............................................... xvii

CHAPTERS

1 INTRODUCTION ............................................... 1

2 CONSERVATION LAWS FOR COMPRESSIBLE MULTICOMPONENT REACTING FLOWS ........................................ 7

2.1 CONSERVATION OF MASS AND CHEMICAL SPECIES .... 7

2.1.1 STATIC VARIABLES FOR A MIXTURE ............... 7

2.1.2 CONSERVATION OF SPECIES AND MASS ........... 8

2.1.3 MOLECULAR DIFFUSION OF SPECIES ............... 9

2.1.4 CHEMICAL KINETICS .................................. 12

2.2 EQUATION OF MOTION ................................... 13

2.2.1 THE EQUATION OF STATE ............................ 13

2.2.2 MOLECULAR DIFFUSION OF MOMENTUM ........... 13

2.3 ENERGY EQUATION ........................................... 14

2.3.1 INTERNAL ENERGY, ENTHALPY AND SPECIFIC HEATS 14

2.3.2 MOLECULAR DIFFUSION OF HEAT AND ENTHALPY 15

2.3.3 TEMPERATURE EVALUATION .......................... 16

2.4 SUMMARY OF SYSTEM OF EQUATIONS .................... 16
3 LARGE EDDY SIMULATION ........................................... 18
3.1 FILTERING THE NAVIER-STOKES EQUATIONS ............... 20
  3.1.1 CONSERVATION OF MASS .................................. 22
  3.1.2 CONSERVATION OF CHEMICAL SPECIES .................. 23
  3.1.3 EQUATION OF MOTION .................................... 23
  3.1.4 ENERGY EQUATION ........................................ 25
  3.1.5 SPECIAL ATTENTION FOR W5 AND IDEAL GAS EQUATION . 25
3.2 EXPLICIT SUB-GRID MODELING ................................. 27
  3.2.1 CLOSING THE FILTERED EQUATION OF MOTION ........ 27
    3.2.1.1 THE SMAGORINSKY MODEL .......................... 27
    3.2.1.2 THE YOSHIZAWA MODEL ............................ 28
    3.2.1.3 THE SELECTIVE STRUCTURE FUNCTION MODEL .... 29
  3.2.2 CLOSING THE FILTERED ENERGY EQUATION .............. 30
  3.2.3 CLOSING THE FILTERED SPECIES EQUATION ............. 30
3.3 Implicit LES .................................................. 30
  3.3.1 MWN ANALYSIS ........................................... 31
  3.3.2 THE INTERACTION BETWEEN NUMERICAL ERRORS AND EXPLICIT SUB-GRID MODELING ............. 34
4 THE WENO SOLVER ................................................ 36
  4.1 TRANSFORMATION BETWEEN CONSERVATIVE, PRIMITIVE AND CHARACTERISTIC FORMS ................. 37
  4.2 CONSERVATIVE FORM FOR DIFFERENTIAL OPERATOR ....... 38
  4.3 WENO SCHEME ................................................ 39
    4.3.1 ENO APPROXIMATION .................................. 40
    4.3.2 1D WENO RECONSTRUCTION PROCEDURE ............... 41
    4.3.3 FLUX SPLITTING ....................................... 42
    4.3.4 SOLUTION FOR SYSTEM OF EQUATIONS ............... 43
  4.4 TIME DISCRETIZATION ....................................... 44
    4.4.1 RKp TVD TIME STEPPING ............................... 44
    4.4.2 CALCULATING THE TIME STEP .......................... 45
4.5 BOUNDARY CONDITIONS ................. 46
  4.5.1 LODI SYSTEM ...................... 46
  4.5.2 FREE NON-REFLECTING BCS ........ 48
  4.5.3 PHYSICAL NON-REFLECTING INLET BOUNDARY CONDITIONS .......... 48
    4.5.3.1 SUBSONIC FORCED NON-REFLECTING INLET .................. 48
    4.5.3.2 SUPERSONIC FORCED NON-REFLECTING INLET ................. 49

4.6 SOME SPECIFICITIES ................... 49
  4.6.1 THE LARROUTUROU METHOD .......... 49
    4.6.1.1 IDEA OF LARROUTUROU .......... 49
    4.6.1.2 APPLICATION TO WENO SOLVER .... 50
  4.6.2 STRUCTURE OF THE CODE .......... 50
  4.6.3 THE QUESTION OF $D_{H0}$ .......... 50
  4.6.4 THE METRICS ..................... 52
  4.6.5 PARALLEL IMPLEMENTATION .......... 53
  4.6.6 LES STRATEGIES .................. 54

5 NUMERICAL EXPERIMENTS ................. 55
  5.1 NON-REACTING CASE .................. 56
    5.1.1 TEST CASE DESCRIPTION .......... 56
    5.1.2 PHYSICAL AND NUMERICAL PARAMETERS .... 58
    5.1.3 FLOW STRUCTURE ................. 63
    5.1.4 STATISTICAL RESULTS ............ 70
    5.1.5 VARIATION OF VELOCITY, SPECIES MASS FRACTION AND PRESSURE .... 76
    5.1.6 IS THE DNS LEVEL REACHED ? ....... 86
    5.1.7 CONCLUSIONS .................... 87

  5.2 REACTING CASE ...................... 88
    5.2.1 TEST CASE DESCRIPTION .......... 88
    5.2.2 INLET FIELD .................... 90
      5.2.2.1 SIMPLE PROFILE ............... 91
<table>
<thead>
<tr>
<th>Chapter</th>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>H</td>
<td>COMPRESSIBLE TEMPORAL MIXING LAYER A PRIORI TESTS</td>
<td>153</td>
</tr>
<tr>
<td>I</td>
<td>COMPUTATIONAL HOURS</td>
<td>157</td>
</tr>
</tbody>
</table>
### LIST OF TABLES

#### TABLES

<table>
<thead>
<tr>
<th>Table</th>
<th>Description</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Table 5.1</td>
<td>Physical parameters for the non-reacting air/H₂ jet. In bold, data from Eggers [32], other values are computed.</td>
<td>58</td>
</tr>
<tr>
<td>Table 5.2</td>
<td>Ratios of approximate Kolmogorov scale and Taylor microscale to the minimum grid spacing.</td>
<td>61</td>
</tr>
<tr>
<td>Table 5.3</td>
<td>Coefficients which are used for the inlet velocity profile.</td>
<td>61</td>
</tr>
<tr>
<td>Table 5.4</td>
<td>The slope and fictitious center of inverse normalized time averaged velocity.</td>
<td>76</td>
</tr>
<tr>
<td>Table 5.5</td>
<td>The slope and fictitious center of inverse normalized time averaged hydrogen mass fraction.</td>
<td>76</td>
</tr>
<tr>
<td>Table 5.6</td>
<td>LAERTE inlet hyperbolic tangent profile coefficients for equation 5.6.</td>
<td>92</td>
</tr>
<tr>
<td>Table 5.7</td>
<td>LAERTE experimental conditions.</td>
<td>94</td>
</tr>
<tr>
<td>Table 5.8</td>
<td>Ratios of approximate Taylor micro-scale and Kolmogorov scale to the minimum grid spacing.</td>
<td>95</td>
</tr>
<tr>
<td>Table A.1</td>
<td>Thermal conductivity coefficients for second order polynomial fit (i.e. equation 2.50).</td>
<td>128</td>
</tr>
<tr>
<td>Table A.2</td>
<td>Coefficients of viscosity for second order polynomial fit (i.e. equation 2.37).</td>
<td>128</td>
</tr>
<tr>
<td>Table A.3</td>
<td>Power law fitted diffusion coefficients for P₀(=1bar), T (i.e. equation 2.21).</td>
<td>129</td>
</tr>
<tr>
<td>Table B.1</td>
<td>Thermodynamic property data format.</td>
<td>130</td>
</tr>
<tr>
<td>Table B.2</td>
<td>Thermodynamic properties. (Burcat [14]).</td>
<td>131</td>
</tr>
<tr>
<td>Table C.1</td>
<td>Efficiency of the species for the ONERA scheme.</td>
<td>132</td>
</tr>
<tr>
<td>Table C.2</td>
<td>H₂/air Chemical reaction mechanism by ONERA.</td>
<td>133</td>
</tr>
<tr>
<td>Table G.1</td>
<td>Coefficients cᵣᵢ for k=1 to 6.</td>
<td>148</td>
</tr>
</tbody>
</table>
Table G.2  Coefficients $d_r$ for $k=1$ to 6. .................................................. 149

Table H.1  D.N.S. physical and numerical parameters for the O$_2$/N$_2$ pair. .............. 154
Table H.2  D.N.S physical and numerical parameters for the O$_2$/H$_2$ pair. .............. 155
Table I.1  Relative cost of LES and MILES, non-reacting and reacting. .................. 157
LIST OF FIGURES

FIGURES

Figure 1.1 Leduc 010, 1949, 680 km/h. ........................................ 2
Figure 1.2 The rocket-powered X-15 (M=6.7 in 1967). .................. 2
Figure 1.3 Schematic of a scramjet [3]. ...................................... 2
Figure 1.4 Specific impulse vs. flight conditions for various engine concepts. Red : H₂, Blue : Kerosene ......................... 3
Figure 1.5 Flight envelop of different engine concepts. .................. 3
Figure 1.6 The X43 in the NASA LaRC M=10 wind tunnel. ............ 4
Figure 1.7 The Boeing X51-A. .................................................. 4
Figure 1.8 The CAD view of the LEA vehicle. .............................. 4
Figure 1.9 The LEA flight test mission. ...................................... 4

Figure 3.1 Linear Modified Wave Number without (top) and with (bottom) non-linear WENO weight. Left : real part, right : imaginary part. (a) 3rd order WENO, (b) 5th order WENO, (c) 7th order WENO, (d) 9th order WENO, (e) 11th order WENO, (f) 1st order upwind FD, (g) 2nd order upwind FD ..................... 33
Figure 3.2 Interaction indicator eq. (3.89) for C = 0, 0.1, 1 and 10. Left: real part, right: imaginary part. .......................... 35

Figure 4.1 Transformations between different forms for the Euler system of equations. 38
Figure 4.2 5th order WENO stencil demonstration. ....................... 40
Figure 4.3 ECBC subsonic flux directions ................................. 47
Figure 4.4 ECBC supersonic flux directions ............................ 47
Figure 4.5 Direction of the waves for subsonic and supersonic conditions for both u > 0 (black arrows) and u < 0 (red arrows) cases. .......... 47
Figure 4.6  Schematic diagram of structure of the code. .......................... 51
Figure 4.7  Coordinate transformation for clustering the grid .................... 52
Figure 4.8  A. Speedup curves for WENO code on clusters VARGAS and Phoebus, B. 2D MPI cartesian topology. .......................... 53

Figure 5.1  Schematic diagram of Eggers jet. ........................................ 57
Figure 5.2  Shear layer convective velocity and Mach numbers[8]. ............. 59
Figure 5.3  Inlet velocity profile for the non-reacting air/H_2 Eggers jet. ....... 59
Figure 5.4  Grid for the Eggers jet of resolution 32×32×128 .................... 60
Figure 5.5  Inlet velocity profile for the non-reacting air/H_2 Eggers jet. ....... 61
Figure 5.6  Radial distribution of confinement function, samples of generated ‘random’ field and the noise applied on transverse velocity components. .......... 62
Figure 5.7  Eggers jet results on 32×32×128 grid. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches. .......................... 65
Figure 5.8  Eggers jet results on 64×64×256 grid. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches. .......................... 66
Figure 5.9  Eggers jet result on 128×128×512 grid with reduced Smagorinsky constant. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches. ............ 67
Figure 5.10 Eggers jet results on 128×128×512 grid. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches. .......................... 68
Figure 5.11 Eggers jet results on 256×256×512 grid. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches. .......................... 69
Figure 5.12 Eggers jet : axial velocity decay. Clockwise from top-left : LES_SM, LES_SSF, MILES_NS and MILES.Euler ......................... 70
Figure 5.13 Eggers jet : transverse velocity profiles at x=0.06, 0.11, 0.18 and 0.29 m from the exit. Resolution 128×128×512 .......................... 71
Figure 5.14 Eggers jet: transverse velocity profiles at \( x = 0.06, 0.11, 0.18 \) and 0.29 m from the exit. Resolution 256x256x512 .............................. 72

Figure 5.15 Eggers jet: \( \text{H}_2 \) mass fraction axial decay. Clockwise from top-left: LES\_SM, LES\_SSF, MILES\_NS and MILES\_EULER .............................. 73

Figure 5.16 Definitions for inverse normalized time averaged quantity. ................. 73

Figure 5.17 Variables for inverse normalized average quantity. ......................... 74

Figure 5.18 Variation of the inverse normalized quantities \( Q^* \) on the axis of turbulent jet. 74

Figure 5.19 The inverse normalized velocity \( Q^*_U \) (left) and the inverse normalized \( \text{H}_2 \) mass fraction \( Q^*_Y_{\text{H}_2} \) (right) for 64x64x256, 128x128x512 and 256x256x512 resolutions (from upper to lower). .............................. 75

Figure 5.20 Instantaneous axial velocity component on the jet axis at 256x256x512. 77

Figure 5.21 Eggers jet: Resolved turbulent kinetic energy, resolution 128x128x512. 79

Figure 5.22 Eggers jet: Nitrogen mass fraction variance, resolution 128x128x512. 79

Figure 5.23 Eggers jet: Resolved turbulent kinetic energy, resolution 256x256x512. 79

Figure 5.24 Eggers jet: Nitrogen mass fraction variance, resolution 256x256x512. 79

Figure 5.25 Sub-figures demonstrating from top: instantaneous velocity over symmetry axis, sampled time variation and spectra of sampled data of 128x128x512 grid solutions (velocity in m/s and frequency in kHz). 80

Figure 5.26 Sub-figures demonstrating from top: instantaneous velocity over symmetry axis, sampled time variation and spectra of sampled data of 256x256x512 grid solutions (velocity in m/s and frequency in kHz). 81

Figure 5.27 Sub-figures demonstrating from top: instantaneous hydrogen mass fraction over symmetry axis, sampled time variation and spectra of sampled data of 128x128x512 grid solutions (frequency in kHz). 82

Figure 5.28 Sub-figures demonstrating from top: instantaneous hydrogen mass fraction over symmetry axis, sampled time variation and spectra of sampled data of 256x256x512 grid solutions (frequency in kHz). 83

Figure 5.29 Sub-figures demonstrating from top: instantaneous pressure over symmetry axis, sampled time variation and spectra of sampled data of 128x128x512 grid solutions (pressure in Pa and frequency in kHz). 84
Figure 5.30 Sub-figures demonstrating from top: instantaneous pressure over symmetry axis, sampled time variation and spectra of sampled data of 256×256×512 grid solutions (pressure in Pa and frequency in kHz). 85

Figure 5.31 Eddy-viscosity ratio, LES_SM. 86

Figure 5.32 Eddy-viscosity ratio, LES_SSF. 87

Figure 5.33 Schema of the LAERTE experiment. 88

Figure 5.34 Experimental data available. From left to right: temperature, velocity and OH molar concentration. 89

Figure 5.35 Initial field data taken from RANS solutions Davidenko [22]; from left to right radial profiles for temperature, velocity and species concentrations. 90

Figure 5.36 Simplified initial field data deduced from RANS solutions by Davidenko [22]; from left to right radial profiles for temperature, velocity and species concentrations. 91

Figure 5.37 Inlet velocity profile for the reacting air/H₂ LAERTE jet. 92

Figure 5.38 Initial field data deduced from Davidenko RANS solutions [22] including shock patterns at inlet; from left to right radial profiles for temperature, velocity and species concentrations. 93

Figure 5.39 0D constant pressure tests for stoichiometric H₂/air (left) and stoichiometric vitiated H₂/air (right) mixtures. The dashed lines refer to the stable solutions and the solid lines to the unstable ones. 95

Figure 5.40 0D constant volume tests for stoichiometric H₂/air (left) and stoichiometric vitiated H₂/air (right) mixtures. The dashed lines refer to the stable solutions and the solid lines to the unstable one. 96

Figure 5.41 Fuel/oxidizer mixture fraction distribution. 98

Figure 5.42 Instantaneous iso-surfaces of the stoichiometric mixture fraction, temperature contours and water mass fraction contours on the symmetry planes for 64×64×256 grid. From top to bottom: LES_SSF, MILES_NS, MILES_EULER. 100

Figure 5.43 Instantaneous iso-surfaces of the stoichiometric mixture fraction, temperature contours and water mass fraction contours in the symmetry planes for 128×128×512 grid. From top to bottom: LES_SSF, MILES_NS, MILES_EULER. 101
Figure 5.44 Instantaneous iso-surfaces of the stoichiometric mixture fraction, temperature contours and water mass fraction contours in the symmetry planes for 256×256×1024 grid. From top to bottom: LES_SSF, MILES_NS, MILES_EULER.

Figure 5.45 Instantaneous axial velocity component on the jet axis and in the main shear region. MILES_NS, 256×256×1024.

Figure 5.46 Scatter plots of temperature. Top to bottom: LES_SSF, MILES_NS, MILES_EULER. Left to right: 64×64×256, 128×128×512, 256×256×1024.

Figure 5.47 Scatter plots of O2 mass fraction. Top to bottom: LES_SSF, MILES_NS, MILES_EULER. Left to right: 64×64×256, 128×128×512, 256×256×1024.

Figure 5.48 Average profiles of velocity, H2 mass fraction and temperature at the center line of the jet and maximum temperature at the transverse sections along x axis. Left to right: 64×64×256, 128×128×512, 256×256×1024.

Figure 5.49 Tranverse profiles of axial velocity (left) and temperature (right) at x = 210 mm. Top to bottom: 64×64×256, 128×128×512, 256×256×1024.

Figure 5.50 Eddy-viscosity, LES_SSF

Figure 5.51 Flame structure of LAERTE chamber with precise initial field, including strong shock patterns.

Figure 5.52 Pressure contours on the symmetry axis, of LAERTE chamber with precise initial field.

Figure 5.53 Wall pressure axial distribution numerical results and experimental data.

Figure 5.54 Maximum temperature distribution along the jet axis computed over normal planes.

Figure 5.55 Transverse OH concentration profiles at x=0.21 m along the axis.

Figure 5.56 Experimental transverse profiles of axial velocity (left) and temperature (right) at x=10 mm, x=72 mm, x=210 mm and x=288 mm.

Figure H.1 Computational box for the temporal direct numerical simulation

Figure H.2 Subgrid terms (H.1). Relative magnitude: O2/N2.

Figure H.3 same as Figure H.2: O2/H2.

Figure H.4 Subgrid terms (H.3). Relative magnitude: O2/N2.

Figure H.5 same as Figure H.4: O2/H2.
Figure H.6 Subrid terms (H.4). Relative magnitude: O$_2$/N$_2$ . . . . . . . . . . . . . . . 156

Figure H.7 same as Figure H.6: O$_2$/H$_2$ . . . . . . . . . . . . . . . . . . . . . . . . . . . 156
CHAPTER 1

INTRODUCTION

The mankind’s desire to fly higher, faster and farther sustained the development of propulsion systems since the first powered flight of Wright brothers in 1903. In the early years of aviation, René Lorin patented an air-breathing device which compresses the incoming air by its inlet (diffuser) and expands it by its exhaust (nozzle) without any moving parts. This engine was the first ramjet concept. At that time, this concept would not be realized since a ramjet cannot produce thrust at zero speed. Similar independent concepts were developed in different countries.

This initial interest in ramjets seemed to be reduced in favor of the sustained development of gas turbine engine, the concept of which was first patented in 1791, by John Barber (UK patent no. 1833). Sir Frank Whittle (England) ran the W2/700 engine, and in 1936 Von Ohain (Germany) ran his engine with H₂. During the second world war (WWII), German flew the first operational ramjet, ground launched V-1 Bomb Flyer which was launched by a solid propellant booster. The first ramjet powered aircraft Leduc 0.10 (Figure 1.1) which was carried aloft by a mother airplane was launched in 1947, in France. Later, after WWII, operational ground launched ramjets were developed as missiles like Talos in US and VEGA in France.

Access to space gave a big impetus for hypersonic flight. The first hypersonic manned rocket powered aircraft called X-15 (Figure 1.2) was built in 60’s by North American Society on behalf of NASA. The X-15 was flying at ≈15 km and reached Mach 6.7 in 1967. This airplane was used to collect experimental data at hypersonic flight regimes at high altitude and to make component tests. The engines driving high-speed flight vehicles had limitations imposed by propulsion: air-breathing engines (turbojet, ramjet) do not reach hypersonic speeds and rocket
engines have the disadvantage of requiring the carriage of oxidant. The scramjet (supersonic combustion ramjet) development which has begun in 1955s permitted to remove these drawbacks. In a ramjet, the flow is subsonic since there is a throat on the flow path. Multi-shocks at the inlet cause high amount of losses at flow velocities higher than Mach 6 (Figure 1.3). On the contrary, in a scramjet, the flow remains supersonic throughout the combustion process, resulting in more efficient operation at hypersonic speeds of the airplane. Figure 1.4 shows the specific impulse vs operating condition of various engine concepts. Rocket engines specific impulse values are about 400 s which is less compared to scramjet. Figure 1.5 shows the altitude limitation vs Mach number and the so-called flight corridor of scramjets. As seen in the Figure, the optimal altitude for scramjet flight at Mach=6 is around 25 km. United States having the hypersonic test capability of X-15, started the NASA hypersonic research engine HRE (hypersonic research engine) program in 1964, and France launched ESOPE (Etude de
Statoréacteur comme Organe de Propulseur Évolué) program in 1966. However, these attempts terminated due to limited technology and interest focused on integrated rocket ramjet (IRR) technology.

In the mid 1980’s, the NASP (National AeroSpace Plane) program was initiated with the goal of flying a reusable launch vehicle, the X-30, powered by a single stage combined cycle engine to Mach 25. Also, at the same time, the PREPHA (Programme de REcherche en Propulsion Hypersonique Avancée) program was launched in France. Considering the state of technology, the goals of these projects were aggressive; eventually these projects were abandoned leaving extensive experience and significant contributions behind. Research in Europe continued with the JAPHAR project (Joint Airbreathing Propulsion for Hypersonic Application Research) co-directed by ONERA (France) and DLR (Germany). These studies have been conducted from 1997 to 2002 as the preliminary study of a dual-mode ramjet hypersonic vehicle.

At the same time, the Hyper-X program launched in the United States aimed to fly a small scale scramjet powered demonstrator at Mach 10. This target was reached in November 2004 by the X-43A (Figure 1.6). Later, the X-51 Waverider program was initiated to demonstrate sustained flight of 300 s with a scramjet powered aircraft. The X-51 was planned to accelerate from Mach 4.7 to beyond Mach 6. The first powered flight test was conducted on 26 May 2010 over 200 s at Mach 5 (Figure 1.7). In Europe, after JAPHAR, the PROMETHEE program and
now LEA (Figure 1.8) program is aimed to develop and launch an experimental hypersonic test vehicle by ONERA-MBDA. The first flight planned to be at 20-30 km altitude from Mach 4 to beyond Mach 8. This experimental vehicle is planned to be delivered to proper altitude and accelerated to the required Mach number by Russian Tu-22 aircraft and Volna launcher (Figure 1.9).

In the present review, some of the hypersonic flight projects were listed. More thorough information about history may be found in references [22, 38, 37, 99, 77, 45, 124].

In a scramjet, the main design considerations are:

- The forebody and the intake act as a diffuser capturing and compressing air with the highest possible efficiency. They contribute to the moment and drag of the vehicle. The flow includes leading bow shock and isentropic or oblique turning shock waves. The transition on the forebody is highly important due to high heating involved.

- The pressure is increased in the isolator which acts as a component of the diffuser.
Oblique shocks penetrating into the combustion chamber are formed in the isolator.

- In the combustion chamber, the fuel is delivered into the flow, mixes and burns in the available chamber length. The (turbulent) mixing and chemical reactions in the supersonic flow conditions have to be complete within less than one millisecond.

- Finally, expansion occurs in the exhaust nozzle and aftbody. The design of the expansion system contributes to the propulsive efficiency and the moments on the vehicle.

The context of this work is the numerical simulation of turbulent mixing and reacting flow in scramjet combustion chamber. Shock waves are inherently present in the flow and require the use of dissipative numerical schemes for stable simulations, whatever the approach retained to model turbulence. Until quite recently, Reynolds averaged calculations (RANS) have been the standard approach to the problem (e.g. [74]). However, turbulence models extended to highly compressible flows and large heat release must be finely tuned, with some uncertainty when extrapolated to different configurations [23, 22]. Direct Numerical Simulation (DNS) of such complex flows is not feasible and may probably never be. LES is a more attractive technique [50] now at hand thanks to the increase in computing power. The purpose of the present thesis is the comparison of the Large Eddy Simulations (LES) and Monotone Integrated Large Eddy Simulation (MILES) numerical approaches in the context of high speed jets. In the former the contribution of the sub-grid motions are modeled based on flow physics and in MILES algorithms designed to satisfy the physical requirements of positivity and causality, in effect have a matching sub-grid model built in [11].

At ICARE the studies related to this topic were initiated by the thesis of Lardjane [80] in which 1D and 2D analysis of numerical methods for turbulent mixing in high density ratio binary systems have been conducted. Later, in the thesis by Gougeon [59], 2D numerical simulations are developed for compressible reacting flows. In the present study, the 2D simulations are extended to full 3D. Closure models for the filtered set of equations are applied for LES and different LES strategies are compared.

The thesis is organized as follows. Chapter 2 presents the governing equations for multicomponent compressible reacting flows, with emphasis on the thermodynamic and transport models. In chapter 3, physical LES equations are detailed; the explicit SGS modeling strategies are explained and dissipation and dispersion characteristics of different numerical approaches
are discussed. In the same chapter, a simple 1D linear analysis of the interaction between
the WENO scheme and SGS model is also proposed. Chapter 4 is devoted to numerics. The
details of the numerical scheme and some aspects of the solver are explained in this chapter.
In chapter 5, both the non-reacting (section 5.1) and reacting (section 5.2) test cases and the
results are presented. Finally, conclusions of this study are given in chapter 6.
CHAPTER 2

CONSERVATION LAWS FOR COMPRESSIBLE
MULTICOMPONENT REACTING FLOWS

In this study, high speed non-reacting and reacting flows are simulated. The cases are high Reynolds number flows near atmospheric conditions, hence the Knudsen number is small and the continuum assumption holds. The system is assumed to be in local thermodynamic equilibrium, having a single temperature. The Navier-Stokes equations for three-dimensional multi-species flows are considered as the governing equations. Perfect gas equation of state is applied for the relation between pressure, temperature and density.

The description of coupled, time dependent, partial differential equations governing the convective flow, diffusion of species and source terms for chemical reactions is presented in this chapter. This set of equations is well known and can be found in many popular textbooks [78, 98, 104, 125, 128]. The purpose of this chapter is to state precisely the notations and definitions used in this study and to establish references for the following chapters.

2.1 CONSERVATION OF MASS AND CHEMICAL SPECIES

2.1.1 STATIC VARIABLES FOR A MIXTURE

Let consider an homogeneous mixture of $N_{sp}$ species $\alpha = 1, ..., N_{sp}$. The density of the mixture may be computed as the sum of partial densities $\rho_{\alpha}$

$$\rho(p, T, Y_{\alpha}) = \sum_{\alpha=1}^{N_{sp}} \rho_{\alpha}(p, T) \quad (2.1)$$

Species mass fractions $Y_{\alpha}$ are
\[ Y_\alpha = \frac{\rho_\alpha}{\rho} ; \quad \sum_{\alpha = 1}^{N_{sp}} Y_\alpha = 1 \]  
(2.2)

Species can also be characterized with their molar concentrations and mole fractions. Molar concentration of species \( \alpha \) in the mixture is expressed in the following way
\[ C_\alpha = \frac{\rho_\alpha}{M_\alpha} = \frac{\rho Y_\alpha}{M_\alpha} \]  
(2.3)

\( M_\alpha \) is the molar mass of the species \( \alpha \). The overall molar concentration of the mixture is
\[ C = \sum_{\alpha = 1}^{N_{sp}} C_\alpha \]  
(2.4)

Mole fractions can be directly computed from molar concentrations as
\[ X_\alpha = \frac{C_\alpha}{C} ; \quad \sum_{\alpha = 1}^{N_{sp}} X_\alpha = 1 \]  
(2.5)

A simple relation between mass fractions, mole fractions and molar mass is
\[ X_\alpha M_\alpha = Y_\alpha M \]  
(2.6)

Average molar mass of the mixture may be computed using both variables \( X_\alpha \) and \( Y_\alpha \) as
\[ M = \frac{\rho}{C} = \sum_{\alpha = 1}^{N_{sp}} X_\alpha M_\alpha = \left( \sum_{\alpha = 1}^{N_{sp}} Y_\alpha \right)^{-1} \]  
(2.7)

### 2.1.2 CONSERVATION OF SPECIES AND MASS

In an infinitesimal control volume, the change of species mass fractions, for chemically reacting flows, is due to convection, diffusion of species and source terms of reactions.
\[ \frac{\partial \rho Y_\alpha}{\partial t} + \frac{\partial \rho Y_\alpha (u_j + V_{\alpha j})}{\partial x_j} = \dot{\omega}_\alpha \]  
(2.8)

In this relation \( u_j \) is the convective velocity, \( V_{\alpha j} \) is the diffusion velocity of species \( \alpha \) in direction \( x_j \) and \( \dot{\omega}_\alpha \) is the source term for species \( \alpha \). The diffusion velocity in equation (2.8) may be used to express the species mass flux
\[ J_{\alpha j} = \rho Y_\alpha V_{\alpha j} \]  
(2.9)

The species conservation equation (2.8) turns out to be
\[ \frac{\partial \rho Y_\alpha}{\partial t} + \frac{\partial \rho u_j Y_\alpha}{\partial x_j} = -\frac{\partial J_{\alpha j}}{\partial x_j} + \dot{\omega}_\alpha \]  
(2.10)
Both the sum of the source terms $\dot{\omega}_\alpha$ and species mass flux $J_{\alpha j}$ must be zero to satisfy conservation of mass.

$$\sum_{\alpha=1}^{N_{sp}} \dot{\omega}_\alpha = 0 \quad ; \quad \sum_{\alpha=1}^{N_{sp}} Y_\alpha V_{\alpha j} = 0$$

The sum of species equations gives conservation of mass. The change of mass in the control volume is due to convection

$$\frac{\partial \rho}{\partial t} + \frac{\partial \rho u_j}{\partial x_j} = 0$$

### 2.1.3 MOLECULAR DIFFUSION OF SPECIES

The species diffusion velocity $V_{\alpha j}$ in equation (2.8) can be expressed according to kinetic theory of gases [16, 36, 44, 66].

$$V_{\alpha j} = -\sum_{\beta=1}^{N_{sp}} \left( \frac{D_{\alpha \beta}^{T} \partial \ln T}{\partial x_j} + D_{\alpha \beta} d_{\beta j} \right)$$

The first term on the r.h.s. is the thermal diffusion also known as the Soret effect [95]. In reacting flows, since temperature gradients take large values, Soret effect may be significant. However, in this study, it is neglected for simplicity.

In the second term on the r.h.s., $d_{\alpha j}$ is the diffusion driving force of species $\alpha$ in direction $x_j$. This term can be derived by elementary analysis [104, 128].

$$d_{\alpha j} = \frac{\partial X_\alpha}{\partial x_j} + (X_\alpha - Y_\alpha) \frac{\partial \ln p}{\partial x_j} + \frac{\rho}{p} \sum_{\beta=1}^{N_{sp}} Y_\beta Y_\alpha (b_{\beta j} - b_{\alpha j})$$

$D_{\alpha \beta}$ is the coefficient for the inter-diffusion of species $\alpha$ and $\beta$ in the presence of any number of other species. The diffusion coefficients take the name “binary diffusion coefficients” with notation $D_{\alpha \beta}$, when there exists two species only. $D_{\alpha \beta}$ can be computed algebraically [16, 66]. In equation (2.13) the diffusion velocities are explicit. However, the multicomponent diffusion coefficients $D_{\alpha \beta}$ are not (as $N_{sp} > 2$). They can be approximated in terms of Sonine polynomial expansion as a function of the primitive variables and binary diffusion coefficients. A system of equations for the coefficients needs to be solved [36]. This procedure is explained in detail by Dixon-Lewis [27] and Coffee & Heimerl [18]. This task is computationally expensive.

An alternative method can be derived using the dual formulation of equation (2.13), neglecting
the Soret effect.

\[ d_{\alpha j} = - \sum_{\beta=1}^{N_{sp}} \Delta_{\alpha \beta} V_{\alpha j} \]  \hspace{1cm} (2.15)

The dual multicomponent diffusion coefficients \( \Delta_{\alpha \beta} \) are not known but can be approximated using the first term of a Sonine polynomial expansion. The Stefan-Maxwell diffusion equations are then obtained [55, 95].

\[ d_{\alpha j} = \sum_{\beta=1}^{N_{sp}} \frac{X_\alpha X_\beta}{D_{\alpha \beta}} V_{\beta j} - \left( \sum_{\beta=1}^{N_{sp}} \frac{X_\alpha X_\beta}{D_{\alpha \beta}} \right) V_{\alpha j} \]  \hspace{1cm} (2.16)

\( D_{\alpha \beta} \) denotes the usual binary diffusion coefficient for two component mixing. The difficulty of computing diffusion coefficients, \( D_{\alpha \beta} \) (for \( N_{sp} > 2 \)) is avoided in this formulation. Even so, solving this system of equations is still computationally expensive [54].

In this work, Fick’s law is applied for computing diffusion velocities. Fick’s law provides close approximations at low computational cost when compared to the two methods that are explained above [25, 35, 65]. The diffusion coefficient \( D_{\alpha m} \) is assumed to be the proportionality factor between the diffusion velocity of each species and the gradient of species mole fraction in the diffusion direction.

\[ V_{\alpha j} = - \frac{D_{\alpha m}}{X_\alpha} X_{\alpha, j} \]  \hspace{1cm} (2.17)

This formulation can be obtained from the Stefan-Maxwell equation (2.16) assuming the body force per unit mass \( b_{\alpha j} \) to be equal for each species and the barotropic (pressure gradient driven) diffusion is negligible [128].

The multicomponent diffusion coefficient of each species in the mixture is computed applying the simple approximation suggested by Hirschfelder-Curtiss [1, 66].

\[ D_{\alpha m} = \frac{1 - Y_\alpha}{\sum_{\beta=1}^{N_{sp}} X_\beta / D_{\alpha \beta}} \]  \hspace{1cm} (2.18)

This approximation is obtained as follows :

- Relation between \( D_{\alpha \beta} \) and \( D_{\alpha m} \) is obtained using the following algebraic relation with the physical constraints \( \sum_\beta Y_\beta D_{\beta \alpha} = 0 \) for \( \alpha = 1, ..., N_{sp} \) to solve the Stefan-Maxwell system.

\[ \Delta D = I - \frac{Y \otimes U}{< U, Y >} \]  \hspace{1cm} (2.19)

where \( U = T \{ 1, 1, 1, ..., 1 \} \). Both \( \Delta \) and \( D \) are assumed to be symmetric and positive definite on \( U^\perp \) and \( Y^\perp \) respectively.
Here, the dual diffusion coefficients are approximated as the first term of the Sonine series expansion

$$\Delta_{\alpha\beta} = \frac{X_\alpha X_\beta}{D_{\alpha\beta}}$$  \hspace{1cm} (2.20)

- Later, this relation is substituted in equation (2.13) assuming pressure is constant, body force per unit mass for each species are equal and thermal diffusion is negligible.

- The approximation (2.18) is obtained substituting $D_{\alpha\beta}$ into Fick’s Law (2.17).

The binary ($D_{\alpha\beta} = D_{\beta\alpha}$) diffusion coefficients for species $\alpha$ and $\beta$ are defined in terms of temperature and pressure. The values are computed applying power law interpolation over the CHEMKIN model [73].

$$D_{\alpha\beta} = \frac{1}{P}a_{\alpha\beta0}T^{b_{\alpha\beta}} \hspace{1cm} \text{(m}^2/\text{s})$$ \hspace{1cm} (2.21)

The coefficients $a_{\alpha\beta0}$ and $b_{\alpha\beta}$ for the variation of $D_{\alpha\beta}$ with temperature at atmospheric pressure are tabulated in appendix A. The subscript 0 stands for atmospheric conditions.

The species mole fractions are considered as independent variables since the diffusion velocities are approximated using Fick’s law (2.17). However, this method causes violation of global mass conservation [54, 97]. The sum of species equations by definition must necessarily result in conservation of mass, in conservative formulation. This condition is not satisfied for Fick’s assumption, unless all diffusion coefficients are equal.

One approach for this problem is discarding one of the species equations from the system. The mass fraction of this final species is calculated fulfilling the sum of the species mass fractions to unity. Also, there are many reasons for the existence of inaccuracies in the solution of Navier-Stokes equations. This last species is chosen to be a diluent in order to absorb the inconsistencies. It is chosen as $N_2$ in the present work since it is an inert species. However, in case of diffusion flames all the species are deficient either in the fuel or oxidizer side. Since the mass fractions of each species are critical at shear layer, this method will result in loss of accuracy. Another approach for preventing the violation of conservation of mass is introducing an artificial correction velocity for all species, which forces the sum of diffusion fluxes to be zero.

$$V_j^c = -\sum_{\beta=1}^{N_{sp}} Y_\beta V_{\beta j}$$ \hspace{1cm} (2.22)

The correction velocity is related to the molecular diffusion. It is more convenient to write it on the right hand side of equation (2.10). The corrected mass flux is written as

$$J_{\alpha j}^c = \rho Y_\alpha \left(V_{\alpha j} + V_j^c\right)$$ \hspace{1cm} (2.23)
So, the final form of the species conservation equation is

\[
\frac{\partial \rho Y_\alpha}{\partial t} + \frac{\partial (\rho Y_\alpha u_j)}{\partial x_j} = -f_{\alpha j}^e + \dot{\omega}_\alpha
\]  

(2.24)

### 2.1.4 CHEMICAL KINETICS

The source term in (2.24) is calculated assuming finite rate chemistry, using Arrhenius laws for the reaction rates. The system of \(N_{\text{reac}}\) reversible chemical reactions occurring in the flow may be symbolized as follows

\[
\sum_{\alpha=1}^{N_{\text{reac}}} \nu'_{\alpha j} A_\alpha \rightleftharpoons \sum_{\alpha=1}^{N_{\text{reac}}} \nu''_{\alpha j} A_\alpha \quad j = 1, ..., N_{\text{reac}}
\]

(2.25)

where \(A_\alpha\) is the chemical symbol of species \(\alpha\), and \(\nu'\) and \(\nu''\) are the stoichiometric coefficients of reactants and products. The chemical scheme gives for each reversible reaction \(j\), the forward \(K_{fj}\) and backward \(K_{bj}\) reaction rates, as the coefficients \((A, b, E_{\text{act}})\) of the Arrhenius law

\[
K = AT^b \exp\left(-\frac{E_{\text{act}}}{RT}\right)
\]

(2.26)

These coefficients are related by the equilibrium constant \(K_{ej}\) of the reaction

\[
K_{ej} = \frac{K_{fj}}{K_{bj}} = \left(\frac{P_{\text{atm}}}{RT}\right)^{\sum_{\beta=1}^{N_{\text{reac}}} \nu'_{\beta j} Y_{\beta}^0} \left(\frac{\Delta S_j^0}{RT} - \frac{\Delta H_j^0}{RT}\right)
\]

(2.27)

In (2.26) \(E_{\text{act}}\) is the activation energy of the reaction, and \(R = 8314.51\) (J/K.kmol) is the universal gas constant. The production rate of species \(\alpha\) due to the \(N_{\text{reac}}\) reversible reactions is computed with the following relation.

\[
\dot{\omega}_\alpha = M_\alpha \sum_{j=1}^{N_{\text{reac}}} (\nu''_{\alpha j} - \nu'_{\alpha j}) \left[ K_{fj} \prod_{\beta=1}^{N_{\text{reac}}} \left(\frac{\rho Y_\beta}{M_\beta}\right)^{\nu'_{\beta j}} - K_{bj} \prod_{\beta=1}^{N_{\text{reac}}} \left(\frac{\rho Y_\beta}{M_\beta}\right)^{\nu''_{\beta j}} \right]
\]

(2.28)

Detailed chemistry has a valuable effect on flow simulation. However, each additional species increases the number of equations and the amount of data to handle. The reaction of H\(_2\)/air mixtures which may be considered to be well known [65] was studied in this work. The 7 species, 14 reactions mechanism of ONERA, which was tested by Davidenko [22], is used for the calculations. The mechanism with coefficients for equation (2.26) is tabulated in appendix B.
The momentum equation, which is the same for reacting and non-reacting cases, represents the dynamics of the flow, in the Navier-Stokes system

\[
\frac{\partial \rho u_i}{\partial t} + \frac{\partial}{\partial x_j} \left( \rho u_i u_j + p \delta_{ij} \right) = \rho f_i + \frac{\partial \tau_{ij}}{\partial x_j} \tag{2.29}
\]

Body forces \( f_i \) are neglected in this work. Pressure \( p \) is calculated applying Dalton’s law

\[
p = \sum_{\alpha=1}^{N_{sp}} p_{\alpha} \tag{2.30}
\]

### 2.2.1 THE EQUATION OF STATE

Each species satisfies the equation of state for ideal gases

\[
p_{\alpha} = \rho_{\alpha} \frac{R}{M_{\alpha}} T = \rho_{\alpha} r_{\alpha} T \tag{2.31}
\]

\( r_{\alpha} = R/M_{\alpha} \) (J.kg\(^{-1}\).K\(^{-1}\)) is the gas constant of species \( \alpha \). The mixture constant and its equation of state are derived using equations (2.7), (2.30) and (2.31).

\[
r = R \sum_{\alpha=1}^{N_{sp}} \frac{Y_{\alpha}}{M_{\alpha}} \quad ; \quad p = \rho RT \sum_{\alpha=1}^{N_{sp}} \frac{Y_{\alpha}}{M_{\alpha}} = \rho r T \tag{2.32}
\]

### 2.2.2 MOLECULAR DIFFUSION OF MOMENTUM

The stress tensor of fluid flow \( \sigma_{ij} \), in terms of the inviscid \( (p:\text{pressure}) \) and the viscous \( (\tau_{ij}) \) contributions, is

\[
\sigma_{ij} = -p \delta_{ij} + \tau_{ij} \tag{2.33}
\]

The viscous stress for a Newtonian fluid (the stress is linearly proportional to the strain rate) is assumed to be symmetric and may be written as

\[
\tau_{ij} = \lambda \delta_{ij} u_{k,k} + 2\mu S_{ij}^* \quad ; \quad S_{ij}^* = \frac{1}{2} \left( \frac{\partial u_i}{\partial x_j} + \frac{\partial u_j}{\partial x_i} \right) \tag{2.34}
\]

Stokes assumption which states that pressure forms the trace and viscous part forms the deviatoric part of the stress tensor \( (\tau_{ii} = 0) \) is applied, resulting in \( \lambda = -2/3\mu \)

\[
\tau_{ij} = \mu S_{ij} = \mu \left( u_{i,j} + u_{j,i} - \frac{2}{3} u_{k,k} \delta_{ij} \right) \tag{2.35}
\]
\( S_{ij} \) is one half of the rate of strain tensor \( S_{ij} \), for an incompressible flow. The viscosity \( \mu \) of the mixture is computed as a function of mole fractions and partial viscosities applying Wilke’s empirical formula [127]

\[
\mu = \sum_{\alpha=1}^{N_{sp}} \frac{X_{\alpha} \mu_{\alpha}}{\sum_{\beta=1}^{N_{sp}} X_{\beta} \phi_{\alpha\beta}} ; \quad \phi_{\alpha\beta} = \left( \frac{1 + \sqrt{\mu_{\alpha} / \mu_{\beta}}} { \mu_{\alpha} / \mu_{\beta}^{1/4} } \right)^2 \sqrt{8 \left( 1 + \frac{M_{\beta}}{M_{\alpha}} \right)}
\] (2.36)

Molecular viscosity \( \mu_{\alpha} \) of species \( \alpha \) are computed using second order polynomials depending on temperature which were generated over the CHEMKIN model

\[
\mu_{\alpha} = \mu_{\alpha 0} + \mu_{\alpha 1} T + \mu_{\alpha 2} T^2 \quad \text{(kg/m.s)}
\] (2.37)

The coefficients \( \mu_{\alpha 0}, \mu_{\alpha 1} \) and \( \mu_{\alpha 2} \) for each species \( \alpha \) are tabulated in appendix A.

### 2.3 ENERGY EQUATION

The energy equation may be written in terms of total energy, internal energy, enthalpy or temperature. Total energy is chosen as the variable of the energy equation since this is the only variable to write the energy equation in conservation form. In total energy, the potential part is neglected

\[
e_t = e + \frac{1}{2} u_i u_i = h - \frac{p}{\rho} + k
\] (2.38)

Here, \( e \) is the internal energy, \( h \) is the enthalpy and \( k \) is the kinetic energy of the mixture per unit mass. The total energy equation for an infinitesimal control volume, neglecting the work done by body forces and assuming there is no heat source term (i.e. electric spark, magnetic energy ...) is as follows

\[
\frac{\partial p e_t}{\partial t} + \frac{\partial}{\partial x_j} \left[ u_j (p e_t + p) \right] = -\frac{\partial q_j}{\partial x_j} + \frac{\partial \tau_{ij} u_i}{\partial x_j}
\] (2.39)

#### 2.3.1 INTERNAL ENERGY, ENTHALPY AND SPECIFIC HEATS

Internal energy per unit mass \( e_\alpha \), enthalpy per unit mass \( h_\alpha = e_\alpha + p_\alpha / \rho_\alpha \), and the specific heats \( C_{p\alpha} \) and \( C_{v\alpha} \) of species \( \alpha \) depend on temperature only, according to the perfect gas assumption

\[
d e_\alpha = C_{v\alpha}(T) dT \quad \text{(2.40)}
\]
\[
d h_\alpha = C_{p\alpha}(T) dT \quad \text{(2.41)}
\]
The enthalpy of species $\alpha$ is obtained by integration of equation (2.41) from a reference temperature $T_0$. This represents the summation of the sensible enthalpy $h_{s\alpha}$ and of the enthalpy of formation $\Delta h^0_{f,\alpha}$ where superscript $0$ represents reference conditions

$$h_{\alpha}(T) = h_{s\alpha}(T) + \Delta h^0_{f,\alpha} = \int_{T_0}^{T} C p_{\alpha}(\theta)d\theta + \Delta h^0_{f,\alpha}$$  \hspace{1cm} (2.42)

The specific heats of species satisfy Mayer relation

$$r_{\alpha} = C p_{\alpha}(T) - C v_{\alpha}(T)$$  \hspace{1cm} (2.43)

and the ratio of specific heats depends on temperature

$$\gamma_{\alpha}(T) = \frac{C p_{\alpha}(T)}{C v_{\alpha}(T)}$$  \hspace{1cm} (2.44)

Specific heats of the mixture are approximated with mass weighted formulas

$$C p(Y_{\alpha}, T) = \sum_{\alpha=1}^{N_{sp}} C p_{\alpha}(T)Y_{\alpha} ; \quad C v(Y_{\alpha}, T) = \sum_{\alpha=1}^{N_{sp}} C v_{\alpha}(T)Y_{\alpha}$$  \hspace{1cm} (2.45)

Specific heats $C p_{\alpha}$ are tabulated as fourth order polynomials (CHEMKIN format tables), for two different temperature ranges of ($T_{min}$-1000) and (1000-$T_{max}$). The data for specific heats at constant pressure $C p_{\alpha}$ (J/kg.K) and enthalpy of formation $\Delta h^0_{f,\alpha}$ taken from Burcat polynomial database [14], are presented in Appendix B.2. The speed of sound in the mixture is

$$c^2 = \gamma r T = \gamma \frac{p}{\rho}$$  \hspace{1cm} (2.46)

The internal energy and enthalpy of the mixture are computed by mass weighted averaging

$$e = \sum_{\alpha=1}^{N_{sp}} e_{\alpha} Y_{\alpha} ; \quad h = \sum_{\alpha=1}^{N_{sp}} h_{\alpha} Y_{\alpha}$$  \hspace{1cm} (2.47)

The final form of the total energy in equation (2.39) is

$$e_t = \sum_{\alpha=1}^{N_{sp}} \left( \Delta h^0_{f,\alpha} + \int_{T_0}^{T} C p_{\alpha}(\theta)d\theta \right) Y_{\alpha} - r T + \frac{1}{2} u_i u_i$$  \hspace{1cm} (2.48)

### 2.3.2 MOLECULAR DIFFUSION OF HEAT AND ENTHALPY

Kinetic theory suggests the heat flux expression $\dot{q}_{ji}$, in direction $x_j$ as

$$\dot{q}_{ji} = \sum_{\alpha=1}^{N_{sp}} h_{\alpha} J_{ij}^\alpha - \lambda T_{ij} - p \sum_{\alpha=1}^{N_{sp}} D_{ij}^\alpha d_{ij}$$  \hspace{1cm} (2.49)
where $d_{\alpha j}$ is the diffusion driving force of specie $\alpha$ given in equation (2.14). The terms of equation (2.49), from left to right, are:

- Partial enthalpy flux of specie $\alpha$.
- Heat transfer by conduction computed by Fourier law. $\lambda$ is the coefficient of thermal conductivity. It is approximated like $\mu$ in equation (2.36). Also, the coefficient of partial conductivity $\lambda_\alpha$ of species $\alpha$ is computed over CHEMKIN model as

$$
\lambda_\alpha = \lambda_{\alpha 0} + \lambda_{\alpha 1} T + \lambda_{\alpha 2} T^2 \quad \text{(W/m.K)}
$$

(2.50)

The constants $\lambda_{\alpha 0}$, $\lambda_{\alpha 1}$ and $\lambda_{\alpha 2}$ are tabulated in appendix A.

- The third term, which is assumed to be negligible compared to conductive heat flux, refers to the heat transfer that is induced by concentration gradients (Dufour effect) [36].

### 2.3.3 TEMPERATURE EVALUATION

Since the species enthalpy $h_\alpha$ in total energy depends on temperature non-linearly, analytical derivation is not possible. Temperature field may be computed from the total energy $e_t$, iteratively. Newton-Raphson iteration is applied for temperature calculations [59, 80].

$$
f(T) = e_t - \left( \sum_{\alpha=1}^{Nsp} h_\alpha Y_\alpha - rT + \frac{1}{2} \mu_i u_i \right)
$$

(2.51)

Given a value for $e_t$, temperature is iterated for convergence of $f(T)$ to zero

$$
T^{it+1} = T^{it} - \frac{f(T^{it})}{f'(T^{it})}
$$

(2.52)

where $it$ is the iteration step. The temperature iteration is initialized with the value from the previous step.

### 2.4 SUMMARY OF SYSTEM OF EQUATIONS

In summary, the system equations which serves as the basis for the simulations presented in this work is

$$
\frac{\partial \vec{U}}{\partial t} + \frac{\partial \vec{F}}{\partial x} + \frac{\partial \vec{G}}{\partial y} + \frac{\partial \vec{H}}{\partial z} = \vec{V} + \vec{S}
$$

(2.53)
The vector of dependent variables $\tilde{U}$, the convective flux vectors $\tilde{F}$, $\tilde{G}$, $\tilde{H}$, the vector of viscous stress $\tilde{V}$ and the vector of source terms $\tilde{S}'$ are given as follows

\[
\tilde{U} = \begin{bmatrix}
\rho \\
\rho u \\
\rho v \\
\rho w \\
\rho e_t \\
\rho Y_1 \\
\vdots \\
\rho Y_{N_{sp}-1}
\end{bmatrix}, \quad \tilde{F} = \begin{bmatrix}
\rho u \\
\rho u^2 + p \\
\rho u v \\
\rho u w \\
(p e_t + p)u \\
\rho u Y_1 \\
\vdots \\
\rho u Y_{N_{sp}-1}
\end{bmatrix}, \quad \tilde{G} = \begin{bmatrix}
\rho v \\
\rho v u \\
\rho v^2 + p \\
\rho v w \\
(p e_t + p)v \\
\rho v Y_1 \\
\vdots \\
\rho v Y_{N_{sp}-1}
\end{bmatrix}, \quad \tilde{H} = \begin{bmatrix}
\rho w \\
\rho w u \\
\rho w v \\
\rho w w \\
(p e_t + p)w \\
\rho w Y_1 \\
\vdots \\
\rho w Y_{N_{sp}-1}
\end{bmatrix}
\]

\[
\tilde{V} = \begin{bmatrix}
0 \\
0 \\
0 \\
0 \\
\tau_{xx,xx} + \tau_{xy,y} + \tau_{xz,z} \\
\tau_{yx,xx} + \tau_{yy,y} + \tau_{yz,z} \\
\tau_{zx,xx} + \tau_{zy,y} + \tau_{zz,z} \\
-\rho_{x,xx} + (\rho u_{xx})_x + (\rho v_{xx})_x + (\rho w_{xx})_x \\
-\rho_{y,xy} + (\rho u_{xy})_y + (\rho v_{xy})_y + (\rho w_{xy})_y \\
-\rho_{z,zx} + (\rho u_{zx})_z + (\rho v_{zx})_z + (\rho w_{zx})_z \\
-J^c_{1,xx} - J^c_{1,yy} - J^c_{1,zz} \\
\vdots
\end{bmatrix}
\]

\[
\tilde{S}' = \begin{bmatrix}
0 \\
0 \\
0 \\
0 \\
0 \\
\omega_a \\
\vdots \\
\omega_{N_{sp}-1}
\end{bmatrix}
\] (2.54)
CHAPTER 3

LARGE EDDY SIMULATION

Turbulent flow problems in the context of this thesis are at high Reynolds numbers and the range of flow scales is fairly wide. Resolution of all scales in realistic or industrial configurations is impossible and it seems to remain so in near future despite the drastic improvement in computational capabilities. Large Eddy Simulation (LES) is suitable and affordable for the simulation of the problems concerned.

Before putting forward the arguments that support the idea of LES one may ask the question: what is being resolved during LES calculations? As Pope [107] does, a clear distinction between real and resolved flow fields have to be carried out. When a field of any turbulent flow variable $U(x,y,z,t)$ is considered, the resolved field of this variable $W(x,y,z,t)$ is the numerical solution of the set of filtered Navier-Stokes equations over a grid. In the context of LES, $W$ is intended to be statistically related to the filtered field $\bar{U}$ which is obtained by applying a low pass filter of characteristic width $\Delta$ to $U$. As the filter width is reduced down to the smallest scales (Kolmogorov microscale $\eta$), the statistics of the solution obtained by applying an accurate numerical method and proper boundary conditions converge to a condition where all energy containing contribution may be represented by $W$. This is the DNS level which is obtained when the oscillations with smaller wavelengths than the grid cut-off have no energy. The statistics at DNS level match those of $\bar{U}$. The resolved variables for LES are commonly considered as "filtered" forms of the solution at the DNS level, in a priori tests. This filter is not precisely defined for numerical schemes having dissipation property, linked to the resolving efficiency. In appropriate LES the statistics achieve an intermediate asymptote as $\Delta$ is in the inertial sub-range.

After four decades of intensive developments [87, 109, 110, 115] and from a theoretical point of view, LES is now a well-established technique for incompressible flows. Intricate interac-
tions between numeric and sub-grid modeling are well understood in academic configurations [43, 52, 76]. Multilevel/multiresolution approaches like Detached-Eddy Simulations (DES) have been developed (see [111] for review), allowing robust and efficient simulations for industrial configurations. Nevertheless, certain aspects of LES such as those in complex physics-shocked flows, multi-component, multiphase, reacting flows, or in complex geometries, remain unclear despite some outstanding achievements [9, 13]. Following S.B. Pope [107], one should first make the distinction between pure physical LES and pure numerical LES.

In the former, filtered equations are solved for the large scales of motion, and the effect of the unresolved turbulent scales, or sub-grid scales (SGS) on the resolved ones is taken into account via a physically sound explicit sub-grid model. Higher-order non-dissipative numerics are needed to avoid interactions between numerical errors and sub-grid modeling. Solving the filtered Navier-Stokes equations in Fourier space with spectral methods is ideal for academic flows. In this case, the low-pass filter is a sharp Fourier cutoff, and spectral sub-grid eddy viscosity closures can be derived from analytical theories [17, 96]. This ideal situation is forcefully limited to homogeneous turbulence in cubic computational domains. High accuracy can also be achieved in physical space on more general configurations (though still very simple) with higher-order centered compact finite difference schemes [83] that include Structure Function models, either standard [96], selective [21] or filtered [30]. Nevertheless, none of these methods can handle sharp gradients or shocked flows without any numerical filtering or artificial viscosity, which already raises the question of competition between numerics and models. If multi-species, reacting variable density flows are considered, the situation is more complicated.

In pure numerical LES, shock-capturing methods are used to solve the Navier-Stokes or Euler equations without any explicit filtering. There is no explicit sub-grid model: small scale fluctuations are damped by the numerical diffusion. This approach which is denoted -highly controversial in the combustion community- is called implicit LES (ILES), or Monotone Integrated LES (MILES) [60, 91]. High-resolution methods are required to avoid a significant impact of the numerics on the turbulent flow statistics. An evaluation of the effects of numerical diffusion on the DNS and LES of compressible turbulence (non-reacting) has been done by E. Garnier [46, 47] for mainly academic demonstrational flows. Several shock-capturing schemes were tested, and the 5th order WENO scheme [69, 88, 112] was found to perform well. This approach has been widely used since. A recent successful MILES has been re-
ported for the simulation of multicomponent Richtmyer-Meshkov turbulent mixing carried out with the Euler equations [119], surprisingly, there are very few attempts to apply this technique to reacting flows [15, 117].

Intermediate between pure physical and pure numerical LES are all other situations, where both explicit sub-grid modeling and robust dissipative schemes are used—although this was not recommended by E. Garnier [47]. Many, if not all, LES of practical (or industrial) interest are carried out that way on irregular or even unstructured grids, with old RANS solvers upgraded for SGS modeling. Hybrid methods have also been developed (e.g. [31, 56]), where the flow solver switches from a second or a fourth-order central scheme away from discontinuities to a shock-capturing scheme, depending on an empirically designed stiffness sensor. Whatever the method, in most of practical physical LES, some SGS terms are systematically neglected, or crudely modeled. As such, irregular grids induce a second-order commutation error [53], and the physically sound part of the SGS model that mimics the small-scale mixing and scalar dissipation rate interacts with the numerical diffusion: both (grid and SGS model) smooth out the flow field. In summary, firstly, filtered conservative variables are introduced which are so-called resolved variables. Then, the set of governing equations satisfied by filtered variables are presented. Finally, two different physical LES modeling strategies are explained, and the dissipation and dispersion characteristics of different schemes for numerical LES are discussed.

3.1 FILTERING THE NAVIER-STOKES EQUATIONS

A filter operation makes the distinction between the large scales \( \bar{f} \) and the small scales \( f' \) of a continuous function \( f \), i.e. \( f = \bar{f} + f' \). It is considered that the projection of function \( f \) on a coarse grid acts as an implicit filtering operation, due to the Nyquist cut-off wavelength.

The filtered part for a homogeneous filter may be standardly defined as

\[
\bar{f}(x) = \int_{\Omega} G_{\Delta}(x-\xi) f(\xi) d\xi
\]  

(3.1)

where \( x \) and \( \xi \) are coordinates in the domain \( \Omega \) and \( \Delta \) is the “filter width” associated with the filter kernel \( G_{\Delta} \) which necessarily satisfies
\[ \int_{\Omega} G_\Delta(\xi) \, d\xi = 1 \quad (3.2) \]

Filtering and derivative commute if the filter is homogeneous [53, 122].

The multicomponent compressible LES equations are similar to the mono-species form given by C.G. Speziale and co-workers [34, 116], but thermodynamics is different. As in the mono-species case, filtered multicomponent equations can be cast in different ways, leading to different SGS modeling strategies ([84, 122, 123] and Garnier et al. [48] for a review). The derivations done in several ways are presented in the literature with more or less details [56, 104, 121]. The way to display LES mathematically is to define the resolved variables (computable quantities) then state the governing equations with separate resolved and SGS terms. This separation enables introduction of models for SGS terms. Primary computable quantities are the resolved field variables \( \vec{W} \) advanced in time.

\[ \vec{W} = T \{ W_1, W_2, \ldots, W_{5+N_p} \} \quad (3.3) \]

In the compressible formulation, if a Reynolds-like decomposition is introduced, filtering the continuity equation (2.12) will lead to an uncomputable variable \( \overline{\rho u_i} \).

\[ u_i = \overline{u_i} + u_i' \quad \rho = \overline{\rho} + \rho' \quad \Rightarrow \quad \overline{\rho u_i} = \overline{\rho} \overline{u_i} + (\overline{\rho u_i} - \overline{\rho} \overline{u_i}) \rightarrow \text{Leonard term} \]
\[ + (\overline{\rho'} u_i' + \rho' \overline{u_i'}) \rightarrow \text{cross term} \]
\[ + \rho' u_i' \rightarrow \text{Reynolds term} \quad (3.4) \]

The Reynolds term in RANS formulation and all three extra terms in LES formulation are non-zero so that \( \overline{\rho u_i} \neq \overline{\rho u_i} \). Favre introduced the concept of mass weighted filtering for compressible RANS equations [39, 40].

\[ u_i = \overline{u_i} + u_i'' \quad \overline{u_i} = \frac{\overline{\rho u_i}}{\overline{\rho}} \quad (3.5) \]

Favre variables are only mathematical definitions which eliminate some of the SGS terms in the LES equations. Despite the loss of physics, Favre variables introduce mathematical simplicity. Theoretically, the set of NS equations (2.53) are filtered and Favre variables are introduced in these equations. The variables obtained solving the filtered set of equations also
include numerical errors.

\[
\begin{bmatrix}
\bar{p} \\
\bar{\rho} u \\
\bar{\rho} v \\
\bar{\rho} w \\
\bar{\rho} e_t \\
\bar{\rho} Y_1 \\
\vdots \\
\bar{\rho} Y_{N_{\rho}-1}
\end{bmatrix} = 
\begin{bmatrix}
p \\
p u \\
p v \\
p w \\
p e_t \\
p Y_1 \\
\vdots \\
p Y_{N_{\rho}-1}
\end{bmatrix}
\]

Given the \( W^j \)'s one can obtain directly the Favre-filtered values \( \bar{u}_i, \bar{e}_t \) and \( \bar{Y}_a \) which are secondary computable quantities. However, temperature and pressure fields are also needed during the simulation. Temperature may be deduced from \( W_5 \) and pressure through ideal gas equation of state. Particular attention will be given on this term at the end of this section.

Filtered equations with Favre variables are presented hereafter. The filtered quantities \( \bar{\phi} \) or \( \tilde{\phi} \) are split into the sum of a part \( \hat{\phi} \) computable out of primary or secondary quantities plus an uncomputable part \( \tilde{\phi} \) (SGS).

\[
\bar{\phi} = \hat{\phi} + \tilde{\phi} \tag{3.6}
\]

### 3.1.1 CONSERVATION OF MASS

Consider the continuity equation (2.12) in conservative form, filtered with a linear filter which may commute with derivatives

\[
\frac{\partial \bar{\rho}}{\partial t} + (\bar{\rho} \bar{u}_j)_j = 0 \tag{3.7}
\]

The convective term \( (\bar{\rho} \bar{u}_j)_j \) is uncomputable so Favre variable is introduced for a straightforward filtered equation (3.7)

\[
\frac{\partial \bar{W}_1}{\partial t} + (\bar{\rho} \bar{u}_j)_j = 0 \tag{3.8}
\]

Since Favre filtering does not commute with derivatives, Favre variables are introduced after commuting the “bar-filter” with partial differential operator.
3.1.2 CONSERVATION OF CHEMICAL SPECIES

The species conservation equation (2.24) after filtering operation is
\[
\frac{\partial \rho Y_\alpha}{\partial t} + \left(\rho u_j Y_\alpha\right)_j = -J_{\alpha j} + \dot{\omega}_\alpha
\]  
(3.9)

Introducing Favre variables into (3.9) yields
\[
\frac{\partial W_{S\alpha}}{\partial t} + \left(\bar{\rho} \bar{u}_j \bar{Y}_\alpha\right)_j = -J_{\alpha j}^c + \dot{\omega}_\alpha - \left[\bar{\rho} \left(\bar{u}_j Y_\alpha - \bar{u}_j \bar{Y}_\alpha\right)\right]_j
\]  
(3.10)

The filtered corrected species mass flux term can be split into computable and uncomputable components, as explained in relation (3.6)
\[
J_{\alpha j} = J_{\alpha j}^c + J_{\alpha j}^u
\]  
(3.11)

The resolved corrected mass flux $J_{\alpha j}^c$ of species $\alpha$ in the direction $x_j$ is computed from equations (2.17)- (2.23) using resolved variables.
\[
\begin{align*}
J_{\alpha j}^c &= \bar{\rho} \left(D_{m} \bar{X}_{\alpha,j} \frac{M_\alpha}{M} + \bar{V}_j \bar{Y}_\alpha\right) \\
J_{\alpha j}^u &= \rho \left(D_{m} \bar{X}_{\alpha,j} \frac{M_\alpha}{M} + \bar{V}_j \bar{Y}_\alpha\right) - \bar{\rho} \left(D_{m} \bar{X}_{\alpha,j} \frac{M_\alpha}{M} + \bar{V}_j \bar{Y}_\alpha\right)
\end{align*}
\]  
(3.12)

So the filtered species conservation equation reads
\[
\frac{\partial W_{S\alpha}}{\partial t} + \left(\bar{\rho} \bar{u}_j \bar{Y}_\alpha\right)_j = -J_{\alpha j}^c + \dot{\omega}_\alpha + \left(C_{1\alpha j} + C_{2\alpha j}\right)_j + C_{3\alpha}
\]  
(3.14)

where
\[
\begin{align*}
C_{1\alpha j} &= -\bar{\rho} \left(\bar{u}_j Y_\alpha - \bar{u}_j \bar{Y}_\alpha\right) \\
C_{2\alpha j} &= -J_{\alpha j}^u \\
C_{3\alpha} &= \bar{\dot{\omega}}_\alpha
\end{align*}
\]  
(3.15)

In a priori LES tests presented in appendix H it has been shown that the $C_{2\alpha j}$ term is generally negligible. On the other hand, the $C_{3\alpha}$ term is the cornerstone of reacting LES [42, 80, 81].

3.1.3 EQUATION OF MOTION

Filtering equation of motion (2.29) and commuting the filter with partial derivative gives
\[
\frac{\partial \rho u_i}{\partial t} + \left(\bar{\rho} \bar{u}_i \bar{u}_j\right)_j = -\bar{p}_j + \bar{\tau}_{ij,j}
\]  
(3.18)
Filtered viscous stress and pressure need special attention.

In the viscous stress, viscosity is not constant

\[ \mu = \mu(Y, T) \] (3.19)

As a result, the filtered viscous stress is uncomputable and may be split into uncomputable and computable parts

\[ \tau_{ij} = \hat{\tau}_{ij} + \bar{\tau}_{ij} \] (3.20)
\[ \hat{\tau}_{ij} = \mu \hat{S}_{ij} \quad ; \quad \bar{\mu} = \mu(\hat{Y}, \hat{T}) \] (3.21)
\[ \bar{\tau}_{ij} = \mu \bar{S}_{ij} - \bar{\mu} \bar{S}_{ij} \] (3.22)

where \( \hat{T} \) is the computable temperature (will be explained in detail in the section for \( W_5 \)) and the filtered strain tensor for compressible flow is

\[ \bar{S}_{ij} = \bar{u}_{i,j} + \bar{u}_{j,i} - \frac{2}{3} \bar{u}_{k,k} \delta_{ij} \] (3.23)

Since the pressure is calculated using the temperature via the equation of state, it has an uncomputable part like the temperature. Specific to multicomponent flows, the uncomputable filtered pressure reads

\[ \bar{p} = \bar{p} + \bar{\rho} \] (3.24)
\[ \bar{\rho} = \bar{\rho} \hat{T} = \rho \left( R \sum_{a=1}^{N_a} \frac{1}{W_a} \hat{Y}_a \right) \hat{T} \] (3.25)
\[ \bar{\rho} = \bar{p} (\hat{T} - \bar{T}) \] (3.26)

Considering (3.20) and (3.24), the Favre variable introduced equation of motion yields

\[ \frac{\partial W_{1+}}{\partial t} + \left( \bar{p} \bar{u}_i \bar{u}_j + \bar{\rho} \delta_{ij} \right)_{,i} = \hat{\tau}_{ij} + \left( A_{1ij} + A_{2ij} + A_3 \delta_{ij} \right)_{,j} \] (3.27)

\[ A_{1ij} = -\bar{\rho} \left( \bar{u}_i \bar{u}_j - \bar{u}_i \bar{u}_j \right) \] (3.28)
\[ A_{2ij} = \hat{\tau}_{ij} \] (3.29)
\[ A_3 = -\bar{\rho} \] (3.30)

The \( A_{1ij} \) term is the most important one and is common to mono-species compressible LES equations. It reduces to Germano’s central moments [51] in the incompressible limit. The \( A_{2ij} \) term appears because the filtered viscous stress is uncomputable as explained and neglected in most compressible LES [122, 123]. It was shown in \textit{a priori} tests presented in appendix H that the \( A_{3,i} \) term may overcome \( A_{1ij,j} \) in some cases. This is never considered in any multicomponent, non-reacting or reacting LES.
3.1.4 ENERGY EQUATION

Filtering and commuting the filter with derivative for the energy equation (2.39) gives

$$\frac{\partial \bar{\rho} \bar{e}_i}{\partial t} + (\bar{\rho} u_j) \frac{\partial \bar{e}_i}{\partial x_j} + (\bar{\rho} u_j) \frac{\partial \bar{u}_i}{\partial x_j} = -\tau_{ij} - (\tau_{ij} u_i)$$  \hspace{1cm} (3.31)

Then, introducing Favre variables in the filtered total energy equation (3.31) yields

$$\frac{\partial W_5}{\partial t} + \left[ (\bar{\rho} \bar{e}_i + \bar{\rho}) \bar{u}_j \right] = \left( \bar{u}_i \tau_{ij} + \bar{u}_i \tilde{\tau}_{ij} \right) - \bar{q}_j + \left( B_{1j} + B_{2j} + B_{3j} + B_{4j} \right)$$  \hspace{1cm} (3.32)

$$B_{1j} = -\bar{\rho} \left( \bar{e}_i \bar{u}_j - \bar{\tau}_{ij} \right)$$  \hspace{1cm} (3.33)

$$B_{2j} = -\left( \bar{\rho} u_j - \bar{\rho} \bar{u}_j \right)$$  \hspace{1cm} (3.34)

$$B_{3j} = \bar{u}_i \tilde{\tau}_{ij} - \bar{u}_i \tilde{\tau}_{ij}$$  \hspace{1cm} (3.35)

$$B_{4j} = -\bar{q}_j$$  \hspace{1cm} (3.36)

Equation (3.32) and term $B_{4j}$ involve respectively the resolved and SGS parts of the uncomputable filtered heat flux. As explained in section 2.3.3, temperature can not be deduced directly but it is computed out of $e_i$

$$\bar{q}_j = \bar{\dot{q}}_j + \bar{q}_j$$  \hspace{1cm} (3.37)

$$\bar{\dot{q}}_j = -\bar{\lambda} \bar{T}_{,j} + \sum \bar{h}_a \bar{f}_{aj}$$  \hspace{1cm} (3.38)

$$\bar{q}_j = -\left( \bar{\lambda} \bar{T}_{,j} - \bar{\lambda} \bar{T}_{,j} \right) + \sum \left( \bar{h}_a \bar{f}_{aj} - \bar{h}_a \bar{f}_{aj} \right)$$  \hspace{1cm} (3.39)

In (3.38), $\bar{\lambda}$ is uncomputable and is obtained the same way as (3.19).

3.1.5 SPECIAL ATTENTION FOR $W_5$ AND IDEAL GAS EQUATION

The total energy is used for construction of the conservative variable in the energy equation. Temperature and pressure are computed from this term. Hence the $W_5$ term deserves particular attention.

$$\bar{\rho} \bar{e}_i = \rho \left( e + \frac{1}{2} u_i u_i \right)$$

$$= \bar{\rho} \bar{h} - \bar{\rho} + \bar{\rho} k$$  \hspace{1cm} (3.40)
The filtered pressure, the Favre-filtered static enthalpy and the Favre-filtered kinetic energy per unit mass are uncomputable.

\[ \rho e_t = \tilde{\rho}e_t = \tilde{\rho} \hat{e}_t + \tilde{\rho} \check{e}_t \]  
\[ \tilde{\rho} \hat{e}_t = \tilde{\rho} \hat{h} - \hat{p} + \tilde{\rho} \hat{k} \]  
\[ \tilde{\rho} \check{e}_t = \tilde{\rho} \check{h} - \check{p} + \tilde{\rho} \check{k} \]  

(3.41)  
(3.42)  
(3.43)

The split uncomputable pressure is given in equation (3.24). The Favre-filtered static enthalpy is written as

\[ \tilde{h} = \hat{h} + \check{h} \]  
\[ \hat{h} = \sum \hat{h}_a \tilde{Y}_a \]  
\[ \check{h} = \sum \left( \tilde{h}_a Y_a - \hat{h}_a \tilde{Y}_a \right) \]  

(3.44)  
(3.45)  
(3.46)

where

\[ \hat{h}_a = \Delta h_0 + \int_{T_0}^{\hat{T}} C_{p_{ol}}(\theta) d\theta \]  

(3.47)

The Favre-filtered kinetic energy is:

\[ \tilde{k} = \hat{k} + \check{k} \]  
\[ \hat{k} = \frac{1}{2} \tilde{u}_i \tilde{u}_i \]  
\[ \check{k} = \frac{1}{2} (\tilde{u}_i \tilde{u}_i - \tilde{u}_i \tilde{u}_i) \equiv k_{sgs} \]  

(3.48)  
(3.49)  
(3.50)

In equation (3.25), and at the upper integration limit in (3.47), the resolved temperature \( \hat{T} \) has been used instead of the Favre-filtered temperature \( \tilde{T} \). Given numbers for the \( W_j^s \), \( \tilde{T} \) could theoretically be obtained by finding iteratively (e.g. via a Newton-Raphson procedure) the value which produces a posteriori the proper \( W_5 \) value from equation (3.40). In practice, since (3.43) is seldom (if ever) explicitly modeled in multicomponent LES, only the analytic expression of the computable part \( \tilde{\rho} \hat{e}_t \), equation (3.42), is used to find the resolved temperature \( \hat{T} \) such that

\[ W_5 = \tilde{\rho} \left[ \sum a \left( \Delta h_0 + \int_{T_0}^{\hat{T}} C_{p_{ol}}(\theta) d\theta \right) \tilde{Y}_a - \tilde{T} \hat{k} + \frac{1}{2} \tilde{u}_i \tilde{u}_i \right] \]  

(3.51)

So, unless the uncomputable total energy per unit volume \( \tilde{\rho} \hat{e}_t \) is properly modeled, the Favre-filtered temperature is uncomputable.
3.2 EXPLICIT SUB-GRID MODELING

In the present study, as in most physical compressible/multicomponent LES, the SGS terms which do not emanate from convective parts of hyperbolic governing equations \( A_{2ij}, A_3, B_{3j}, B_{4j} \) and \( C_{2ai} \) will not be explicitly modeled. One can consider that they are simply neglected, or put together with the dominant dynamic terms \( A_{1ij}, B_{1j} \) and \( C_{1ai} \). For the main \( A_{1ij} \) sub-grid term in equation (3.27), a classical sub-grid eddy-viscosity assumption is made for the deviatoric part of the SGS tensor

\[
A_{1ij}^D = A_{1ij} - \frac{1}{3} A_{1kk} \delta_{ij} = \mu_{sgs}' S_{ij}
\]  

so that

\[
\bar{\rho} (\bar{u}_i \bar{u}_j - \bar{u}_i \bar{u}_j) - \frac{2}{3} \bar{\rho} k_{sgs} \delta_{ij} = -\mu_{sgs}' S_{ij}
\]  

Two SGS models are implemented: the Smagorinsky (SM) model [115] extended to compressible flows, and the Selective Structure Function (SSF) model [21, 96].

3.2.1 CLOSING THE FILTERED EQUATION OF MOTION

3.2.1.1 THE SMAGORINSKY MODEL

The first known application of the idea of physical LES is by Joseph Smagorinsky [115]. The turbulent viscosity \( \mu_{sgs}' \) is dimensionally equal to \( \bar{\rho} u l \). The velocity may be approximated by assuming the energy dissipation is equal to the energy transferred to small scales, with no backscattering.

\[
\epsilon \approx \bar{\rho} \frac{u^3}{l} \approx \frac{u^3}{\Delta}
\]  

In this relation, the characteristic length is taken as the filter width \( \Delta \)

\[
\mu_{sgs}' = \bar{\rho} \left( \frac{\Delta \epsilon}{\rho} \right)^{1/3} \Delta
\]  

The main assumption for the model states that production \( P_{k_{sgs}} \) is equal to dissipation for the sub-grid kinetic energy

\[
P_{k_{sgs}} = A_{1ij}^D \bar{u}_i \bar{u}_j = \mu_{sgs}' S_{ij} \bar{u}_i \bar{u}_j
\]  

When one substitute the production into equation (3.55) for dissipation, the turbulent viscosity reads

\[
\mu_{sgs}' = \bar{\rho}^{2/3} \left( \mu_{sgs}' S_{ij} \bar{u}_i \bar{u}_j \right)^{1/3} \Delta^{4/3}
\]  

27
The usual form of the compressible SM model is [34, 49]

\[
\mu'_{sgs} = \rho(C_S \Delta^2 |\tilde{S_*}|) \tag{3.58}
\]

\[
|\tilde{S_*}| = \left( 2 \tilde{S_{ij}} \tilde{S_{ij}} \right)^{1/2} \tag{3.59}
\]

It is common to apply the equation (3.59) for construction of SM model which is only valid for incompressible case. However, considering the sub-grid scale production term \(A_{ij}^D \tilde{\mu}_{i,j} \) [56], the traceless closure (3.53) leads rather to

\[
|\tilde{S_*}| = \left( \tilde{S_{ij}} \tilde{S_{ij}} \right)^{1/2} \tag{3.60}
\]

This is the expression implemented in the present study. In the SM model, the filter width, or turbulence resolution length scale \( \Delta \), is multiplied by the model constant. The product \( C_S \Delta \) can be written as \( \lambda \Delta c \) where \( \Delta c = (\Delta x_1 \Delta x_2 \Delta x_3)^{1/3} \) is the geometrical mean local grid resolution. Various derivations which result in different values for the constant \( C_S \) are possible (p.587 in [106]). In the numerical experiments presented in section 5.1, simulations with the SM model are done for \( \lambda^2 = 0.01 \) and \( \lambda^2 = 0.02 \). Taking the constant \( C_S = 0.1 \), this may be interpreted as varying the numerical accuracy from \( \Delta/h = 1 \) to \( \Delta/h = \sqrt{2} \). This is the usual range for physical LES using dissipative numerical schemes (question 7 in [107]).

### 3.2.1.2 THE YOSHIZAWA MODEL

The isotropic part

\[
A'_{1ij} = \frac{1}{3} A_{1kk} \delta_{ij} = -\frac{2}{3} \rho k_{sgs} \delta_{ij} \tag{3.61}
\]

can be represented by the Yoshizawa model [129] as

\[
\bar{\rho} k_{sgs} = \bar{\rho} C_I \Delta^2 |\tilde{S_*}|^2 \tag{3.62}
\]

However, it may be neglected if the sub-grid Mach number \( M_{sgs}^2 = \bar{\rho} k_{sgs} / \bar{\gamma} p \) is small [86, 34]. The sub-grid Mach number will be verified \textit{a priori} later in section 5.1.5.
3.2.1.3 THE SELECTIVE STRUCTURE FUNCTION MODEL

The SSF model is based on the standard structure function (SF) model \[85\] for $\mu_{sgs}^j$ in $A_{ij}^D$. In the SF model, the second order velocity structure function $F_2(\vec{x}, \Delta)$ is used

$$F_2(\vec{x}, \Delta x) = \left\langle \left\| \vec{\tilde{u}}(\vec{x}) - \vec{\tilde{u}}(\vec{x} + \vec{r}, t) \right\|^2 \right\rangle_{||\vec{r}||=\Delta x}$$ \hspace{1cm} (3.63)

In practice, the velocity structure function is computed over differences in the resolved velocity field at adjacent grid points. The SF on uniform grid $\Delta x_i = \text{Const.}$ in direction $x_i$ with unit vector $\vec{e}_i$ is

$$F_2^{(i)} = \left\| \vec{\tilde{u}}(\vec{x}) - \vec{\tilde{u}}(\vec{x} + \Delta x_i \vec{e}_i) \right\|^2 + \left\| \vec{\tilde{u}}(\vec{x}) - \vec{\tilde{u}}(\vec{x} - \Delta x_i \vec{e}_i) \right\|^2$$ \hspace{1cm} (3.64)

The model may be modified for non-uniform rectilinear grid \[86\]

$$F_2(\vec{x}, \Delta) = \frac{1}{6} \sum_{i=1}^{3} F_2^{(i)} \left( \frac{\Delta c}{\Delta x_i} \right)^{2/3}$$ \hspace{1cm} (3.65)

The SF model gives good results for flows in which turbulence in small scales is close to isotropy, but it is too dissipative for shear layers. The SSF model is an improved version of SF model\[21\]. It allows to switch-off the model in regions where sharp gradients are present but where the flow is not three-dimensional enough. This is typically the case in the core region of high-speed jets, where transition to turbulence has not yet occurred. The measure of three dimensionality is chosen as the angle $\theta$ between the local vorticity and the space averaged value of the vorticity over the six closest neighboring points. The SSF model reads

$$\mu_{sgs}^j = f_{\theta_0}(\theta) C_{SF} \Delta \vec{p} \left[ F_2(\vec{x}, \Delta) \right]^{1/2}$$ \hspace{1cm} (3.66)

where

$$f_{\theta_0}(\theta) = \begin{cases} 1 & \text{if } \theta \geq \theta_0 = 20^\circ \\ 0 & \text{else} \end{cases}$$ \hspace{1cm} (3.67)

The model constant is taken as

$$C_{SF} = 0.105 C_K^{-3/2} = 0.0634$$ \hspace{1cm} (3.68)

where $C_K = 1.4$ is the Kolmogorov constant. The choice of the $C_{SF}$ value is discussed in \[21, 30, 31, 85\].
3.2.2 CLOSING THE FILTERED ENERGY EQUATION

In the filtered energy equation, the sum of the major sub-grid terms $B_1 j + B_2 j$ is modeled as a whole with a gradient assumption, using a constant $Pr_{sgs} = \mu_{sgs} C_p/\lambda_{sgs}$ number.

\[
B_1 j + B_2 j = -\frac{\mu_{sgs} C_p}{Pr_{sgs}} \hat{T}_{-j} = -\lambda_{sgs} \hat{T}_{-j}
\]

(3.69)

in which the thermal conductivity for energy is connected to the eddy viscosity through unity $Pr_{sgs}$.

3.2.3 CLOSING THE FILTERED SPECIES EQUATION

The same is done for the $C_{1\alpha j}$ term of the species equations, equation (3.14), using a constant sub-grid Schmidt number $Sc_{sgs}$.

\[
C_{1\alpha j} = -\frac{\mu_{sgs}}{Sc_{sgs}} \bar{Y}_{\alpha,j} = -\rho D_{sgs} \bar{Y}_{\alpha,j}
\]

(3.70)

This is quite a crude -although commonly used- modeling compared, for example, to the dynamic methodology proposed in [56] which closes the major part (3.50) of (3.43) leading to $\hat{T} \approx \bar{T}$. However, it is sufficient to assess the aim of this study, which is to compare LES and MILES for high speed shear flows. In this state of mind, it will simply taken that $Pr_{sgs} = Sc_{sgs} = 1$.

Modeling the sub-grid chemical source term $C_{3\alpha}$ is the cornerstone of LES of reacting flows. Most of the models for non-premixed combustion rely on a conserved scalar like the mixture fraction, for which a filtered (joint)probability density function (FPDF) has to be prescribed (usually a $\beta$-function). Examples are flamelet models [20], progress-variable models [101]) and conditional moment closure (CMC) models [75]. These models also require the variance of the mixture fraction and/or its dissipation rate at the smallest turbulent scales, which is difficult to achieve in implicit LES [103]. Hence, in the framework of the present study, the $C_{3\alpha}$ reaction term is left to the numerical diffusion.

3.3 Implicit LES

In the simulations, a set of non-linear PDEs is being solved. In order to satisfy stability and monotonicity of solutions including discontinuities, specially designed schemes are applied.
These schemes, by their nature, cause the waves speed and amplitude of the solution to deviate from theoretical exact solution. In “numerical LES” approach, the deformation due to the numerical scheme, linked to the resolving efficiency of the derivation scheme, contributes as SGS motions. This argument is equivalent to what is indicated by Boris: “monotone CFD algorithms readily have built-in filter, and corresponding built-in SGS model” [12]. The numerical deformation of wave speeds and amplitudes due to the numerical schemes are dispersive and dissipative errors respectively. The dissipative and dispersive errors may be separated analytically, applying the modified wave number (MWN) spectral analysis. This analysis may be done analytically for linear numerical schemes.

3.3.1 MWN ANALYSIS

Consider the one-dimensional scalar function $F_k(x)$, $x \in [0, 2\pi]$ for simplicity, containing a single Fourier mode $k$. On a regular grid $x_i = i\Delta x$, $\Delta x = 2\pi/N$, $i = 0, \ldots, N$, $k$ must stay within the numerical cutoff $|k| \leq N/2$, $|k\Delta x| \leq \pi$ to avoid aliasing errors. The discrete projection of $F_k(x)$ is

$$F_k(x_i) = \widehat{F}_k e^{jkx_i} \equiv F_{k,i} \quad (3.71)$$

**Linear MWN analysis**

A linear scheme produces a numerical derivative at the single mode $k$

$$\frac{\partial F_k}{\partial x} \bigg|_{i} = \frac{\delta F_k^{L}}{\delta x} = \frac{\delta \widehat{F}_k}{\delta x}(k)e^{jkx_i} = jk'(k)\widehat{F}_k e^{jkx_i} \quad (3.72)$$

where $k'(k\Delta x)$ is the MWN of the scheme, whose real and imaginary parts are associated respectively with dispersive and dissipative errors. The analysis of response produced by some linear schemes on mode $k$ may be demonstrated as follows.

- Second order centered scheme

$$\frac{\partial F_k}{\partial x} \bigg|_{i} = F_{k,i+1} - F_{k,i-1} \quad \frac{N/2}{2 \Delta x} \sum_{k=-N/2}^{N/2} \widehat{F}_k \left( e^{jk\Delta x} - e^{-jk\Delta x} \right) e^{jkx_i}$$

$$k'(k\Delta x) = \frac{\sin (k\Delta x)}{\Delta x} \quad (3.73)$$
\begin{itemize}
  \item First order upwind scheme
  \[
  \frac{\partial F_k}{\partial x} \bigg|_{i} = \frac{F_{k,i} - F_{k,i-1}}{\Delta x} = \sum_{k=-N/2}^{N/2} \hat{F}_k \left( \frac{1 - e^{-jk\Delta x}}{\Delta x} \right) e^{-jkx_i}
  \]
  \[
k'(k\Delta x) = \sin \left( \frac{k\Delta x}{\Delta x} \right) - j \left( \frac{1 - \cos \left( \frac{k\Delta x}{\Delta x} \right)}{\Delta x} \right)
  \tag{3.74}
  \]
  \item Second order upwind scheme
  \[
  \frac{\partial F_k}{\partial x} \bigg|_{i} = \frac{3F_{k,i} - 4F_{k,i-1} + F_{k,i-2}}{2\Delta x} = \sum_{k=-N/2}^{N/2} \hat{F}_k \left( \frac{3 - 4e^{-jk\Delta x} + e^{-2jk\Delta x}}{2\Delta x} \right) e^{jkx_i}
  \]
  \[
k'(k\Delta x) = \frac{1}{2\Delta x} \left[ 4\sin \left( \frac{k\Delta x}{\Delta x} \right) - \sin \left( \frac{2k\Delta x}{\Delta x} \right) \right] - \frac{j}{2\Delta x} \left[ 3 - 4\cos \left( \frac{k\Delta x}{\Delta x} \right) + \cos \left( \frac{2k\Delta x}{\Delta x} \right) \right]
  \tag{3.75}
  \]
\end{itemize}

**Non-linear MWN analysis**

In this study, the non-linear WENO procedure, which will be detailed in the next chapter, is applied to compute reconstructed vectors \( \hat{\vec{F}}_{i+1/2} \). Separation of dissipative and dispersive errors of non-linear schemes is not possible analytically. However, the numerical error analysis which will be presented in this section, may be done for any scheme.

In the present case of WENO, at a point \( \{x_i = i\Delta x\} \) of a structured cartesian grid, the conservative approximation to the first derivative at fifth-order

\[
\left. \frac{\partial \hat{F}}{\partial x} \right|_i = \hat{F}_{i+1/2} - \hat{F}_{i-1/2} + O(\Delta x^5)
\tag{3.76}
\]

applies in smooth regions of the flow. The following spectral analysis of the non-linear WENO scheme (4.14) was introduced by N. Lardjane [80] and further developed by S. Pirozzoli [102].

A non-linear scheme will produce a response at other frequencies which may be separated applying the modified wave number (MWN) analysis numerically.

\[
\left. \frac{\delta F_k}{\delta x} \right|_i^{NL} = \sum_{n=-N/2}^{N/2} \frac{\delta F_k}{\delta x} (n) e^{jn\xi_i}
\tag{3.77}
\]

The MWN \( k'_x \) of the non-linear scheme is introduced, such that

\[
\left. \frac{\delta F_k}{\delta x} \right|_i^{NL} = jk'_x \hat{F}_k e^{jkx_i}
\tag{3.78}
\]

in which one can separate the contribution to mode \( k \) from the contribution to other modes

\[
jk'_x \hat{F}_k e^{jkx_i} = \frac{\delta \hat{F}_k}{\delta x} (k) e^{jkx_i} + \sum_{n\neq k} \frac{\delta \hat{F}_k}{\delta x} (n) e^{jn\xi_i}
\tag{3.79}
\]
By analogy with (3.72), \( k'_L(k) = k'_N(k) + k'_{NL}(k) \) is written with

\[
k'_L(k) = -j \frac{\delta F_L}{F_k \delta x} (k)
\]  

(3.80)

and

\[
k'_{NL}(k) = -j \sum_{n=-N/2}^{N/2} \frac{\delta F_k(n)e^{i(n-k)x_i}}{F_k n}
\]  

(3.81)

The linear MWN (3.80) has been computed on a grid of size \( N = 1024 \), without (smooth function) and with the non-linear WENO weights. In the former case, the WENO scheme simply becomes linear. Figure 3.1 displays the real and imaginary parts of the scaled linear MWN, \( \omega'(\omega) = \Delta x k'_L(k\Delta x) \), for each case. The effective spectral properties of the WENO scheme are very different from the theoretical ones, and the peaks intentionally kept on the curves show that the stiffness sensors may be put in default for some specific frequencies on a given grid. An analysis of the non-linear MWN (3.81) can be found in [80].

Figure 3.1: Linear Modified Wave Number without (top) and with (bottom) non-linear WENO weight. Left: real part, right: imaginary part. (a) 3rd order WENO, (b) 5th order WENO, (c) 7th order WENO, (d) 9th order WENO, (e) 11th order WENO, (f) 1st order upwind FD, (g) 2nd order upwind FD
3.3.2 THE INTERACTION BETWEEN NUMERICAL ERRORS AND EXPLICIT SUB-GRADE MODELING

The following 1D Fourier analysis simply demonstrates the interaction of SGS model with numerical errors. As the MWN analysis, this analysis may be conducted for any numerical scheme. Consider the viscous linear wave equation

$$\frac{\partial u}{\partial t} + c \frac{\partial u}{\partial x} = \nu \frac{\partial^2 u}{\partial x^2}$$  \hspace{1cm} (3.82)

whose dispersion relation is $\Omega(k) = kc - j\nu k^2$ for a single Fourier mode $u(x,t) = \hat{u}(k,t)e^{ikx}$.

A “physical LES” for equation (3.82) may be considered, in which the advection term is computed with a WENO scheme, and the diffusion term with a central FD scheme for first order derivative, applied twice. Introducing the scaled MWN of both schemes, the numerical LES dispersion relation can be written

$$\Omega_{num}^{LES}(k) = kc \left( \frac{\omega'_L}{\omega} \right) - j\nu'_{sgs} k^2 \left( \frac{\omega'_{FD}}{\omega} \right)^2$$  \hspace{1cm} (3.83)

where $\omega'_{FD}(\omega) = \frac{1}{6}(8\sin(\omega) - \sin(2\omega))$ for the 4th order central FD scheme.

In 1D, the Smagorinsky model reads

$$\nu_{sgs}^{SM} = \sqrt{2} C_S^2 \Delta x \frac{\partial u}{\partial x} \approx \sqrt{2} C_S^2 \Delta x \omega'_{FD} |\hat{u}|$$  \hspace{1cm} (3.84)

The second order structure function in the SSF model is

$$F_2(x,t) = \langle \|u(x,t) - u(x+r,t)\|^2 \rangle_{|r|=\Delta x} \approx \Delta x^2 \frac{\partial u}{\partial x} |_{FD}^2$$  \hspace{1cm} (3.85)

hence

$$\nu_{sgs}^{SSF} \approx C_{SF} \Delta x \omega'_{FD} |\hat{u}|$$  \hspace{1cm} (3.86)

Setting $C = C_{SGS}|\hat{u}|/c$ with $C_{SGS} = \sqrt{2} C_S^2$ (SM model) or $C_{SGS} = C_{SF}$ (SSF model) yields

$$\frac{1}{kc} \Omega_{num}^{LES}(\omega) = \frac{\omega'_L}{\omega} - j\frac{\omega'_{FD}^2}{\omega} C$$  \hspace{1cm} (3.87)

to be compared with

$$\frac{1}{kc} \Omega_{exact}^{LES}(\omega) = 1 - j\omega^2 C$$  \hspace{1cm} (3.88)

for exact derivatives. The departure from 1 (resp. 0) of the real (resp. imaginary) part of the ratio

$$\eta = \frac{\Omega_{num}^{LES}}{\Omega_{exact}^{LES}}$$  \hspace{1cm} (3.89)
indicates the frequency range, depending on the value of $C$, for which numerical errors begin to interact with the sub-grid model. For a given frequency $\omega \neq 0$, when $C = 0$, $\eta = \omega' / \omega$, and when $C \rightarrow \infty$, $\eta \rightarrow \left(\omega'_{FD} / \omega\right)^3$. Figure 3.2 shows the real and imaginary parts of (3.89) for the 5th order WENO scheme, associated with the 4th order central FD scheme. About 20% of the frequencies that the grid can capture are not affected by numerics.

Figure 3.2: Interaction indicator eq. (3.89) for $C = 0, 0.1, 1$ and 10. Left: real part, right: imaginary part.
CHAPTER 4

THE WENO SOLVER

The vector representation of the Navier-Stokes equations for multi-species flows (2.53) is given by

\[
\frac{\partial \tilde{U}}{\partial t} + \frac{\partial \tilde{F}}{\partial x} + \frac{\partial \tilde{G}}{\partial y} + \frac{\partial \tilde{H}}{\partial z} = \tilde{V} + \tilde{S}
\]  

(4.1)

In this chapter, \(\tilde{\cdot}\) indicates the conservative variables, and not Favre variables.

\[
\tilde{U} = ^T \begin{bmatrix} \rho, \rho u, \rho v, \rho w, \rho e_t, \rho Y_1, \ldots, \rho Y_{N_{sp}-1} \end{bmatrix}
\]  

(4.2)

The Euler equations (\(\tilde{V} = \tilde{0}, \tilde{S} = \tilde{0}\)) are a set of hyperbolic PDEs in each plane \((x, t), (y, t)\) and \((z, t)\)

\[
\frac{\partial \tilde{U}}{\partial t} + \frac{\partial \tilde{F}}{\partial x} + \frac{\partial \tilde{G}}{\partial y} + \frac{\partial \tilde{H}}{\partial z} = \tilde{0}
\]  

(4.3)

In order to illustrate the WENO method, consider a 1D hyperbolic problem

\[
\frac{\partial \tilde{U}}{\partial t} + \frac{\partial \tilde{F}}{\partial x} = \tilde{0}
\]  

(4.4)

The finite difference spatial derivative of the conservative flux (\(\tilde{F}\)) is computed to advance the solution in time. Use of the conservative form guarantees discontinuities to move at the correct speed satisfying Rankine-Hugoniot conditions. WENO method with Lax Friedrichs flux splitting is applied for ensuring stability of the scheme. The 1D system (4.4) is diagonalized then flux splitting and WENO reconstruction are applied on characteristic fields. Derivatives at grid points are computed using reconstructed fluxes at cell interfaces for each direction, separately.

In this chapter, firstly transformations between conservative \(\tilde{U}\), primitive \(\hat{U}\) and characteristic \(\hat{W}\) variables are explained. Then, the conservative finite difference scheme is defined. In
the third part, the WENO reconstruction with Lax-Friedrichs flux splitting algorithm is explained. The time integration and boundary conditions are presented in sections 4.4 and 4.5. Finally, some specificities of the numerical tool are listed.

4.1 TRANSFORMATION BETWEEN CONSERVATIVE, PRIMITIVE AND CHARACTERISTIC FORMS

The notations for transformations between conservative, primitive and characteristic forms of the EULER system, in $x$ direction is presented in this section. The system in conservative form is written as

$$\frac{\partial \tilde{\mathbf{U}}}{\partial t} + [\tilde{A}] \frac{\partial \tilde{\mathbf{U}}}{\partial x} = \mathbf{0} \quad \text{where} \quad [\tilde{A}]_{ij} = \frac{\partial \tilde{F}_i}{\partial \tilde{U}_j}$$ (4.5)

Conservative variables are transformed into primitive form using matrix $[P] = \frac{\partial \tilde{\mathbf{U}}}{\partial \mathbf{U}}$

$$\frac{\partial \mathbf{U}}{\partial t} + [A] \frac{\partial \mathbf{U}}{\partial x} = \mathbf{0} \quad \text{where} \quad [A] = [P^{-1}] [\tilde{A}] [P]$$ (4.6)

The vector of primitive variables is

$$\mathbf{U} = \mathbf{T} [\rho, u, v, w, T, Y_1, ..., Y_{N_{sp} - 1}]$$ (4.7)

Matrix $[A]$ is diagonalized with left and right eigenmatrices, $[L]$ and $[R]$ respectively.

$$[\Lambda] = [L] [A] [R]$$ (4.8)

Explicitly, the terms of the system with size $(N_{sp} + 4)$ with the eigenvalues ordered $\tilde{\lambda} = \mathbf{T} [u - c, u, u, u + c, ..., u]$ are as follows

$$[L] = \mathbf{T} [\tilde{l}^{(1)}, \tilde{l}^{(2)}, ..., \tilde{l}^{(N_{sp} + 4)}]$$ (4.9)

where $\tilde{l}^{(i)}$ are the left eigenvectors of primitive jacobian matrix $[A]$ (Figure 4.1) related to the eigenvalue $\lambda^{(i)}$

$$\tilde{l}^{(i)} [A] = \lambda^{(i)} \tilde{l}^{(i)} \quad \text{for} \quad i = 1, ..., N_{sp} + 4$$ (4.10)

Diagonalization is done on primitive form since it is difficult to obtain $[\tilde{L}]$ and $[\tilde{R}]$ which are left and right eigenmatrices of the conservative jacobian matrices $[\tilde{A}]$.

$$\frac{\partial \tilde{W}}{\partial t} + [\Lambda] \frac{\partial \tilde{W}}{\partial x} = \mathbf{0} \quad \text{where} \quad \tilde{W} = [L] \tilde{U}$$ (4.11)

The definitions and notations are summarized in Figure 4.1 The transformation matrices
Figure 4.1: Transformations between different forms for the Euler system of equations.

$[\tilde{R}]$ and $[\tilde{L}]$ are given in appendix D. Although the derived transformation matrices are of size $N_{sp} + 5$, the mass fraction of the last species $Y_{N_{sp}}$ is computed out of the others. The same derivation for the characteristic boundary conditions is conducted for $N_{sp} + 4$ system of equations, presented in appendix E.

### 4.2 CONSERVATIVE FORM FOR DIFFERENTIAL OPERATOR

The objective of the finite difference spatial discretization scheme is to compute the spatial derivative of any function. $f$ is the flux function of a non-linear PDE

$$\frac{\partial u}{\partial t} + \frac{\partial f}{\partial x} = 0$$

$$u(x, 0) = u_0(x)$$

Then the solution may be advanced in time with an integration scheme like a TVD Runge Kutta method. Consider pointwise function $f \equiv f(x_i)$ defined on a grid $(x_0, ..., x_i, ..., x_N)$. A conservative approximation for equation (4.12) is of the form

$$\frac{df}{dx}|_{x_i} = \frac{1}{\Delta x} \left( \hat{f}_{i+1/2} - \hat{f}_{i-1/2} \right)$$

where,

$$\hat{f}_{i+1/2} = \hat{f}_{i+1/2}(f_{i-1}, ..., f_{i+1}) \quad i = 0, ..., N$$

(4.13)
and \( s \) are positive integers. Conservative approximation for derivative of flux function yields discontinuities in the solution to move at physically correct speed [126]. However, stability is not ensured by conservative schemes.

### 4.3 WENO SCHEME

The importance of the numerical solutions to the Navier-Stokes equations increases due to improvements enabling solutions for real problems of concern. One side of this improvement is in computational capacity. The growth in understanding of PDE theory and development of new numerical techniques, is the other side. Godunov’s “order barrier theorem” is one of the milestones for numerical solution of flow problems including discontinuities [57], in 1959. After this early work, in the 80’s Lax, Van Leer, Harten and many others published on approximate solution methods for the Riemann problem [108, 120]. Harten presented his work on total variation diminishing (TVD) schemes. This work opens the door for higher order “monotonicity preserving” schemes that do not contravene Godunov’s theorem [61]. Later, Harten introduced the “essentially non-oscillatory” (ENO) scheme with his colleagues [62]. ENO scheme is based on the choice of the smoothest shifted stencil for flux reconstruction near discontinuities. ENO schemes are successful for shock capturing. Shu and Osher suggested TVD Runge-Kuta time discretization and application of ENO procedure directly on flux values rather than on the variables [113]. These modifications enhance the accuracy of ENO schemes for lower cost. Later, in a second paper they introduced ENO-Roe, ENO-LLF (local Lax-Friedrichs flux splitting) methods for improvement [113]. Weighted essentially non oscillatory scheme (WENO) which is applied in the numerical method of this thesis is developed by Liu, Osher and Chan [88]. Recently, modifications to improve the accuracy of WENO method, in the literature. Mapped WENO schemes achieving \((2r-1)^{th}\) order accuracy near discontinuities [64], more accurate band-width optimized WENO scheme [93, 118] and low cost WENO-Z schemes [10] are some examples. These improvements are not considered, since the scope is not related to achieve higher accuracy.

The key idea in WENO is constructing flux values using the weighted average of approximate flux values at cell interfaces. Normalized weights are determined depending on the smoothness of the ENO approximations at shifted sub-stencils shown in Figure 4.2.

In the WENO procedure, \( \hat{f}_{i+1/2} \) in equation (4.13) is reconstructed using ENO approximations
on different sub-stencils \((S_{i}^{(r)})\) in Figure 4.2) including \(x_{i+1/2}\). The overall stencil \((S_{i})\) size for \(k^{th}\) order reconstruction of \(\hat{f}_{i+1/2}\) by WENO scheme is \(2k - 1\). Weighted average of \(k\) number of approximations \(\hat{f}_{i+1/2}^{(r)}\) for \(r = 0, ..., k - 1\) gives reconstructed flux

\[
\hat{f}_{i+1/2} = \sum_{r=0}^{k-1} w_{r} \hat{f}_{i+1/2}^{(r)} \tag{4.14}
\]

As an example for the fifth order WENO reconstruction Figure 4.2 the approximations \(\hat{f}_{i+1/2}^{(r)}\) are computed on sub-stencils \(S_{i}^{(0-2)}\) of size \(k\) (Figure 4.2).

\[
\hat{f}_{i+1/2} = \hat{f}_{i+1/2}^{(0)} w_{0} + \hat{f}_{i+1/2}^{(1)} w_{1} + \hat{f}_{i+1/2}^{(2)} w_{2} \tag{4.15}
\]

and \(\hat{f}_{i+1/2}^{(r)}\) is obtained from \(k^{th}\) order ENO approximation.

![Figure 4.2: 5th order WENO stencil demonstration.](image)

4.3.1 ENO APPROXIMATION

\(\hat{f}_{i+1/2}^{(r)}\) are values of the interpolating polynomials, the coefficients of which are derived on following grid, cells and grid points

\[
a = x_{1/2} < x_{3/2} < \ldots < x_{N-1/2} < x_{N+1/2} = b
\]

\[
I_{i} = [x_{i-1/2}, x_{i+1/2}]; \quad x_{i} = \frac{1}{2}(x_{i-1/2} + x_{i+1/2})
\]

At the cell \(I_{i}\), \(\hat{f}_{i+1/2}^{(r)}\) which is \(k^{th}\) order accurate approximation is based on stencil \(S_{i}^{(r)}\) of length \(k\) cells, \(r\) cells to the right, and \(s\) cells to the left, with \(r + s + 1 = k\)

\[
S_{i}^{(r)} \equiv \{I_{i-r}, \ldots, I_{i+s}\} \tag{4.17}
\]
The approximations are constrained by the integral equality for conservative schemes

\[
\bar{f}_j \equiv \frac{1}{\Delta x_j} \int_{x_{j-1/2}}^{x_{j+1/2}} f(\xi) d\xi \equiv \frac{1}{\Delta x_j} \int_{x_{j-1/2}}^{x_{j+1/2}} \hat{f}^{(r)}(\xi) d\xi \quad j = i - r, ..., i + s
\] (4.18)

There exists a unique polynomial approximation of degree \(k - 1\), on nodes of stencil \(S_i^{(r)}\) matching \(f_j\) values in the sub-stencil. So, there exists linear set of coefficients \(c_{r,j}\) depending on left shift for \(\hat{f}^{(r)}_{i+1/2}\).

\[
\hat{f}^{(r)}_{i+1/2} = \sum_{j=0}^{k-1} c_{r,j} f_{i-r+j}
\] (4.19)

The linear coefficients which are derived starting from the Lagrange form of approximate polynomials by Shu are

\[
c_{r,j} = \left( \sum_{m=j+1}^{k} \sum_{l=0}^{k-m} \prod_{q=0}^{k} \frac{x_{i+r-q+m} - x_{i-r-q+1}}{x_{i+r-q+m} - x_{i-r-q+1}} \right) \Delta x_{i-r+j}
\] (4.20)

If the grid is uniform, the coefficients do not depend on \(i\) (\(\Delta x_i = \Delta x\))

\[
c_{r,j} = \sum_{m=j+1}^{k} \sum_{l=0}^{k-m} \prod_{q=0}^{k} \frac{r - q + 1}{m - l}
\] (4.21)

The constants \(c_{r,j}\) are listed in Table at appendix G for \(k = 1, ..., 6\).

### 4.3.2 1D WENO RECONSTRUCTION PROCEDURE

The reconstruction procedure to compute \(\hat{f}_{i+1/2}\) using \(f_i\) is as follows.

- At \(x_{i+1/2}\), \(k\) approximate values \(\hat{f}^{(r)}_{i+1/2}\) of \(k^{th}\) order accuracy are computed over sub-stencil \(S_i^{(r)}\).

\[
\hat{f}^{(r)}_{i+1/2} = \sum_{j=0}^{k-1} c_{r,j} f_{i-r+j} \quad ; \quad S_i^{(r)} = \{x_{i-r}, ..., x_{i-r+k-1}\}
\] (4.22)

- The reconstruction is done by weighted averaging of \(\hat{f}^{(r)}_{i+1/2}\) as seen in equation (4.14).

- The normalized weights \(w_r\) are related to smoothness. \(w_r\) are used to increase the contribution of the approximation at the sub-stencil \(S_i^{(r)}\) where interpolated polynomial is smoother.

\[
w_r = \frac{\alpha_r}{\sum_{s=0}^{k-1} \alpha_s} \quad r = 0, ..., k - 1 \quad \alpha_r = \frac{d_r}{(\epsilon + \beta_r)^2}
\] (4.23)
\( \beta_r \) is called the smoothness indicator which is a measure of discontinuity (stiffness) of the solution. \( \epsilon \) is a positive small coefficient to avoid division by zero. The ideal weights \( d_r \) which generate \((2k-1)^{th}\) order upwind scheme are tabulated in appendix G for \( k = 1, \ldots, 6 \). If the function is smooth in the stencil \( S_i^{(r)} \) the value of the smoothness indicator \( \beta_r = O(\Delta x^2) \) so that \( w_r = O(1) \). When there exists a discontinuity in the stencil \( \beta_r = O(1) \) so that \( w_r = O(\Delta x^4) \). The numerical formulation suggested by Shu is presented in appendix G for \( k = 1, \ldots, 6 \).

### 4.3.3 FLUX SPLITTING

Application of Lax-Friedrichs flux splitting is suggested by Shu to enhance the stability characteristics of ENO [114]. The same enhancement is extended to WENO schemes. In ICASE report, Shu states that the flux splitting algorithm is cost effective and improves stability characteristics [112]

\[
f(u) = f^+(u) + f^-(u)
\]

where for Lax-Friedrichs flux splitting,

\[
f^\pm(u) = \frac{1}{2}(f(u) \pm \alpha u) \quad \alpha = \max_u \left| f'(u) \right|
\]

The range of relevant \( u \) for taking maximum may be chosen at the overall domain or locally as the stencil \( S_i \). Depending on the choice, the method is named “local LF” or “global LF”.

In application, WENO procedure is applied on split fluxes \( f_{i+1/2}^+ \) and \( f_{i+1/2}^- \). Note that, in the \( c_{rj} \) Table which is presented in appendix G, \( r \) takes the value \(-1\). The transformation of coefficients \( \tilde{c}_{rj} = c_{r-1,j} \) is for \( \tilde{f}_{i+1/2}^+ \) as, \( c_{rj} \) is for \( \tilde{f}_{i+1/2}^- \). After WENO procedure, fluxes are reconstructed

\[
f_{i+1/2} = \tilde{f}_{i+1/2}^+ + \tilde{f}_{i+1/2}^-
\]

Lax-Friedrichs flux splitting is the less expensive but one of the most dissipative flux splitting algorithms. However, as the order of the scheme increases, this dissipation effect decreases [2].
4.3.4 SOLUTION FOR SYSTEM OF EQUATIONS

It is explained in the introduction of this chapter, that the 1D flux derivative is computed in $x$ direction and this formulation is used for $y$ and $z$ directions. Application of WENO scheme directly to the coupled system of equations is possible. However, this is reported to cause spurious oscillations for higher order approximations [112]. 1D characteristic system of variable is

$$\frac{\partial \vec{W}}{\partial t} + [\tilde{L}] \frac{\partial \vec{F}}{\partial x} = \vec{0} ; \quad [\tilde{L}] \frac{\partial \vec{F}}{\partial x} = \frac{\partial \tilde{g}}{\partial x}$$

(4.27)

which may be written in component form as

$$\frac{\partial W_k}{\partial t} + \frac{\partial g_k}{\partial x} = 0 ; \quad k = 1, ..., N_{sp} + 5$$

(4.28)

where $\tilde{g}$ is the flux for the characteristic system. The overall procedure of WENO scheme for flux derivative computation having the conservative variables $\vec{\tilde{U}}_i$ and the flux values $\vec{\tilde{F}}_i$ at grid points, is as follows.

- Transformation matrices from conservative to characteristic variables $[\tilde{R}]$, $[\tilde{L}]$ and $[\Lambda]$, are constructed at $x_{i+1/2}$. Matrices are given analytically in appendix D.

$$[\tilde{R}] = [\tilde{R}](\vec{U}_{i+1/2}) \quad [\tilde{L}] = [\tilde{L}](\vec{U}_{i+1/2}) \quad [\Lambda] = [\Lambda](\vec{U}_{i+1/2})$$

(4.29)

- These matrices are assumed to be constant in the stencil of concern $S_i$. The cell interface values of the primitive variables $\vec{U}_{i+1/2}$ in equations (4.29) are deduced from conservative variables $\vec{\tilde{U}}_{i+1/2}$ which are computed by simple averaging.

$$\vec{\tilde{U}}_{i+1/2} = \frac{1}{2}(\vec{\tilde{U}}_i + \vec{\tilde{U}}_{i+1})$$

(4.30)

- Having obtained the left eigenvectors $\vec{\tilde{l}}$ (lines of left eigenmatrice $[\tilde{L}]$) the characteristic system flux values are computed.

$$g_k = \sum_{p=1}^{s+N_{sp}} \hat{r}_p^{(k)} \vec{F}_p = \lambda^{(k)} W_k$$

$$k = 1, ..., N_{sp} + 5$$

(4.31)

- Lax-Friedrichs flux splitting is applied on $g_k$ as

$$g_k = g^+_k + g^-_k$$

(4.32)
\[ g_k^\pm = \frac{1}{2} \left[ g_k(\bar{W}) \pm \alpha_k W_k \right] \]
\[ \alpha_k = \max \left| \frac{\partial g_k}{\partial W_k} \right| = \max |\lambda^k| \]  \hspace{1cm} (4.33)

- WENO procedure is applied to obtain reconstructed fluxes at cell interfaces \( \hat{g}_{k,i+1/2}^\pm \)

\[ \hat{g}_{k,i+1/2}^{\pm(r)} = \sum_{j=0}^{k-1} C_j \hat{g}_{k,i-r+j} \]

\[ \hat{g}_{k,i+1/2}^{\pm} = \sum_{r=0}^{k-1} \omega_r \hat{g}_{k,i+1/2}^{\pm(r)} \]  \hspace{1cm} (4.34)

- Back transformation is applied from characteristic to conservative flux.

\[ \hat{F}_{i+1/2}^\pm = \left[ \hat{R} \right]_{i+1/2} \hat{g}_{i+1/2}^\pm \]  \hspace{1cm} (4.35)

- Finally, the overall reconstructed numerical fluxes are evaluated as

\[ \hat{F}_{i+1/2} = \hat{F}_{i+1/2}^++\hat{F}_{i+1/2}^- \]  \hspace{1cm} (4.36)

### 4.4 TIME DISCRETIZATION

Having computed the flux derivative in space, the solution is advanced using explicit Runge-Kutta time discretization.

#### 4.4.1 RK\(_p\) TVD TIME STEPPING

General form of Runge-Kutta methods to solve initial value problem of ODE: \( u_t = L(u) \), which is obtained from spatial discretization of PDE: \( u_t = f(u)_x \), is

\[ u^{(p)} = \sum_{k=0}^{p-1} (\alpha_{pk} u^{(k)} + \Delta t \beta_{pk} L(u^{(k)})), \quad p = 1, ..., m \]

\[ u^{(0)} = u^0, \quad u^{(m)} = u^{n+1} \]  \hspace{1cm} (4.37)

Considering an ODE (in time), the total variation \( TV(u) = \sum_j |u_{j+1} - u_j| \) for first order in time Euler forward stepping, does not increase under the restriction

\[ \Delta t \leq \Delta t_{FE} \]  \hspace{1cm} (4.38)
Time integration is stable provided that $\Delta t \leq c \Delta t_{FE}$ where $c$ is the CFL coefficient. It is proved, higher order explicit Runge-Kutta time discretization (eq. 4.37) is TVD under the CFL-like condition [113]

$$c = \min_{i,k} \frac{\alpha_{ik}}{\beta_{ik}} \quad \alpha_{ik} > 0, \beta_{ik} > 0$$ (4.39)

The third order TVD Runge Kutta method with $c = 1$ is

$$u^{(1)} = u^n + \Delta t L(u^n)$$

$$u^{(2)} = \frac{3}{4} u^n + \frac{1}{4} u^{(1)} + \frac{1}{4} \Delta t L(u^{(1)})$$

$$u^{n+1} = \frac{1}{3} u^n + \frac{2}{3} u^{(2)} + \frac{2}{3} \Delta t L(u^{(2)})$$ (4.40)

4.4.2 CALCULATING THE TIME STEP

A non dimensional CFL condition is used to calculate the time step for time integration. CFL value less than unity guaranties the stability of the time integration.

$$\Delta t_{\text{max}} = \frac{C \Delta x_{\text{min}}}{|\lambda_{x}|_{\text{max}}}$$ (4.41)

where $\lambda_x$ is the set of eigenvalues of the system ($\lambda_x = \{u + c, u, u - c\}$) that correspond to the wave speed. In application, three time steps considering stability of hydrodynamics, viscous terms and chemical reaction mechanism are determined. The minimum time step for stability of time integration is chosen.

$$\Delta t = \min (\Delta t_{\text{hydro}}, \Delta t_{\text{visc}}, \Delta t_{\text{chem}})$$ (4.42)

The hydrodynamic time step is computed applying CFL condition [70] as

$$\Delta t_{\text{hydro}} = \frac{C}{|\lambda_{x}|_{\text{max}} + |\lambda_{y}|_{\text{max}} + |\lambda_{z}|_{\text{max}}}$$ (4.43)

The CFL number is chosen to be 0.7 for the computations. The time step determined for viscous stability criteria is computed (for Navier-Stokes solutions) as

$$\Delta t_{\text{visc}} = \frac{1}{2\mu \left( \frac{1}{\Delta x} + \frac{1}{\Delta y} + \frac{1}{\Delta z} \right)}$$ (4.44)

$\Delta x$, $\Delta y$ and $\Delta z$ are the grid spacing in directions $x$, $y$ and $z$.

The chemical source terms are highly non-linear. Maximum time step for the reacting cases ($\Delta t_{\text{chem}}$) to prevent instability due to mechanism is determined conducting 0D, constant volume H$_2$/air mixture ignition tests. These tests are presented with the results since they are specific to the reaction mechanism and the conditions.
4.5 BOUNDARY CONDITIONS

Boundary conditions (BC) are of critical importance in numerical simulations. Indeed they are necessary to implement the physical requirements of the problem and to prevent unphysical wave reflections at the boundaries of the domain. BCs must assure the system to be “well-posed”. The system constrained by the BCs must reveal a unique outcome. The first way of constraining boundaries is direct implementation of physical values at the boundaries of the domain. This forcing is applied repeatedly at every time step. Secondly, “ghost cells” which are virtual cells prolonged outside at the domain boundaries may be utilized like boundary nodes. The presence of ghost cells is unavoidable for the spatial discretization stencils of nodes at the boundaries. However, forcing all conditions will generate unphysical waves to reflect at the boundaries. As a result, a method based on 1D characteristic analysis along the direction perpendicular to the boundary is applied. The first systematic work on non-reflecting inlet, outlet, slip and no-slip wall boundary conditions implemented on subsonic and supersonic inviscid flow problems is published by Kevin Thompson [79], to the author’s knowledge. The system for derivation of BC’s is called “local one-dimensional inviscid” (LODI) system of equations. Thompson refers to Hedstrom, who developed characteristic boundary conditions for one-dimensional, nonlinear and hyperbolic PDE solutions, on rectangular domain [63]. Thompson states his work is application of Hedstrom’s method on multidimensional Euler equations on non-rectangular domains with transformed coordinates. Later, Poinsot and Lele presented LODI system BC’s modified for Navier-Stokes equations [105]. Baum, Poinsot and Thévenin expanded the BC’s to multispecies reacting cases for gases with realistic thermodynamic properties [5]. Explanation of applied BCs and details of application are expressed in two classifications. The physical BCs are the ones implementing known physical properties at the boundaries. When the physically definite conditions are not enough to constrain the equations at the boundaries, numerical BCs are applied. The LODI system for application of BCs on characteristic basis is as follows.

4.5.1 LODI SYSTEM

When the number of definite physical BCs are not enough for well posedness of the solution, BCs may be applied on wave amplitude variations $\tilde{L}$ in characteristic form. The idea is based on the fact the solution is not only governed by the state of the problem but the waves entering
to the domain also [79].

One-dimensional (normal direction to the boundary), inviscid, characteristic form is used for \( \mathcal{L} \) formulation. The effect of waves in tangential directions and viscous terms on the boundaries are neglected for the test cases studied. This type of BCs are called Euler Characteristic Boundary Conditions (ECBC).

\[
\frac{\partial \vec{W}}{\partial t} + \vec{L} = \vec{0} \quad ; \quad \vec{L} = [\Lambda] \frac{\partial \vec{W}}{\partial x} = [\Lambda] [L] \frac{\partial \vec{U}}{\partial x} \quad (4.45)
\]

\( \vec{L} \) vector is written explicitly as

\[
\mathcal{L}_i = \lambda^{(i)} \sum_{j=1}^{N_{sp}+4} \kappa_j \frac{\partial U_j}{\partial x} \\
i = 1, ..., N_{sp} + 4 \quad (4.46)
\]

The components of \( \mathcal{L} \) in \( x, y \) and \( z \) are presented in appendix F. Once all components of wave amplitude variations are computed, time derivatives of conservative variables to advance in time, may be obtained applying transformation. Firstly, it is necessary to determine the direction of the waves related to \( \mathcal{L}_i \)'s, as explained in Figure 4.5 depending on

- nature of the flow, supersonic or subsonic
- direction of the flow, \( u < 0 \) or \( u > 0 \)

![Figure 4.3: ECBC subsonic flux directions](image)

![Figure 4.4: ECBC supersonic flux directions](image)

![Figure 4.5: Direction of the waves for subsonic and supersonic conditions for both \( u > 0 \) (black arrows) and \( u < 0 \) (red arrows) cases.](image)

Amplitude variations \( \mathcal{L}_i \) of the waves which leave the domain are computed applying equation (4.46), using one-sided derivatives based on interior data. \( \mathcal{L}_i \)'s of waves which enter the
domain can not be computed this way. Depending of the type of boundary, these amplitude
variations are

- either set to zero: numerical BC.
- or related to outgoing waves by physical restrictions applied on time derivatives of
  primitive variables: physical BC.

\[
[R] \frac{\partial \tilde{W}}{\partial t} + [R] \tilde{L} = \tilde{0}
\]

\[
\frac{\partial \tilde{U}}{\partial t} + \tilde{d} = \tilde{0} \quad ; \quad \tilde{d} = [R] \tilde{L} 
\]  

(4.47)

This system which relates wave amplitude variations to time derivatives of primitive variables
is called local one-dimensional inviscid (LODI) system. Detailed formulations of \( \tilde{L} \) and
\( \tilde{d} \) for each direction \( x, y \) and \( z \) are presented in appendix F. Having explained the basis
and definitions for ECBC, specific implementations depending on the type of BC will be
presented.

4.5.2 FREE NON-REFLECTING BCS

At free boundaries none of the primitive variables may be implemented explicitly. \( \tilde{L}_i \)'s for
incoming waves are equated to zero.

4.5.3 PHYSICAL NON-REFLECTING INLET BOUNDARY CONDITIONS

Physical non-reflecting boundary conditions are applied at \( x = 0 \) inlet only. The incoming
wave amplitude variations for subsonic and supersonic case are computed as follows.

4.5.3.1 SUBSONIC FORCED NON-REFLECTING INLET

All characteristic waves, except \( \tilde{L}_1 \) related to eigenvalue \( (u - c) \), enter the domain. \( \tilde{L}_1 \) is
computed from interior points. Other wave amplitude variations are expressed in terms of \( \tilde{L}_1 \)
solving the LODI system (4.47). The physical BC for a subsonic inlet at \( x = 0 \) are constant
velocity components \((u = u_{inlet}, v = 0, w = 0)\), temperature \((T = T_{inlet})\) and mass fractions
\( \{Y_1, Y_2, ..., Y_{Nsp}\} = \text{Const} \). The wave amplitude variations \( L_i \) are deduced equating time derivatives of these primitive variables which are presented in appendix F.2 to zero.

\[
\begin{align*}
L_2 &= 0 \\
L_3 &= 0 \\
L_4 &= -2L_1 \\
L_{\alpha+3} &= 0 \quad \alpha = (2, ..., N_{sp} - 1) \\
L_{N_{sp}+3} &= -\frac{\rho}{T}L_4 \\
L_{N_{sp}+4} &= L_1
\end{align*}
\] (4.48)

### 4.5.3.2 SUPersonic FORCED NON-REFLECTING INLET

In supersonic flow case, all the waves enter into the domain. All time derivatives of primitive variables are equated to \textit{zero}.

### 4.6 SOME SPECIFICITIES

#### 4.6.1 THE LARROUTUROU METHOD

As mentioned early in section 2.1, numerical simulation of multi-species flows with the conservative form of equations poses difficulty to keep species mass fractions positive. This problem is especially encountered for reacting flows involving large numbers of species. The positivity problem which has been the subject of many works [4, 82] are related to the resolution of Navier-Stokes equations in conservation form. Larrouturou method is applied for preserving positivity of the species mass fractions.

##### 4.6.1.1 IDEA OF LARROUTUROU

The mass fraction of species \( \alpha \) is computed as the ratio of two conservative variables

\[
Y_{\alpha,i} = \frac{\rho_i Y_{\alpha,i}}{\rho_i} \quad \alpha = 1, ..., N_{sp} - 1
\] (4.49)

Due to the numerical scheme employed, the mass fractions obtained do not respect the discrete maximum principle \((0 \leq Y_{\alpha} \leq 1)\). Larrouturou proposed to use the same discrete mass flux.
for continuity \( f_{i+1/2}^{\rho} \) and species mass fraction \( f_{i+1/2}^{\alpha} \) fluxes

\[
f_{i+1/2}^{\alpha} = f_{i+1/2}^{\rho} \begin{cases} 
Y_{\alpha,i} & \text{if } u_{i+1/2} > 0 \\
Y_{\alpha,i+1} & \text{if } u_{i+1/2} < 0 
\end{cases}
\]  

(4.50)

This modification guaranties maximum principle with suitable CFL condition [82].

### 4.6.1.2 APPLICATION TO WENO SOLVER

Larrouturou suggests to advance the numerical solution for whole system as coupled flow and species equations, then apply the method for species mass fluxes \( \hat{f}_{i+1/2}^{\alpha} \). It is intended to keep the accuracy of approximation at the same order for both flow and species variables. The method is applied as follows

\[
\hat{f}_{i+1/2}^{\alpha} = \hat{f}_{i+1/2}^{\rho} \begin{cases} 
\hat{Y}_{\alpha,i+1/2} & \text{if } \hat{f}_{i+1/2}^{\rho} > 0 \\
\hat{Y}_{\alpha,i+1/2}^{+} & \text{if } \hat{f}_{i+1/2}^{\rho} < 0 
\end{cases}
\]  

(4.51)

The conservation of mass as sum of species mass fractions (\( \sum_{\alpha=1}^{N_{sp}} \hat{Y}_{\alpha,i+1/2} = 1 \)) is satisfied applying the same weights for all species. It corresponds to the maximum of all the weights of WENO reconstruction for the species.

### 4.6.2 STRUCTURE OF THE CODE

The structure for single processor case is presented in Figure (4.6). In parallel case each processor follows this task and communicates at each time step for ghost cells on interior faces.

### 4.6.3 THE QUESTION OF \( D_{H0} \)

The energy is expressed in terms of the total energy which is computed with a reference to temperature \( T_0 \) that is chosen as \( T_0 = 298.15K \). This the energy variable which is zero at this reference temperature \( T = T_0 \), is called sensible energy. The conservative variable for energy equation is given in (2.48)
Main program
- init_parallel
- initialisation
- solver
- output

Numerical fluxes
- viscous fluxes
- reaction source terms
- Euler fluxes on x dr.
- Euler fluxes on y dr.
- Euler fluxes on z dr.

Type of simulation
- explicit SGS term
- viscosity

Spatial derivatives
- boundaries
- single time step
- explicit SGS term
- viscous flux

Boundary conditions
- NSCBC @ x=0
- NSCBC @ x=L
- NSCBC @ y=0
- NSCBC @ y=L
- NSCBC @ z=0
- NSCBC @ z=L

Communication
- fill ghost cells + inlet
- corners + edges
- evaluate primitive
- eigen values comp.
- numerical flux

Init parallel
- read parameters
- read initial field
- init communication

Solver
- calculate time step
- advance in time
- check time
- write results

Output
- read initial field
- init communication

Time step
- compute primitive
- consen
- trans. matrices
- WENO base
- final assembling

Rk
- U_{i+1/2} = \frac{1}{2} (U_i + U_{i+1})
- U'_{i+1/2} = \frac{1}{2} (U_i + U_{i+1})
- WENO base rout
- WENO base rout
- R
- U_{i+1/2}

Figure 4.6: Schematic diagram of structure of the code.
\[
\rho e_t = \rho \sum_{a=1}^{N_{sp}} \left( \int_0^T C p_a(\theta) d\theta \right) Y_a - p + \frac{1}{2} \rho u_i u_i + \rho D_{H0} \tag{4.52}
\]

with
\[
\rho D_{H0} = \rho \sum_{a=1}^{N_{sp}} \left( \Delta h_{f,a}^0 - \int_0^T C p_a(\theta) d\theta \right) Y_a
\]

Since solution is concerned with variation in total energy, it is suitable to carry on calculations for sensible energy, adding and subtracting \(\rho D_{H0}\) to \(\rho e_t\) every time step.

### 4.6.4 THE METRICS

At some regions of the domain where the flow is turbulent and mixing and reactions take place, high resolution is necessary. However there are regions where the solution is smooth, so there is no need for high resolution. The grid is clustered introducing geometric transformation in each direction in order to reduce the computational cost.

![Coordinate transformation for clustering the grid](image)

\[
x = L \frac{X}{\sqrt{1 - X^2}}, \quad \frac{dX}{dx} = \frac{L^2}{(X^2 + L^2)^{3/2}} \tag{4.53}
\]

The capital \(X\) stands for numerical coordinates, \(x\) is for physical coordinates and \(L\) is a constant for the ratio of minimum physical to numerical grid spacing. Spatial derivatives are transformed by chain rule.

\[
\frac{\partial}{\partial x} = \frac{\partial}{\partial X} \frac{dX}{dx} \tag{4.54}
\]

The filtering operation explained in chapter 3 does not commute with differential operator. It is shown that the commutation error is significant when the truncation error is higher than second order [53]. In this work the commutation errors are assumed to be negligible, since
the clustering is mostly effective at the region of smooth solution [92]. It is also applied by Lardjane [80] and Gougeon [59].

### 4.6.5 PARALLEL IMPLEMENTATION

Parallelization of the code is achieved by domain decomposition in all directions. Standard MPI routines for cartesian topology are used for parallelization.

The parallel efficiency may be represented in terms of scalability curves in Figure 4.8.A. Speed-up is calculated dividing wall time for a task for sequential processing by wall time for the same task processed with N processors.

\[
S_N = \frac{T_{1 \text{ proc}}}{T_{N \text{ procs}}}
\]  

(4.55)

The speed-up curves presented are for clusters VARGAS/IDRIS and Phoebus/CCRC (Centre de Calcul en région Centre).

- Phoebus: 42 nodes of 4 Intel XEON E5450 3.00 Ghz cores with 25Gb for each node.
- Vargas:
  - 112 nodes of 32 core Power 6 4.7 Ghz core with 128 Gb memory
  - 84 nodes and 256 Gb memory 28 nodes.

AR: the ratio of long edge to short edge for the square prism sub-domain blocks. AR is an indicator of ratio of communication (at surfaces) to computation (within volume) of each node.
4.6.6 LES STRATEGIES

The set of governing LES equations (3.8)(3.27)(3.32)(3.14) can be written in the following form, including the contribution of the sub-grid scales

\[
\frac{\partial \tilde{U}}{\partial t} + \frac{\partial \tilde{F}}{\partial x} + \frac{\partial \tilde{G}}{\partial y} + \frac{\partial \tilde{H}}{\partial z} = \tilde{V} + \tilde{r} + \tilde{S} \tag{4.56}
\]

where \( \tilde{F}, \tilde{G}, \\text{and} \ \tilde{H} \) are the inviscid fluxes in the \( x, y \) and \( z \) direction, respectively, \( \tilde{V} \) is the vector of viscous fluxes, \( \tilde{S} \) is the vector of chemical source terms, and \( \tilde{r} \) represents the vector of explicit sub-grid terms. The different approaches are characterized by:

- **LES_SM**: \( \tilde{V} \neq \tilde{0} \); \( \tilde{r} \): SM model
- **LES_SSF**: \( \tilde{V} \neq \tilde{0} \); \( \tilde{r} \): SSF model
- **MILES_NS**: \( \tilde{V} \neq \tilde{0} \); \( \tilde{r} = \tilde{0} \)
- **MILES_EULER**: \( \tilde{V} = \tilde{0} \); \( \tilde{r} = \tilde{0} \)

In the non-reacting cases, \( \tilde{S} \neq \tilde{0} \), and in the reacting ones,

\[
\tilde{S} = \set{0, 0, 0, 0, \hat{\omega}_\alpha; \alpha = 1, \ldots, N_{sp}}
\]
In this thesis, experiments are presented focused on the following questions by S.B. Pope [107]: “Is LES a physical model, a numerical procedure or a combination of both?” and “How are different LES models to be appraised?”. Different LES strategies are evaluated and compared for the numerical simulation of high-speed non-reacting and reacting air/H₂ jets. Widely studied simple turbulent jet case is chosen as the flow prototype. The complexities like multi-injection or chamber geometry are excluded. The focus is shifted to the behavior of the numerical approaches on the main problems of mixing, transition and interaction of turbulence with chemistry [28].

The simple turbulent jet is a good flow prototype for the evaluation of MILES and LES, indeed;

- High-speed compressible flows do usually involve shocks. Consequently, it is mandatory to apply dissipative numerical methods for the simulation of these flows.

- The time scales are at the order less than one millisecond. Considering that the turbulent mixing and chemical reaction are included in the problem, the parameters questioned in the simulation (potential core length, ignition length ...) are sensitive to the molecular transport.

The conditions for the test cases are based on the experimental data published in the literature. The non-reacting case is taken from the experiments of J.M. Eggers [32], and the reacting one is from the experiments conducted at LAERTE supersonic combustion chamber of ONERA [19, 90]. These free shear flows are selected at first because Kelvin-Helmholtz instabilities, primarily responsible for large scale turbulent mixing, are not much influenced by diffusion,
and because large scale turbulent mixing is determinant for non-premixed combustion which occurs at the molecular level due to micro-mixing or diffusion of reactants. The non-reacting test case isolates the first aspect and the reacting one includes both aspects.

In both cases, complete inlet profiles, necessary for constructing a simulation, are not available. Also, in the available data significant inconsistencies are reported. Therefore, approximate profiles that include assumptions are applied for the inlet conditions. It is pragmatically assumed that LES method (of dissipative schemes) is not sensitive to the accuracy of the inlet conditions as much as RANS approach does [22].

Numerical simulations are performed using a 5th order WENO scheme\(^1\), with resolutions ranging from \(32 \times 32 \times 128\) to \(256 \times 256 \times 1024\), with (LES) and without (MILES) explicit sub-grid model. MILES simulations are carried out for the Navier-Stokes equations and for the Euler equations, i.e. without any molecular transport terms. LES simulations include molecular transport terms.

The sensitivities of different numerical approaches to initial and boundary conditions are also assessed. It is intended to obtain experience in solving high speed reacting flows like those existing in scramjet engines, beyond the purpose of qualitative comparison.

In this chapter, LES solutions of non-reacting and reacting compressible H\(_2\)/air jets for different approaches are presented. In each case, first the flow problem is explained in detail including available experimental data and parameters for numerical simulations. Later, results showing interpretation of flow structure for straightforward visualization and statistical results for more thorough analysis are presented. The simulation results are also compared with the published experimental data.

5.1 NON-REACTING CASE

5.1.1 TEST CASE DESCRIPTION

The first simulated test case is a non-reacting transonic round H\(_2\) jet which is in a co-flowing supersonic round air jet. This coaxial mixing experiment was conducted in 1971 by Eggers [32]. The simple schematic diagram of the experiment is shown in Figure 5.1. The internal

\(^1\) Although WENO schemes are not monotone, MILES will be used to designate numerical LES in this thesis.
and external diameters of hydrogen injector are 11.6 mm and 12.7 mm, respectively. The
diameter of the co-flowing air nozzle is 152 mm. Mixing occurs in unconfined region, so the
flow static pressure is close to atmospheric pressure. The dimensions of the flow domain to
be simulated are $L_y \times L_z \times L_x = 70 \times 70 \times 700 \text{ mm}^3$, $x$ being the axial direction of the jet
starting from the injector exit. The inner shear layer is assumed not to be affected by the outer
shear layer for the region of interest.

Primary variables which have been measured by Eggers are pitot pressure, total temperature

\[ P_{\text{tot}} = \frac{(y + 1)}{2 M^2} \left[ \frac{\gamma + 1}{2 \gamma M^2 - (\gamma - 1)} \right]^{\frac{1}{\gamma - 1}} \]

and from the basic isentropic relations for a subsonic flow. The local values of the total
temperature are computed knowing the total temperatures in the pure H$_2$ and in the pure air
streams, using an energy balance as

\[ T_{\text{tot}} = \frac{C_p H_2 Y_{H_2} T_{\text{tot,H}_2} + C_p \text{air}(1 - Y_{H_2}) T_{\text{tot,air}}}{C_p H_2 Y_{H_2} + C_p \text{air}(1 - Y_{H_2})} \]

Figure 5.1: Schematic diagram of Eggers jet.
The static temperature was then deduced from the total temperature and the Mach number. Finally, the velocity was obtained from the local Mach number and the speed of sound. Eggers indicated a [-16%, +4%] uncertainty in the integrated H2 mass flow rate over transverse planes. The reliability of the velocity data is also affected by experimental uncertainty. The results of this experiment are frequently used in the literature for the validation of compressible RANS solvers [22, 28, 33].

### 5.1.2 PHYSICAL AND NUMERICAL PARAMETERS

Physical parameters of the flow are gathered in Table 5.1. The subscript jet stands for H2 jet conditions. Rej in the Table is the Reynolds number per meter. Data taken from report of Eggers are written in bold. The other parameters of the initial field are computed assuming a constant static pressure of 100 kPa and using the thermodynamic properties explained in chapter 2. A guess-and-try technique is used to find the static temperature that gives the experimental total temperature, knowing the Mach number in both streams. The characteristics of the flow may be estimated depending on parameters as density (ρjet/ρco-flow) and velocity (Ujet/Uco-flow) ratios. The convective Mach number Mc is suggested by Bogdanoff [8] for parametrization of compressibility. It is the Mach number of the free streams adjacent to the shear layer, relative to the coherent structures velocity Uc (Figure 5.2). The convection velocity Uc is given as

\[
U_c = \frac{c_2}{c_1 + c_2} \frac{U_1 + c_1 U_2}{U_1 + c_1 U_2}
\]  

Table 5.1: Physical parameters for the non-reacting air/H2 jet. In bold, data from Eggers [32], other values are computed.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>H2 jet</th>
<th>air co-flow</th>
</tr>
</thead>
<tbody>
<tr>
<td>U (m/s)</td>
<td>1074</td>
<td>394</td>
</tr>
<tr>
<td>Tstat/Ttot (K)</td>
<td>260/300 6/12</td>
<td>222/300 3/2</td>
</tr>
<tr>
<td>Pstat/Ptot (kPa)</td>
<td>100/167</td>
<td>100/285</td>
</tr>
<tr>
<td>Mach</td>
<td>0.886</td>
<td>1.32</td>
</tr>
<tr>
<td>µ (kg.m⁻¹.s⁻¹)</td>
<td>0.878 10⁻⁵</td>
<td>1.687 10⁻⁵</td>
</tr>
<tr>
<td>ρ (kg.m⁻³)</td>
<td>0.093</td>
<td>1.563</td>
</tr>
<tr>
<td>Rej (1/m)</td>
<td>11.6 10⁶</td>
<td>36.6 10⁶</td>
</tr>
<tr>
<td>Rejet</td>
<td>1.34 10⁵</td>
<td></td>
</tr>
<tr>
<td>Mc</td>
<td>0.44</td>
<td></td>
</tr>
</tbody>
</table>
Figure 5.2: Shear layer convective velocity and Mach numbers[8].

The shear layer convective Mach number is calculated by equating the convective Mach numbers of adjacent flows assuming constant specific heat ratio ($\gamma_1 = \gamma_2$)

$$\frac{U_1 - U_c}{c_1} = \frac{U_c - U_2}{c_2} \Rightarrow M_c = \frac{U_1 - U_2}{c_1 + c_2}$$

(5.4)

As in the current case of low convective Mach numbers, the assumption for the equality of $M_{c_1}$ and $M_{c_2}$ is relatively accurate [100]. It is important to note that convective Mach number is one of the parameters for characterization of jet but not the only one. It is shown by Fediou and Lardjane that temperature and density ratio differentiate the characteristics of binary temporal shear layer even if the convective Mach number is constant [41].

It is interesting to have an estimate, or at least an order of magnitude of the range of turbulent scales to be resolved. Large scales of the flow are produced by initial Kelvin-Helmholtz inflexional instability at the jet/co-flow interface, as illustrated in Figure 5.3. From experimental velocity profiles, the end of the potential core is at about 5.5 jet diameters, i.e. $x - x_0 \approx 64$ mm. At this location, the estimated shear layer width $\delta(x) = \delta_0 + \delta'(x - x_0)$ is about 8mm.
(Figure 11 (a) of [32]), giving the scale of the largest eddies. This value corresponds to a growth rate \( \delta' \approx 0.12 \), coherent with the literature [26] for compressible, high density ratio shear layers. The velocity scale of the largest eddies is typically half the velocity difference, \( \Delta U/2 = 170 \) m/s. Taking an average viscosity and an average density for the values listed in Table 5.1, one finds the large-scale Reynolds number as

\[
Re_{(\Delta U/2, \delta)} = \frac{\pi \Delta U/2 \delta}{\mu} \approx 88000 \tag{5.5}
\]

Hence, the Kolmogorov scale is about \( \eta \approx \delta \times Re_{(\Delta U/2, \delta)}^{-3/4} \approx 0.0015 \) mm and the Taylor microscale is about \( \lambda \approx \delta \times Re_{(\Delta U/2, \delta)}^{-1/2} \approx 0.027 \) mm.

The grid resolution is indicated as \( N_y \times N_z \times N_x \). Simulations are performed for 4 different grids namely 32\( \times \)32\( \times \)128, 64\( \times \)64\( \times \)256, 128\( \times \)128\( \times \)512 and 256\( \times \)256\( \times \)512. In the latter case, the computational domain is shortened to half length, i.e. \( L_x = 350 \) mm. The grid is refined in the transverse direction toward the jet centerline, using analytical metrics explained previously in section 4.6.4 with \( z \in [-L_z/2, +L_z/2] \) for \( Z \in [-1, +1] \). The size parameter \( L \) is chosen such that the ratio of the longest to the shortest cell is 5. The grid is also refined in the axial direction toward the inlet, with a ratio of 6. The 2D views of 32\( \times \)32\( \times \)128 resolution grid are shown in Figure 5.4. For the fine grid, the shortest grid cell is \( \Delta x_{min} = 0.323 \) mm and \( \Delta y_{min} = \Delta z_{min} = 0.160 \) mm. Assuming \( \Delta \approx \Delta y_{min} \), gives \( \Delta/\eta \approx 100 \) and \( \Delta/\lambda \approx 6 \). Hence, even on the fine grid, the simulation is far from a DNS. The numerical cut-off is probably in the inertial range. The approximations of the scales for all grid resolutions are listed in Table 5.1.2.

The inlet velocity profiles for all grids are shown in Figure 5.5. The profiles are fitted to the
Table 5.2: Ratios of approximate Kolmogorov scale and Taylor microscale to the minimum grid spacing.

<table>
<thead>
<tr>
<th>Grid</th>
<th>$\Delta x_{min}$</th>
<th>$\Delta y_{min}$</th>
<th>$\Delta/\eta$</th>
<th>$\Delta/\lambda$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$32 \times 32 \times 128$</td>
<td>$0.303 \times 10^{-2}$</td>
<td>$0.132 \times 10^{-2}$</td>
<td>880</td>
<td>49</td>
</tr>
<tr>
<td>$64 \times 64 \times 256$</td>
<td>$0.151 \times 10^{-2}$</td>
<td>$0.612 \times 10^{-3}$</td>
<td>408</td>
<td>23</td>
</tr>
<tr>
<td>$128 \times 128 \times 512$</td>
<td>$0.754 \times 10^{-3}$</td>
<td>$0.322 \times 10^{-3}$</td>
<td>215</td>
<td>12</td>
</tr>
<tr>
<td>$256 \times 256 \times 512$</td>
<td>$0.323 \times 10^{-3}$</td>
<td>$0.161 \times 10^{-3}$</td>
<td>107</td>
<td>6</td>
</tr>
</tbody>
</table>

Eggers experimental results using hyperbolic tangent equation 5.6

$$f = \frac{f_{\text{max}}}{2} \left(1 + \tanh \left[A \left(\frac{B r C}{E} + \frac{1}{r C - D}\right)\right]\right)$$

$$r = \left(y^2 + z^2\right)^{1/2}$$

(5.6)

The coefficients for the velocity profiles (co-flow/jet) and for the air mass fraction profile are listed in Table 5.3. Air is assumed to be composed of $N_2$ and $O_2$ in the atmospheric concentration ratio. Hydrogen mass fraction is computed full-filling the sum of all species mass fractions to unity. While adopting the hyperbolic tangent profiles the following issues are considered:

- The $H_2$ jet and air jet profile must be distinct at pitot region of nozzle. As there exists high density and velocity ratio. Otherwise, air at the speed of hydrogen jet may cause significant momentum injection at shear layer.

- At the pitot region, as the gradients of inlet profiles are high, the number of nodes

![Figure 5.5: Inlet velocity profile for the non-reacting air/H$_2$ Eggers jet.](image-url)
must be more than the WENO sub-stencil size. Otherwise the scheme will smooth the profiles and even out the effect of pitot at inlet.

This slight modification of the initial field causes some error in the volumetric flow rate which is $6 \times 10^{-2}$ (m$^3$/s) (computed by trapezoidal rule) for the experimental profiles. The error is at most 10% for the 64x64x256 grid and around 5% for the other resolutions. This is within the uncertainty limits of Eggers data.

In order to trigger the transition to turbulence, a random noise is applied for the $y$ and $z$ velocity components. First, for each line 20 random values are generated using the portable random number generator presented in the literature [68, 89]. By applying back Fourier transform a function is obtained using these random numbers. The noise which is confined by Gaussian distribution in the H$_2$ central jet, vanishes in the air co-flow (Figure 5.6) The normalized weight of the Gaussian distribution Conf is computed as

$$\text{Conf} = a e^{-\frac{r^2}{2bL^2}}$$

(5.7)

The amplitude $a$ and parameter $b$ controlling the Gaussian shape are respectively taken to be 10 m/s and 0.04. $L$ is the size of the window for Gaussian shape (diagonal length of the YZ surface where the bruit is applied.). No other tunable parameter is introduced in the initial field. Non-reflecting boundary conditions are applied for open boundaries.

At the co-flow velocity, one crossing of the full computational domain takes 1.8 ms. At the three lowest resolutions, time averaging of the flow variables are computed for 0.7 ms after the simulation is allowed to progress for 2 ms from the initial field. Simulation with the finer

![Figure 5.6: Radial distribution of confinement function, samples of generated 'random' field and the noise applied on transverse velocity components.](image)
grid (half domain) is progressed for 0.82 ms before averaging for 0.15 ms only, due to the high computational cost (see appendix I).

5.1.3 FLOW STRUCTURE

In Figures 5.7 - 5.11, results of LES_SM, LES_SSF, MILES_NS and MILES_EULER simulations are shown from top to bottom. Vertical and horizontal contour maps are the projections of the instantaneous pressure fields and the instantaneous H\textsubscript{2} mass fraction fields in the corresponding symmetry planes, respectively. For a better visualization, y and z scales are magnified by a factor of two. These figures allow a straightforward comparison of the behavior of LES and MILES at different grid resolutions. Mixing and pressure fluctuations may be observed from the hydrogen mass fraction and the pressure contours, respectively.

The structure of the flow is displayed at all resolutions as the Q-criterion [29, 67] iso-surface which is commonly used for coherent structure identification. Q-criterion is the measure of domination of the strain $S^{*}_{ij}$ by the rotation $\Omega_{ij}$ which are the symmetric and the antisymmetric parts of the velocity gradient tensor, respectively.

$$Q = \frac{1}{2} (\Omega_{ij} \Omega_{ij} - S^{*}_{ij} S^{*}_{ij})$$  \hspace{1cm} (5.8)

where

$$\Omega_{ij} = \frac{1}{2} (u_{i,j} - u_{j,i})$$  \hspace{1cm} (5.9)

Q-criterion is also related to the Lagrangian of pressure for incompressible formulation, by the well known Poisson equation.

$$Q = -\frac{1}{\rho} \Delta p$$  \hspace{1cm} (5.10)

Results for $32\times32\times128$ grid resolution (Figure 5.7):

The SM model introduces excessive (more than physical) amount of sub-grid viscosity. This results in the damping of flow shear and also delays transition to downstream locations where the stretching of the grid may even out the linear instabilities. These solutions are not considered to be physical since the flow field is not resolved.

Results for $64\times64\times256$ grid resolution (Figure 5.8):

Although the spatial resolution is still not adequate at this grid level, the obtained results are more amended for physical interpretation. The difference of the LES_SM result compared to
the other solutions is apparent. Coherent structures for LES_{SSF} solution are of larger intensity since the explicit sub-grid model damps the small perturbations. For the LES_{SSF} solution, the large scale structures disappear approximately at the end of concentration potential core. MILES_{EULER} and MILES_{NS} simulations generate similar results. In the MILES_{NS} solution the light turbulent structures are further damped out. Disturbances which are present in the pressure field are associated with the flow instabilities can be called as Mach waves [94]. Hydrogen mixing starts to be observed at this resolution. The concentration potential core lengths for MILES_{EULER}, MILES_{NS} and LES_{SSF} are close.
Figure 5.7: Eggers jet results on 32x32x128 grid. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches.
Figure 5.8: Eggers jet results on 64x64x256 grid. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches.
Results for 128×128×512 grid resolution (Figure 5.10): The solutions at the initial core of the jet for MILES_{EULER}, MILES_{NS} and LES_{SSF} are identical at 128×128×512 resolution. After the transition, the structures displayed as Q-criterion iso-surfaces are damped along the axial direction earlier in the LES_{SSF} solution. The sensor (3.67) initially switch off the SSF model in the initial core of the jet where the gradients are naturally high and later, when the flow is 3D enough, model is switched on to diffuse turbulence effectively. Concentration potential core lengths of all numerical approaches are identical and shorter than the ones for 64×64×256 grid solution. LES_{SM} solution is observed to be more dissipative at this resolution. This solution is more comparable with lower resolution of 64×64×256 grid MILES_{EULER}, MILES_{NS} solutions in Figure 5.8. Simulation for lower Smagorinsky constant (half) is conducted for understanding the effect of the SM model (Figure 5.9). The solution at the initial core region became identical to the solutions for other numerical approaches and coherent structures propagate downstream for the LES_{SM} solution which is sensitive to the Smagorinsky constant.

Results for 256×256×512 grid resolution (Figure 5.11):

Figures at fine grid resolution are almost identical to the solutions for 128×128×512. The LES_{SM} solution is closer to the solutions of the other approaches however the flow field at the beginning of the shear layer is still very much affected by the model. The pressure waves are more clustered compared to the results obtained with lower grid resolutions.

Figure 5.9: Eggers jet result on 128×128×512 grid with reduced Smagorinsky constant. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches.
Figure 5.10: Eggers jet results on 128×128×512 grid. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches.
Figure 5.11: Eggers jet results on 256×256×512 grid. Instantaneous structure of the flow visualized using iso-Q. Pressure and Hydrogen mass fraction in the symmetry planes for different approaches.
5.1.4 STATISTICAL RESULTS

The center-line, time-averaged, $x$-velocity distribution (axial decay) is shown in Figure 5.12, for the different simulations. Clockwise from top-left, sub-figures display LES_SM, LES_SSF, MILES_NS and MILES_EULER results for different resolutions. The lowest resolution $32 \times 32 \times 128$ is physically inadequate for all numerical approaches. Increasing the resolution to $64 \times 64 \times 256$ improves slightly the results, but the grid is still too coarse to capture the initial development of shear instabilities. They are damped by the numerical dissipation, and the transition is delayed far downstream. The worst case is observed for LES_SM (top-left) because the Smagorinsky model, can not distinguish the mean flow gradients from turbulent structures. Hence, the model is active and introduces additional dissipation in the laminar, high-shear, jet entrance region. Increasing again the grid resolution up to $128 \times 128 \times 512$ improves dramatically the results. Except for LES_SM with $\lambda^2 = 0.02$ which is still too dissipative, all the simulations show convergence toward the experimental results. MILES_EULER (bottom-left) and MILES_NS (bottom-right) are very close to each other, but MILES_NS is a little closer to the experiment at the end of the computational domain. The LES with explicit
SSF sub-grid model does not show any improvement compared to the MILES. At the highest resolution 256×256×512, results are further closer to the experimental data and grid convergence is almost achieved, even with MILES_EULER although there is no physical cutoff in this calculation. As mentioned above, the curves are a little wavy because time statistics are not perfectly converged. Transverse velocity profiles at four specific downstream locations investigated by Eggers are shown on the 128×128×512 and 256×256×512 grids in Figures 5.13 and 5.14 respectively. With the exception of LES_SM with $\lambda^2 = 0.02$, all numerical approaches under-predicts qualitatively, the spreading of the jet. On the finer grid 256×256×512 (Figure 5.14), results are further closer to the experimental data.

![Figure 5.13: Eggers jet : transverse velocity profiles at $x = 0.06, 0.11, 0.18$ and 0.29 m from the exit. Resolution 128×128×512](image)

Figure 5.15 shows the hydrogen mass fraction along the centerline. Although the agreement is not so good as for the dynamic field, most of the previous conclusions can still be observed again, here. One may have to keep in mind the uncertainty in the experimental data as well.

In order to evaluate the axial variation of a quantity $Q(x, y, z, t)$, it is appropriate to look at the variation of time averaged variable ($\bar{Q}(x, y, z)$) at the centerline ($Q_0(x)$) (Figure 5.17). The subscripts 0 and $\infty$ are respectively the centerline and far-field average values of the quantity.
Figure 5.14: Eggers jet: transverse velocity profiles at \( x = 0.06, 0.11, 0.18 \) and 0.29 m from the exit. Resolution 256×256×512 (Figure 5.16). \( x_0 \) is the fictitious center of the jet. Considering the above definitions the definition for an inverse normalized time averaged quantity \([106]\) at the centerline is

\[
Q_* = \frac{Q_{jet} - Q_\infty}{Q_0(x) - Q_\infty} \approx \frac{x - x_0}{B_Q d} \tag{5.11}
\]

\( B_Q \) is the spreading rate of quantity after the initial region of the jet. The variations of the inverse of quantities eventually appear to show a linear trend as shown in Figure 5.18. The axial normalized inverse of velocity and hydrogen mass fraction profiles are shown in Figure 5.19 for all grids except 32×32×128 grid. The \( B \) and \( x_0 \) values for the axial velocity and for the hydrogen mass fraction are listed respectively in Tables 5.4 and 5.5. These results show that at low resolution of 64×64×256, the potential core estimated far downstream and the spreading rate are larger. At low resolution the gradients of the inlet profile are smoother since there are few points at the pitot region of the inlet. High numerical diffusion, because of the very low resolution, enlarges the spreading angle. It is observed in the profiles shown on Figure 5.19 that the Smagorinsky model is not able to resolve most physical properties related to the problem. The velocity decay for MILES_EULER is one of the fastest which has an
early plateau around \(x=0.5\) m. The hydrogen concentration shows similar behavior. At higher resolutions of \(128 \times 128 \times 512\) and \(256 \times 256 \times 1024\) the simulations give similar results except for the Smagorinsky model. The Smagorinsky model causes transition delay by smoothing the gradients which are present in the initial region. However the velocity decay slopes of the solutions of Smagorinsky model are close to the MILES\_NS and LES\_SSF results. At \(64 \times 64 \times 256\) Smagorinsky gives unrealistic results as stated earlier in section 5.1.3. At high
resolution all cases give similar behavior as shown in Figure 5.19 and tabulated data (Table 5.4).

The H₂ mass fraction potential core values for all cases are smaller than the velocity potential core values, by a factor of 4 ± 2. The diffusion of momentum is more than the diffusion of species so that the overall Schmidt Sc number is less than unity.
Figure 5.19: The inverse normalized velocity $Q_u$ (left) and the inverse normalized H$_2$ mass fraction $Q_y$ (right) for 64$\times$64$\times$256, 128$\times$128$\times$512 and 256$\times$256$\times$512 resolutions (from upper to lower).
The variation of velocity, hydrogen mass fraction and pressure are computed and stored once these flow statistics have reached steady state, for different axial locations of the jet flow field.

The numerical probe locations are indicated over the instantaneous velocity contours on the symmetry axis (figures 5.25 and 5.26), the time variation of velocity at these locations and spectra of velocity data are also given on the same figures.

In the coarser grid, core flow tends to persist for a longer time downstream the computational domain. In figure 5.25, the coherent structures in the flow are more pronounced in MILES_NS and LES_SSF. Amplitude variation of oscillations for MILES EULER is large especially in the jet core region. This leads to an earlier end up of the velocity potential core before reaching two third of the computational domain. The end of velocity potential core for MILES_NS solution is in the second half of the domain. LES_SSF results are similar to MILES_NS results. The velocity spectra of LES_SSF and MILES NS also carry the same slope characteristics. This is also true for the solution on high resolution grid given in figure 5.26. The velocity data of MILES_EULER shows high amplitude fluctuations which is also
reflected in the corresponding velocity spectra. In LES_SM solution (figure 5.25) higher rate of dissipation is observed and oscillations are damped. Whenever the resolution increases, this effect disappears.

The passage frequency of linear instability waves are recognized in time variation figures. The size of the coherent structures is approximately 8 mm based on the observation of instantaneous velocity histogram along the shear region (figure 5.20). The convective velocity of these structures \( U_c \) is calculated to be approximately \( \sim 518.2 \) m/s (equation 5.3). So the passage frequency of the linear instabilities is roughly \( \sim 70 \) kHz in accordance with the peak frequencies of the velocity spectra displayed in figures 5.26.

Next two figures 5.27 and 5.28 relate to the hydrogen mass fraction time variations for two highest resolutions of \( 128\times128\times512 \) and \( 256\times256\times512 \). The concentration related results which are obtained by MILES_NS and LES_SSF are similar for both resolutions. Whereas the concentration potential core for MILE.Euler is shorter. It persists longer in LES_SM results at \( 128\times128\times512 \) grid resolution. Higher resolution results also show that the jet core persists for larger distance in LES_SSF and MILES_NS. Larger structures are present for LES_SM and MILES.Euler results.

Low resolution LES_SM does not capture the pressure wave fluctuations which do appear for higher resolution shown in figure 5.30. The LES SSF, MILES.Euler and MILES NS show

![Figure 5.20: Instantaneous axial velocity component on the jet axis at 256×256×512.](image)
a higher much resolved Mach wave clustering near the jet inlet. As a result, relatively smaller eddies and smaller fluctuations are present in the flow domain. The velocity spectra for all high resolution solutions look alike.

In figures for pressure (figure 5.29 and 5.30) there exists two peaks in time variation spectra. The frequency of the first peak is at the order of the linear instability passage frequency. This is related to the Mach waves radiated by the growth and decay of linear instability waves [94]. The frequency of these waves is the same for all numerical approaches.

The time series taken are used to compute second order moments for the resolved velocity field \( \tilde{u}_i \) and the Nitrogen mass fraction \( \tilde{Y}_{N_2} \), at the probe locations. The resolved kinetic energy \( \hat{k}(\vec{x}, t) \) in (3.48) written using the Reynolds decomposition, e.g. \( \tilde{u}_i = \langle \tilde{u}_i \rangle + \tilde{u}'_i \) is

\[
\hat{k}(\vec{x}, t) = \frac{1}{2} \langle \tilde{u}_i \rangle \langle \tilde{u}_i \rangle + \langle \tilde{u}'_i \rangle + \frac{1}{2} \tilde{u}'_i \tilde{u}'_i
\] (5.12)

where \( \langle \tilde{\cdot} \rangle \) represents time average. The values of numerically resolved turbulent kinetic energy \( \hat{k}' = \frac{1}{2} \langle \tilde{u}'_i \tilde{u}'_i \rangle \) and the Nitrogen mass fraction variance \( \langle \hat{Y}'_{N_2} \rangle \) are plot in Figures 5.21-5.22 and Figures 5.23-5.24, respectively for 128\( \times \)128\( \times \)512 and 256\( \times \)256\( \times \)512 grid solutions. Decay of turbulent kinetic energy observed in Figure 5.21. As the resolution increase the results make more sense. MILES produces higher second order statistics than LES in the early development of turbulent processes (probe at \( x = 0.11 \) m, Figures 5.23 and 5.24). Mixing in LES_SM, associated with the peak in \( \langle \hat{Y}'_{N_2} \rangle \), is delayed downstream compared to other simulations. Since small-scales are dissipated by the SM model, large energy-containing eddies dominate in LES_SM. Hence, both \( \hat{k}' \) and \( \langle \hat{Y}'_{N_2} \rangle \) overtake the values of other solutions in the downstream region of the jet. The peak turbulent Mach number \( M_{RANS} = (\tilde{p}\hat{k}'/\gamma \bar{p})^{1/2} \) is about 0.1 in the H\(_2\) jet. This result justifies the assumption that the isotropic part (3.61) of the SGS term (3.28) is negligible since the isotropic term can be written as \( A_{ij}^i = \gamma M_{sgs}^2 \bar{p} \) [34].
Figure 5.21: Eggers jet: Resolved turbulent kinetic energy, resolution 128×128×512.

Figure 5.22: Eggers jet: Nitrogen mass fraction variance, resolution 128×128×512.

Figure 5.23: Eggers jet: Resolved turbulent kinetic energy, resolution 256×256×512.

Figure 5.24: Eggers jet: Nitrogen mass fraction variance, resolution 256×256×512.
Figure 5.25: Sub-figures demonstrating from top: instantaneous velocity over symmetry axis, sampled time variation and spectra of sampled data of 128x128x512 grid solutions (velocity in m/s and frequency in kHz).
Figure 5.26: Sub-figures demonstrating from top: instantaneous velocity over symmetry axis, sampled time variation and spectra of sampled data of 256×256×512 grid solutions (velocity in m/s and frequency in kHz).
Figure 5.27: Sub-figures demonstrating from top: instantaneous hydrogen mass fraction over symmetry axis, sampled time variation and spectra of sampled data of 128×128×512 grid solutions (frequency in kHz).
Figure 5.28: Sub-figures demonstrating from top: instantaneous hydrogen mass fraction over symmetry axis, sampled time variation and spectra of sampled data of 256×256×512 grid solutions (frequency in kHz).
Figure 5.29: Sub-figures demonstrating from top: instantaneous pressure over symmetry axis, sampled time variation and spectra of sampled data of 128×128×512 grid solutions (pressure in Pa and frequency in kHz).
Figure 5.30: Sub-figures demonstrating from top: instantaneous pressure over symmetry axis, sampled time variation and spectra of sampled data of 256×256×512 grid solutions (pressure in Pa and frequency in kHz).
5.1.6 IS THE DNS LEVEL REACHED?

The question that naturally arises is to know if the explicit sub-grid model is still active at the finest grid resolution. An appropriate numerical indicator is the eddy-viscosity ratio $\mu_{\text{sg}}/\mu$. In a free shear flow, one can consider that for $\mu_{\text{sg}}/\mu \lesssim 1$, the model is no more active. This would not be the case in the boundary layer for a wall-bounded flow where the flow exhibits large gradients. Figures 5.31 and 5.32 show respectively the instantaneous distribution of $\mu_{\text{sg}}/\mu$ for LES$_{\text{SM}}$ and LES$_{\text{SSF}}$. At resolutions $64 \times 64 \times 256$ and $128 \times 128 \times 512$, the models are very pronounced in both LES$_{\text{SM}}$ and LES$_{\text{SSF}}$. The effect of the laminarity sensor (3.67) in the SSF model is apparent, and the model is globally less dissipative than the Smagorinsky model, although instantaneous and local peak values may be higher. At the highest resolution $256 \times 256 \times 512$, the SM model is still active whereas the SSF model is almost vanished. Although the DNS limit is far from being reached as the ratio of minimum grid spacing to Taylor microscale is 6 (Table 5.1.2). This means that the numerical dissipation of the WENO scheme is responsible for most of the sub-grid modeling on the finer grid.

![Figure 5.31: Eddy-viscosity ratio, LES_SM.](image)

Figure 5.31: Eddy-viscosity ratio, LES_SM.
5.1.7 CONCLUSIONS

The main conclusion for this non-reacting test-case is that conventional LES does not show any superiority compared to MILES. The Smagorinsky model deteriorates the results due to its over-dissipative behavior. The SSF model is less dissipative, but does not improve the results, whatever the grid resolution is. MILES\textsubscript{EULER} and MILES\textsubscript{NS} give almost similar results, the molecular dissipation being far below the numerical one. This means that the computational effort involved in highly accurate transport models is simply wasted, when shock capturing schemes are used to solve the compressible Navier-Stokes equations. The results of MILES\textsubscript{EULER} surprisingly seem to show grid convergence toward the experimental data although no physical cutoff is present in the simulation. This is an open question for further research.
5.2 REACTING CASE

5.2.1 TEST CASE DESCRIPTION

Having investigated the mixing problem of a compressible H$_2$/air jet, the experimental results from LAERTE combustion chamber of ONERA [50, 86] are chosen as the reacting flow test case. This test facility is designed for the fundamental study of supersonic combustion and is intended to develop background data material on hypersonic air-breathing propulsion.

The experimental setup consists of:

- an initial section of the air pre-heater and the supersonic air nozzle for co-flow,
- the supersonic injector for nitrogen or fuel (H$_2$),
- the combustion chamber.

A sketch demonstrating the setup is given in Figure 5.33.

Co-flowing air is first heated by a primary heater up to 850 K then a secondary hydrogen flame heater is used to elevate the temperature up to 1850 K. Before entering into the chamber, the co-flow is enriched with oxygen to the atmospheric concentration (%21). The supersonic air nozzle accelerates the flow to Mach 2. The vitiation of the co-flow with water, by the flame heater, is around %16 (molar fraction). Total flow rate of vitiated air is 0.65 kg/s and the static pressure is around 80 kPa. Hydrogen is injected through a round supersonic injector of inner and outer diameters of 6 mm and 10 mm, respectively. The fuel flow rate is 6.2 g/s at Mach 2 and the static pressure is also around 80 kPa. The combustion chamber starts with a 45x45 mm
constant section part of 370 mm then the section diverges at an angle of 1.15° equal at upper and lower surfaces for 500 mm along the axis. Optical access is provided on the sides of the chamber using silica windows for measurements.

Since the injector and the air nozzle are subject to elevated temperatures, they are cooled with water. The experiment can be conducted for short durations at the order of seconds (15 s for H₂ combustion [22]).

Available experimental results which are obtained within the combustion chamber are listed below [19]:

- pressure measurements are taken at different axial locations on both upper and lower wall surfaces.
- radial temperature profiles are measured at at \( x = 10 \text{mm}, x = 72 \text{mm}, x = 210 \text{mm} \) and \( x = 288 \text{mm} \).
- radial OH concentration distribution is measured by planar laser induced florescence (PLIF) technique at \( x = 210 \text{mm} \).
- velocity measurements are taken applying LDA or PIV methods at \( x = 10 \text{mm}, x = 210 \text{mm} \) and \( x = 288 \text{mm} \).
- the ignition position is reported to be roughly at \( x = 17 \text{cm} \)

The data available are shown in Figure 5.34. In the present study, only the first constant section part is simulated.

![Figure 5.34: Experimental data available. From left to right: temperature, velocity and OH molar concentration.](image-url)
Computations are performed over a domain with the dimensions 45×45×350 mm³. The domain is the section of the combustion chamber located after the end of hydrogen injector. The following assumptions are applied on the computational domain for the sake of simplicity.

- Slip wall (symmetric) boundary conditions are applied on the sides of the domain, to avoid the necessity of resolving wall boundary layers.
- Divergent section at the last 13 mm of the computational domain is not considered.

The purpose is not to reproduce strictly the experimental data numerically, so simplifications are applied despite their significant effect.

5.2.2 INLET FIELD

There exists no detailed quantitative experimental flow field measurement at the inlet. It is reasonable to fit a hyperbolic tangent profile for the velocity of the pure hydrogen injection that satisfies the mass flow rate measurement. However, the initial field of variables for the co-flow, especially the species mass fractions are not easy to predict. Davidenko simulated the hydrogen heater and the co-flow nozzle in his work in order to generate inlet BCs for the combustion chamber problem [22]. 1D profiles (Figure 5.35) for the radial distribution of the primitive variables, at the inlet of the combustion chamber are taken from these solutions.

This RANS simulation was conducted with the initial flow composition of 0.542 kg/s air, 0.1 kg/s O₂ and 8 g/s H₂ at total temperature of 850 K at the inlet of the hydrogen heater. Calculated profiles by Davidenko for the inlet field are in good agreement with the total temperature
data measured very near the inlet, located at x=10 mm (shown in Figure 5.35). Also, the estimated co-flow inlet oxygen concentration is at atmospheric level (%21), showing agreement with reported experimental data [24]. Therefore, it is assumed that the inlet conditions applied in this section are adequate.

Two set of initial fields are deduced from the results of Davidenko. The simulations with first simple profile intended to the comparison of numerical approaches. The second inlet profile is more complex, including the radial velocity and pressure profiles. These complexities induce strong shocks within the flow which are reflected at the walls and affect the flowfield. Sensitivity of the solution to the inlet field conditions is investigated with this improved inlet.

5.2.2.1 SIMPLE PROFILE

Using a simplified initial field is generated fitting hyperbolic tangent profiles over the given initial conditions, (equation 5.6) similar to the Eggers jet. The walls are subject to sudden temperature change and also the wall boundary layer can have significant effect on the flow inside. Therefore, Davidenko applied various conditions for the walls. The profiles considered in this study are corresponding to adiabatic wall boundary conditions for the cooled nozzle and hydrogen injector. However, to be able to include wall effects a high resolution is necessary around such flow domain. As an alternative, symmetric boundary conditions are applied on wall at low resolution and the velocity gradients at the wall boundaries are removed. As such, the profiles are modified to be normal at the wall boundaries. Also, the radial component of the velocity and the pressure profiles which lead to strong shock patterns are neglected. This also allows to single out the analysis of turbulent flame without occurrence of shock.

Figure 5.36: Simplified initial field data deduced from RANS solutions by Davidenko [22]; from left to right radial profiles for temperature, velocity and species concentrations.
turbulence interaction effects.

The simple profiles of temperature, velocity and species concentrations (except radicals) are shown in Figure 5.36. The chemical composition of the vitiated air co-flow also includes radicals (not shown in Figure 5.36): \( Y_{\text{OH}} = 2.285 \times 10^{-4} \), \( Y_{\text{O}} = 1.8 \times 10^{-4} \) which have an important effect on ignition.

The coefficients for hyperbolic tangent profiles used in equation 5.6 for the jet and the co-flow are given in Table 5.6. In the Figure 5.37, the jet core region inlet velocity profiles are shown with different resolutions. The shear layer is shown to be highly resolved especially with the fine grid.

Table 5.6: LAERTE inlet hyperbolic tangent profile coefficients for equation 5.6.

<table>
<thead>
<tr>
<th>var</th>
<th>( U_{\text{jet}} )</th>
<th>( U_{\text{co-flow}} )</th>
<th>( Y_{\text{air}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( f_{\text{max}} )</td>
<td>1970</td>
<td>1400</td>
<td>1</td>
</tr>
<tr>
<td>A</td>
<td>9</td>
<td>0.8</td>
<td>12</td>
</tr>
<tr>
<td>B</td>
<td>10</td>
<td>0.75</td>
<td>1</td>
</tr>
<tr>
<td>C</td>
<td>10</td>
<td>-1.33</td>
<td>-1.11</td>
</tr>
<tr>
<td>D</td>
<td>1</td>
<td>-1</td>
<td>0</td>
</tr>
<tr>
<td>E</td>
<td>1</td>
<td>-1</td>
<td>-1</td>
</tr>
</tbody>
</table>

Figure 5.37: Inlet velocity profile for the reacting air/H\(_2\) LAERTE jet.

### 5.2.2.2 IMPROVED PROFILE

The simulations are conducted with more realistic conditions that also include radial velocity and pressure profiles at the inlet. These additional profiles generate strong shocks through the flow. The resolution of the wall shear layer is not tractable even with the fine grid spacing so the velocity decay on the walls is not included in these simulations. The additional radial flow field and pressure profiles computed by Davidenko normal to the axis of the jet is shown on the set of Figures 5.38. The concentrations of the radicals (not shown in the Figure) in the co-flow are taken to be: \( Y_{\text{OH}} = 2.285 \times 10^{-4} \), \( Y_{\text{O}} = 1.8 \times 10^{-5} \) and \( Y_{\text{H}} = 8.2 \times 10^{-7} \).
Figure 5.38: Initial field data deduced from Davidenko RANS solutions [22] including shock patterns at inlet; from left to right radial profiles for temperature, velocity and species concentrations.
5.2.3 NUMERICAL PARAMETERS

5.2.3.1 FLOW PARAMETERS

The flow conditions at the inlet of the chamber are summarized in Table 5.7.

Table 5.7: LAERTE experimental conditions.

<table>
<thead>
<tr>
<th></th>
<th>H₂ jet</th>
<th>air co-flow</th>
</tr>
</thead>
<tbody>
<tr>
<td>U (m/s)</td>
<td>1970</td>
<td>1336</td>
</tr>
<tr>
<td>Tₜₚₜ/Tₜₜₜ (K)</td>
<td>160/300</td>
<td>1200/1850</td>
</tr>
<tr>
<td>Pₚₚₚₜ/Pₜₜₜ (kPa)</td>
<td>80/680</td>
<td>80/720</td>
</tr>
<tr>
<td>Mach</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>(\dot{q} ) (g/s)</td>
<td>6.2</td>
<td>650</td>
</tr>
<tr>
<td>(\mu ) (kg.m⁻¹.s⁻¹)</td>
<td>0.729 10⁻⁵</td>
<td>0.729 10⁻⁵</td>
</tr>
<tr>
<td>(\rho ) (kg.m⁻³)</td>
<td>0.104</td>
<td>0.283</td>
</tr>
<tr>
<td>Reₖ</td>
<td>2.05 10⁵</td>
<td></td>
</tr>
<tr>
<td>Mₖ</td>
<td>0.39</td>
<td></td>
</tr>
</tbody>
</table>

The variables that are written in bold are taken from experiments while the remaining ones are calculated.

5.2.3.2 SPATIAL PARAMETERS

Grid resolutions are 64×64×256, 128×128×512 and 256×256×1024 with the same clustering characteristics of grid nodes as for the Eggers jet (equation (4.53)). The size of the smallest cell for the fine grid is \(\Delta x_{\text{min}} = 0.216 \text{ mm} \) and \(\Delta y_{\text{min}} = \Delta z_{\text{min}} = 0.110 \text{ mm} \). Assuming the size of the large scales to be approximately 5 mm in the mixing region, a calculation similar to equation (5.5) gives \(Re(\Delta U/2, \delta) \approx 1300\), a Kolmogorov scale \(\eta \approx 0.023 \text{ mm}\) and a Taylor micro-scale \(\lambda \approx 0.14 \text{ mm}\). The resolution characteristics for all grids are listed in Table 5.8. Comparing non-reacting Eggers and reacting LAERTE test cases, it is seen that the flow field is more resolved for reacting test case on the fine grids.
Table 5.8: Ratios of approximate Taylor micro-scale and Kolmogorov scale to the minimum grid spacing.

<table>
<thead>
<tr>
<th>Grid</th>
<th>$\Delta x_{\text{min}}$</th>
<th>$\Delta y_{\text{min}}$</th>
<th>$\Delta/\eta$</th>
<th>$\Delta/\lambda$</th>
</tr>
</thead>
<tbody>
<tr>
<td>64×64×256</td>
<td>0.865×10^{-2}</td>
<td>0.450×10^{-3}</td>
<td>20</td>
<td>3.2</td>
</tr>
<tr>
<td>128×128×512</td>
<td>0.430×10^{-3}</td>
<td>0.223×10^{-3}</td>
<td>10</td>
<td>1.6</td>
</tr>
<tr>
<td>256×256×512</td>
<td>0.226×10^{-3}</td>
<td>0.110×10^{-3}</td>
<td>5</td>
<td>0.8</td>
</tr>
</tbody>
</table>

5.2.3.3 TIME PARAMETERS

The chemical source terms are non-linear, so that it is not possible to deduce analytical restriction for time step. The maximum time step constraint $\Delta t_{\text{chem}}$ is validated by numerical 0D tests. Since the current problem is in more or less atmospheric conditions, it is proper to conduct the tests for constant pressure ($dp/dt = 0$). Time derivative of pressure is related to the enthalpy

$$\rho \frac{dh}{dt} = \frac{dp}{dt}$$

(5.13)

When the enthalpy is written explicitly, it yields

$$\rho C_p \frac{\partial T}{\partial t} = - \sum_{\alpha=1}^{N_p} \left( \Delta h_{\alpha}^0 + \int_{T_0}^{T} C_{p,\alpha}(\theta) d\theta \right) \dot{\omega}_{\alpha}$$

(5.14)

0D tests are conducted by iterating the time step for pure stoichiometric and vitiated stoichiometric ($Y_{H_2,O}=0.1124$) mixtures. Initial pressure and temperature values are taken to be respectively 80 kPa and 1500 K. Third order Runge-Kutta method is applied for time integration. The variation of temperature and species mass fractions are shown in Figure 5.39.

Figure 5.39: 0D constant pressure tests for stoichiometric H$_2$/air (left) and stoichiometric vitiated H$_2$/air (right) mixtures. The dashed lines refer to the stable solutions and the solid lines to the unstable ones.
The dashed lines refer to stable solutions. The solid lines refer to the unstable solutions. The maximum time step for both pure and vitiated H$_2$-air mixtures is $\Delta t_{chem} = 1.3 \times 10^{-7}$.

However, the 0D calculations with constant volume conditions are more severe than constant pressure. Also, it is more representative of the flow enclosed with walls including pressure jumps due to shocks. 0D constant volume ($\frac{dv}{dt} = 0$) conditions refer to constant energy condition

$$\frac{dv}{dt} = 0 \Rightarrow \frac{de}{dt} = 0 \quad (5.15)$$

When the enthalpy is written explicitly, it yields

$$\rho C_v \frac{dT}{dt} = - \sum_{\alpha=1}^{N_g} \left( \Delta h_\alpha^0 + \int_{T_0}^{T} C p_\alpha(\theta) d\theta - \frac{RT}{M_\alpha} \right) \dot{\omega}_\alpha \quad (5.16)$$

0D constant volume tests (Figure 5.40) are conducted with the same conditions of constant pressure tests. Both constant volume computations for stoichiometric H$_2$/air (left) and stoichiometric vitiated H$_2$/air (right) mixtures are stable with the maximum time step constraint of $\Delta t_{chem} = 0.6 \times 10^{-7}$ s. Considering that $\Delta t_{chem}$ reduces by increasing initial pressure and temperature, the chemical constraint for time step is chosen to be $\Delta t_{chem} = 0.2 \times 10^{-7}$ s [58].

In application, at high resolution of $256 \times 256 \times 1024$, $\Delta t_{hydro}$ which is computed out of CFL condition, is less than the chemical time step limit.
5.2.4 RESULTS

5.2.4.1 RESULTS FOR SIMPLE INLET PROFILE

The LAERTE chamber simulation results, with the simple inlet field (section 5.2.2.1), are presented in this section. In summary, the inlet radial velocity and the inlet pressure profiles, estimated numerically by Davidenko for the experiment, are not applied for the simulations that are presented here. All the inlet profiles are smoothed and mass fraction of the radical O is taken to be more than the numerical estimations by one order. Because of poor performance of Smagorinsky model in non-reacting case, only LESSSF, MILES_NS and MILES.Euler are considered.

5.2.4.1.1 FLAME STRUCTURE

The flame structure is analyzed from the fuel/oxidizer mixture fraction

\[ \tilde{z} = \frac{Z - Z_{\text{co-flow}}}{Z_{\text{jet}} - Z_{\text{co-flow}}} \]  

(5.17)

\[ Z = s Y_{\text{H}_2} - Y_{O_2} \]  

(5.18)

\[ Z \] is the first of the three Schwab-Zeldovitch variables for a single-step reaction (e.g. [104] p.84), and \( s = Y_{O_2} / Y_{\text{H}_2} |_{st} = 8 \) is the mass stoichiometric ratio for the H\(_2\)/O\(_2\) chemistry. Hence, in the pure H\(_2\) central jet

\[ Y_{\text{H}_2}^{\text{jet}} = 1 \ ; \ Y_{O_2}^{\text{jet}} = 0 \ ; \ Z^{\text{jet}} = 8 \ ; \ z = 1 \]  

(5.19)

and in the vitiated air co-flow

\[ Y_{\text{H}_2}^{\text{co-flow}} = 0 \ ; \ Y_{O_2}^{\text{co-flow}} = 0.2447 \ ; \ Z_{\text{co-flow}} = -Y_{O_2}^{\text{co-flow}} \ ; \ z = 0 \]  

(5.20)

Introducing boundary conditions (5.19)(5.20) for \( Z \) in equation 5.17, and making use of the flame equivalence ratio \( \phi_0 \) yields

\[ \phi_0 = s \frac{Y_{\text{H}_2}^{\text{jet}}}{Y_{O_2}^{\text{jet}}} = 32.69 \]  

(5.21)

\( z \) obtained by mixing the same amount mass of fluid taken in the fuel stream and in the oxidizer stream with incorporated equation 5.17 becomes

\[ z = s \left( \frac{Y_{\text{H}_2}}{Y_{O_2}} \right) - \left( \frac{Y_{O_2}}{Y_{O_2}^{\text{co-flow}}} \right) + 1 \]  

\[ 1 + \phi_0 \]  

(5.22)
The flame front is supposed to be where $Y_{H_2} = Y_{O_2} = 0$, i.e. at stoichiometric locations where

$$z_{st} = \frac{1}{1 + \phi_0} = 0.0297 \quad (5.23)$$

This value follows from the assumption of a unitary Lewis number in both fuel and oxidizer streams.

The distribution of fuel/oxidizer mixture fraction is showing over $128 \times 128 \times 512$ resolution MILES_NS solution data to have an idea of the distribution, in Figure 5.41. In Figures 5.42-

![Figure 5.41: Fuel/oxidizer mixture fraction distribution.](image)

5.44, 3D instantaneous iso-surface of the mixture fraction ($z_{st}$) at the stoichiometric value are plotted for LES_SSF, MILES_NS and MILES_EULER on different grid resolutions. The mixture fraction definition given in equation 5.22 is derived for one-step infinitely fast chemistry.

When finite rate chemistry is considered, other passive scalar derivations may be constructed using

- the atomic mass fractions [6, 7]

$$z = \frac{0.5(Z_H - Z_{H,O})/W_H - (Z_O - Z_{O,O})}{0.5(Z_{H,f} - Z_{H,O})/W_H - (Z_{O,f} - Z_{O,O})/W_O} \quad (5.24)$$

where $Z$ is the elemental mass fraction

- the mass fraction of an inert species, if there exists any. In current study, $N_2$ is the inert species.

$$z = \frac{Y_{N_2} - Y_{N_2}|_{jet}}{Y_{N_2}|_{co-flow} - Y_{N_2}|_{jet}} \quad (5.25)$$

A lifted diffusion flame can then be traced using passive scalar definition based on atomic mass fractions (equation 5.24). However, since the co-flow includes combustion products,
equations 5.24, 5.25 and 5.41 are giving similar stoichiometric iso-surfaces at the shear layer. The first definition is (equation 5.22) kept for mixture fraction definition for visualization.

Vertical and horizontal contour maps are the projections of the instantaneous temperature field and water mass fractions in the corresponding symmetry planes, respectively.

Increasing the grid resolution produces a more wrinkled flame. But for a given resolution, there are apparently no major differences between LES and MILES. Some subtle differences are present however. The solution at coarse grid of 64×64×256 (Figure 5.42) is reasonable, even though the gradients at shear region of inlet profiles are poorly resolved. Although there is no molecular transport except for the numerical diffusion the MILES_EULER solution gives qualitatively similar results when compared to MILES_NS and LES_SS. The flame region is designated as the warm color areas in the temperature and water mass fraction contour plots. It consists of a diffusion layer surrounding the reactive layer, embedded in the stoichiometric surface. The reactive layer at the jet/co-flow interface is wrinkled by large-scale turbulent eddies, which bring in contact reactants and mix them at the turbulent level. During that time, the finite-rate chemistry proceeds.
Figure 5.42: Instantaneous iso-surfaces of the stoichiometric mixture fraction, temperature contours and water mass fraction contours on the symmetry planes for 64×64×256 grid. From top to bottom: LES_SSF, MILES_NS, MILES_EULER.
Figure 5.43: Instantaneous iso-surfaces of the stoichiometric mixture fraction, temperature contours and water mass fraction contours in the symmetry planes for 128×128×512 grid. From top to bottom: LES_SSF, MILES_NS, MILES_EULER.
Figure 5.44: Instantaneous iso-surfaces of the stoichiometric mixture fraction, temperature contours and water mass fraction contours in the symmetry planes for $256 \times 256 \times 1024$ grid. From top to bottom: LES_SSF, MILES_NS, MILES_EULER.
5.2.4.1.2 STATISTICAL RESULTS

In the MILES,NS 256×256×1024 simulation, the turbulent time scale can be estimated from Figure 5.45. The Figure shows the instantaneous distribution of $U_x$ along two longitudinal lines one located on the jet axis ($r=0$ mm) and the other in the shear region ($r=3.2$ mm, wake of jet nozzle). At the end of the potential core, the turbulent length scale deduced from the mean distance between major peaks is about 1 cm or less. Applying the so-called Taylor hypothesis with a mean velocity of about 1500 m/s gives a turbulent time scale $\tau_t \approx 5.10^{-6}$ s. Figures 5.39 and 5.40 show the time evolution of the species mass fractions and the temperature in a premixed stoichiometric 0D air/H$_2$ flame at constant volume and at constant pressure conditions, respectively. In both cases the ignition delay is around $3.10^{-5}$s, and the time for the reaction to complete is $\tau_c \approx 5.10^{-5}$s. Hence, there is a strong interaction between chemistry and turbulence in a thick flame region since the estimated the Damköhler number is around

\[
Da = \frac{\tau_t}{\tau_c} \approx 0.1
\]  

(5.26)

As in the non-reacting case, the grid spacing is mostly influencing the amount of turbulent mixing. The finer is the grid spacing, the lower is the sub-grid scale dissipation. This is either present as numerical dissipation in MILES or numerical and explicit dissipation in LES.

Figure 5.45: Instantaneous axial velocity component on the jet axis and in the main shear region. MILES,NS, 256×256×1024.
Increased grid resolution also leads to more finer turbulent structures in the flow field. The residual numerical diffusion combines with molecular transport -except in MILES_EULER- to finally shape the mixture fraction across the diffusion layer and control the reaction rate.

Further insight can be gained from the scatter-plots of Figures 5.46 and 5.47. The most dissipative simulation is the top-left sub-figure and the less dissipative one is the bottom-right sub-figure. In the scatter plots the mixture fraction definition is chosen as the normalized nitrogen mass fraction (equation 5.25). In order to indicate the position of points on scatter plots relative to the flame front, color that represent an index related to the water concentration is used

\[
c_{\text{H}_2\text{O}} = \left(\frac{Y_{\text{H}_2\text{O}} - Y_{\text{H}_2\text{O}}|_{\text{jet}}}{Y_{\text{H}_2\text{O}}|_{\text{max}} - Y_{\text{H}_2\text{O}}|_{\text{jet}}}\right)\left(\frac{Y_{\text{H}_2\text{O}} - Y_{\text{H}_2\text{O}}|_{\text{co-flow}}}{Y_{\text{H}_2\text{O}}|_{\text{max}} - Y_{\text{H}_2\text{O}}|_{\text{co-flow}}}\right)
\] (5.27)

\(c_{\text{H}_2\text{O}}\) is an index indicating the relative mass fraction of water which is the main product of chemical reactions. By definition \(c_{\text{H}_2\text{O}}\) takes the value 1 in the flame and 0 elsewhere. The points colored in green correspond to flow regions close to the flame front. Finite-rate, reversible full chemistry is clearly felt since the flame structure departs strongly from infinitely fast, irreversible one-step chemistry (straight lines). The temperature mixing line is curved and is below the theoretical linear mixing line (Figure 5.46). This is due to the cold hydrogen stream at 160K (see Table 5.7) which has a very high heat capacity (13200 J/kg.K at 160K) when compared to that of air (1170 J/kg.K at 1200K). The dark blue color points are at high mixture fraction region, indicate the initial core, far from the flame front. When hydrogen mixes with surrounding air, it takes heat from the oxidizer stream and lowers the mixture temperature below the self-ignition limit (\(\approx 1000\text{K}\) at stoichiometry).

On the coarse grid (left columns of Figures 5.46 and 5.47), the flame structure is almost the same for all simulations. The points are clustered close to the equilibrium states, indicating a relatively fast chemistry compared to large-eddies turbulent time scale (higher Damköhler number). Few points are close to the mixing line at \(z \approx z_{\text{st}}\) indicating “mixed is burned” behavior: the high level of numerical diffusion artificially brings reactants together at the molecular level and reactions are completed.

On the intermediate grid 128x128x512 (central columns), a stronger departure from irreversible infinitely fast chemistry is observed. Diagrams are more “filled” because turbulent mixing is more intense and the reactive mixture can be found in various intermediate states combining reactants and products, especially in LES_SSF due to the explicit sub-grid turbu-
lent diffusion. More points can also be found near stoichiometry where reaction does not occur (high $Y_{O_2}$, low $T$) due to a “lack of time”. Physical LES and MILES are close to each other indicating that the contributions of the explicit sub-grid model and the numerical diffusion in MILES are at the same level.

On the fine grid, the scatter plots for the flame are slightly different. More points can be found close the mixing lines, mainly in the $H_2$ stream ($z \rightarrow 1$) where $O_2$ is present without reaction, because of the low temperature. This is due to fast and efficient turbulent mixing relative to the chemistry. This is mainly true for the MILES_EULER simulation (bottom-right sub-figure in Figure 5.47) which differs clearly from LES_SSF and MILES_NS : the low level of numerical diffusion and the absence of viscous model restrict the small scale molecular mixing, hence the reaction rates are reduced.

The velocity, temperature and $H_2$ mass fractions of the average fields at the center line and
the maximum temperature value on transverse planes along x axis are plotted for different resolutions in Figure 5.48. The average profiles are approaching each other for high grid resolutions. Both velocity potential core and concentration potential core lengths are longer for low resolution of $64 \times 64 \times 512$, regardless of numerical approach. At this initial region of the flow shear layer instabilities are mostly responsible of mixing and transition to turbulence. At low resolution since the sharp gradients are smoothed due to large amount of dissipation, transition and mixing are delayed. At low resolutions high level oscillations are noticed at the location where jet is entering the computational domain. This may be attributed to the sharpness of the oblique shock forming immediately around the inlet region. At the two higher resolutions potential core lengths for the solutions are the same (four sub-figures top right). The dynamics of the flow is resolved for $128 \times 128 \times 512$ and higher resolution. Although the resolution is increased, the potential core length remains the same.
The maximum temperature distribution along the jet axis on the normal planes are plotted to denote the ignition location along the axis as the temperature jump (bottom sub-figures in 5.48). As the resolution is increased, the jump in the maximum temperature for MILES_EULER slightly deviates to an upstream axial position (bottom right sub-figure in 5.48). As diffusion is enhanced with molecular transport and physical LES, this jump can

Figure 5.48: Average profiles of velocity, H\textsubscript{2} mass fraction and temperature at the center line of the jet and maximum temperature at the transverse sections along x axis. Left to right : 64×64×256, 128×128×512, 256×256×1024.
be expected to be earlier for MILES_NS and LES_SSF. However, further analysis shows that after the occurrence of ignition, the penetration of the high temperature zones associated with transverse diffusion is prevented by the cold hydrogen flow close to the inlet. Taking into account the fact that the molecular transport enhances cooling, the temperature jump rise along the axis is delayed. Temperature jumps for MILES_NS and LES_SSF are exactly at the same axial positions for all resolutions.

Radial profiles of axial velocity and temperature at $x = 210$ mm are shown in Figure 5.49, with experimental measurements of ONERA. For these measurements, temperature is obtained with CARS (Coherent Anti-Stokes Raman Scattering) and velocity with the LDA technique. The computed velocity that is overestimated compared to experiments, is probably because particles cannot follow perfectly the fast and low-density flow. Computations underestimate the mean experimental temperature in the center because the jet is flapping and the points of measurement are alternatively in the central jet or in the surrounding reaction zone. Also, the high temperature wall boundary layer which is not considered may lead temperature to increase in the actual flow.
Figure 5.49: Tranverse profiles of axial velocity (left) and temperature (right) at $x = 210$ mm. Top to bottom: $64\times64\times256$, $128\times128\times512$, $256\times256\times1024$. 
Figure 5.50 displays the instantaneous distribution of sub-grid eddy-viscosity ratio $\mu'/\mu$ in the vertical symmetry plane, for the different resolutions. On the 256×256×1024 grid, the SSF model is barely active within the flame. The turbulent oscillations which emanate from the inlet noise clustered at the jet center are laminarized while passing through the flame surface. The SSF model is not active outside the flame.

**5.2.4.2 RESULTS FOR IMPROVED INLET PROFILE**

In the previous part the behavior of numerical approaches MILES and LES are investigated using a simplified initial field with weak shocks which is deduced from simulation results of Davidenko for the LAERTE chamber [22]. In this section, the improved inlet conditions explained in section 5.2.2.2 are applied. The inlet field is improved adding the pressure and radial velocity profiles calculated in the work by Davidenko. So that strong shock patterns which do emanate from the inlet are included in the solution. Also, the concentration of rad-
ical O is reduced to the level of numerical estimations by Davidenko whereas in previous section 5.2.4.1 O mass fraction were calculated as \((1 - \sum Y_{a \neq O})\). The effect of radical O mass fraction is showing itself as a late ignition. MILES\,EULER and MILES\,NS simulations are conducted at \(128 \times 128 \times 512\) and \(256 \times 256 \times 1024\) resolutions. LES\,SSF simulation is not performed since it is shown that it converges to the same statistics with MILES\,NS for reacting and non-reacting flows.

5.2.4.2.1 FLAME STRUCTURE

Flame structure for the LAERTE solution is shown as the mixture fraction stoichiometric value iso-surfaces together with temperature and water concentration contours on the corresponding symmetry axis, in Figure 5.51. When compared with the previous simulations of simple BCs (presented in section 5.2.4.1), MILES\,EULER has larger structures especially in the second half of the computational domain. The MILES\,EULER solution is more wrinkled compared to the MILES\,NS solution. Considering both temperature and water mass fraction contours, it is observed that ignition occurs at a later axial location as the radical O concentration in the co-flow is less than the value in previous simulations with a simplified initial field. Instantaneous pressure contours on the axis of the jet are shown on Figure 5.52. The resolved shock patterns are almost the same for both MILES\,NS and MILES\,EULER solutions. As the stoichiometric iso-surfaces are passing across the shock, the corresponding surface is contracted. Thin layers of quenching can be observed on temperature and water mass fraction contours which occur at these contraction regions.

When comparing medium and high resolution solutions, it is seen that ignition occurs at the same axial location. Higher temperatures occur after ignition. More pronounced quenching effect is also seen. This may be due to the increase in the sharpness of the shocks on the high resolution grid. The shock patterns for the medium and high resolution solutions look alike. Even though, in the solution on high resolution grid, much sharper shocks are resolved. As a result, the radial extent of higher pressure zones along the oblique shocks is increased. The turbulent structures in the first diamond shape shock structure near the inlet are also more pronounced and sharper.
Figure 5.51: Flame structure of LAERTE chamber with precise initial field, including strong shock patterns.
Figure 5.52: Pressure contours on the symmetry axis, of LAERTE chamber with precise initial field.
5.2.4.2 STATISTICAL RESULTS

The experimental pressure measurements over the chamber wall are compared with time averaged numerical results on Figure 5.53. The statistical results of these simulations are also compared to the numerical results obtained by RANS and adaptive mesh refinement (AMR) methods. The AMR simulation results are taken from Eude. Since wall boundaries are not considered in the MILES_NS and MILES_EULER computations, the experimental pressure growth is larger than that computed. As in RANS calculations these wall boundary conditions are considered, agreement with the experimental data is apparent. The experimental pressure growth is reasonably higher due to velocity decay on the boundaries. There exists quite good agreement in shock reflection positions despite the stretched grid at the boundaries.

In the Figure 5.54, maximum temperature distribution over YZ planes along the \(x\) axis is shown. Although results of MILES_NS MILES_EULER and AMR are similar for pressure (Figure 5.53), they differ with each other for maximum temperature. The ignition location which is denoted as the temperature jump in this Figure 5.54, occurs earlier in MILES_NS solution. This delay is also observed in flame structure Figures. In the LAERTE experiment the ignition point is given roughly at 10 diameters along the axis form the jet entrance. This location is in good agreement with the axial temperature profile jump location computed by MILES_EULER MILES_NS and AMR methods.
The radial distribution of OH molar concentration measurements are compared with the numerical results in Figure 5.55. Both AMR and MILES solutions give quantitatively good agreement with the experimental results. On the other hand RANS simulation results overestimate OH concentrations.

Both transverse velocity and temperature distributions at different axial locations on the symmetry axis of the domain are shown on Figure 5.56. At the far-field \((x = 288 \text{ mm})\) both set of results deviate from the experimental measurements. This may be attributed at the neglect of the temperature and the velocity gradients near the wall. Furthermore, there may exist some errors in the measurements due to flame flapping and insufficient transport of PIV particles in low density flows. When comparing Figure 5.49 (simplified inlet field) and Figure 5.56 it is seen that transverse profiles of average temperature and velocity are not much affected at the far-field by the change of inlet field conditions. Despite the reduced scalar concentration \((Y_o)\) and radial inlet pressure and velocity profiles which generate strong shocks, the transverse profiles are identical to the results of the previous section.

5.2.5 CONCLUSIONS

In this section, reacting tests conducted for the LAERTE chamber, are simulated. At first, a simplified initial field is used for the comparative evaluation of the numerical approaches. Later, a more precise initial condition is applied in order to enhance flow physics. In conclu-
Figure 5.56: Experimental transverse profiles of axial velocity (left) and temperature (right) at $x=10$ mm, $x=72$ mm, $x=210$ mm and $x=288$ mm.

sion for the first case, flow statistics converge to physical results as the resolution is increased for both MILES_NS and LES_SSF computations. However at high resolution MILES_EULER
results differ from those of MILES_NS and LES_SSF. The achieved LES asymptotes for the statistics also explained in the paper by Pope are similar to these two approaches. The more precise inlet conditions are applied in the second case. Both MILES_NS and MILES_EULER simulation results are in good agreement. Even if the inlet profiles are improved, the average field results are similar to the solutions for simplified inlet profiles. Present results are also compared with the AMR results with precise initial conditions and they are in good agreement.
CHAPTER 6

CONCLUSIONS

In this study three dimensional computations are performed for reacting and non-reacting compressible H₂/air jets. In the computations, fifth order WENO scheme is used. The main objective is to investigate the respective performance of LES and MILES numerical approaches in the context of high speed reacting flows. It was also the intention of this work to determine the sensitivities of a high speed reacting jet flow simulations to various physical models. In the non-reacting case, the time step is limited by the CFL condition. In the reacting case the time step is limited to $2 \times 10^{-8}$ s as dictated by the chemistry. The relative cost of the different simulations is presented at Table I.1 in appendix I.

The following conclusions are derived from the simulation of non-reacting Eggers jet [71]:

- The Smagorinsky model is found to be over-dissipative, hence inhibiting the initial development of instabilities. The three-dimensionality sensor in the Selective Structure Function model fixes the problem.

- In particular, LES solutions do not show superiority compared to MILES, Navier-Stokes or Euler, considering the statistics related to the mixing and the dynamics of the flow.

- In the non-reacting case, calculation of the molecular transport model in MILES-NS represents an extra cost around 18%. The cost of introducing the Smagorinsky model in LES-SM is hardly felt because the velocity derivatives required in the model are already computed in viscous terms. The SSF model is a little more expensive due to the calculation of velocity differences and of the three-dimensionality sensor, (equation 3.67), but is also negligible.
• MILES_EULER simulations are showing grid convergence although there exists no physical cutoff in the simulation. This is an open question for further research.

In the reacting LAERTE chamber case, the main conclusions are [72]:

• The trend is the same as in the Eggers jet, but depending on the grid resolution, chemistry can be faster or slower relatively to the flow dynamics.

• One of the most sensitive parameter to the molecular transport is the ignition point. When comparing the maximum temperature change on the normal planes along the axis, a slight shift of temperature jump is observed at the order of jet diameter between EULER and NS solutions.

• When comparing MILES_NS and LES_SSF average field results, no significant differences are found. This similarity is valid even in the scatter plots of temperature (Figure 5.46) and O₂ mass fraction (Figure 5.47) which are representing the interactions at the molecular level.

• The high sensitivity to the inlet profiles of RANS method is not found in LES.

• The 7 species, 14 reactions chemistry in the reacting MILES_EULER represent 85% of the WENO effort, although the routines for source terms are thoroughly optimized.

In conclusion, the non-linear WENO scheme is self-adapting to the flow gradients, and a balance is achieved between the explicit sub-grid scale hyperviscosity in physical LES and the numerical dissipation of the scheme. In a physical LES on a coarse grid, the proper modeling of the sub-grid chemical source term is important, but the molecular transport is not. In a numerical LES however, a molecular viscous cut-off in the simulation is mandatory. The choice of whether to introduce or not an explicit sub-grid model in the simulation is not a matter of computational cost. MILES-Navier-Stokes performed with “clever” dissipative numerics provides almost the same grid-independent flow statistics as physical LES. Quoting S.B. Pope’s conclusion [107] “The most that an LES calculation can hope to achieve, is to obtain an accurate estimate of the intermediate asymptote [...]”. This is achieved for the two finest grids used in this study, with and without explicit sub-grid model.
REFERENCES


APPENDIX A

TRANSPORT COEFFICIENTS

In this appendix the constants for the curve fits of viscosity, thermal conductivity versus temperature and binary diffusion coefficients versus temperature and pressure are tabulated.

Table A.1: Thermal conductivity coefficients for second order polynomial fit (i.e. equation 2.50).

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Table A.2: Coefficients of viscosity for second order polynomial fit (i.e. equation 2.37).

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Table A.3: Power law fitted diffusion coefficients for $P_0$ (=1bar), T (i.e., equation 2.21).

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APPENDIX B

THERMODYNAMIC DATA

The thermodynamic data used in the calculation are presented in Table B.2, following the format given in Table B.1. It is assumed that the pressure is not elevated so thermodynamic properties are function of temperature only. The data are the coefficients of fitted polynomials on the temperature functions of heat capacities $C_{p\alpha}$ and $C_{v\alpha}$ of species $\alpha$.

$$\frac{C_{p\alpha}}{R} = C_{p0\alpha} + TC_{p1\alpha} + C_{p2\alpha}T^2 + C_{p3\alpha}T^3 + C_{p4\alpha}T^4 \quad (B.1)$$

Enthalpy $H^0/RT$ and entropy $S^0/RT$ of species are computed using this data. There exists two sets of coefficients referring to two different polynomials for low temperature $[T_{min} - 1000K]$ and high temperature $[1000K - T_{max}]$ intervals. The coefficients $C_{p5}$ and $C_{p6}$ are used to compute the enthalpy $\Delta H^0$ and the entropy $\Delta S^0$ of system.

Table B.1: Thermodynamic property data format.

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<td>0.04920308E+02</td>
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<td>0.02421032E-04</td>
</tr>
<tr>
<td>0.01602843E-07</td>
<td>0.03890696E-11</td>
<td>0.02914764E+06</td>
<td>0.02963995E+02</td>
</tr>
<tr>
<td>O₂</td>
<td>300</td>
<td>5000</td>
<td>31.99880</td>
</tr>
<tr>
<td>0.03697578E+02</td>
<td>0.06135197E-02</td>
<td>-0.01258842E-02</td>
<td>0.01775281E-09</td>
</tr>
<tr>
<td>0.01233930E+05</td>
<td>0.03189166E+02</td>
<td>0.03212936E+02</td>
<td>0.01127486E-01</td>
</tr>
<tr>
<td>0.01313877E-07</td>
<td>-0.08768554E-11</td>
<td>-0.01005249E+05</td>
<td>0.06034738E+02</td>
</tr>
<tr>
<td>OH</td>
<td>300</td>
<td>5000</td>
<td>17.00734</td>
</tr>
<tr>
<td>0.02882730E+02</td>
<td>0.01013974E-01</td>
<td>-0.02276877E-05</td>
<td>0.02174684E-09</td>
</tr>
<tr>
<td>0.03886888E+05</td>
<td>0.05595712E+02</td>
<td>0.03637266E+02</td>
<td>0.01850910E-02</td>
</tr>
<tr>
<td>0.02387203E-07</td>
<td>-0.08431442E-11</td>
<td>0.03606782E+05</td>
<td>0.01358860E+02</td>
</tr>
<tr>
<td>H₂O</td>
<td>300</td>
<td>5000</td>
<td>18.01528</td>
</tr>
<tr>
<td>0.02672146E+02</td>
<td>0.03056293E-01</td>
<td>-0.08730260E-05</td>
<td>0.01209966E-08</td>
</tr>
<tr>
<td>0.02989921E+06</td>
<td>0.06828178E+02</td>
<td>0.03386842E+02</td>
<td>0.03474982E-01</td>
</tr>
<tr>
<td>0.06968581E-07</td>
<td>-0.02506588E-10</td>
<td>-0.03020811E+06</td>
<td>0.02590233E+02</td>
</tr>
<tr>
<td>N₂</td>
<td>300</td>
<td>5000</td>
<td>28.01348</td>
</tr>
<tr>
<td>0.02926640E+02</td>
<td>0.01487977E-01</td>
<td>-0.05684761E-05</td>
<td>0.01009704E-08</td>
</tr>
<tr>
<td>0.09227977E+04</td>
<td>0.05980528E+02</td>
<td>0.03298677E+02</td>
<td>0.01408240E-01</td>
</tr>
<tr>
<td>0.05641515E-07</td>
<td>-0.02444855E-10</td>
<td>-0.01020900E+05</td>
<td>0.03950372E+02</td>
</tr>
</tbody>
</table>
APPENDIX C

CHEMICAL KINETICS

The chemical kinetics data used for source term calculation are presented in Table C.2. The rate constant $K_r$ is computed applying Arrhenius law:

$$K_r = A_r T^{b_r} \exp \left( \frac{E_r}{RT} \right)$$

where for the $r^{th}$ reaction:

- $A_r$ is the pre-exponential factor.
- $E_r$ is the activation energy.
- $b_r$ constant for pre-exponential factor $T^{b_r}$.

The efficiencies of the species included in the H$_2$/Air chemical kinetics mechanism is:

Table C.1: Efficiency of the species for the ONERA scheme.

<table>
<thead>
<tr>
<th></th>
<th>H</th>
<th>H$_2$</th>
<th>H$_2$O</th>
<th>O</th>
<th>OH</th>
<th>O$_2$</th>
<th>N$_2$</th>
<th>Ar</th>
</tr>
</thead>
<tbody>
<tr>
<td>ONERA</td>
<td>1.0</td>
<td>2.5</td>
<td>12</td>
<td>1.0</td>
<td>1.0</td>
<td>1.00</td>
<td></td>
<td>1.0</td>
</tr>
</tbody>
</table>
Table C.2: H\textsubscript{2}/air Chemical reaction mechanism by ONERA.

<table>
<thead>
<tr>
<th>r</th>
<th>Reaction</th>
<th>A\textsubscript{r} (cm.mol.s)</th>
<th>b\textsubscript{r}</th>
<th>E\textsubscript{r} (kJ.mol\textsuperscript{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>H\textsubscript{2} + O\textsubscript{2} → OH + OH</td>
<td>1.70E+13</td>
<td>0.0</td>
<td>199.9122</td>
</tr>
<tr>
<td>2</td>
<td>OH + OH → H\textsubscript{2} + O\textsubscript{2}</td>
<td>4.03E+10</td>
<td>0.3168</td>
<td>121.0074</td>
</tr>
<tr>
<td>3</td>
<td>H + O\textsubscript{2} → OH + O</td>
<td>1.99E+14</td>
<td>0.0</td>
<td>70.3043</td>
</tr>
<tr>
<td>4</td>
<td>O + OH → H + O\textsubscript{2}</td>
<td>8.93E+11</td>
<td>0.3383</td>
<td>-0.9778</td>
</tr>
<tr>
<td>5</td>
<td>H\textsubscript{2} + OH → H\textsubscript{2}O + H</td>
<td>1.02E+08</td>
<td>1.6</td>
<td>13.8008</td>
</tr>
<tr>
<td>6</td>
<td>H + H\textsubscript{2}O → H\textsubscript{2} + OH</td>
<td>7.96E+08</td>
<td>1.528</td>
<td>77.3248</td>
</tr>
<tr>
<td>7</td>
<td>H\textsubscript{2} + O → OH + H</td>
<td>5.12E+04</td>
<td>2.67</td>
<td>26.3016</td>
</tr>
<tr>
<td>8</td>
<td>OH + H → H\textsubscript{2} + O</td>
<td>2.70E+04</td>
<td>2.649</td>
<td>18.6212</td>
</tr>
<tr>
<td>9</td>
<td>OH + OH → H\textsubscript{2}O + O</td>
<td>1.51E+09</td>
<td>1.14</td>
<td>0.4142</td>
</tr>
<tr>
<td>10</td>
<td>H\textsubscript{2}O + O → OH + OH</td>
<td>2.22E+10</td>
<td>1.089</td>
<td>71.6144</td>
</tr>
<tr>
<td>11</td>
<td>H + OH + M → H\textsubscript{2}O + M</td>
<td>2.21E+22</td>
<td>-2.0</td>
<td>0.0</td>
</tr>
<tr>
<td>12</td>
<td>H\textsubscript{2}O + M → H + OH + M</td>
<td>8.94E+22</td>
<td>-1.835</td>
<td>496.7304</td>
</tr>
<tr>
<td>13</td>
<td>H + H + M → H\textsubscript{2} + M</td>
<td>9.79E+16</td>
<td>-0.6</td>
<td>0.0</td>
</tr>
<tr>
<td>14</td>
<td>H\textsubscript{2} + M → H + H + M</td>
<td>5.09E+16</td>
<td>-0.3624</td>
<td>433.2265</td>
</tr>
</tbody>
</table>
APPENDIX D

TRANSFORMATION MATRICES FOR SYSTEM OF $N_S p + 5$ EQUATIONS

The following matrices are used for finite difference characteristic discretization. The vector of conservative variables is:

$$\vec{\tilde{U}} = [\rho, \rho u, \rho v, \rho w, \rho e_1, \rho e_2, \rho Y_1, \ldots, \rho Y_{N_{sp}}]^T$$

and the vector of primitive variables is considered as:

$$\vec{U} = [\rho, u, v, w, T, Y_1, \ldots, Y_{N_{sp}}]^T$$

Here the matrices for transformation between conservative variables to characteristic variables are presented, for the spatial discretization $\tilde{R}$ and $\tilde{L}$. These can be directly obtained diagonalizing the conservative jacobian matrices $\tilde{A}$ but the easier way is using the relations $\tilde{R} = PR$ and $\tilde{L} = LP^{-1}$.

with

1. $K = \frac{1}{2} u_i u_i$
2. $\epsilon_i = h_{i\alpha} - r_{i\alpha} T$
3. $\epsilon = \sum_{\alpha=1}^{N_{sp}} h_\alpha Y_\alpha - r T$
\[
\tilde{R} = \begin{pmatrix}
\rho & \rho & 0 & 0 & \rho & \rho \frac{r-c}{r} & \ldots & \ldots & \ldots & \rho \frac{r-c}{r} \\
\rho(u-c) & \rho u & 0 & 0 & \rho(u+c) & \rho u \frac{r-c}{r} & \ldots & \ldots & \ldots & \rho u \frac{r-c}{r} \\
\rho v & \rho v & \rho & 0 & \rho v & \rho v \frac{r-c}{r} & \ldots & \ldots & \ldots & \rho v \frac{r-c}{r} \\
\rho w & \rho w & 0 & \rho & \rho w & \rho w \frac{r-c}{r} & \ldots & \ldots & \ldots & \rho w \frac{r-c}{r} \\
\rho e_1 + P - \rho uc & \rho e_1 - \frac{P}{r} & \rho v & \rho w & \rho e_1 + P + \rho uc & \rho e_1 \frac{r-c}{r} + \rho (e_1 - \epsilon) & \ldots & \ldots & \ldots & \rho e_1 \frac{r-c}{r} + \rho (e_{N_{\text{sp}}} - \epsilon) \\
\rho Y_1 & \rho Y_1 & 0 & 0 & \rho Y_1 & \rho Y_1 \frac{r-c}{r} + \rho & \rho Y_1 \frac{r-c}{r} & \ldots & \ldots & \rho Y_1 \frac{r-c}{r} \\
\rho Y_2 & \rho Y_2 & 0 & 0 & \rho Y_2 & \rho Y_2 \frac{r-c}{r} & \ldots & \ldots & \ldots & \rho Y_2 \frac{r-c}{r} \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ldots & \ldots & \ldots & \vdots \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ldots & \ldots & \ldots & \vdots \\
\rho Y_{N_{\text{sp}}} & \rho Y_{N_{\text{sp}}} & 0 & 0 & \rho Y_{N_{\text{sp}}} & \rho Y_{N_{\text{sp}}} \frac{r-c}{r} & \ldots & \ldots & \rho Y_{N_{\text{sp}}} \frac{r-c}{r} - 1 & \rho Y_{N_{\text{sp}}} \frac{r-c}{r} + \rho \\
\end{pmatrix}
\] (D.1)
\[
\bar{L} = \begin{pmatrix}
\frac{c^2 + \gamma uc + \gamma(\gamma-1)(K-\epsilon)}{2\gamma pc^2} & -\frac{(y-1)u-c}{2pc^2} & -\frac{(y-1)v}{2pc^2} & -\frac{(y-1)w}{2pc^2} & \frac{y-1}{2pc^2} & \frac{(y-1)(e_1) + (r_1-r)T}{2pc^2} & \ldots & \ldots & \ldots & \frac{(y-1)(e_{Nsp}) + (r_{Nsp}-r)T}{2pc^2} \\
\frac{y-1}{\gamma pc^2}(c^2 - \gamma(K-\epsilon)) & -\frac{(y-1)u}{\rho c^2} & -\frac{(y-1)v}{\rho c^2} & -\frac{(y-1)w}{\rho c^2} & \frac{y-1}{\rho c^2} & \frac{(y-1)(e_1) + (r_1-r)T}{\rho c^2} & \ldots & \ldots & \ldots & \frac{(y-1)(e_{Nsp}) + (r_{Nsp}-r)T}{\rho c^2} \\
-\frac{\frac{v}{\rho^2}}{2} & 0 & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 \\
-\frac{\frac{w}{\rho^2}}{2} & 0 & \frac{1}{\rho} & 0 & 0 & 0 & \ldots & \ldots & 0 \\
\frac{c^2 - \gamma uc + \gamma(\gamma-1)(K-\epsilon)}{2\gamma pc^2} & -\frac{(y-1)u+c}{2pc^2} & -\frac{(y-1)v}{2pc^2} & -\frac{(y-1)w}{2pc^2} & \frac{y-1}{2pc^2} & \frac{(y-1)(e_1) + (r_1-r)T}{2pc^2} & \ldots & \ldots & \ldots & \frac{(y-1)(e_{Nsp}) + (r_{Nsp}-r)T}{2pc^2} \\
-\frac{\frac{\gamma_1}{\rho^2}}{2} & 0 & 0 & 0 & 0 & \frac{1}{\rho} & 0 & \ldots & 0 \\
-\frac{\frac{\gamma_2}{\rho^2}}{2} & 0 & 0 & 0 & 0 & 0 & \frac{1}{\rho} & \ldots & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \ddots \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \ddots \\
-\frac{\frac{Y_{Nsp}}{\rho}}{2} & 0 & \ldots & \ldots & \ldots & \ldots & \ldots & \ldots & 0 & \frac{1}{\rho}
\end{pmatrix}
\]
APPENDIX E

TRANSFORMATION MATRICES FOR SYSTEM OF $N_{sp}+4$ EQUATIONS

Matrices for the transformation between conservative, primitive and characteristic forms of governing equations are presented in this appendix. The derivations for these matrices were held over equations for $N_{sp}-1$ species.

E.1 TRANSFORMATION MATRICES BETWEEN THE PRIMITIVE VARIABLES AND THE CONSERVATIVE VARIABLES

The vector of conservative variable is:

$$\vec{\tilde{U}} = T [\rho, \rho u, \rho v, \rho w, \rho e_t, \rho Y_1, \ldots, \rho Y_{N_{sp}-1}]$$

and primitive variable vector is considered as:

$$\vec{U} = T [\rho, u, v, w, T, Y_1, \ldots, Y_{N_{sp}-1}]$$
The transformation matrices of the conservative variables to primitive variables and vice versa are given below respectively $P$ and $P^{-1}$.

$$
P = \begin{pmatrix}
1 & 0 & 0 & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 \\
u & \rho & 0 & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 \\
v & 0 & \rho & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 \\
w & 0 & 0 & \rho & 0 & 0 & 0 & \ldots & \ldots & 0 \\
P_{5,1} & \rho u & \rho v & \rho w & \rho C_v & P_{5,6} & P_{5,7} & \ldots & \ldots & P_{5,N_{sp}+4} \\
y_1 & 0 & 0 & 0 & 0 & \rho & 0 & \ldots & \ldots & 0 \\
y_2 & 0 & 0 & 0 & 0 & \rho & 0 & \ldots & \ldots & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots & \vdots \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & 0 \\
y_{N_{sp}-1} & 0 & 0 & 0 & 0 & 0 & 0 & \ldots & 0 & \rho
\end{pmatrix}
$$

with

- $P_{5,1} = h - rT + 1/2u_i u_i$
- $P_{5,5+a} = \rho[h_a - h_{N_{sp}} - (r_a - r_{N_{sp}})T]$ avec $a = 1,...,N_{sp}-1$

$$
P^{-1} = \begin{pmatrix}
1 & 0 & 0 & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 \\
-\nu/\rho & 1/\rho & 0 & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 \\
-\nu/\rho & 0 & 1/\rho & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 \\
-\nu/\rho & 0 & 0 & 1/\rho & 0 & 0 & 0 & \ldots & \ldots & 0 \\
P_{5,1} & \frac{\nu}{\rho C_v} & \frac{\nu}{\rho C_v} & \frac{\nu}{\rho C_v} & \frac{1}{\rho C_v} & P_{5,6} & P_{5,7} & \ldots & \ldots & P_{5,N_{sp}+4} \\
-\nu_1/\rho & 0 & 0 & 0 & 0 & 1/\rho & 0 & \ldots & \ldots & 0 \\
-\nu_2/\rho & 0 & 0 & 0 & 0 & 1/\rho & 0 & \ldots & \ldots & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots & \vdots \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & 0 \\
-\nu_{N_{sp}-1}/\rho & 0 & 0 & 0 & 0 & 0 & 0 & \ldots & 0 & 1/\rho
\end{pmatrix}
$$

avec

- $P_{5,1}^{-1} = \frac{1}{\rho C_v}[r_{N_{sp}} T - h_{N_{sp}} + 1/2u_i u_i]$
- $P_{5,5+a}^{-1} = \frac{1}{\rho C_v}[(r_a - r_{N_{sp}})T - (h_a - h_{N_{sp}})-]$ avec $a = 1,...,N_{sp}-1$
E.2 TRANSFORMATION MATRICES BETWEEN THE PRIMITIVE VARIABLES AND THE CHARACTERISTIC VARIABLES

The system for multispecies compressible flow equations may be identified with the matrices $A$. Considering the $x$ direction, the matrix $A$ associated with the primitive variables defined in the previous section is

$$A = \begin{pmatrix}
  u & \rho & 0 & 0 & \cdots & \cdots & \cdots & \cdots & 0 \\
  \frac{RT}{\mu_M} & u & 0 & 0 & \frac{R}{M} & A_{2,6} & A_{2,7} & \cdots & A_{2,N_{sp}+4} \\
  0 & 0 & u & 0 & \cdots & \cdots & \cdots & 0 \\
  0 & 0 & 0 & u & 0 & \cdots & \cdots & \vdots \\
  0 & \frac{RT}{C_vM} & 0 & 0 & u & 0 & \cdots & \vdots \\
  0 & 0 & 0 & \ddots & \ddots & \ddots & \ddots & \vdots \\
  \vdots & \ddots & \ddots & 0 & u & 0 & \vdots \\
  \vdots & \ddots & \ddots & 0 & u & 0 \\
  0 & \cdots & \cdots & \cdots & 0 & 0 & 0 & u
\end{pmatrix}$$

with $A_{2,5+i} = R(1/M_\alpha - 1/M_\alpha) i=1,\ldots,N_{sp} - 1$. 

139
The jacobian for primitive variables $A$ is diagonalized using the derivation of left and right eigenvector matrices $R$ and $L$ (this definition is implemented for the derivation of characteristic boundary conditions § 4.5.1) as $A = RAL$. Diagonal matrix of eigenvalues $\Lambda$ is as follows:

$$
\begin{pmatrix}
    u - c & 0 & 0 & \ldots & \ldots & \ldots & \ldots & 0 \\
    0 & u & 0 & \vdots & & & & \vdots \\
    \vdots & 0 & u & 0 & \vdots & & & \vdots \\
    \vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\
    \vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \ddots & 0 \\
    \vdots & 0 & u & 0 & \vdots & & & \vdots \\
    0 & \ldots & \ldots & \ldots & \ldots & \ldots & 0 & u + c \\
\end{pmatrix}
$$

Matrices for right and left eigenvectors are given below

$$
L = \begin{pmatrix}
    \frac{Cp}{c} & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 & 1 & \frac{Cp}{c} \\
    \frac{Cp}{c} & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 & 0 & \frac{Cp}{c} \\
    0 & 1 & 0 & 0 & 0 & \ldots & \ldots & 0 & 0 & 0 \\
    0 & 0 & 1 & 0 & 0 & \ldots & \ldots & 0 & 0 & 0 \\
    1 & 0 & 0 & 1 & 1 & \ldots & \ldots & 1 & 0 & 1 \\
    0 & 0 & 0 & \frac{-r}{(r_1 - Nsp)\gamma} & 0 & \ldots & \ldots & 0 & \frac{-r}{(r_1 - Nsp)\gamma} & 0 \\
    0 & 0 & 0 & \frac{-r}{(r_2 - Nsp)\gamma} & 0 & \ldots & \ldots & 0 & \frac{-r}{(r_2 - Nsp)\gamma} & 0 \\
    \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots & \vdots & \vdots \\
    \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots & \vdots & \vdots \\
    0 & 0 & 0 & 0 & \ldots & \ldots & 0 & \frac{-r}{(Nsp - r_1 - Nsp)\gamma} & 0 & 0 \\
\end{pmatrix}
$$

$$
R = \begin{pmatrix}
    \frac{(\gamma - 1)}{\gamma} & L & -\frac{c}{2c_p} & 0 & 0 & (\frac{(\gamma - 1)}{2\gamma}) & (\frac{2 - \gamma}{2\gamma}) & \frac{r_{1Nsp}}{\gamma} T & (\frac{2 - \gamma}{2\gamma}) & \frac{r_{2Nsp}}{\gamma} T & \ldots & \ldots & (\frac{2 - \gamma}{2\gamma}) & \frac{r_{Nsp - 1 - Nsp}}{\gamma} T \\
    0 & 0 & 1 & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 & 0 \\
    0 & 0 & 0 & 1 & 0 & 0 & 0 & \ldots & \ldots & 0 & 0 \\
    0 & 0 & 0 & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 & 0 \\
    0 & 0 & 0 & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 & 0 \\
    0 & 0 & 0 & \frac{c}{2c_p} & 0 & 0 & 0 & \ldots & \ldots & 0 & \frac{r_{1Nsp}}{\gamma} T & \frac{r_{2Nsp}}{\gamma} T & \ldots & \ldots & \frac{r_{Nsp - 1 - Nsp}}{\gamma} T \\
    0 & \ldots & \ldots & \ldots & \ldots & \ddots & \ddots & \ldots & \ddots & \ddots & 0 & \ldots & \ldots & \ldots & 0 \\
    0 & \ldots & \ldots & \ldots & \ldots & \ddots & \ddots & \ldots & \ddots & \ddots & 0 & \ldots & \ldots & \ldots & 0 \\
    0 & \ldots & \ldots & \ldots & \ldots & \ddots & \ddots & \ldots & \ddots & \ddots & 0 & \ldots & \ldots & \ldots & 0 \\
    0 & \ldots & \ldots & \ldots & \ldots & \ddots & \ddots & \ldots & \ddots & \ddots & 0 & \ldots & \ldots & \ldots & 0 \\
    0 & \ldots & \ldots & \ldots & \ldots & \ddots & \ddots & \ldots & \ddots & \ddots & 0 & \ldots & \ldots & \ldots & 0 \\
    0 & \ldots & \ldots & \ldots & \ldots & \ddots & \ddots & \ldots & \ddots & \ddots & 0 & \ldots & \ldots & \ldots & 0 \\
\end{pmatrix}
$$
E.3 TRANSFORMATION MATRICES BETWEEN THE CONSERVATIVE VARIABLES AND THE CHARACTERISTIC VARIABLES

Here the matrices for transformation between conservative variables and characteristic variables are presented, for the spatial discretization $\tilde{R}$ et $\tilde{L}$. These can be directly obtained diagonalizing the conservative jacobian matrices $\tilde{A}$ but the easier way is using the relations $R = PR$ and $L = LP^{-1}$. 
\[
\bar{R} = \begin{pmatrix}
\frac{\rho C_p}{c} & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 & 1 & \frac{\rho C_p}{c} \\
\frac{\rho C_p}{c}(u - c) & 0 & 0 & 0 & 0 & \ldots & \ldots & 0 & u & \frac{\rho C_p}{c}(u + c) \\
\frac{\rho C_p}{c} Y & \rho & 0 & 0 & 0 & \ldots & \ldots & 0 & v & \frac{\rho C_p}{c} Y \\
\frac{\rho C_p}{c} W & \rho & 0 & 0 & 0 & \ldots & \ldots & 0 & w & \frac{\rho C_p}{c} W \\
\frac{\rho C_p}{c}(h + K - uc) Y_1 & \rho v & \rho w & R_{S,4} & R_{S,5} & \ldots & \ldots & R_{S,Nsp+3} & h - r \frac{h_1 - h_{Nsp}}{r_{1-Nsp}} + K & \frac{\rho C_p}{c}(h + K + uc) Y_1 \\
\frac{\rho C_p}{c} Y_2 & 0 & 0 & -\frac{pr}{(r_1-r_{Nsp})F} & 0 & \ldots & \ldots & 0 & Y_1 - r \frac{r_1 - r_{Nsp}}{r_{1-Nsp}} & \frac{\rho C_p}{c} Y_2 \\
\vdots & \vdots & \vdots & \vdots & 0 & \ddots & \ddots & \vdots & \vdots & \vdots \\
\vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots & \vdots & \vdots \\
\frac{\rho C_p}{c} Y_{Nsp-1} & 0 & 0 & 0 & \ldots & \ldots & 0 & -\frac{pr}{(r_{Nsp-1}-r_{Nsp})F} & Y_{Nsp-1} & \frac{\rho C_p}{c} Y_{Nsp-1} \\
\end{pmatrix}
\]
\[ \mathbf{L} = \begin{pmatrix} R_{1,1}^{-1} & \gamma (y-1) u & y (y-1) v & u (y-1) & y^{-1} & R_{1,6}^{-1} & R_{1,7}^{-1} & \ldots & R_{1,N_{sp}+4}^{-1} \\ -\frac{u}{p} & 0 & \frac{1}{p} & 0 & 0 & 0 & 0 & \ldots & 0 \\ -\frac{w}{p} & 0 & 0 & \frac{1}{p} & 0 & 0 & 0 & \ldots & 0 \\ \frac{c}{p} & \frac{c}{p} & \frac{c}{p} & \frac{c}{p} & \frac{c}{p} & h_1-h_{sp} & \frac{h_2-h_{sp}}{p} & \ldots & \frac{h_{N_{sp}+1}-h_{sp}}{p} \\ \frac{(r_2-r_{sp})}{p r} Y_2 & 0 & 0 & 0 & 0 & 0 & \frac{r_2-r_{sp}}{p r} T & 0 & \ldots & 0 \\ : & : & : & : & : & : & : & \ldots & : \\ : & : & : & : & : & : & : & \ldots & : \\ : & : & : & : & : & : & : & \ldots & : \\ \frac{(r_{N_{sp}+1}-r_{sp})}{p r} Y_{N_{sp}+1} & 0 & 0 & 0 & 0 & 0 & 0 & \ldots & 0 & \frac{(r_{N_{sp}+1}-r_{sp})}{p r} T \\ \frac{r-r_{sp}}{r} \frac{-h_{sp}+\delta}{C_p T} & \frac{u}{C_p T} & \frac{w}{C_p T} & \frac{\gamma-1}{2 C_p} & \frac{1}{C_p T} & -\frac{r_1-r_{sp}}{r} & \frac{h_1-h_{sp}}{C_p T} & \ldots & \frac{h_{N_{sp}+1}-h_{sp}}{C_p T} \\ R_{N_{sp}+4,1}^{-1} & \frac{r(r-1)}{4 p C_p} & \frac{(y-1)}{2 p C_p} & \frac{(y-1)}{2 p C_p} & \frac{(y-1)}{2 p C_p} & R_{N_{sp}+4,6}^{-1} & R_{N_{sp}+4,7}^{-1} & \ldots & R_{N_{sp}+4,N_{sp}+4}^{-1} \end{pmatrix} \]
with

- \( K = 1/2u_iu_i \)

- \( R_{5,\alpha+3} = pC_p \frac{\rho r(h_a - h_{Nsp})}{T(r_a - r_{Nsp})} \)

- \( R^{-1}_{1,1} = \frac{\gamma_{Nsp} \frac{T}{y^{-1}} \sum_{a=2}^{Nsp} (r_{Nsp} - r_a) Y_a - h_{Nsp} + K - u_c}{2pC_p} \)

- \( R^{-1}_{4,1} = \frac{1}{pC_p} \left[ \frac{T}{y^{-1}} \sum_{a=2}^{Nsp} (r_{Nsp} - r_a) Y_a - h_{Nsp} + K \right] \)

- \( R^{-1}_{1,\alpha+5} = \frac{\gamma T(r_a - r_{Nsp}) - (y-1)(h_a - h_{Nsp})}{2pC_p} \) with \( \alpha = 1, \ldots, N_{2p} - 1 \)

- \( R^{-1}_{N_{1p},4,1} = \frac{\gamma_{Nsp} \frac{T}{y^{-1}} \sum_{a=2}^{Nsp} (r_{Nsp} - r_a) Y_a - h_{Nsp} + K - u_c}{2pC_p} \)

- \( R^{-1}_{N_{1p},4,\alpha+5} = R^{-1}_{1,\alpha+5} \) with \( \alpha = 1, \ldots, N_{2p} - 1 \)
APPENDIX F

WAVE AMPLITUDES AND LODI SYSTEM IN TERMS OF
PRIMITIVE VARIABLES

LODI system wave amplitudes in x direction:

\[ L_1 = \frac{u - c}{2 \rho C p} \left( \frac{\partial P}{\partial x} - \rho c \frac{\partial u}{\partial x} \right) \]

\[ L_2 = u \frac{\partial v}{\partial x} \]

\[ L_3 = u \frac{\partial w}{\partial x} \]

\[ L_4 = u T \left[ \frac{1}{\rho} \left( \frac{1}{c^2} \frac{\partial P}{\partial x} \right) - \left( \frac{r_1 - r_{Nsp}}{r} \right) \frac{\partial Y_1}{\partial x} \right] \]

\[ L_{\alpha+3} = -u T \left( \frac{r_\alpha - r_{Nsp}}{r} \right) \frac{\partial Y_\alpha}{\partial x} ; \quad \alpha = (2, ..., N_{sp} - 1) \]

\[ L_{N_{sp}+3} = u \left( \frac{\partial p}{\partial x} - \frac{\partial P}{\partial x} \right) \]

\[ L_{N_{sp}+4} = \frac{u + c}{2 \rho C p} \left( \frac{\partial P}{\partial x} + \rho c \frac{\partial u}{\partial x} \right) \]  \hspace{1cm} (F.1)

LODI system primitive derivatives in x direction:

\[ d_1 = \frac{\partial \rho}{\partial t} = \frac{\rho C_p}{c^2} (L_1 + L_{N_{sp}+4}) + L_{N_{sp}+3} \]

\[ d_2 = \frac{\partial u}{\partial t} = \frac{C_p}{c} (L_{N_{sp}+4} - L_1) \]

\[ d_3 = \frac{\partial v}{\partial t} = L_2 \]

\[ d_4 = \frac{\partial w}{\partial t} = L_3 \]

\[ d_5 = \frac{\partial T}{\partial t} = L_1 + L_4 + \sum_{\alpha=2}^{N_{sp}-1} L_{\alpha+3} + L_{N_{sp}+4} \]

\[ d_6 = \frac{\partial Y_1}{\partial t} = - \left( \frac{r}{r_1 - r_{Nsp}} \right) \left( \frac{L_{N_{sp}+3}}{\rho} + \frac{L_4}{T} \right) \]

\[ d_{5+\alpha} = \frac{\partial Y_\alpha}{\partial t} = - \left( \frac{r}{r_\alpha - r_{Nsp}} \right) \frac{L_{\alpha+3}}{T} ; \quad \alpha = (2, ..., N_{sp} - 1) \]  \hspace{1cm} (F.2)
LODI system wave amplitudes in y direction:

\[ L_1 = \frac{v - c}{2 \rho C_p} \left( \frac{\partial P}{\partial y} - \rho \frac{\partial v}{\partial y} \right) \]

\[ L_2 = \frac{\partial u}{\partial y} \]

\[ L_3 = v T \left[ 1 \left( \frac{1}{\rho} \frac{\partial P}{\partial y} - \frac{\partial \rho}{\partial y} \right) - \left( \frac{r_1 - r_{N_p}}{r} \right) \frac{\partial Y_1}{\partial y} \right] \]

\[ L_4 = v \frac{\partial w}{\partial y} \]

\[ L_{\alpha+3} = -v T \left( \frac{r_\alpha - r_{N_p}}{r} \right) \frac{\partial Y_{\alpha}}{\partial y} \quad : \alpha = (2, ..., N_{sp} - 1) \]

\[ L_{N_{sp}+3} = \left( \frac{\partial \rho}{\partial y} - \frac{\partial P}{\partial y} \right) \left( \frac{1}{\rho^2} \right) \]

\[ L_{N_{sp}+4} = \frac{v + c}{2 \rho C_p} \left( \frac{\partial P}{\partial y} + \rho \frac{c}{c} \frac{\partial v}{\partial y} \right) \tag{F.3} \]

LODI system primitive derivatives in y direction:

\[ d_1 = \frac{\partial \rho}{\partial t} = \frac{\rho C_p}{c^2} (L_1 + L_{N_{sp}+4}) + L_{N_{sp}+3} \]

\[ d_2 = \frac{\partial u}{\partial t} = L_2 \]

\[ d_3 = \frac{\partial v}{\partial t} = \frac{C_p}{c} (L_{N_{sp}+4} - L_1) \]

\[ d_4 = \frac{\partial w}{\partial t} = L_4 \]

\[ d_5 = \frac{\partial T}{\partial t} = L_1 + L_3 + \sum_{\alpha=2}^{N_{sp}-1} L_{\alpha+3} + L_{N_{sp}+4} \]

\[ d_6 = \frac{\partial Y_1}{\partial t} = -\left( \frac{r}{r_1 - r_{N_p}} \right) \left( \frac{L_{N_{sp}+3}}{\rho} + \frac{L_3}{T} \right) \]

\[ d_{5+\alpha} = \frac{\partial Y_{\alpha}}{\partial t} = -\left( \frac{r}{r_\alpha - r_{N_p}} \right) \frac{L_{\alpha+3}}{T} \quad \alpha = (2, ..., N_{sp} - 1) \tag{F.4} \]
LODI system wave amplitudes in $z$ direction:

$$\begin{align*}
L_1 &= \frac{w - c}{2 \rho C_p} \left( \frac{\partial P}{\partial z} - \rho_c \frac{\partial w}{\partial z} \right) \\
L_2 &= w T \left[ \frac{1}{\rho} \left( \frac{1}{c^2} \frac{\partial P}{\partial z} - \frac{\rho}{c} \frac{\partial w}{\partial z} \right) - \left( \frac{r_1 - r_{N_p}}{r} \right) \frac{\partial Y_1}{\partial z} \right] \\
L_3 &= \frac{\partial u}{\partial z} \\
L_4 &= \frac{\partial v}{\partial z} \\
L_{\alpha+3} &= -w T \left( \frac{r_1 - r_{N_p}}{r} \right) \frac{\partial Y_\alpha}{\partial z} ; \quad \alpha = (2, ..., N_{sp} - 1) \\
L_{N_{sp}+3} &= w \left( \frac{\partial P}{\partial z} - \frac{\partial P}{\partial z} \frac{1}{c^2} \right) \\
L_{N_{sp}+4} &= \frac{w + c}{2 \rho C_p} \left( \frac{\partial P}{\partial z} + \rho c \frac{\partial u}{\partial z} \right)
\end{align*}$$

(F.5)

LODI system primitive derivatives in $z$ direction:

$$\begin{align*}
d_1 &= \frac{\partial \rho}{\partial t} = \frac{\rho C_p}{c^3} (L_1 + L_{N_{sp}+4}) + L_{N_{sp}+3} \\
d_2 &= \frac{\partial w}{\partial t} = \frac{C_p}{c} (L_{N_{sp}+4} - L_1) \\
d_3 &= \frac{\partial u}{\partial t} = L_3 \\
d_4 &= \frac{\partial v}{\partial t} = L_4 \\
d_5 &= \frac{\partial T}{\partial t} = L_1 + L_2 + \sum_{\alpha=2}^{N_{sp}-1} L_{\alpha+3} + L_{N_{sp}+4} \\
d_6 &= \frac{\partial Y_1}{\partial t} = -\left( \frac{r}{r_1 - r_{N_p}} \right) \left( \frac{L_{N_{sp}+3}}{\rho} + \frac{L_2}{T} \right) \\
d_{5+\alpha} &= \frac{\partial Y_\alpha}{\partial t} = -\left( \frac{r}{r_1 - r_{N_p}} \right) \frac{L_{\alpha+3}}{T} ; \quad \alpha = (2, ..., N_{sp} - 1)
\end{align*}$$

(F.6)
### WENO COEFFICIENTS

Table G.1: Coefficients $c_{rj}$ for $k=1$ to 6.

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<th>$j = 1$</th>
<th>$j = 2$</th>
<th>$j = 3$</th>
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<td>7/60</td>
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<td>19/20</td>
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<td>-1/30</td>
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<td>-163/60</td>
<td>79/20</td>
<td>-71/20</td>
<td>49/20</td>
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</table>
The coefficients $\beta$ necessary for computation of the smoothness indicators $\beta_r$, for different stencils presented for reconstruction for $k = 1$ to 6.

$k=1$
There exist single stencil for reconstruction : $S_0 = (i)$.

$$\beta_0 = f_i f_i$$

$k=2$
There exist two stencils for reconstruction : $S_0 = (i, i + 1)$ $S_1 = (i - 1, i)$.

$$\beta_0 = (f_{i+1} - f_i)^2$$

$$\beta_1 = (f_i - f_{i-1})^2$$

$k=3$
There exist three stencils for reconstruction : $S_0 = (i, i + 1, i + 2)$ $S_1 = (i - 1, i, i + 1)$ $S_2 = (i - 2, i - 1, i)$.

$$\beta_0 = \frac{13}{12} (f_i - 2 f_{i+1} + f_{i+2})^2 + \frac{1}{4} (3 f_i - 4 f_{i+1} + f_{i+2})^2$$

$$\beta_1 = \frac{13}{12} (f_{i-1} - 2 f_i + f_{i+1})^2 + \frac{1}{4} (f_{i-1} - f_{i+1})^2$$

$$\beta_2 = \frac{13}{12} (f_{i-2} - 2 f_{i-1} + f_i)^2 + \frac{1}{4} (f_{i-2} - 4 f_{i-1} + 3 f_i)^2$$

$k=4$
There exist four stencils for reconstruction : $S_0 = (i, i + 1, i + 2, i + 3)$ $S_1 = (i - 1, i, i + 1, i + 2, i + 3)$

<table>
<thead>
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<th>K</th>
<th>r=0</th>
<th>r=1</th>
<th>r=2</th>
<th>r=3</th>
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<td>5/77</td>
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Table G.2: Coefficients $d_r$ for $k=1$ to 6.
2) \( S_2 = (i - 2, i - 1, i, i + 1) \quad S_3 = (i - 3, i - 2, i - 1, i) \). 

\[
\beta_0 = 2107f_i^2 - 9402f_{i+1}f_i + 7042f_{i+2}f_i - 1854f_{i+3}f_i + 11003f_{i+4}^2 - 17246f_{i+2}f_{i+1} \\
+ 4642f_{i+3}f_{i+1} + 7043f_{i+2}^2 - 3882f_{i+3}f_{i+2} + 547f_{i+3}^2 \\
\beta_1 = 547f_{i-1}^2 - 2522f_{i-1}f_i + 1922f_{i+1}f_{i-1} - 494f_{i+2}f_{i-1} + 3443f_i^2 - 5966f_{i-1}f_i \\
+ 1602f_{i+2}f_i + 2843f_{i+1}^2 - 1642f_{i+2}f_{i+1} + 267f_{i+1}^2 \\
\beta_2 = 267f_{i-2}^2 - 1642f_{i-2}f_{i-1} + 1602f_{i-2}f_{i-1} - 494f_{i+1}f_{i-2} + 2843f_{i-1}^2 - 5966f_{i-2}f_{i-1} \\
+ 1922f_{i+1}f_{i-1} + 3443f_i^2 - 2522f_{i+1}f_i + 547f_{i+1}^2 \\
\beta_3 = 547f_{i-3}^2 - 3882f_{i-3}f_{i-2} + 4642f_{i-1}f_{i-3} - 1854f_{i-3}f_{i-3} + 7043f_{i-2}^2 - 17246f_{i-2}f_{i-2} \\
+ 7042f_{i-2}f_{i-2} + 11003f_{i-2}^2 - 9402f_{i-1}f_{i-2} + 2107f_i^2 \\
\]

\[K=5\]

There exist five stencils for reconstruction: 

\( S_0 = (i, i + 1, i + 2, i + 3, i + 4) \quad S_1 = (i - 1, i, i + 1, i + 2, i + 3) \quad S_2 = (i - 2, i - 1, i, i + 1, i + 2) \quad S_3 = (i - 3, i - 2, i - 1, i, i + 1) \quad S_4 = (i - 4, i - 3, i - 2, i - 1, i). \)

\[
\beta_0 = 107918f_i^2 - 649501f_{i+1}f_i + 758823f_{i+2}f_i - 411487f_{i+3}f_i + 86329f_{i+4}f_i \\
+ 1020563f_{i+1}^2 - 2462076f_{i+2}f_{i+1} + 1358458f_{i+3}f_{i+1} - 288007f_{i+4}f_{i+1} + 1521393f_{i+2}^2 \\
- 1704396f_{i+3}f_{i+2} + 364863f_{i+4}f_{i+2} + 482963f_{i+3}^2 - 208501f_{i+4}f_{i+3} + 22658f_{i+4}^2 \\
\beta_1 = 22658f_{i-1}^2 - 140251f_{i-1}f_i + 165153f_{i+1}f_{i-1} - 88297f_{i+2}f_{i-1} + 18079f_{i+3}f_{i-1} \\
+ 242723f_i^2 - 611976f_{i+1}f_i + 337018f_{i+2}f_i - 70237f_{i+3}f_i + 406293f_{i+1}^2 \\
- 464976f_{i+2}f_{i+1} + 99213f_{i+3}f_{i+1} + 138563f_{i+4}f_{i+1} - 60871f_{i+2}f_{i+3} + 6908f_{i+4}^2 \\
\beta_2 = 6908f_{i-2}^2 - 51001f_{i-2}f_{i-2} + 67923f_{i-2}f_{i-2} - 38947f_{i+1}f_{i-2} + 8209f_{i+2}f_{i-2} \\
+ 104963f_i^2 - 299076f_{i+1}f_i + 179098f_{i+2}f_{i+1} - 38947f_{i+2}f_{i+2} + 231153f_i^2 \\
- 299076f_{i+1}f_i + 67923f_{i+2}f_{i+1} + 104963f_{i+3}f_{i+1} - 51001f_{i+2}f_{i+3} + 6908f_{i+2}^2 \\
\beta_3 = 6908f_{i-3}^2 - 60871f_{i-2}f_{i-3} + 99213f_{i-1}f_{i-3} - 70237f_{i+1}f_{i-3} + 18079f_{i+2}f_{i-3} \\
+ 138563f_{i-2}^2 - 464976f_{i-1}f_{i-2} + 337018f_{i-2}f_{i-2} - 88297f_{i+1}f_{i-2} + 406293f_{i-1}^2 \\
- 611976f_{i-1}f_{i-1} + 165153f_{i+1}f_{i-1} + 242723f_i^2 - 140251f_{i+2}f_{i-1} + 22658f_{i+3}^2 \\
\beta_4 = 22658f_{i-4}^2 - 208501f_{i-3}f_{i-4} + 364863f_{i-2}f_{i-4} - 288007f_{i-1}f_{i-4} + 86329f_{i-4}^2 \\
+ 482963f_{i-3}^2 - 1704396f_{i-2}f_{i-3} + 1358458f_{i-1}f_{i-3} - 411487f_{i-2}f_{i-3} + 1521393f_{i-2}^2 \\
- 2462076f_{i-1}f_{i-2} + 758823f_{i-2}f_{i-2} + 1020563f_{i-1}^2 - 649501f_{i-1}f_i + 107918f_i^2 \]
There exist six stencils for reconstruction: $S_0 = (i, i + 1, i + 2, i + 3, i + 4, i + 5)$, $S_1 = (i - 1, i + 1, i + 2, i + 3, i + 4)$, $S_2 = (i - 2, i - 1, i + 1, i + 2, i + 3)$, $S_3 = (i - 3, i - 2, i - 1, i + 1, i + 2)$, $S_4 = (i - 5, i - 4, i - 3, i - 2, i - 1, i)$, $S_5 = (i - 4, i - 3, i - 2, i - 1, i)$.

\[
\beta_0 = 6150211 f_{i-1}^2 - 47460464 f_{i+1} f_i + 76206736 f_{i+2} f_i - 63394124 f_{i+3} f_i + 27060170 f_{i+4} f_i 
- 4712740 f_{i+5} f_i + 94851237 f_{i+1}^2 - 311771244 f_{i+2} f_{i+1} + 262901672 f_{i+3} f_{i+1} - 113206788 f_{i+4} f_{i+1} 
+ 19834350 f_{i+5} f_{i+1} + 260445372 f_{i+2}^2 - 444003904 f_{i+3} f_{i+2} + 192596472 f_{i+4} f_{i+2} 
- 33918804 f_{i+5} f_{i+2} + 190757572 f_{i+3}^2 - 166461044 f_{i+4} f_{i+3} + 29442256 f_{i+5} f_{i+3} 
+ 36480687 f_{i+4}^2 = -12950184 f_{i+5} f_{i+4} + 1152561 f_{i+5}^2
\]

\[
\beta_1 = 1152561 f_{i-1}^2 - 9117992 f_{i-1} f_{i+1} + 14742480 f_{i+1} f_{i-1} - 12183636 f_{i+2} f_{i-1} + 5134574 f_{i+3} f_{i-1} 
- 880548 f_{i+1} f_{i-1} + 19365967 f_{i}^2 - 65224244 f_{i+1} f_{i} + 55053752 f_{i+2} f_{i} - 23510468 f_{i+3} f_{i} 
+ 4067018 f_{i+4} f_{i} + 56662212 f_{i+1}^2 - 97838784 f_{i+2} f_{i+1} + 42405032 f_{i+3} f_{i+1} - 7408908 f_{i+4} f_{i+1} 
+ 43093692 f_{i+2}^2 - 37913324 f_{i+3} f_{i+2} + 6694608 f_{i+4} f_{i+2} + 8449957 f_{i+3}^2 - 3015728 f_{i+4} f_{i+3} 
+ 271779 f_{i+4}^2
\]

\[
\beta_2 = 271779 f_{i-2}^2 - 2380800 f_{i-1} f_{i-2} + 4086352 f_{i-2} f_{i-1} - 3462252 f_{i+1} f_{i-2} + 1458762 f_{i+2} f_{i-2} 
- 245620 f_{i+1} f_{i-2} + 5653317 f_{i-1}^2 - 20427884 f_{i-1} f_{i-1} + 17905032 f_{i+1} f_{i-1} - 7727988 f_{i+2} f_{i-1} 
+ 1325006 f_{i+3} f_{i-1} + 19510972 f_{i}^2 - 35817664 f_{i+1} f_{i} + 15929912 f_{i+2} f_{i} - 2792660 f_{i+3} f_{i} 
+ 17195652 f_{i+1}^2 - 15880404 f_{i+2} f_{i+1} + 2863984 f_{i+3} f_{i+1} + 3824847 f_{i+2} + 1429976 f_{i+3} f_{i+2} 
+ 139633 f_{i+3}^2
\]

\[
\beta_3 = 139633 f_{i-3}^2 - 1429976 f_{i-2} f_{i-3} + 2863984 f_{i-1} f_{i-3} - 2792660 f_{i} f_{i-3} + 1325006 f_{i+1} f_{i-3} 
- 245620 f_{i+2} f_{i-3} + 3824847 f_{i-1}^2 - 15880404 f_{i-1} f_{i-1} + 15929912 f_{i+2} f_{i-1} - 7727988 f_{i+3} f_{i-2} 
+ 1458762 f_{i+2} f_{i-2} + 17195652 f_{i-1}^2 - 35817664 f_{i-1} f_{i-1} + 17905032 f_{i+2} f_{i-1} - 3462252 f_{i+3} f_{i-1} 
+ 19510972 f_{i}^2 - 20427884 f_{i+1} f_{i} + 4086352 f_{i+2} f_{i} + 5653317 f_{i+1}^2 - 2380800 f_{i+2} f_{i+1} 
+ 271779 f_{i+2}^2
\]

\[
\beta_4 = 271779 f_{i-4}^2 - 3015728 f_{i-3} f_{i-4} + 6694608 f_{i-2} f_{i-4} - 7408908 f_{i-1} f_{i-4} + 4067018 f_{i} f_{i-4} 
- 880548 f_{i+1} f_{i-4} + 8449957 f_{i-2}^2 - 37913324 f_{i-2} f_{i-2} + 42405032 f_{i-3} f_{i-2} - 23510468 f_{i} f_{i-3} 
+ 5134574 f_{i+1} f_{i-3} + 43093692 f_{i-1}^2 - 97838784 f_{i-1} f_{i-1} + 55053752 f_{i+2} f_{i-1} - 12183636 f_{i+3} f_{i-1} 
+ 56662212 f_{i-2}^2 - 65224244 f_{i-1} f_{i-2} + 14742480 f_{i+1} f_{i-2} + 19365967 f_{i}^2 - 9117992 f_{i+1} f_{i} 
+ 1152561 f_{i+1}^2
\]

(G.1)
\( \beta_5 = 1152561 f_{i-5}^2 - 12950184 f_{i-4} f_{i-4} + 29442256 f_{i-3} f_{i-4} - 33918804 f_{i-2} f_{i-4} \\
+ 19834350 f_{i-1} f_{i-4} - 4712740 f_{i} f_{i-4} + 36480687 f_{i-4}^2 - 166461044 f_{i-3} f_{i-3} \\
+ 192596472 f_{i-2} f_{i-3} - 113206788 f_{i-1} f_{i-3} + 27060170 f_{i} f_{i-3} + 190757572 f_{i-2}^2 \\
- 444003904 f_{i-2} f_{i-2} + 262901672 f_{i-1} f_{i-2} - 63394124 f_{i} f_{i-2} + 260445372 f_{i-2}^2 \\
- 311771244 f_{i-1} f_{i-1} + 76206736 f_{i} f_{i-1} + 94851237 f_{i-1}^2 - 47460464 f_{i} f_{i} + 6150211 f_i^2 \)
APPENDIX H

COMPRESSIBLE TEMPORAL MIXING LAYER A PRIORI TESTS

The \textit{a priori} tests which were conducted by Lardjane and Fedioun, for the sub-grid terms of NS equations in the framework of a temporal binary mixing layer are presented in this appendix.

The effect of physical parameters (density ratio, temperature, compressibility...) on the following sub-grid terms which were derived filtering the multicomponent compressible NS equations in convective form was investigated.

![Figure H.1: Computational box for the temporal direct numerical simulation](image)

Figure H.1: Computational box for the temporal direct numerical simulation
• filtered equation of motion

\begin{align*}
A_{1i} &= -[\rho(\bar{u}_i \bar{u}_j - \bar{u}_i \bar{u}_j)]_{ij} \\
A_{2i} &= (\mu \bar{S}_{ij} - \bar{\mu} \bar{S}_{ij})_{ij} \\
A_{3i} &= -[\rho \left( \bar{r}T - \bar{r}T \right)]_{ij} \quad \text{(H.1)}
\end{align*}

• filtered energy equation

\begin{align*}
B_1 &= \left( \bar{\alpha}_T - \bar{\alpha}_T \right)_{ij} \\
B_2 &= -\left( \bar{p}\bar{u}_{ij} - \bar{p}\bar{u}_{ij} \right) \\
B_3 &= \bar{u}_{ij} \bar{\bar{u}}_{ij} - \bar{u}_{ij} \bar{\bar{u}}_{ij} \quad \text{(H.3)}
\end{align*}

• filtered species equation

\begin{align*}
C_{\alpha 1} &= -[\rho \left( \bar{\bar{Y}}_{\alpha i} \bar{u}_j - \bar{\bar{Y}}_{\alpha i} \bar{u}_j \right)]_{ij} \\
C_{\alpha 2} &= [\rho \left( \bar{D}_{\alpha m} \bar{\bar{Y}}_{\alpha ij} - \bar{D}_{\alpha m} \bar{\bar{Y}}_{\alpha ij} \right)]_{ij} \quad \text{(H.4)}
\end{align*}

The field of sub-grid terms was rebuilt out of D.N.S. flow variables by means of explicit S.O.C.F. box filtering of flow variables [53]. The DNS solutions were obtained applying 6th order compact scheme for spatial discretization and 3rd order time integration and accurate transport model. The physical and numerical parameters for several DNS computations are gathered in Tables H.1 and H.2. The sub-grid terms were quantified as r.m.s. of transverse profiles obtained averaging variables in homogeneous streamwise x and spanwise y directions. The relative magnitudes of sub-grid terms are presented in Figures H.2 - H.7. It is seen that, when the index \(i\) corresponds to the cross-flow direction, \(A_{3i}\) becomes non-negligible when the \(N_2\) stream is heated relatively to the \(O_2\) stream and may overcome \(A_{1ij}\) in some cases.

Table H.1: D.N.S. physical and numerical parameters for the \(O_2/N_2\) pair.

<table>
<thead>
<tr>
<th>(O_2/N_2)</th>
<th>#1</th>
<th>#2</th>
<th>#3</th>
<th>#4</th>
<th>#5</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Re_{ref})</td>
<td>100</td>
<td>100</td>
<td>200</td>
<td>400</td>
<td>100</td>
</tr>
<tr>
<td>(M_e)</td>
<td>0.293</td>
<td>0.586</td>
<td>0.880</td>
<td>0.293</td>
<td>0.246</td>
</tr>
<tr>
<td>(U (\text{m/s}))</td>
<td>±100</td>
<td>±200</td>
<td>±300</td>
<td>±100</td>
<td>±100</td>
</tr>
<tr>
<td>(T_2/T_1 (K))</td>
<td>300/300</td>
<td>300/300</td>
<td>300/300</td>
<td>300/300</td>
<td>600/300</td>
</tr>
<tr>
<td>(Lx\times Ly\times Lz)</td>
<td>40×40×80</td>
<td>same as #1</td>
<td>40×40×40</td>
<td>28×28×40</td>
<td>same as #1</td>
</tr>
<tr>
<td>(Nx\times Ny\times Nz)</td>
<td>181×181×245</td>
<td>same as #1</td>
<td>255×255×257</td>
<td>same as #3</td>
<td>same as #1</td>
</tr>
</tbody>
</table>

154
Table H.2: D.N.S physical and numerical parameters for the O$_2$/H$_2$ pair.

<table>
<thead>
<tr>
<th>O$_2$/H$_2$</th>
<th>#1</th>
<th>#2</th>
<th>#3</th>
<th>#4</th>
<th>#5</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Re_{ref}$</td>
<td>100</td>
<td>100</td>
<td>200</td>
<td>200</td>
<td>100</td>
</tr>
<tr>
<td>$M_c$</td>
<td>0.121</td>
<td>0.243</td>
<td>0.607</td>
<td>0.121</td>
<td>0.113</td>
</tr>
<tr>
<td>$U$ (m/s)</td>
<td>±100</td>
<td>±200</td>
<td>±500</td>
<td>±100</td>
<td>±100</td>
</tr>
<tr>
<td>$T_2/T_1$ (K)</td>
<td>300/300</td>
<td>300/300</td>
<td>300/300</td>
<td>300/300</td>
<td>600/300</td>
</tr>
<tr>
<td>$L_x \times L_y \times L_z$</td>
<td>33\times33\times60</td>
<td>20\times20\times60</td>
<td>same as #2</td>
<td>same as #1</td>
<td>same as #2</td>
</tr>
<tr>
<td>$N_x \times N_y \times N_z$</td>
<td>199\times199\times485</td>
<td>same as #1</td>
<td>same as #1</td>
<td>same as #1</td>
<td>same as #1</td>
</tr>
</tbody>
</table>
Figure H.2: Subrid terms (H.1). Relative magnitude: $O_2/N_2$

Figure H.3: same as Figure H.2: $O_2/H_2$

Figure H.4: Subrid terms (H.3). Relative magnitude: $O_2/N_2$

Figure H.5: same as Figure H.4: $O_2/H_2$

Figure H.6: Subrid terms (H.4). Relative magnitude: $O_2/N_2$

Figure H.7: same as Figure H.6: $O_2/H_2$
### Computational Hours

Table I.1: Relative cost of LES and MILES, non-reacting and reacting.

<table>
<thead>
<tr>
<th>Method</th>
<th>Non-Reacting</th>
<th>Reacting</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>wall time</td>
<td>CPU/step</td>
</tr>
<tr>
<td>MILES_EULER</td>
<td>14238 s</td>
<td>508.4 s</td>
</tr>
<tr>
<td>MILES_NS</td>
<td>16581 s</td>
<td>599.2 s</td>
</tr>
<tr>
<td>LES_SSF</td>
<td>16482 s</td>
<td>625.1 s</td>
</tr>
<tr>
<td>LES_SM</td>
<td>15680 s</td>
<td>603.6 s</td>
</tr>
</tbody>
</table>