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# **MOLECFIT Pipeline User Manual**

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### **Change Record**

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| 2.4     | 03/05/2012 | Sect. 1, 2.2, 4.4.2, 6.5,<br>and 7          | Discussing telluric absorption correction,<br>Figure 1 modified                                    |
| 2.3     | 21/03/2012 | Sect. 4 and 5.5.3                           | Voigt profile approximation added  |
| 2.2     | 16/03/2012 | Sect. 5                                     | Updating information concerning GDAS profiles<br>Comment about width zero components (convolution) |
| 2.1     | 23/11/2011 | Sect. 4 and 5.5.3                           | Adding comments of Alain Smette  |
| 2.0     | 06/04/2011 | All   | Second version   |
| 1.0     | 28/02/2011 | All   | First version  |



### Contents

| 1 | Intro | oduction                                       | 5  |
|---|-------|--|----|
|   | 1.1   | Scope  | 5  |
|   | 1.2   | Acknowledgements                               | 5  |
|   | 1.3   | Stylistic conventions                          | 6  |
|   | 1.4   | Notational Conventions                         | 6  |
| 2 | Rela  | ated Documents                                 | 7  |
|   | 2.1   | Applicable Documents                           | 7  |
|   | 2.2   | Reference Documents                            | 7  |
| 3 | Defi  | nitions, Acronyms and Abbreviations            | 9  |
| 4 | Ove   | rview  | 11 |
|   | 4.1   | The Project                                    | 11 |
|   | 4.2   | Algorithm                                      | 12 |
|   | 4.3   | Major Changes between MOLECFIT v1.5.9 and v3.x | 16 |
|   |       | 4.3.1 Driver                                   | 16 |
|   |       | 4.3.2 Plot Files                               | 16 |
|   |       | 4.3.3 ASCII Data Files                         | 16 |
|   |       | 4.3.4 OpenMP                                   | 16 |
|   | 4.4   | Major Changes between MOLECFIT v3.x and v4.0   | 16 |
|   |       | 4.4.1 Removed Parameters                       | 16 |
|   |       | 4.4.2 Changed Parameters                       | 17 |
|   | 4.5   | Instrument Specific MOLECFIT Recipes           | 17 |
| 5 | Quio  | ck Start                                       | 19 |
|   | 5.1   | The MOLECFIT Pipeline Recipes                  | 19 |
|   | 5.2   | Running the MOLECFIT Pipeline Recipes          | 19 |
|   |       | 5.2.1 Getting Started with <i>EsoRex</i>       | 19 |
|   | 5.3   | Running MOLECFIT using <i>EsoRex</i>           | 20 |
|   |       | 5.3.1 Environment Variables                    | 20 |



|  | 5.3.2   | Input spectrum FITS header keywords   | 21  |
|--|---|---|---|
|  | 5.3.3   | Executing molecfit_model  | 21  |
|  | 5.3.4   | Executing molecfit_calctrans  | 22  |
|  | 5.3.5   | Executing molecfit_correct  | 23  |
| Rec  | ipe Ref   | erence  | 24  |
|  | -   |   |   |
| ••••   |   |   |   |
| 0.2  | 6.2.1   |   |   |
|  | 6.2.2   | ·   |   |
| 6.3  |   |   |   |
|  | 6.3.1   |   |   |
|  | 6.3.2   | Calctrans Recipe  |   |
|  | 6.3.3   |   | 56  |
| 6.4  | Input S   | Spectrum  | 59  |
| 6.5  | Output  | t Products  | 59  |
|  |   |   |   |
|  |   |   |   |
| Gui  |   | NOLECFIT 1.5.9 Users  | 60  |
| <b>Gui</b><br>7.1  |   | <b>IOLECFIT 1.5.9 Users</b> tion Format   |   |
|  | Execu   |   | 60  |
| 7.1  | Execut<br>Param   | tion Format   | 60<br>60  |
| 7.1<br>7.2   | Execut<br>Param<br>Input C  | tion Format   | 60<br>60<br>61  |
| <ol> <li>7.1</li> <li>7.2</li> <li>7.3</li> <li>7.4</li> </ol>                           | Execut<br>Param<br>Input C<br>Plot Fi   | tion Format   | 60<br>60<br>61  |
| <ol> <li>7.1</li> <li>7.2</li> <li>7.3</li> <li>7.4</li> </ol>                           | Execut<br>Param<br>Input C<br>Plot Fi<br>Extra I  | tion Format   | 60<br>60<br>61<br>62<br>62  |
| <ol> <li>7.1</li> <li>7.2</li> <li>7.3</li> <li>7.4</li> <li>7.5</li> </ol>              | Execut<br>Param<br>Input C<br>Plot Fi<br>Extra I<br>Simple                                      | tion Format   | <ul> <li>60</li> <li>60</li> <li>61</li> <li>62</li> <li>62</li> <li>65</li> </ul>  |
| <ol> <li>7.1</li> <li>7.2</li> <li>7.3</li> <li>7.4</li> <li>7.5</li> <li>7.6</li> </ol> | Execut<br>Param<br>Input C<br>Plot Fi<br>Extra I<br>Simple                                      | tion Format   | <ul> <li>60</li> <li>60</li> <li>61</li> <li>62</li> <li>62</li> <li>65</li> <li>67</li> </ul>  |
| <ol> <li>7.1</li> <li>7.2</li> <li>7.3</li> <li>7.4</li> <li>7.5</li> <li>7.6</li> </ol> | Execut<br>Param<br>Input C<br>Plot Fi<br>Extra I<br>Simple<br>Old Pa                            | tion Format   | <ul> <li>60</li> <li>60</li> <li>61</li> <li>62</li> <li>62</li> <li>65</li> <li>67</li> <li>67</li> </ul>  |
| <ol> <li>7.1</li> <li>7.2</li> <li>7.3</li> <li>7.4</li> <li>7.5</li> <li>7.6</li> </ol> | Execut<br>Param<br>Input C<br>Plot Fi<br>Extra I<br>Simple<br>Old Pa<br>7.7.1                   | tion Format   | <ul> <li>60</li> <li>60</li> <li>61</li> <li>62</li> <li>62</li> <li>65</li> <li>67</li> <li>67</li> <li>67</li> </ul>  |
| <ol> <li>7.1</li> <li>7.2</li> <li>7.3</li> <li>7.4</li> <li>7.5</li> <li>7.6</li> </ol> | Execut<br>Param<br>Input C<br>Plot Fi<br>Extra I<br>Simple<br>Old Pa<br>7.7.1<br>7.7.2          | tion Format   | <ul> <li>60</li> <li>60</li> <li>61</li> <li>62</li> <li>62</li> <li>65</li> <li>67</li> <li>67</li> <li>67</li> <li>67</li> </ul>  |
| <ol> <li>7.1</li> <li>7.2</li> <li>7.3</li> <li>7.4</li> <li>7.5</li> <li>7.6</li> </ol> | Execut<br>Param<br>Input C<br>Plot Fi<br>Extra I<br>Simple<br>Old Pa<br>7.7.1<br>7.7.2<br>7.7.3 | tion Format   | <ul> <li>60</li> <li>60</li> <li>61</li> <li>62</li> <li>62</li> <li>65</li> <li>67</li> <li>67</li> <li>67</li> <li>67</li> <li>68</li> </ul>  |
|  | <ul><li>6.1</li><li>6.2</li><li>6.3</li><li>6.4</li></ul>                                       | 5.3.3<br>5.3.4<br>5.3.5<br><b>Recipe Ref</b><br>6.1 The St<br>6.2 The .1<br>6.2.1<br>6.2.1<br>6.2.2<br>6.3 The Pa<br>6.3.1<br>6.3.2<br>6.3.3<br>6.4 Input S | 5.3.3       Executing molecfit_model         5.3.4       Executing molecfit_calctrans         5.3.5       Executing molecfit_correct         6.1       The SOF files         6.2       The .rc files         6.2.1       molecfit_model default .rc parameters         6.2.2       molecfit_correct default .rc parameters         6.3       The Parameters         6.3.1       Model Recipe         6.3.2       Calctrans Recipe         6.3.3       Correct Recipe         6.4       Input Spectrum |



|    |       | 7.7.7     | Line Spread Functions   | 69 |
|----|-------|-----------|---|----|
|    |       | 7.7.8     | Ambient Parameters  | 70 |
|    |       | 7.7.9     | Instrument Parameters   | 70 |
|    |       | 7.7.10    | Atmostpheric Profiles   | 71 |
| 8  | The   | Model     |   | 72 |
|    | 8.1   | Atmos     | pheric profiles and meteorological data                         | 72 |
|    |       | 8.1.1     | MIPAS profiles  | 72 |
|    |       | 8.1.2     | GDAS profiles   | 77 |
|    |       | 8.1.3     | ESO Meteo Monitor   | 77 |
|    |       | 8.1.4     | Processing of ESO Meteor Monitor data, GDAS, and MIPAS profiles | 79 |
|    | 8.2   | Radiat    | ive transfer code   | 81 |
|    |       | 8.2.1     | Line File / Line-By-Line Radiative Transfer Model (LNFL/LBLRTM) | 82 |
|    |       | 8.2.2     | aer line database   | 83 |
|    | 8.3   | Molecu    | ular spectra  | 85 |
|    | 8.4   | Therm     | al emission by telescope  | 85 |
|    | 8.5   | Adapta    | ation of model to input spectrum                                | 86 |
|    |       | 8.5.1     | The continuum   | 86 |
|    |       | 8.5.2     | The wavelength solution   | 86 |
|    |       | 8.5.3     | The resolution  | 87 |
| 9  | Insta | allation  |   | 89 |
| 10 | Cod   | o Porfo   | ormance   | 90 |
| 10 | 000   |           |   |    |
| 11 | Tips  | and tr    | icks  | 91 |
| A  | Exp   | ert Fitti | ing   | 93 |
|    | A.1   | Mappi     | ng Multiple Extension Data                                      | 93 |
|    | A.2   | Mappi     | ng Ranges to Chips  | 94 |
|    | A.3   | Expert    | Range Specific Continuum Modelling                              | 94 |
|    | A.4   | Expert    | Chip Specific Wavelength Correcting                             | 95 |
|    | A.5   | Expert    | Mode Initial Fitting Values                                     | 95 |



|        | А.6                | Expert  | Analysis of the BEST_FIT_PARAMETERS.fits FILE                  | 96   |
|--------|--------------------|---|--|--|
|        |                    | A.6.1   | The value of the Status variable                               | 96   |
|        |                    | A.6.2   | Wavelength correction  | 97   |
|        |                    | A.6.3   | Continuum  | 98   |
|        | A.7                | Scaling   | g the transmission function to the PWV of the science spectrum | 98   |
|        | A.8                | Quality   | of the correction parameter QC MEAN_ABS_DEV MAX                | 99   |
| в      | Para               | ameters   |  | 100  |
|        | B.1                | Molecf  | it Parameter Name Changes: MOLECFIT_MODEL                      | 100  |
|        | B.2                | Molecf  | it Parameter Name Changes: MOLECFIT_CALCTRANS                  | 103  |
|        | B.3                | Molecf  | it Parameter Name Changes: MOLECFIT_CORRECT                    | 103  |
|        | B.4                | molec:  | fit_model SOF Tag names  | 104  |
|        | B.5                | molec:  | fit_calctrans SOF Tag Names                                    | 105  |
|        | B.6                | molec:  | fit_correct SOF Tag Names                                      | 105  |
|        | B.7                | Molecf  | it Output Files  | 106  |
|        |                    | B.7.1   | molecfit_model Output Files                                    | 106  |
|        |                    | B72   | molecfit_calctrans Output Files                                | 107  |
|        |                    | BE  |  |  |
|        |                    |   | molecfit_correct Output Files                                  |  |
| с      | Mole               |   | molecfit_correct Output Files                                  |  |
| C<br>D |                    | B.7.3<br>ecular S   | molecfit_correct Output Files                                  | 107  |
| _      |                    | B.7.3<br>ecular S   | molecfit_correct Output Files                                  | 107<br>108<br>114  |
| _      | Mair               | B.7.3<br>ecular S<br>ntenanc<br>Mainte  | molecfit_correct Output Files                                  | 107<br>108<br>114  |
| _      | Mair               | B.7.3<br>ecular S<br>ntenanc<br>Mainte<br>D.1.1   | molecfit_correct Output Files                                  | 107<br><b>108</b><br><b>114</b><br>114<br>114                                    |
| _      | Mair               | B.7.3<br>ecular S<br>ntenanc<br>Mainte<br>D.1.1<br>D.1.2  | molecfit_correct Output Files                                  | 107<br><b>108</b><br><b>114</b><br>114<br>114<br>114                             |
| _      | Mair               | B.7.3<br>ecular S<br>ntenanc<br>Mainte<br>D.1.1<br>D.1.2<br>D.1.3                                       | molecfit_correct Output Files                                  | 107<br><b>108</b><br><b>114</b><br>114<br>114<br>114<br>115                      |
| _      | <b>Maiı</b><br>D.1 | B.7.3<br>ecular S<br>ntenanc<br>Mainte<br>D.1.1<br>D.1.2<br>D.1.3<br>D.1.4                              | molecfit_correct Output Files                                  | 107<br><b>108</b><br><b>114</b><br>114<br>114<br>114<br>115<br>115               |
| _      | <b>Maiı</b><br>D.1 | B.7.3<br>ecular S<br>ntenanc<br>Mainte<br>D.1.1<br>D.1.2<br>D.1.3<br>D.1.4<br>License                   | molecfit_correct Output Files                                  | 107<br><b>108</b><br><b>114</b><br>114<br>114<br>114<br>115<br>115<br>117        |
| _      | <b>Maiı</b><br>D.1 | B.7.3<br>ecular S<br>ntenanc<br>Mainte<br>D.1.1<br>D.1.2<br>D.1.3<br>D.1.4<br>License<br>D.2.1          | molecfit_correct Output Files                                  | 107<br><b>108</b><br><b>114</b><br>114<br>114<br>115<br>115<br>117<br>117        |
| _      | <b>Maiı</b><br>D.1 | B.7.3<br>ecular S<br>ntenanc<br>Mainte<br>D.1.1<br>D.1.2<br>D.1.3<br>D.1.4<br>License<br>D.2.1<br>D.2.2 | molecfit_correct Output Files                                  | 107<br><b>108</b><br><b>114</b><br>114<br>114<br>115<br>115<br>117<br>117<br>118 |



| Doc. Number:  | VLT-MAN-ESO- |
|---------------|--------------|
|               | 19550-5772   |
| Doc. Version: | 4.4.2        |
| Released on:  | 2025-02-11   |
| Page:         | 8 of 120     |



### 1 Introduction

MOLECFIT is a software tool for correcting atmospheric absorption features in astronomical observations, based on fitting synthetic transmission spectra to the astronomical data. It can also estimate molecular abundances, especially the water vapour content of the Earth's atmosphere.

Correcting for these telluric absorption features usually requires supplementary calibration data which are very expensive to obtain in terms of telescope time. In addition, the observation scheduling flexibility is restricted as these data usually have to be taken immediately before or after the science observations to which the correction will be applied, due to the high variability of the telluric absorption which depends on the state and chemical composition of the atmosphere at the time of observations. A tool for telluric absorption correction such as MOLECFIT, that is based on synthetic transmission spectra calculated by a radiative transfer code does not require these supplementary calibration data if the science data can be fitted directly. This can save a significant amount of valuable telescope time and as such increase the instrumental efficiency.

As useful as MOLECFIT is, it has some limitations and observation of a telluric standard star may be better suited in some conditions. For instance, MOLECFIT might perform poorly with:

- large number of intrinsic features,
- little or no continuum emission,
- low signal-to-noise ratio,
- large airmass observations with high water vapor content.

The tool can be run from a command-line or more conveniently through a GUI (see the MOLECFIT Reflex Tutorial [RD3]).

#### 1.1 Scope

This document describes the MOLECFIT package version 4.4.2. The most recent version of this document may be found on the VLT Instrument Pipelines MOLECFIT Downloads page, at <a href="https://www.eso.org/sci/software/pipelines/molecfit/molecfit-pipe-recipes.html">https://www.eso.org/sci/software/pipelines/molecfit/molecfit-pipe-recipes.html</a>.

The examples on running individual pipeline recipes in this manual use the esorex command and manually created list of input files. Several interfaces to automatically organise the data, create the list of input files and execute the pipeline recipes in the proper sequence are available, see the *https://www.eso.org/pipelines* for details.

#### 1.2 Acknowledgements

MOLECFIT was developed for ESO by a team of astronomers at the Institute for Astro- and Particle Physics at the University of Innsbruck on the base of an IDL prototype developed by Alain Smette, Hannes Horst, Hugues Sana, Alexandre Gallenne. It is one of three sky modeling and correction tools provided in the framework of the Austrian in-kind contribution to ESO.



More recently, with version 3, it has been refactored by the ESO Pipelines team, and divided into the user interface component, and its underlying TELLURICCORR library with the goal to be easily integrated into the various instrument workflows. Development and maintenance of MOLECFIT is now carried out by the AAO Software Pipelines Team.

If you use MOLECFIT to produce a scientific publication, we ask that you please use the following references:

- Smette, A. and Sana, H. and Noll, S. and Horst, H. and Kausch, W. and Kimeswenger, S. and Barden, M. and Szyszka, C. and Jones, A. M. and Gallenne, A. and Vinther, J. and Ballester, P. and Taylor, J. 2015, A&A 576, A77<sup>1</sup>
- Kausch, W. and Noll, S. and Smette, A. and Kimeswenger, S. and Barden, M. and Szyszka, C. and Jones, A. M. and Sana, H. and Horst, H. and Kerber, F. 2015, A&A 576, A78<sup>2</sup>
- Molecfit Pipeline Team, MOLECFIT Pipeline User Manual v4.4.2, 2023. ESO VLT-MAN-ESO-19550-5772

#### **1.3 Stylistic conventions**

Throughout this document the following stylistic conventions are used:

- **bold** in text sections for commands and other user input which has to be typed as shown *italics* in the text and example sections for parts of the user input which have to be replaced with real contents
- teletype in the text for FITS keywords, program names, file paths, and terminal output, and as the general style for examples, commands, code, etc

In example sections expected user input is indicated by a leading shell prompt.

In the text **bold** and *italics* may also be used to highlight words.

#### 1.4 Notational Conventions

Hierarchical FITS keyword names, appearing in the document, are given using the dot-notation to improve readability. This means, that the prefix "HIERARCH ESO" is left out, and the spaces separating the keyword name constituents in the actual FITS header are replaced by a single dot.

<sup>&</sup>lt;sup>1</sup>Bibtex reference: https://ui.adsabs.harvard.edu/abs/2015A%26A...576A..77S/exportcitation <sup>2</sup>BibTex reference: https://ui.adsabs.harvard.edu/abs/2015A%26A...576A..78K/exportcitation



### 2 Related Documents

#### 2.1 Applicable Documents

| [AD                      | 1] VLT D<br>Contr             | nata Flow System Specifications for Pipeline and Quality   | VLT-SPE-ESO-19600-1233   |
|--------------------------|-------------------------------|--|--|
| [AD<br>[AD<br>[AD<br>[AD | 2] Data<br>3] DFS F<br>4] ESO | Flow for VLT Instruments Requirement Specification<br>Pipeline & Quality Control — User Manual<br>DICB — Data Interface Control Document<br>non Pipeline Library User Manual | VLT–SPE–ESO–19000–1618/2.0<br>VLT–MAN–ESO–19500-1619<br>GEN–SPE–ESO–19400–0794<br>VLT–MAN–ESO–19500–2720 |
| 2.2                      | Referen                       | ce Documents   |  |
|                          |                               |  |  |
|                          | [RD01]                        | Detailed Specification Document  | VLT-SPE-ESO-19550-5769   |
|                          | [RD02]                        | DR06 User Manual   | VLT-MAN-ESO-19550-5286   |
|                          | [RD03]                        | Reflex MOLECFIT Tutorial   | VLT-MAN-ESO-19550-5928   |
|                          | [RD04]                        | Moré, J.J., Garbow, B.S., & Hillstrom, K.E. 1980,  |  |
|                          |                               | User Guide for MINPACK-1,  | - 111  |
|                          | [RD05]                        | Argonne National Laboratory Report ANL-80-74, Argonne Clough, S.A, Shephard, M.W., Mlawer. E.J., et al. 2005,  | e, III.  |
|                          | [IID00]                       | J. Quant. Spectrosc. Radiat. Transfer, 91, 233   |  |
|                          | [RD06]                        | Noll, S., Kausch, W., Barden, M., et al. 2012,   |  |
|                          | [11200]                       | A&A, 543, A92  |  |
|                          | [RD07]                        | VLT Astronomical Site Monitor, ASM Data, User Manual   | VLT-MAN-ESO-17440-1773   |
|                          | [RD08]                        | Anu Dudhia, private communication, 2009.   |  |
|                          | [RD09]                        | SM-01 User Manual,   | VLT-MAN-ESO-19550-5770   |
|                          | [RD10]                        | Niro, F., Jucks, K., & Hartmann, JM. 2005,   |  |
|                          |                               | J. Quant. Spectrosc. Radiat. Transfer, 95, 469   |  |
|                          | [RD11]                        | Clough, S.A., Iacono, M.J., & Moncet, JL. 1992,  |  |
|                          |                               | J. Geophys. Res., 97, 15761  |  |
|                          | [RD12]                        | Masuda, K., Takashima, T., & Takayama, Y. 1988,  |  |
|                          |                               | Remote Sens. Environ., 24, 313   |  |
|                          | [RD13]                        | Wu, X. & Smith, L. 1997,   |  |
|                          |                               | Appl. Opt., 36, 2609   |  |
|                          | [RD14]                        | Patat, F., Moehler, S., O'Brien, K., at al. 2011,  |  |
|                          |                               | A&A, 527, A91  |  |
|                          | [RD15]<br>[RD16]              | SM-03 Science Report,<br>Modigliani. A., Goldoni, P., Royer, F., et al. 2010,  | VLT-TRE-ESO-19550-5774   |
|                          | נהטוסן                        | Proc. SPIE, 7737, 773728   |  |
|                          | [RD17]                        | LBLRTM, http://rtweb.aer.com/lblrtm_frame.htm  | ٦  |
|                          | [RD18]                        | CMPFIT, http://www.physics.wisc.edu/~craigm/i  |  |
|                          | [RD19]                        | AER, http://www.physics.wisc.cdu/ cruigm/i   | , captionic  |
|                          | [RD20]                        | HITRAN, http://www.cfa.harvard.edu/HITRAN/   |  |
|                          | [RD21]                        | NOAA, http://www.arl.noaa.gov/READYamet.php  |  |



- [RD22] GDAS, http://ready.arl.noaa.gov/gdas1.php
- [RD23] GRIB, http://nomads.ncep.noaa.gov/txt\_descriptions/
  - fast\_downloading\_grib.shtml
- [RD24] RFM, http://www.atm.ox.ac.uk/RFM/
- [RD25] LBLRTM\_instructions,
  - http://irina.eas.gatech.edu/Lab\_5560/lblrtm/lblrtm\_inst.html
- [RD26] PWV, http://www.eso.org/observing/dfo/quality/GENERAL/PWV/HEALTH/
- [RD27] CDIAC, http://cdiac.ornl.gov/
- [RD28] LBLRTM FAQ, http://rtweb.aer.com/docs/aqFAQ\_LBLRTM.pdf



### 3 Definitions, Acronyms and Abbreviations

| AER<br>AOPP | Atmospheric and Environmental Research Inc.<br>Atmospheric, Oceanic and Planetary Physics, Oxford University, UK |
|-------------|--|
| ARL         | Air Resources Laboratory   |
| ASCII       | American Standard Code for Information Interchange   |
| CalibDB     | Calibration Database   |
| CPL         | Common Pipeline Library  |
| CCD         | Charge Coupled Device  |
| CRIRES      | CRyogenic high-resolution InfraRed Echelle Spectrograph  |
| DFS         | Data Flow System   |
| DRS         | Data Reduction System  |
| EMM         | ESO Meteo Monitor  |
| Envisat     | Environmental Satellite  |
| ESO         | European Southern Observatory  |
| EsoRex      | ESO Recipe Execution Tool  |
| FITS        | Flexible Image Transport System  |
| FOV         | Field Of View  |
| FORS        | Focal Reducer/low dispersion Spectrograph  |
| FWHM        | full width at half-maximum   |
| GDAS        | Global Data Assimilation System  |
| GRIB        | GRIdded Binary   |
| GUI         | Graphical User Interface   |
| HITRAN      | HIgh-resolution TRANsmission molecular absorption database   |
| HTML        | Hypertext Markup Language  |
| IDL         | Interactive Data Language  |
| KMOS        | K-band Multi Object Spectrograph   |
| LBLRTM      | Line-by-line Radiative Transfer Model  |
| LNFL        | Line File  |
| LSF         | Line Spread Function   |
| MIPAS       | Michelson Interferometer for Passive Atmospheric Sounding  |
| MIR         | mid-infrared   |
| NCEP        | National Centers for Environmental Prediction  |
| NIR         | near-infrared  |
| NOAA        | National Oceanic and Atmospheric Administration  |
| OB          | Observation Block  |
| pixel       | picture element (of a raster image)  |
| PSF         | Point Spread Function  |
| PWV         | precipitable water vapour  |
| QC          | Quality Control  |
| RFM         | Reference Forward Model  |
| RMS         | root mean square   |
| RPG         | Radiometer Physics GmbH  |
| SDP         | Science Data Product   |
| SOF         | Set Of Frames  |



| TBD   | To be defined                                |
|-------|--|
| TBC   | To be confirmed                              |
| VISIR | VLT Imager and Spectrometer for mid InfraRed |
| VLT   | Very Large Telescope ]                       |
| WCS   | World Coordinate System                      |



### 4 Overview

This document is organised as follows:

Section 4 gives a brief overview of the project and the incorporated algorithms. Section 5 contains a description on how to run the code, the required input parameter file, and the output files. Users of MOLECFIT version 1.59 may wish to read Section 7, which describes how the user interaction with the package has changed since the v3.0 overhaul. In Section 8 the atmospheric model and its adaptation to the observed spectrum are described in detail. Section 9 provides information on the installation procedures, and tips and tricks are outlined in Section 11. Finally, the appendices provide a reference, and information for the advanced user.

#### 4.1 The Project

Ground-based astronomical observations suffer from emission and absorption processes in the atmosphere which deteriorate the quality of the obtained data. At wavelengths longer than  $2 - 2.5 \,\mu m$ , where thermal radiation from molecules in the lower atmosphere dominates, the amount of this perturbing background radiation can determine whether scientific observations are feasible at all. For this wavelength regime, it is crucial to be able to estimate the intensity of the atmospheric emission in advance. Such a prediction requires good knowledge of the column densities of atmospheric constituents that significantly contribute to the greenhouse effect. The most important molecules are water ( $H_2O$ ), carbon dioxide ( $CO_2$ ), methane ( $CH_4$ ), nitrous oxide  $(N_2O)$ , and ozone  $(O_3)$ . In particular, a good knowledge of the water abundance, i.e. air humidity, is essential, since H<sub>2</sub>O is the main contributor to the IR atmospheric spectra. Moreover, it is much more variable than the other important species. Due to this variability of molecular abundances, the removal of telluric absorption features from astronomical spectra depends on observations of telluric standard stars with relatively smooth continua, at similar time and airmass as the scientific target. Even if those observations are available, the telluric absorption correction becomes tricky in the near- and mid-IR, where wavelength regions with negligible atmospheric absorption are hardly present. In this case, a reliable determination of the shape of the unabsorbed continuum is very difficult if an interpolation approach is used. Therefore, a realistic model of the atmospheric absorption for given observing conditions makes the telluric absorption correction more reliable, and reduces the number of required observations of telluric standard stars.

For a periodic monitoring of the abundances of crucial atmospheric constituents as well as a high-quality correction of telluric absorption features in astronomical spectra a fast, user-friendly, and reliable software tool is needed. MOLECFIT provides such a tool. Thanks to the advances made in modelling the Earth's atmosphere, it is now possible to compute realistic atmospheric emission and absorption spectra. Such model spectra can be fitted to observed atmospheric spectra with relative deviations of only a few percent. The MOLECFIT pipeline is a tool designed to fit such atmospheric models and derive reliable estimates of the atmosphere's convolution impact on observed spectra. The incorporated model spectra are generated using the radiative transfer code LBLRTM [RD17]. The basic input for the radiative transfer code is an atmospheric profile, which is created from a standard atmosphere (containing information on height, pressure, temperature, and chemical composition for general tropical environments up to 120 km), modelled meteorological GDAS data (containing pressure, temperature, and relative humidity for the region of the observing site for elevations up to  $\approx 25 \text{ km}$ ), and on-site meteorological measurements by the ESO Meteo Monitor (EMM). Using iterative techniques, the input atmospheric profile is varied to obtain a model spectrum that fits the scientific input spectrum (see Section 4.2 for more details). In this process, it is also necessary to optimise the scaling, wavelength grid, and resolution of the



model. To reliably compute radiance spectra, grey body radiation from the telescope itself also has to be taken into account.

The use of MOLECFIT as part of the data reduction process can significantly improve the efficiency of astronomical instruments working in the thermal IR such as CRIRES or VISIR. Since strong molecular absorption affects all observable wavelength regimes except for the visual, many ESO instruments could benefit from an improved correction of telluric features in spectra of astronomical targets.

The MOLECFIT package relies on the following third-party code and input files:

- CMPFIT: The C version of the least-squares fitting library mpfit by C. Markwardt [RD18] based on the FORTRAN fitting routine MINPACK-1 by Moré et al. [RD04]
- The radiative transfer code LBLRTM v12.11. This publicly available software is developed within the Radiative Transfer Working Group of Atmospheric and Environmental Research Inc. (AER); see also Clough et al. [RD05], [RD19], and [RD17]) for more details. LBLRTM can handle all molecules incorporated in a line parameter database, e.g. HITRAN [RD20], and offers a wide range of possibilities for adjusting input parameters. An additional part of this software is the LNFL, which provides the required line information based on the line parameter database.
- A line database of molecular parameters: Currently, the line parameter list aer\_v\_3.8.1.2 delivered with the LBLRTM v12.11 package is included in the MOLECFIT package. This line parameter list is built from HITRAN 2016 [RD20] and contains some updates (see [RD17] and Appendix D.1.3 for more details).
- An atmospheric profile valid for the time of observation: The profile is created using standard atmospheric profiles, on-site measurements by the EMM [RD07], and GDAS data (a product of the NCEP (National Centers for Environmental Prediction) model, created by the Air Resources Laboratory (ARL) of the National Oceanic and Atmospheric Administration (NOAA), see [RD19], [RD21], [RD22]). Such atmospheric profiles include pressure, temperature, and humidity for a number of layers of the atmosphere, at any point on the Earth for all recent dates (back to December 2004), on a 3h-grid basis.
- Perl scripts get\_inv.pl and get\_grib.pl from the NCEP NOMAD server [RD23] for fast download of GDAS data in the GRIB2 (GRIdded Binary 2) format.

#### 4.2 Algorithm

Block diagrams of the basic functionality of the software are shown in Figures 4.1 and 4.2 providing, respectively, a general overview of the workflow, and a more detailed view of the molecfit\_model component.

First, the code reads the science spectrum from a FITS table or FITS image, and an ASCII driver file containing the user input parameters. If available, ESO keywords including EMM data are directly taken from FITS header information. A single atmospheric profile is compiled from data provided by three sources: a standard atmospheric profile for a given climate zone, an appropriate GDAS model profile for the time of the observation and telescope site, and the corresponding ground-based EMM measurements (see Section 8.1 for more details). Input for the radiative transfer model is the resulting merged atmospheric profile (with a possible pre-selection of relevant molecules) and the target airmass at the time of observation (see Section 8.2). To match the observed spectrum, the code adapts the atmospheric spectrum (either transmission or radiation; see Section 8.3) further



| Doc. Number:  | VLT-MAN-ESO- |
|---------------|--------------|
|               | 19550-5772   |
| Doc. Version: | 4.4.2        |
| Released on:  | 2025-02-11   |
| Page:         | 17 of 120    |



**Figure 4.1:** Overview of the software workflow. It shows the input and output for the three executables molecfit\_model, molecfit\_calctrans, and molecfit\_correct and the connection between them.



## MOLECFIT Pipeline User Manual

| Doc. Number:  | VLT-MAN-ESO- |
|---------------|--------------|
|               | 19550-5772   |
| Doc. Version: | 4.4.2        |
| Released on:  | 2025-02-11   |
| Page:         | 18 of 120    |
|               |              |



Figure 4.2: Workflow of the molecfit\_model routine.



by flux scaling, wavelength grid correction, and convolution with a suitable instrument profile (see Section 8.5). For a radiance spectrum, the contribution of thermal emission from the telescope can also be taken into account (see Section 8.4).

The central component of the algorithm is the comparison/fitting of the calculated and the input science spectrum by means of mpfit [RD18]. The  $\chi^2$  minimisation procedure of this routine is based on the Levenberg-Marquardt technique (see Moré et al., [RD04]), an iterative search algorithm characterised by gradient-controlled jumps in parameter space. Since this technique is prone to finding local minima, reasonable starting values and constraints for the fit parameters are required. mpfit checks whether the desired fit quality is reached. If this is not the case, it changes fit parameters in an appropriate way to search for a better  $\chi^2$ . Each function call causes a new calculation of the sky model. For a change of the molecular abundances, the profiles of molecules can be scaled by simple factors, which represent a subset of the fit parameters provided to mpfit. The other parameters are coefficients of polynomials for continuum scaling, coefficients of Chebyshev polynomials for the object of the sector of the wavelength solution, the FWHM of boxcar, Gaussian, and Lorentzian kernels that are used to build a realistic instrumental profile, and the telescope emissivity if a radiance spectrum is computed. When a satisfactory fit is reached after several iterations of the  $\chi^2$  minimisation procedure, the code writes the best-fit spectrum, atmospheric profile, and fit parameters to output files. The best-fit molecular profiles are integrated to get molecular columns in ppmv. The molecfit\_model recipe also computes the water vapour content of the atmosphere in mm. The atmospheric abundances are part of a special output summary file (see Section 6.5).

The fit parameters are defined in the input .rc file (see Section 6.2). Dynamically setting fit flags to include and exclude parameters from the fitting procedure allows the code to find a reasonable solution in a relatively fast and robust way. In detail, the molecfit\_model recipe follows a plan consisting of six steps by default:

- Step 1: scaling of the continuum
- Step 2: wavelength and resolution fit
- · Step 3: rescaling of the continuum
- Step 4: fitting of the column density for each molecule
- · Step 5: joint continuum, wavelength, and resolution fit
- Step 5: fit of all components (molecules column density, continuum, wavelength, and resolution)

For the telluric absorption correction, molecfit\_calctrans must be used with an input set of frames (SOF) which includes the name of the file with the best-fit parameters obtained by molecfit\_model. The molecfit\_calctrans recipe calculates the atmospheric transmission for the full wavelength range of the input spectrum and corrects this spectrum using this function. These calculations are separated from the fitting procedure, since the run of a radiative transfer code for a wide wavelength range is very time consuming. This is particularly critical if the fit is optimised by several code runs with different input parameters. For this reason, the fitting procedure should be performed for several well-defined narrow wavelength ranges that can be provided either as part of the input .rc file or in an independent FITS file (see Section 6.3). Very wide fit ranges are also not recommended due to the probable failure of the polynomial continuum fit. The narrow wavelength range of CRIRES allows one to fit the full spectrum at once. The molecfit\_calctrans recipe writes a FITS table with the model transmission function, the telluric absorption corrected spectrum, and a quality flag.

Finally, molecfit\_correct corrects the spectra provided as a list in its input file, preserving their format.



#### 4.3 Major Changes between MOLECFIT v1.5.9 and v3.x

A major overhaul and refactoring of the MOLECFIT package was carried out in 2018-2020, which included changes which are backward incompatible to v1.5.9. We encourage users to move to the latest supported version, currently v4.4.2.

For a detailed listing of parameter name and other user input changes between the two versions, please see Section 7.

#### 4.3.1 Driver

MOLECFIT 1.5.9 consisted of a set of standalone executables. These have been replaced with a set of *EsoRex* recipes to be processed from the *EsoRex* environment. As a consequence changes have been made to certain parameter names, parameter value formats, and how input files are declared in order to be compliant with *EsoRex* pipeline standards. Further details of the differences of the parameter formats are covered in Section 7.

#### 4.3.2 Plot Files

MOLECFIT versions 3.x and later do not support the 1.5.9 plot file creation options, as *EsoRex* and *Reflex* offer other support methods to graph results.

#### 4.3.3 ASCII Data Files

All data files must be in FITS format – i.e. ASCII files are not supported. This is in line with *EsoRex* pipeline standards.

#### 4.3.4 OpenMP

MOLECFIT 3.x and later support OpenMP and will try to utilise multi cores by performing the fitting process on multiple independent ranges in parallel. The maximum number of threads to use in the executions is by default set to 1 (i.e. no parallel processing), but can be overridden by the value of the environment variable OMP\_NUM\_THREAD if that has been set.

#### 4.4 Major Changes between MOLECFIT v3.x and v4.0

#### 4.4.1 Removed Parameters

The following parameters have been removed:

- CONT\_BARY\_RV (not yet implemented)
- FIT\_CHIP[1234] (no longer used)



- FIT\_RANGE[1234] (no longer used)
- LBLRTM\_REF\_LAT (use LATITUDE\_VALUE or value in LATITUDE\_KEYWORD)
- LBLRTM\_H1 (use ELEVATION\_VALUE or value in ELEVATION\_KEYWORD)
- MASK\_BINARY (no longer used)
- MASK\_THRESHOLD (no longer used)
- OPENMP\_THREADS (no longer used)
- OUTPUT\_NAME (no longer used)
- OUTPUT\_PATH (no longer used)
- WLC\_REF (no longer used)

#### 4.4.2 Changed Parameters

The name or meaning of the following parameters have changed:

- CONTINUUM\_N (see Tab. 6.1)
- CONTINUUM\_CONST (see Tab. 6.1)
- FIT\_CONTINUUM (see Tab. 6.1)
- FIT\_WLC (see Tab. 6.1)
- LNFL\_LINE\_DB=aer\_v\_3.8.1.2 (updated from aer\_v\_3.8)
- MAP\_REGIONS\_TO\_CHIP is new (see Tab. 6.1)
- WAVELENGTHS\_FRAME is now WAVELENGTH\_FRAME
- WL\_GTO\_MICRON is now WLG\_TO\_MICRON
- USE\_ONLY\_INPUT\_PRIMARY\_EXTENSION is now USE\_ONLY\_INPUT\_PRIMARY\_DATA

#### 4.5 Instrument Specific MOLECFIT Recipes

The following pipelines now have their own independent MOLECFIT recipes:

- FORS
- KMOS
- X-Shooter



These streamlined recipes have been designed to make it easier to run MOLECFIT on the respective pipeline data products with several instrument defaults already configured.

Additional instruments will be added as they are incorporated into their pipelines.

For more information on these recipes and how to run them, please see the respective pipeline user manuals.



### 5 Quick Start

This chapter provides a brief description of the MOLECFIT pipeline recipes, their usage and execution.

#### 5.1 The MOLECFIT Pipeline Recipes

The MOLECFIT pipeline consists of the following three recipes:

- 1. molecfit\_model: determines the parameters that best fit the data in user selected include regions;
- 2. molecfit\_calctrans: determines the telluric transmission spectrum over the full spectral range of the input spectrum;
- 3. molecfit\_correct: corrects the list of input spectra using the transmission spectrum provided by molecfit\_calctrans.

Since 2022, similar MOLECFIT recipes are now available for some specific instruments. Please see Section 4.5 for more details.

#### 5.2 Running the MOLECFIT Pipeline Recipes

#### 5.2.1 Getting Started with EsoRex

*EsoRex* is a command-line tool which can be used to execute the recipes of all standard VLT/VLTI instrument pipelines. With *EsoRex* in your path, the general structure of an *EsoRex* command line is

1> esorex [esorex options] [recipe [recipe options] [sof [sof]...]]

where options appearing before the recipe name are options for *EsoRex* itself, and options given after the recipe name are options which affect the recipe.

All available *EsoRex* options can be listed with the command

1> esorex --help

and the full list of available parameters of a specific recipe can be obtained with the command

1> esorex --help <recipe name>

The output of this command shows as parameter values the current setting, i.e. all modifications from a configuration file or the command line are already applied.

The listing of all recipes known to EsoRex can be obtained with the command



1> esorex --recipes

The last arguments of an *EsoRex* command are the so-called *set-of-frames*. A *set-of-frames* is a simple ASCII file containing a list of input data files for the recipe. Each input file is followed by an unique identifier (frame classification or frame tag), indicating the contents of this file. The input files can be given as a relative or absolute path; however, *EsoRex* allows the use of environment variables so that a common directory prefix can be abreviated. Individual lines may be commented out by using a hash character (#) in the first column. An example of a *set-of-frames* is shown in the following:

```
1> cat molecfit_model.sof
$SOF_DATA/raw/XSH0OTER_NIR_Pipeline_R71_input.fits SCIENCE
$SOF_DATA/static_calib/XSH0OTER_NIR_Pipeline_R71_molecules.fits MOLECULES
$SOF_DATA/static_calib/XSH0OTER_NIR_Pipeline_R71_range_wl_inc.fits WAVE_EXCLUDE
$SOF_DATA/static_calib/XSH0OTER_NIR_Pipeline_R71_range_pix_exc.fits PIXEL_EXCLUDE
```

These *set-of-frames* files will be created by the user using a text editor, for instance. Finally, if more than one *set-of-frames* is given on the command-line *EsoRex* concatenates them into a single *set-of-frames*.

#### 5.3 Running MOLECFIT using EsoRex

MOLECFIT consists of three separate recipes: molecfit\_model, molecfit\_calctrans, and molecfit\_correct, each of which is an unique *EsoRex* recipe to be called from the *EsoRex* environment. If both *EsoRex* and the MOLECFIT recipes have been installed correctly, then by typing:

1> esorex -recipes

molecfit\_model, molecfit\_calctrans, and molecfit\_correct will be listed as available recipes. Each recipe can be called via *EsoRex* as follows:

1> esorex --recipe-config= <Pars2Use.rc> <recipe\_name> <SOF\_filename>

where *Pars2Use.rc* contains the list of processing parameters (covered in detail in Section 6.2). The SOF (Set Of Files) contains an ASCII list of the names of the input data files (which are expected to be in FITS format) with their associated tags (see Section 6.1).

#### 5.3.1 Environment Variables

Prior to any recipe execution the following environment variables can be defined:

| TELLURICCORRDIR      | Defines the installation directory where TELLURICCORR may be   |
|----------------------|--|
|                      | found. If this is not set MOLECFIT will use the installation direc-  |
|                      | tory path defined at compile time.   |
| TELLURICCORRDIR_DATA | Defines the data directory. If this is not set MOLECFIT will use the path of the data directory created at compile time. |
| OMP_NUM_THREADS      | Defines the number of threads to use. If undefined one thread per CPU is used.   |



As mentioned earlier, SOF files can include environment variables in file pathnames – e.g. \$HOME, as well as those listed above.

#### 5.3.2 Input spectrum FITS header keywords

The CDELT1, CDELT2 and CDELT3 FITS header keywords are now *deprecated* in favour of the corresponding FITS header keywords CD1\_1, CD2\_2 and CD3\_3. If your input spectrum is missing the latter keywords, a warning will be raised about the input spectrum. This warning can be resolved by adding these keywords to the FITS header.

#### 5.3.3 Executing molecfit\_model

The first stage of MOLECFIT is to derive a telluric model that best describes the given spectral data. This model is defined by three specific group of parameters:

- 1. A list of atmospheric molecules under consideration for their telluric impact in the spectral range of interest e.g. H<sub>2</sub>O, O<sub>2</sub>, O<sub>3</sub>, CH<sub>4</sub>, etc.
- 2. Atmospheric parameters relevant to the time and location that the input science spectra was captured, e.g. altitude-specific values of pressure, temperature, molecular abundances, etc.
- 3. A remaining set of parameters characterising the observational setup, e.g. various relative scales, background contributions, wavelength corrections, etc.

The first group is declared by the user either as a parameter in the relevant .rc file or in a table of a FITS file specified in the SOF file. The second one is derived via interpolation and combination of data from various sources packaged with the molecfit installation. Initial values for the third group of parameters are also provided in the .rc file or in FITS tables. The bulk of the molecfit\_model process is to determine the molecular abundance for the molecules listed, as well as various of the parameters that characterise the observations.

The modelling recipe of MOLECFIT is called via esorex in the following way:

1> esorex --recipe-config= <Pars2Use.rc> molecfit\_model <Files2Use.sof>

where *Pars2Use.rc* contains the list of processing parameters and, at a minimum, the SOF file contains the pathname of the science data file of interest and is tagged as SCIENCE — e.g. :

\$HOME/raw/CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits SCIENCE

If successful, the recipe execution will deliver the following intermediate FITS files:



| BEST_FIT_MODEL.fits       | Model of the best fit to the data.  |  |
|---------------------------|---|--|
| GDAS.fits                 | Initial atmospheric profiles (height, temperature, relative humidity) as    |  |
|                           | a function of altitude, resulting from the combination of the profiles      |  |
|                           | GDAS_BEFORE.fits, GDAS_AFTER.fits, and EMM.                                 |  |
| GDAS_BEFORE.fits          | GDAS profile before the observation time.                                   |  |
| GDAS_AFTER.fits           | GDAS profile after the observation time.                                    |  |
| ATM_PROFILE_STANDARD.fits | Initial atmospheric profiles, in particular, for all molecules available to |  |
|                           | molecfit_model.   |  |
| ATM_PROFILE_COMBINED.fits | Atmospheric profiles resulting from the combination of                      |  |
|                           | ATM_PROFILE_STANDARD, GDAS, and EMM data, used as initial                   |  |
|                           | value by molecfit_model.  |  |

and the following results files that explicitly define the model, and which are required as inputs to the molecfit\_calctrans recipe:

| ATM_PARAMETERS.fits      | The atmospheric profiles, including the volume mixing ratio for each of the relevant molecules as result of the fitting in molecfit_model. |
|--------------------------|--|
| MODEL_MOLECULES.fits     | The list of molecules used in the model, with boolean flags indicating if  |
|                          | the column densities were to be fitted and the initial values of the column densities relative to the combined atmospheric profile.        |
| BEST_FIT_PARAMETERS.fits | The least-squares fitted parameter values.   |

#### 5.3.4 Executing molecfit\_calctrans

The second stage of MOLECFIT generates the telluric model from the set of parameters determined by the model stage, and derives data for a convolution process that can be used to correct spectra with the derived telluric transmission spectrum.

The calctrans process of molecfit is called via *EsoRex* in the following way:

1> esorex --recipe-config= <Pars2Use.rc> molecfit\_calctans <Files2Use.sof>

Where *Pars2Use.rc* contains the list of processing parameters and, at a minimum, the SOF file contains the pathnames of the FITS files that uniquely define the model and are tagged as follows:

\$MYDATADIR/SCIENCE\_FILE.fits SCIENCE \$MYDATADIR/ATM\_PARAMETERS.fits ATM\_PARAMETERS \$MYDATADIR/MODEL\_MOLECULES.fits MODEL\_MOLECULES \$MYDATADIR/BEST\_FIT\_PARAMETERS.fits BEST\_FIT\_PARAMETERS

If successful, the recipe execution will deliver the following intermediate FITS files:

| LBLRTM_RESULTS.fits | The transmission spectrum as produced by the radiative transfer code  |
|---------------------|---|
|                     | (LBLRTM).   |
| TELLURIC_DATA.fits  | The processed transmission spectrum including, in particular, the convolution by the instrumental line spread function. |



and the following results file that exclusively contains the convoluted transmission spectrum, required as input to the third stage - the molecfit\_correct recipe:

TELLURIC\_CORR.fits The convoluted transmission spectrum.

#### 5.3.5 Executing molecfit\_correct

The final stage of MOLECFIT is to correct input spectra with the assumed telluric contamination, using the convoluted transmission data generated by the molecfit\_calctrans recipe. This MOLECFIT recipe is called via *EsoRex* in the following way:

1> esorex --recipe-config= <Pars2Use.rc> molecfit\_correct <Files2Use.sof>

where *Pars2Use.rc* contains the list of processing parameters and the *SOF* file which, at a minimum, contains the pathnames of the science data to be corrected and the TELLURIC\_CORR.fits file - the convoluted telluric transmission spectrum produced by the molecfit\_calctrans recipe – e.g. :

```
$HOME/raw/CRIRES_HighSNR_Telluric_input_AM1p472.fits SCIENCE
./TELLURIC_CORR.fits TELLURIC_CORR
```

If successful, the recipe execution will deliver the following output FITS file:

SCIENCE\_TELLURIC\_CORR\_CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits

that is the input science frame with the original spectra replaced with that of the corrected spectra.



### 6 Recipe Reference

#### 6.1 The SOF files

SOF files are ASCII files that contain a list of file path locations for specific input data. These are in the form:

file path name 1 tagname 1 file path name 2 tagname 2 . . file path name N tagname N

where file path names may contain environment variables and *tagnames* are specific keywords that declare the file's role in the process e.g. input science spectra, input best-fit parameters, etc. A full list of the SOF *tagnames* are given in Appendix B.4.

There are fundamentally two types of tagged files within MOLECFIT: those which are mandatory, and files which are used for optional parameter substitution. The latter is a convenient way of assigning a structured parameter that is better described as a binary table within an external FITS file instead of a long string declaration in the .rc file. A good example is the wavelength inclusion parameter. In the .rc file this is defined as string in the following format:

WAVE\_INCLUDE = ini1, end1, ..., iniN, endN

which for a long set of ranges can be quite complex. Alternatively, if the parameter value is specified as NULL, i.e.

WAVE\_INCLUDE = NULL

then MOLECFIT will look for a file path name in the SOF file with the tag WAVE\_INCLUDE and, if listed, it will assume that these data exist in the specified file as FITS binary table with columns LOWER\_LIMIT and UPPER\_LIMIT. For example, the following string definition of the wavelength ranges in the molecfit\_model .rc file:

WAVE\_INCLUDE = 4.8417300, 4.8421100, 4.8408000, 4.841320, 4.837330, 4.837950, 4.838740, 4.83949, 4.844310, 4.845210

can be defined by the following parameter setting

WAVE\_INCLUDE = NULL

and the associated line in the SOF file:

./wave\_include.fits WAVE\_INCLUDE

where wave\_include.fits contains the following bin table:



| LOWER_LIMIT        | UPPER_LIMIT        |
|--------------------|--------------------|
| 1D                 | 1D                 |
| 4.84173000000E+00  | 4.842110000000E+00 |
| 4.84080000000E+00  | 4.84132000000E+00  |
| 4.837330000000E+00 | 4.83795000000E+00  |
| 4.838740000000E+00 | 4.83949000000E+00  |
| 4.844310000000E+00 | 4.845210000000E+00 |

This format is identical for the WAVE\_EXCLUDE and PIX\_INCLUDE parameters and SOF tags.

Another less straight forward example is the list of molecules to be included and/or fitted, and their values of relative column. In the .rc files this is defined by three parameters, as shown in this example:

LIST\_MOLEC = H<sub>2</sub>O, CO<sub>2</sub>, O<sub>3</sub>, CO, OCS FIT\_MOLEC = 1, 1, 1, 1, 0 REL\_COL = 0.422, 1.012, 2.808, 0.809

The first line defines the list of molecules to use. The second lists if they are to be fitted (1) or not (0), and the third lists the values of the relative columns to be used, respectively. Any of these three parameters can be parameter substituted, but only by a single tagged file for all three (tagname = MOLECULES). The declared FITS file must contain a binary table that includes the required data in a column with a header of the parameter name being substituted.

For example, the above can be replaced with

```
LIST_MOLEC = NULL
FIT_MOLEC = NULL
REL_COL = NULL
```

with the associated line in the SOF file:

\$FULL\_DIR\_PATH/molecules.fits MOLECULES

where molecules.fits contains the following bin table:

| LIST_MOLEC       | FIT_MOLEC | REL_COL           |
|------------------|-----------|-------------------|
| 3A               | 1J        | 1D                |
| H <sub>2</sub> 0 | 1         | 4.22000000000E-01 |
| CO <sub>2</sub>  | 1         | 1.01200000000E+00 |
| O <sub>3</sub>   | 1         | 2.80800000000E+00 |
| CO               | 1         | 8.09000000000E-01 |
| OCS              | 0         | 1.00000000000E+00 |

Another example that mixes the two approaches:

```
LIST_MOLEC = H_20, CO_2, O_3, CO, OCS
FIT_MOLEC = 1, 1, 1, 1, 0
REL_COL = NULL
```

where molecules.fits need only contain the single column table "REL\_COL".



The most common tag names and their purpose are listed below. For a full list see Appendix B.4.

With regard to molecfit\_model:

| Tag Name      | Туре                   | Purpose                                   |
|---------------|------------------------|---|
| SCIENCE       | Mandatory              | Input science spectrum                    |
| WAVE_INCLUDE  | Parameter Substitution | List of wavelength ranges to be included  |
| WAVE_EXCLUDE  | Parameter Substitution | List of wavelength ranges to be excluded  |
| PIXEL_EXCLUDE | Parameter Substitution | List of pixel ranges to be excluded       |
| MOLECULES     | Parameter Substitution | List of molecules to include in the model |

With regard to molecfit\_calctrans:

| Tag Name            | Туре      | Purpose   |
|---------------------|-----------|---|
| ATM_PARAMETERS      | Mandatory | Atmospheric parameter values as used in the model |
| MODEL_MOLECULES     | Mandatory | List of molecules as used in the model            |
| BEST_FIT_PARAMETERS | Mandatory | Best fit values as used in the model              |

With regard to molecfit\_correct:

| Tag Name      | Туре      | Purpose  |
|---------------|-----------|--|
| TELLURIC_CORR | Mandatory | Telluric convolution data necessary to correct telluric contami- |
|               |           | nated data   |
| TELLURIC_DATA | Mandatory | In case WLC_REF is set to MODEL, as it includes the wave-        |
|               |           | length grid with the telluric lines used as reference            |

**PLEASE NOTE** :molecfit\_correct will only accept IDP compliant transmission products, if the input SCIENCE product is IDP compliant.

#### 6.2 The .rc files

The *EsoRex* .rc files are only necessary in the event of using any parameter values other than the default, and even then only need to list the parameters with the non-default values. Generate an .rc file containing the default values as follows:

1> esorex --create-config= <filename> <recipe\_name>

This can then be used as a template for the parameter settings required by the user.



#### 6.2.1 molecfit\_model default .rc parameters

The command

1> esorex --create-config= Default\_model.rc molecfit\_model

generates a file containing the following default parameters:

```
# File: Default_model.rc
    #
    # Note: This configuration file has been automatically
    #
            generated by the esorex (v3.13.6) program.
    #
    # Date: 04-May-2023 18:28:10
    #
    #
    # --USE_ONLY_INPUT_PRIMARY_DATA
10
    # Value=TRUE implies that only the fits primary contains the input science flux
    # data.
    # Value=FALSE implies that the fits extensions also contains input science
    # flux data.
    USE_ONLY_INPUT_PRIMARY_DATA=FALSE
    # --USE_DATA_EXTENSION_AS_DFLUX
    # Only valid if USE_ONLY_INPUT_PRIMARY_DATA=TRUE. The fits extension index that
    # contains the
   # errors of the science flux data (DFLUX). A value of 0 implies that there is
20
    # no DFLUX.
    USE_DATA_EXTENSION_AS_DFLUX=0
    # --USE_DATA_EXTENSION_AS_MASK
    # Only valid if USE_ONLY_INPUT_PRIMARY_DATA=TRUE. The fits extension index that
    # contains the
    # mask associated with the science flux data. A value of 0 implies that there
    # is no mask data.
    USE_DATA_EXTENSION_AS_MASK=0
30
    # --USE_INPUT_KERNEL
    # If TRUE, use the kernel library if it is provided.
    USE_INPUT_KERNEL=TRUE
    # --MODEL_MAPPING_KERNEL
    # Mapping 'STD_MODEL/SCIENCE' - 'MODEL_KERNEL_LIBRARY' [string with ext_number
    # comma separated (int)] :
    # If set to NULL, check if the TAG[MODEL_MAPPING_KERNEL] FITS BINTABLE values
    # is provided.
    # The FITS BINTABLE have to one column [KERNEL_LIBRARY_EXT].
40
    MODEL_MAPPING_KERNEL=NULL
```



# --LIST\_MOLEC

# List of molecules to be included in the model. Represented as a comma # separated # string of molecule names, e.g. "H20,C02,O3". # If set to NULL, the input TAG[MOLECULES] FITS BINTABLE values have to be # provided # where the FITS BINTABLE specified contains the three columns: # LIST\_MOLEC; FIT\_MOLEC; and REL\_COL. 50LIST\_MOLEC=NULL # --FIT\_MOLEC # List of flags that specify which of the listed molecules are to be fitted for. # Flag=1 implies yes. Flag=0 implies no. Represented as a string of comma # separated # integers in the same order as the listed molecules. For example: if # LIST\_MOLEC="H20,C02,03", then # FIT\_MOLEC="1,0,1" implies that only H2O and O3 should be fitted for. # If set to NULL, the input TAG[MOLECULES] FITS BINTABLE values have to be 60 # provided where the FITS # BINTABLE specified contains the three columns: LIST\_MOLEC; FIT\_MOLEC; and # REL\_COL. FIT\_MOLEC=NULL # --REL\_COL # List of the intial values of fitting of the molecular columns expressed # relatively to the input # ATM profile columns. Represented as a comma separated list of doubles in # the same order as the 70 # listed molecules. For example, if LIST\_MOLEC="H20,CO2,O3", then # REL\_COL="1.0,1.2,0.8" # implies that H2O, CO2 and O3 have initial relative values of 1.0, 1.2 and # 0.8 respectively. # If set to NULL, the input TAG[MOLECULES] FITS BINTABLE values have to be # provided where the FITS # BINTABLE specified contains the three columns: LIST\_MOLEC; FIT\_MOLEC; and # REL\_COL. REL\_COL=NULL 80 # --WAVE\_INCLUDE # Wavelength ranges to be included. Represented as a string of comma separated # doubles in pairs # specifying the start and end wavelengths of a range. The wavelength units # are always in microns. # For example a KMOS sample data in the range of 1.11um to 1.67um may have # WAVE\_INCLUDE="1.773,1.78633,1.79098,1.80434,1.187691,1.189937" to represent # three inclusion regions: # [1.773,1.78633], [1.79098,1.80434] and [1.187691,1.189937]. # If set to NULL, molecfit will check if the TAG[WAVE\_INCLUDE] FITS BINTABLE 90 # values is provided where # the FITS BINTABLE specified has the two columns: LOWER\_LIMIT; and # UPPER\_LIMIT.



WAVE\_INCLUDE=NULL # --WAVE\_EXCLUDE # Wavelength ranges to be excluded. Represented as a string of comma separated # doubles in pairs # specifying the start and end wavelengths of a range. The wavelength units # are always in microns. 100 # as the input science data. For example a KMOS sample data in the range of # 1.11um to 1.67um may have # WAVE\_EXCLUDE="1.773,1.78633,1.79098,1.80434,1.187691,1.189937" to represent # three exclusion regions: # [1.773,1.78633], [1.79098,1.80434] and [1.187691,1.189937]. # If set to NULL, molecfit will check if the TAG[WAVE\_EXCLUDE] FITS BINTABLE # values is provided where # the FITS BINTABLE specified has the two columns: LOWER\_LIMIT; and # UPPER LIMIT. WAVE\_EXCLUDE=NULL 110 # --PIXEL\_EXCLUDE # Pixel ranges to be excluded. Represented as a string of comma separated # integers in pairs specifying the # start and end pixel of a range. For example: # PIXEL\_EXCLUDE="54,128,512,514,1020,1024" represents three # exclusion regions: [54,128], [512,514] and [1020,1024]. # If set to NULL, molecfit will check if the TAG[PIXEL\_EXCLUDE] FITS BINTABLE # values is provided where the # FITS BINTABLE specified has the two columns: LOWER\_LIMIT; and UPPER\_LIMIT. 120 PIXEL\_EXCLUDE=NULL # --TELLURICCORR\_PATH # Installation directory. TELLURICCORR\_PATH=TELLURICCORR\_PARAMETER\_DEFAULT # --TELLURICCORR\_DATA\_PATH # Data directory. TELLURICCORR\_DATA\_PATH=TELLURICCORR\_PARAMETER\_DEFAULT 130 # --TMP\_PATH # Temporary directory. TMP\_PATH=TELLURICCORR\_PARAMETER\_DEFAULT # --SILENT\_EXTERNAL\_BINS # Silent the output of the external binaries. SILENT\_EXTERNAL\_BINS=TRUE # --TRANSMISSION # Type of input spectrum : 0 = Emission(radiance); 1 = Transmission. 140 TRANSMISSION=TRUE # --COLUMN\_LAMBDA # Wavelength column ('NULL' can be used if the file is an image and that



```
# the data are in the primary
     # (data are given by the FITS header keywords [CRVAL1=wave_ini, CD1_1=step])
     # f CD1_1 is absent, then the DEPRECATED CDELT1 keyword will be used.
     COLUMN_LAMBDA=lambda
    # --COLUMN_FLUX
150
     # Flux column.
     COLUMN_FLUX=flux
     # --COLUMN_DFLUX
     # Flux error column (Avoided by writing 'NULL') : 1-sigma error on the flux.
     COLUMN_DFLUX=NULL
     # --COLUMN_MASK
     # Mask column (Avoided by writing 'NULL') : Indicates if a pixel is invalid.
    COLUMN_MASK=NULL
160
     # --DEFAULT_ERROR
     # Default error relative to mean for the case that the error column
     # is not provided.
    DEFAULT_ERROR=0.01
     # --WLG_TO_MICRON
     # Multiplicative factor applied to the wavelength to express is in micron.
     # E.g.: if wavelength is given in nm, the value should be 0.001.
    WLG_TO_MICRON=1.0
170
     # --WAVELENGTH_FRAME
     # Wavelength in vacuum
                                                                 = 'VAC'.
     # Wavelength in air with the observatory reference frame = 'AIR'.
     # Wavelength in vacuum with another reference frame = 'VAC_RV'.
     # Wavelength in air with another
                                                reference frame = 'AIR_RV'.
       (typically the sun or the barycenter of the solar system).
     #
    # In the latter case, the radial velocity of the observatory relative
    # to the external reference frame must be provided in the parameter obs_RV.
    WAVELENGTH_FRAME=VAC
180
     # --OBS_ERF_RV_KEY
     # The radial velocity of the observatory in km/s
     # relative to the external reference frame;
     # It is positive if the distance between the science target and the Earth
     # increases along the line-of-sight to the science target.
     # It must be provided if MF_PARAMETERS_WAVELENGTH_FRAME = 'VAC_RV' or 'AIR_RV'.
    OBS_ERF_RV_KEY=NONE
     # --OBS_ERF_RV_VALUE
190
     # If OBS_ERF_RV_KEYWORD=='NONE' take this value.
     # It must be provided if MF_PARAMETERS_WAVELENGTH_FRAME = 'VAC_RV' or 'AIR_RV'.
     OBS_ERF_RV_VALUE=0.0
     # --CLEAN_MODEL_FLUX
```



```
# Set model flux to 0 for non-fitted pixels.
     CLEAN_MODEL_FLUX=FALSE
     # --FTOL
     # Relative chi-square convergence criterion.
200
     FTOL=1e-10
     # --XTOL
     # Relative parameter convergence criterion.
     XTOL=1e-10
     # --FLUX_UNIT
     # Conversion of fluxes from phot/(s*m2*mum*as2) (emission spectrum only)
     # to flux unit of observed spectrum:
210
     # 0: phot / (s * m<sup>2</sup> * mum * as<sup>2</sup>) [no conversion]
     # 1:
          W / (
                      m^2 * mum * as^2)
     # 2: erg / (s * cm<sup>2</sup> * A * as<sup>2</sup>)
     # 3: mJy / (
                                    as^2)
     # For other units, the conversion factor has to be considered
     # as constant term of the continuum fit.
     FLUX_UNIT=0
     # --FIT_TELESCOPE_BACKGROUND
     # Fit of telescope background -- 1 = yes; 0 = no (emission spectrum only).
220
     FIT_TELESCOPE_BACKGROUND=TRUE
     # --TELESCOPE_BACKGROUND_CONST
     # Initial value for telescope background fit.
     TELESCOPE_BACKGROUND_CONST=0.1
     # --FIT_CONTINUUM
     # Comma deliminated string of flags (1=true, 0=false) for fitting continuum in
     # specific regions.
     # If set to NULL, check if the TAG[WAVE_INCLUDE] points to a FITS BINTABLE
     # with column CONT_FIT_FLAG provided.
230
    FIT_CONTINUUM=1
     # --CONTINUUM_N
     # Polynomial order for continuum model for each region. Presented as a comma
     # deliminated string.
     # If set to NULL, check if the TAG[WAVE_INCLUDE] points to a FITS BINTABLE
     # with column CONT_POLY_ORDER provided.
     CONTINUUM_N=O
     # --CONTINUUM_CONST
240
     # Initial constant term for continuum fit (valid for all fit ranges)
     # [emission spectrum: about 1 for correct flux_unit].
     CONTINUUM_CONST=1.0
     # --MAP_REGIONS_TO_CHIP
     # Comma deliminated string of chip indices that each range is associated with.
```



```
# If set to NULL, check if the TAG[WAVE_INCLUDE] points to a FITS BINTABLE
     # with column MAPPED_TO_CHIP provided.
     MAP_REGIONS_TO_CHIP=1
250
     # --FIT_WLC
     # Flags for including regions in wavelength corrections.
     # If set to NULL, check if the TAG[WAVE_INCLUDE] points to
     # a FITS BINTABLE with column WLC_FIT_FLAG provided.
     FIT_WLC=0
     # --WLC_N
     # Polynomial degree of the refined wavelength solution.
     WLC_N=1
260
     # --WLC_CONST
     # Initial constant term for wavelength adjustment
     # (shift relative to half wavelength range).
     WLC_CONST=0.0
     # --FIT_RES_BOX
     # Fit resolution by Boxcar LSF.
     FIT_RES_BOX=TRUE
     # --RES_BOX
270
     # Initial value for FWHM of Boxcar rel. to slit width
     # at the centre of the spectrum.
     RES_BOX=1.0
     # --FIT_RES_GAUSS
     # Fit resolution by Gaussian.
     FIT_RES_GAUSS=TRUE
     # --RES_GAUSS
     # Initial value for FWHM of the Gaussian in pixels
280
     # at the centre of the spectrum.
     RES_GAUSS=1.0
     # --FIT_RES_LORENTZ
     # Fit resolution by Lorentzian.
     FIT_RES_LORENTZ=TRUE
     # --RES_LORENTZ
     # Initial value for FWHM of the Lorentz in pixels
     # at the centre of the spectrum.
290
     RES_LORENTZ=1.0
     # --KERNMODE
     # Voigtian profile approximation instead of independent Gaussian and
     # Lorentzian?.
     KERNMODE=FALSE
```


```
# --KERNFAC
     # Size of Voigtian/Gaussian/Lorentzian kernel in FWHM.
     KERNFAC=3.0
300
     # --VARKERN
     # Does the kernel size increase linearly with wavelength?.
     VARKERN=FALSE
     # --OBSERVING_DATE_KEYWORD
     # Observing date in years or MJD in days (not string).
     OBSERVING_DATE_KEYWORD=MJD-OBS
     # --OBSERVING_DATE_VALUE
310
     # If OBSERVING_DATE_KEYWORD=='NONE' take this value.
     OBSERVING_DATE_VALUE=-1.0
     # --UTC_KEYWORD
     # UTC in s.
     UTC_KEYWORD=UTC
     # --UTC_VALUE
     # If UTC_KEYWORD=='NONE' take this value.
     UTC_VALUE=-1.0
320
     # --TELESCOPE_ANGLE_KEYWORD
     # Telescope altitude angle in deg.
     TELESCOPE_ANGLE_KEYWORD=ESO TEL ALT
     # --TELESCOPE_ANGLE_VALUE
     # If TELESCOPE_ANGLE_KEYWORD == 'NONE' take this value.
     TELESCOPE_ANGLE_VALUE=90.0
     # --RELATIVE_HUMIDITY_KEYWORD
330
     # Relative humidity in %.
     RELATIVE_HUMIDITY_KEYWORD=ESO TEL AMBI RHUM
     # --RELATIVE_HUMIDITY_VALUE
     # If RELATIVE_HUMIDITY_KEYWORD=='NONE' take this value.
     RELATIVE_HUMIDITY_VALUE=15.0
     # --PRESSURE_KEYWORD
     # Pressure in hPa.
     PRESSURE_KEYWORD=ESO TEL AMBI PRES START
340
     # --PRESSURE_VALUE
     # If PRESSURE_KEYWORD=='NONE' take this value.
     PRESSURE_VALUE=750.0
     # --TEMPERATURE_KEYWORD
     # Ambient temperature in deg C.
     TEMPERATURE_KEYWORD=ESO TEL AMBI TEMP
```



# -- TEMPERATURE\_VALUE 350 # If TEMPERATURE\_KEYWORD=='NONE' take this value. TEMPERATURE\_VALUE=15.0 # --MIRROR\_TEMPERATURE\_KEYWORD # Mirror temperature in deg C. MIRROR\_TEMPERATURE\_KEYWORD=ESO TEL TH M1 TEMP # --MIRROR\_TEMPERATURE\_VALUE # If MIRROR\_TEMPERATURE\_KEYWORD=='NONE' take this value. MIRROR\_TEMPERATURE\_VALUE=15.0 360 # --ELEVATION\_KEYWORD # Elevation above sea level in m (default is Paranal: 2635. m). ELEVATION\_KEYWORD=ESO TEL GEOELEV # --ELEVATION\_VALUE # If ELEVATION\_KEYWORD=='NONE' take this value. ELEVATION\_VALUE=2635.0 # --LONGITUDE\_KEYWORD 370 # Longitude (default is Paranal: -70.4051 deg). LONGITUDE\_KEYWORD=ESO TEL GEOLON # --LONGITUDE\_VALUE # If LONGITUDE\_KEYWORD=='NONE' take this value. LONGITUDE\_VALUE=-70.4051 # --LATITUDE\_KEYWORD # Latitude (default is Paranal: -24.6276 deg). LATITUDE\_KEYWORD=ESO TEL GEOLAT 380 # --LATITUDE\_VALUE # If LATITUDE\_KEYWORD=='NONE' take this value. LATITUDE\_VALUE=-24.6276 # --SLIT\_WIDTH\_KEYWORD # Slit width in arcsec (taken from FITS header if present). SLIT\_WIDTH\_KEYWORD=ESO INS SLIT1 WID # --SLIT\_WIDTH\_VALUE 390 # If SLIT\_WIDTH\_KEYWORD=='NONE' take this value. SLIT\_WIDTH\_VALUE=0.4 # --PIX\_SCALE\_KEYWORD # Pixel scale in arcsec (taken from this file only). PIX\_SCALE\_KEYWORD=NONE # --PIX\_SCALE\_VALUE # If PIX\_SCALE\_KEYWORD=='NONE' take this value.



```
PIX_SCALE_VALUE=0.086
400
     # --REFERENCE_ATMOSPHERIC
     # Reference atmospheric profile. Possible values:
     # - equ.fits (default; equatorial atmosphere, valid for Paranal);
     # - tro.fits (tropical atmosphere);
     # - std.fits (standard atmosphere);
     # - Other file located in :
     #
       ({TELLURICCORR_DATA_PATH}/profiles/mipas/).
     REFERENCE_ATMOSPHERIC=equ.fits
410
     # --GDAS_PROFILE
     # Specify which GDAS profile to use. Possible values:
     # - 'auto', automatic retrieval of the GDAS profiles
     # (P[hPa] HGT[m] T[K] RELHUM[%]) close in time to the
     # observation and in location to the observatory. If
     # the files are not on disk and there is no internet
     # connection, the average profile is taken from
     # share/molecfit/data/profiles/lib corresponding to
    # the month of the observation (GDAS_t0_s1.fits for
    # Dec/Jan, GDAS_t0_s2.fits for Feb/Mar, etc)
420
     # See Sec. 8.1.4 of the molecfit manual for more info.
     # - 'null', use the profile in the SOF with tag GDAS.
     # If there is no profile in the SOF, the behaviour
     # is the same as GDAS_PROF=auto.
     # - 'none', use the average profile taken from
     # share/molecfit/data/profiles/lib corresponding to
     # the month of observation (see 'auto' description).
     # - 'directory/file.fits', use the specified path and
     # filename as the GDAS profile. Either an absolute path
     # (starting with '/') or a relative path may be used,
430
     # however a relative path is preferred, since only
     # the first 40 char of the path and filename are
     # copied to the FITS header.
     GDAS_PROFILE=auto
     # --LAYERS
     # Grid of layer heights for merging ref_atm and GDAS profile.
     # Fixed grid = CPL_TRUE and natural grid = CPL_FALSE.
    LAYERS=TRUE
440
     # --EMIX
     # Upper mixing height in km for considering data of a local meteo station.
     # If emix is below geoelev, rhum, pres, and temp are not used
     # for modifying the corresponding profiles.
    EMIX=5.0
     # -- PWV
     # PWV value in mm for the input water vapor profile.
     # The merged profile composed of ref_atm, GDAS, and local meteo data
    |# will be scaled to this value if pwv > 0 (default: -1 -> no scaling).
450
```



```
PWV=-1.0
     # --LNFL_LINE_DB
     # File name of the line list (must be stored in the directory :
     # ({TELLURICCORR_DATA_PATH}/hitran/).
     LNFL_LINE_DB=aer_v_3.8.1.2
     # --LNFL_LINE_DB_FORMAT
     # Format of the line file: gives the length in terms of characters per line.
460
     LNFL_LINE_DB_FORMAT=100.0
     # --LBLRTM_ICNTNM
     # Continua and Rayleigh extinction [0,1,2,3,4,5].
     LBLRTM_ICNTNM=5
     # --LBLRTM_IAERSL
     # Aerosols [0,1].
     LBLRTM_IAERSL=0
     # --LBLRTM_MPTS
470
     # Number of optical depth values.
     LBLRTM_MPTS=5
     # --LBLRTM_NPTS
     # Number of values for each panel.
     LBLRTM_NPTS=5
     # --LBLRTM_V1
     # Beginning wavenumber value for the calculation.
480
     LBLRTM_V1=1.9
     # --LBLRTM_V2
     # Ending wavenumber value for the calculation.
     LBLRTM_V2=2.4
     # --LBLRTM_SAMPLE
     # Number of sample points per mean halfwidth [between 1 to 4, default=4].
     LBLRTM_SAMPLE=4
     # --LBLRTM_ALFALO
490
     # Average collision broadened halfwidth [cm-1/atm].
     LBLRTM_ALFALO=0.0
     # --LBLRTM_AVMASS
     # Average molecular mass [amu] for Doppler halfwidth.
     LBLRTM_AVMASS=0.0
     # --LBLRTM_DPTMIN
     # Minimum molecular optical depth below which lines will be rejected.
     LBLRTM_DPTMIN=0.0002
500
```



# --LBLRTM\_DPTFAC

```
# Factor multiplying molecular continuum optical depth.
     LBLRTM_DPTFAC=0.001
     # --LBLRTM_TBOUND
     # Temperature of boundary [K].
     LBLRTM_TBOUND=0.0
     # --LBLRTM_SREMIS1
510
     # Emissivity coefficient 1.
     LBLRTM_SREMIS1=0.0
     # --LBLRTM_SREMIS2
     # Emissivity coefficient 2.
     LBLRTM_SREMIS2=0.0
     # --LBLRTM_SREMIS3
     # Emissivity coefficient 3.
     LBLRTM_SREMIS3=0.0
520
     # --LBLRTM_SRREFL1
     # Reflectivity coefficient 1.
     LBLRTM_SRREFL1=0.0
     # --LBLRTM_SRREFL2
     # Reflectivity coefficient 2.
     LBLRTM_SRREFL2=0.0
     # --LBLRTM_SRREFL3
530
     # Reflectivity coefficient 3.
     LBLRTM_SRREFL3=0.0
     # --LBLRTM_MODEL
     # Atmospheric profile [0,1,2,3,4,5,6].
     LBLRTM_MODEL=0
     # --LBLRTM_ITYPE
     # Type of path [1,2,3].
     LBLRTM_ITYPE=3
540
     # --LBLRTM_NOZERO
     # Zeroing of small amounts of absorbers [0,1].
     LBLRTM_NOZERO=0
     # --LBLRTM_NOPRNT
     # Do not print output? [0,1].
     LBLRTM_NOPRNT=0
     # --LBLRTM_IPUNCH
550
     # Write out layer data to TAPE7 [0,1].
     LBLRTM_IPUNCH=0
```



```
# --LBLRTM_RE
     # Radius of earth [km].
     LBLRTM_RE=0.0
     # --LBLRTM_HSPACE
     # Altitude definition for space [km].
     LBLRTM_HSPACE=120.0
560
     # --LBLRTM_H2
     # Upper height limit [km].
     LBLRTM_H2=0.0
     # --LBLRTM_RANGE
     # Length of a straight path from H1 to H2 [km].
     LBLRTM_RANGE=0.0
     # --LBLRTM_BETA
570
     # Earth centered angle from H1 to H2 [degrees].
     LBLRTM_BETA=0.0
     # --LBLRTM_LEN
     # Path length [0,1].
     LBLRTM_LEN=0
     # --LBLRTM_HOBS
     # Height of observer.
     LBLRTM_HOBS=0.0
580
     # --LBLRTM_AVTRAT
     # Maximum Voigt width ratio across a layer.
     LBLRTM_AVTRAT=2.0
     # --LBLRTM_TDIFF1
     # Maximum layer temperature difference at ALTD1 [K].
     LBLRTM_TDIFF1=5.0
     # --LBLRTM_TDIFF2
590
     # Maximum layer temperature difference at ALTD2 [K].
     LBLRTM_TDIFF2=8.0
     # --LBLRTM_ALTD1
     # Altitude of TDIFF1 [km].
     LBLRTM_ALTD1=0.0
     # --LBLRTM_ALTD2
     # Altitude of TDIFF2 [km].
    LBLRTM_ALTD2=0.0
600
     # --LBLRTM_DELV
     # Number of wavenumbers [cm-1] per major division.
```



620

```
LBLRTM_DELV=1.0
     # --EXPERT_MODE
     # If set to true, will check if TAG[INIT_FIT_PARAMETERS] points to a fits file
     # with a bintable of parameter values to use as initial values for the
     # fitting process.
     EXPERT_MODE=FALSE
610
     # --CHIP_EXTENSIONS
     # Flag that determines if image extensions are to be treated as independent
     # science data to be fitted for independently or as CHIP specific subranges
     # of a single observation to be fitted for as a single combined spectrum.
     # Value = TRUE implies to treat as CHIPS to be combined. Value = FALSE
     # implies
     # to treat as independent. [FALSE].
     CHIP_EXTENSIONS=FALSE
     #
     # End of file
    6.2.2 molecfit_correct default .rc parameters
    The command
    1> esorex --create-config= Default_correct.rc molecfit_correct
    generates a file containing the following:
     # File: Default_correct.rc
     #
     # Note: This configuration file has been automatically
             generated by the esorex (v3.13.6) program.
     #
     #
     # Date: 04-May-2023 18:27:29
     #
     #
```

# --USE\_ONLY\_INPUT\_PRIMARY\_DATA 10

```
# Value=TRUE implies that only the fits primary contains the input science flux
# data.
# Value=FALSE implies that the fits extensions also contains input science
# flux data.
USE_ONLY_INPUT_PRIMARY_DATA=FALSE
```

# --USE\_DATA\_EXTENSION\_AS\_DFLUX # Only valid if USE\_ONLY\_INPUT\_PRIMARY\_DATA=TRUE. The fits extension index that # contains the

```
# errors of the science flux data (DFLUX). A value of 0 implies that there is
20
```



30

60

```
# no DFLUX.
    USE_DATA_EXTENSION_AS_DFLUX=0
    # --USE_DATA_EXTENSION_AS_MASK
    # Only valid if USE_ONLY_INPUT_PRIMARY_DATA=TRUE. The fits extension index that
    # contains the
    # mask associated with the science flux data. A value of 0 implies that there
    # is no mask data.
    USE_DATA_EXTENSION_AS_MASK=0
    # --SUPPRESS_EXTENSION
    # Suppress arbitrary filename extension : TRUE (apply) or FALSE (don't apply).
    SUPPRESS_EXTENSION=FALSE
    # --MAPPING_CORRECT
    # Mapping 'SCIENCE' - 'TELLURIC_CORR' [string with ext_number comma separated
    # (int)] :
    # If set to NULL, check if the TAG[MAPPING_CORRECT] FITS BINTABLE value is
    # provided.
    # The FITS BINTABLE have to one column [TELLURIC_CORR_EXT].
40
    MAPPING_CORRECT=NULL
    # --WLC_REF
    # Indicates that the reference for the wavelength calibration :
    # - If it is set to 'DATA', is the input data.
    # - If it is set to 'MODEL', is the output model, and TELLURIC_DATA file is
    #required. This feature is currently available for FITS binary table inputs
    #only.
    WLC_REF=DATA
50
    # --CHIP_EXTENSIONS
    # Flag that determines if image extensions are to be treated as independent
    # science data to be fitted for independently or as CHIP specific subranges
    # of a single observation to be fitted for as a single combined spectrum.
    # Value = TRUE implies to treat as CHIPS to be combined. Value = FALSE
    # implies
    # to treat as independent. [FALSE].
    CHIP_EXTENSIONS=FALSE
    #
    # End of file
```



# 6.3 The Parameters

Here we consider the MOLECFIT parameters in more detail.

### 6.3.1 Model Recipe

Table 6.1 describes the individual model recipe parameters, in the order in which they appear in the .rc file. Description of the LBRTM parameters is available in

molecfit-kit-<version>/molecfit\\_third\\_party-<version>lblrtm/docs/html/ lblrtm\_instructions\_frame.html.

| Parameter                   | Description  |
|-----------------------------|--|
| USE_ONLY_INPUT_PRIMARY_DATA | Indicates where the science data are located.                        |
|                             | Type = Boolean; Default = FALSE                                      |
|                             | Value=TRUE: Only the primary data array contains the input sci-      |
|                             | ence flux data.  |
|                             | Value=FALSE: The FITS extensions also contain input science flux.    |
| USE_DATA_EXTENSION_AS_DFLUX | Specifies the FITS extension index that contains the                 |
|                             | 1-sigma error on the science flux (DFLUX).                           |
|                             | Type = Integer; Default = 0  |
|                             | Only valid if USE_ONLY_INPUT_PRIMARY_DATA = TRUE.                    |
|                             | Value = 0: There is no DFLUX available.                              |
| USE_DATA_EXTENSION_AS_MASK  | The FITS extension index that contains the mask associated with      |
|                             | the science flux data.   |
|                             | Type = Integer; Default = 0  |
|                             | Only valid if USE_ONLY_INPUT_PRIMARY_DATA = TRUE.                    |
|                             | Value = 0: There is no mask data.                                    |
| USE_INPUT_KERNEL            | Type = Boolean; Default = TRUE                                       |
|                             | Value = TRUE: use a user-provided kernel library, in which case      |
|                             | the FITS file with the kernel library must be given in the SOF, with |
|                             | the tag KERNEL_LIBRARY or MODEL_KERNEL_LIBRARY                       |
| MODEL_MAPPING_KERNEL        | Type = String with comma-separated integers that represent the       |
|                             | extension numbers; Default = NULL                                    |
|                             | If USE_INPUT_KERNEL = FALSE, this parameter is unused.               |
|                             | If USE_INPUT_KERNEL = TRUE, value is a list of extensions            |
|                             | to map from the kernel library file (MODEL_KERNEL_LIBRARY or         |
|                             | KERNEL_LIBRARY) to the extensions of the input file (STD_MODEL       |
|                             | or SCIENCE).   |

#### Table 6.1: Model recipe parameters.



| Parameter | Description   |
|-----------|---|
|           | e.g. Assuming an input science file has extensions "1,2,3,4", then    |
|           | Value="2,2,2,2" means that extension 2 of the kernel library is used  |
|           | for extensions 1, 2, 3 and 4 of the input spectrum. i.e. mapping 2    |
|           | to 1, 2 to 2, 2 to 3, and 2 to 4.                                     |
|           | Similarly, Value="1,2,3,4" would map 1 to 1, 2 to 2, 3 to 3, and 4 to |
|           | 4.  |
|           |   |
|           | Description continues on the next page                                |



| continuation of Tab | le <mark>6.1</mark> |
|---------------------|---------------------|
|---------------------|---------------------|

| Parameter    | Description  |
|--------------|--|
|              | If Value = NULL check if the TAG [MODEL_MAPPING_KERNEL] FITS   |
|              | BINTABLE values are provided.  |
|              | The FITS BINTABLE must have one column   |
|              | [KERNEL_LIBRARY_EXT].  |
|              | See Appendix A.1 for more details.   |
| LIST_MOLEC   | List of molecules to be included in the model.   |
|              | Type = (N_val : nmolec). Comma-separated string of molecule  |
|              | names; Default = NULL.   |
|              | e.g. Value="H20,CO2,O3"  |
|              | If Value = NULL, the input TAG [MOLECULES] FITS BINTABLE must  |
|              | be provided and must contain the three columns: [LIST_MOLEC,   |
|              | FIT_MOLEC, REL_COL].   |
| FIT_MOLEC    | List of flags specifying which of the molecules in LIST_MOLEC to fit.  |
|              | Type = (N_val: nmolec). String containing comma-separated flags  |
|              | (1=fit; 0=do not fit) in the same order as the molecules listed in   |
|              | LIST_MOLEC; Default = NULL   |
|              | e.g. if LIST_MOLEC="H20,CO2,O3", then FIT_MOLEC="1,0,1" im-  |
|              | plies that only $H_2O$ and $O_3$ should be fitted.   |
|              | If Value = NULL, the input TAG [MOLECULES] FITS BINTABLE must  |
|              | be provided and must contain the three columns [LIST_MOLEC,  |
|              | FIT_MOLEC, REL_COL].   |
| REL_COL      | List of the initial values of fitting of the molecular column densities  |
|              | expressed relative to the input ATM profile column densities.  |
|              | Type = String containing a list of comma-separated doubles, in the   |
|              | same order as the listed molecules; Default = NULL.  |
|              | e.g. if LIST_MOLEC = "H20,CO2,O3", then REL_COL = "1.0,1.2,0.8"  |
|              | implies that $H_2O$ , $CO_2$ and $O_3$ have initial relative values of 1.0, 1.2,   |
|              | and 0.8 respectively.  |
|              | If Value = NULL, the input TAG[MOLECULES] FITS   |
|              | BINTABLE must be provided and must have three columns  |
|              | [LIST_MOLEC,FIT_MOLEC,REL_COL].  |
| WAVE_INCLUDE | Wavelength ranges to be included.  |
|              | Type = String with comma separated wavelengths (in microns) in   |
|              | pairs of Double; Default = NULL.   |
|              | e.g. WAVE_INCLUDE="1.773,1.78633,1.79098,1.80434,1.187691,   |
|              | 1.189937" specifies the ranges   |
|              | [1.773,1.78633], [1.79098,1.80434], and [1.187691,1.189937].<br>If set to NULL, check if the TAG[WAVE_INCLUDE] FITS BINTABLE |
|              | values are provided. The FITS BINTABLE must have two columns   |
|              | ·  |
|              | [LOWER_LIMIT, UPPER_LIMIT].  |



| Parameter              | Description   |
|------------------------|---|
| WAVE_EXCLUDE           | Wavelength ranges excluded.   |
|                        | Type = String containing a list of comma-separated pairs of wave-           |
|                        | lengths (Doubles in microns); Default = NULL                                |
|                        | e.g. WAVE_EXCLUDE="1.773,1.78633,1.79098,1.80434,1.187691,                  |
|                        | 1.189937" specifies the ranges  |
|                        | [1.773,1.78633], [1.79098,1.80434], [1.187691,1.189937].                    |
|                        | If Value = NULL, check if the TAG[WAVE_EXCLUDE] FITS BINTABLE               |
|                        | values are provided. The FITS BINTABLE must have two columns                |
|                        | [LOWER_LIMIT, UPPER_LIMIT].   |
| PIXEL_EXCLUDE          | Pixel ranges to be excluded.  |
|                        | Type = String containing comma-separated Integer pairs of pixel             |
|                        | numbers; Default = NULL   |
|                        | e.g. PIXEL_EXCLUDE="54,128,512,514,1020,1024"                               |
|                        | specifies the ranges [54,128], [512,514], [1020,1024].                      |
|                        | If Value = NULL, check if the TAG[PIXEL_EXCLUDE] FITS                       |
|                        | BINTABLE values are provided. The FITS BINTABLE must have                   |
|                        | two columns [LOWER_LIMIT, UPPER_LIMIT].                                     |
| TELLURICCORR_PATH      | Installation directory.   |
|                        | Default = TELLURICCORR_PARAMETER_DEFAULT                                    |
| TELLURICCORR_DATA_PATH | Data directory.   |
|                        | Default = TELLURICCORR_PARAMETER_DEFAULT                                    |
| TMP_PATH               | Temporary directory.  |
|                        | Default = TELLURICCORR_PARAMETER_DEFAULT                                    |
| SILENT_EXTERNAL_BINS   | Supress the output of external binaries.                                    |
|                        | Default = TRUE  |
| TRANSMISSION           | Type of input spectrum.   |
|                        | Type = Boolean; Default = TRUE  |
|                        | Value = TRUE: Transmission spectrum;  |
|                        | Value = FALSE: Emission (radiance) spectrum.                                |
| COLUMN_LAMBDA          | Wavelength column.  |
|                        | Default = lambda  |
|                        | Value = 'NULL' is used if the file is an image and the data are in the      |
|                        | primary data array, in which case the wavelength scale is calculated        |
|                        | from the FITS header keywords CRVAL1= <initial wavelength=""> and</initial> |
|                        | CD1_1= <wavelength step="">.</wavelength>                                   |
|                        | Mandatory if the parameter  |
|                        | USE_ONLY_INPUT_PRIMARY_DATA = FALSE.  |
| COLUMN_FLUX            | Flux column.  |
|                        | Default = flux  |
|                        | Set Value = NULL if the file is an image and the data are in the            |
|                        | primary data array.   |
|                        | Mandatory if USE_ONLY_INPUT_PRIMARY_DATA = FALSE.                           |



| continuation of Tab | le <mark>6.1</mark> |
|---------------------|---------------------|
|---------------------|---------------------|

| Parameter        | Description   |
|------------------|---|
| COLUMN_DFLUX     | 1-sigma error on the flux.  |
|                  | Optional parameter. Default = NULL                                      |
|                  | Set Value = NULL if the file is an image.                               |
| COLUMN_MASK      | Mask column; indicates if a pixel is invalid.                           |
|                  | Optional. Default = NULL  |
|                  | Set Value = NULL if the file is an image                                |
| DEFAULT_ERROR    | Default error relative to the mean. Only used if the error column is    |
|                  | not provided.   |
|                  | Default = 0.01  |
| WLG_TO_MICRON    | Multiplicative factor applied to the wavelength to express it in mi-    |
|                  | crometres.  |
|                  | e.g. if the wavelength is given in nm, set Value = 0.001.               |
|                  | Default = 1.0   |
| WAVELENGTH_FRAME | Wavelength reference frame.   |
|                  | Default = VAC   |
|                  | Value = 'VAC': Wavelength in vacuum;                                    |
|                  | Value = 'AIR': Wavelength in air with the observatory reference         |
|                  | frame;  |
|                  | Value = 'AIR_RV': Wavelength in air with the another reference          |
|                  | frame;  |
|                  | Value = 'VAC_RV': Wavelength in vacuum with another reference           |
|                  | frame, typically the sun or the barycenter of the solar system.         |
|                  | In the latter case, the radial velocity of the observatory relative to  |
|                  | the external reference frame must be provided in the parameter          |
|                  | OBS_ERF_RV_VALUE or provided by the keyword given as a param-           |
|                  | eter to OBS_ERF_RV_KEY.   |
| OBS_ERF_RV_KEY   | The radial velocity of the observatory in km/s relative to the external |
|                  | reference frame. It is positive if the distance between the science     |
|                  | target and the Earth increases along the line-of-sight to the science   |
|                  | target.   |
|                  | This parameter must be provided if                                      |
|                  | WAVELENGTH_FRAME = 'VAC_RV' or 'AIR_RV'.                                |
|                  | Default = NONE  |
| OBS_ERF_RV_VALUE | If OBS_ERF_RV_KEY=='NONE' use this value.                               |
|                  | This parameter must be provided if                                      |
|                  | WAVELENGTH_FRAME = 'VAC_RV' or 'AIR_RV'.                                |
|                  | Default = 0.0   |
| CLEAN_MODEL_FLUX | Set model flux to 0 for non-fitted pixels.                              |
|                  | Default = FALSE   |
| FTOL             | Relative chi-square convergence criterion.                              |
|                  | Default = 1e-10   |
| XTOL             | Relative parameter convergence criterion.                               |
|                  | Default = 1e-10   |



| continuation | of Table | 6.1      |
|--------------|----------|----------|
| 001101100011 | 01 10010 | <b>.</b> |

| Parameter                  | Description  |
|----------------------------|--|
| FLUX_UNIT                  | Conversion of fluxes from $phot/(s * m^2 * mum * as^2)$ (emission        |
|                            | spectrum only) to flux unit of observed spectrum:                        |
|                            | Value = 0: $phot/(s * m^2 * mum * as^2)$ [no conversion]                 |
|                            | Value = 1: $W/(m^2 * mum * as^2)$  |
|                            | Value = 2: $erg/(s * cm^2 * A * as^2)$                                   |
|                            | Value = 3: mJy / ( $as^2$ )  |
|                            | For other units, the conversion factor has to be considered as a         |
|                            | constant term of the continuum fit.                                      |
|                            | Default = 0  |
| FIT_TELESCOPE_BACKGROUND   | Specifies whether to include fitting for the telescope background.       |
|                            | Type = Boolean; Default = TRUE.  |
|                            | Value = TRUE: yes;   |
|                            | Value = FALSE: no (fit emission spectrum only).                          |
| TELESCOPE_BACKGROUND_CONST | Initial value for telescope background constant (relevant for emis-      |
|                            | sion spectrum only).   |
|                            | Default = 0.1  |
| FIT_CONTINUUM              | Flag to enable/disable the polynomial fit of the continuum.              |
|                            | Type: String with comma-separated integers of value 0 or 1, with         |
|                            | each number corresponding to a range.                                    |
|                            | Default = "1" for each range - e.g. if there are four ranges, the de-    |
|                            | fault value is FIT_CONTINUUM="1,1,1,1"                                   |
|                            | e.g. If there are 4 ranges, FIT_CONTINUUM="1,1,0,1" requests con-        |
|                            | tinuum to be fitted for ranges 1, 2, and 4, with no continuum fitting    |
|                            | for range 3.   |
|                            | If FIT_CONTINUUM="NULL", the values are read from the                    |
|                            | TAG[WAVE_INCLUDE] FITS BINTABLE from a column named                      |
|                            | CONT_FIT_FLAG.   |
| CONTINUUM_N                | Degree of the polynomial continuum fit to use per range.                 |
|                            | Type = String of comma-separated integers, each giving the order         |
|                            | of the polynomial to be used in the fit (one per range). Default = "1"   |
|                            | e.g. If there are 4 ranges to fit, CONTINUUM_N="3,3,3,3," means that     |
|                            | all four ranges will use an order 3 polynomial to fit the continuum.     |
|                            | If CONTINUUM_N="NULL", the values are instead read from the              |
|                            | TAG[WAVE_INCLUDE] FITS BINTABLE from a column named                      |
|                            | CONT_POLY_ORDER.   |
| CONTINUUM_CONST            | Initial constant term for continuum fit (valid for all fit ranges). Must |
|                            | be set to the same order of magnitude as the flux.                       |
|                            | Default = 1.0  |



| Parameter       | Description   |
|-----------------|---|
| FIT_WLC         | Defines whether a region is used for the refinement of the wave-        |
|                 | length calibration solution.  |
|                 | Type = String containing a comma-separated list of integers of          |
|                 | value 0 or 1.   |
|                 | Default = "1" for each range. If, for example, there are four ranges,   |
|                 | the default value is FIT_WLC="1,1,1,1"                                  |
|                 | Value = 1: the region is used for the refinement;                       |
|                 | Value = 0: the region is not used.                                      |
|                 | If FIT_WLC="NULL", the values are read from the                         |
|                 | TAG[WAVE_INCLUDE] FITS BINTABLE from a column named                     |
|                 | WLC_FIT_FLAG.   |
|                 | Note that we do not explicitly specify that a chip is to be fitted with |
|                 | this parameter, but rather whether the ranges on the chip are to be     |
|                 | fitted.   |
| WLC_N           | Polynomial degree of the refined wavelength solution.                   |
|                 | Default = 1   |
| WLC_CONST       | Initial constant term for wavelength adjustment - i.e. shift relative   |
|                 | to half wavelength range.   |
|                 | Default = 0.0   |
| FIT_RES_BOX     | Fit the width of a Boxcar LSF.  |
|                 | Default = TRUE  |
| RES_BOX         | Initial value for FWHM of Boxcar relative to slit width at the centre   |
|                 | of the spectrum.  |
|                 | Default = 1.0   |
| FIT_RES_GAUSS   | Fit the FWHM of a Gaussian LSF.   |
|                 | Default = TRUE  |
| RES_GAUSS       | Initial value for FWHM of the Gaussian in pixels, at the centre of      |
|                 | the spectrum.   |
|                 | Default = 1.0   |
| FIT_RES_LORENTZ | Fit the FWHM of a Lorentzian LSF.                                       |
|                 | Default = TRUE  |
| RES_LORENTZ     | Initial value for FWHM of the Lorentz in pixels at the centre of the    |
| -               | spectrum.   |
|                 | Default = 1.0   |
| KERNMODE        | Use a Voigtian profile approximation instead of independent Gaus-       |
|                 | sian and Lorentzian.  |
|                 | Default = FALSE   |
| KERNFAC         | Size of Voigtian / Gaussian / Lorentzian kernel in FWHM.                |
|                 | Default = 3.0   |
| VARKERN         | Does the kernel size increase linearly with wavelength?                 |
|                 | Default = FALSE   |
|                 |   |



| Parameter                  | Description   |
|----------------------------|---|
| OBSERVING_DATE_KEYWORD     | Observing date in years or MJD in days (not string).          |
|                            | Default = MJD-OBS   |
|                            | If Value == NONE, OBSERVING_DATE_VALUE will be used instead.  |
| OBSERVING_DATE_VALUE       | If OBSERVING_DATE_KEYWORD=='NONE' use this value.             |
|                            | Default = -1.0  |
| UTC_KEYWORD                | UTC in seconds.   |
|                            | Default = UTC   |
|                            | If Value == NONE, UTC_VALUE will be used.                     |
| UTC_VALUE                  | If UTC_KEYWORD=='NONE' use this value.                        |
|                            | Default = -1.0  |
| TELESCOPE_ANGLE_KEYWORD    | Telescope altitude angle in degrees.                          |
|                            | Default = ESO TEL ALT   |
| TELESCOPE_ANGLE_VALUE      | If TELESCOPE_ANGLE_KEYWORD=='NONE' take this value.           |
|                            | Default = 90.0  |
| RELATIVE_HUMIDITY_KEYWORD  | Relative humidity in %.                                       |
|                            | Default = ESO TEL AMBI RHUM                                   |
| RELATIVE_HUMIDITY_VALUE    | If RELATIVE_HUMIDITY_KEYWORD=='NONE' take this value.         |
|                            | Default = 15.0  |
| PRESSURE_KEYWORD           | Pressure in hPa.  |
|                            | Default = ESO TEL AMBI PRES START                             |
| PRESSURE_VALUE             | If PRESSURE_KEYWORD=='NONE' take this value.                  |
|                            | Default = 750.0   |
| TEMPERATURE_KEYWORD        | Ambient temperature in deg C.                                 |
|                            | Default = ESO TEL AMBI TEMP                                   |
| TEMPERATURE_VALUE          | If TEMPERATURE_KEYWORD=='NONE' take this value.               |
|                            | Default = 15.0  |
| MIRROR_TEMPERATURE_KEYWORD | Mirror temperature. (Relevant for emission spectrum only)     |
|                            | Default = ESO TEL TH M1 TEMP                                  |
| MIRROR_TEMPERATURE_VALUE   | If MIRROR_TEMPERATURE_KEYWORD=='NONE' take this value,        |
|                            | specified in degrees C.                                       |
|                            | Default = 15.0  |
| ELEVATION_KEYWORD          | Elevation above sea level in m (default is Paranal: 2635. m). |
|                            | Default = ESO TEL GEOELEV                                     |
| ELEVATION_VALUE            | If ELEVATION_KEYWORD=='NONE' take this value.                 |
|                            | Default = 2635.0  |
| LONGITUDE_KEYWORD          | Longitude (default is Paranal: -70.4051 deg).                 |
|                            | Default = ESO TEL GEOLON                                      |
| LONGITUDE_VALUE            | If LONGITUDE_KEYWORD=='NONE' take this value.                 |
| LONGTIONS_AUTOR            | Default = $-70.4051$  |
| LATITUDE_KEYWORD           | Latitude (default is Paranal: -24.6276 deg).                  |
|                            | Default = ESO TEL GEOLAT                                      |
|                            |   |



|  | continuation | of Table | 6.1 |
|--|--------------|----------|-----|
|--|--------------|----------|-----|

| Parameter             | Description   |
|-----------------------|---|
| LATITUDE_VALUE        | If LATITUDE_KEYWORD=='NONE' take this value.                          |
|                       | Default = -24.6276  |
| SLIT_WIDTH_KEYWORD    | Slit width in arcsec (taken from FITS header if present).             |
|                       | Default = ESO INS SLIT1 WID   |
| SLIT_WIDTH_VALUE      | If SLIT_WIDTH_KEYWORD=='NONE' take this value.                        |
|                       | Default = 0.4   |
| PIX_SCALE_KEYWORD     | Pixel scale in arcsec (taken from this file only).                    |
|                       | Default = NONE  |
| PIX_SCALE_VALUE       | If PIX_SCALE_KEYWORD=='NONE' take this value.                         |
|                       | Default = 0.086   |
| REFERENCE_ATMOSPHERIC | Reference atmospheric profile. Possible values: - equ.fits (default;  |
|                       | equatorial atmosphere, valid for Paranal); - tro.fits (tropical atmo- |
|                       | sphere); - std.fits (standard atmosphere); - Other file located in    |
|                       | (TELLURICCORR_DATA_PATH/profiles/mipas/).                             |
|                       | Default = equ.fits  |
| GDAS_PROFILE          | Specify which GDAS profile to use. If GDAS_PROFILE=='auto',           |
|                       | automatic retrieval of the GDAS profiles (P[hPa] HGT[km] T[K]         |
|                       | RELHUM[%]) close in time to the observation and in location           |
|                       | to the observatory. If the files are not on disk and there            |
|                       | is no internet connection, the average profile is taken from          |
|                       | share/molecfit/data/profiles/lib corresponding to the month of the    |
|                       | observation (GDAS_t0_s1.fits for Dec/Jan, GDAS_t0_s2.fits for         |
|                       | Feb/Mar, etc) See Sec. 8.1.4 of the molecfit manual for more          |
|                       | info. If GDAS_PROFILE=='null', use the profile in the SOF with        |
|                       | tag GDAS. If there is no profile in the SOF, the behaviour is the     |
|                       | same as GDAS_PROF=auto. If GDAS_PROFILE=='none', use the              |
|                       | average profile taken from share/molecfit/data/profiles/lib corre-    |
|                       | sponding to the month of observation (see 'auto' description). If     |
|                       | GDAS_PROFILE=='directory/file.fits', use the specified path and file- |
|                       | name as the GDAS profile. Either an absolute path (starting with '/') |
|                       | or a relative path may be used, however a relative path is preferred, |
|                       | since only the first 40 char of the path and filename are copied to   |
|                       | the FITS header. The file format must be a FITS binary table with     |
|                       | columns having the names 'press height temp relhum' and units         |
|                       | hPa, km, K and %, respectively.                                       |
|                       | Default = auto  |
| LAYERS                | Grid of layer heights for merging ref_atm and GDAS profile. Fixed     |
|                       | grid = CPL_TRUE and natural grid = CPL_FALSE.                         |
|                       | Default = TRUE  |
| EMIX                  | Upper mixing height in km for considering data of a local meteo       |
|                       | station. If emix is below geoelev, rhum, pres, and temp are not       |
|                       | used for modifying the corresponding profiles.                        |
|                       | Default = 5.0   |



| continuation of Table | 6.1      |
|-----------------------|----------|
| oonanaaalon on habio  | <b>U</b> |

| Parameter           | Description  |
|---------------------|--|
| PWV                 | PWV value in mm for the input water vapor profile. The merged        |
|                     | profile composed of ref_atm, GDAS, and local meteo data will be      |
|                     | scaled to this value if pwv > 0 (default: -1 -> no scaling).         |
|                     | Default = -1.0   |
| LNFL_LINE_DB        | File name of the line list (must be stored in the directory :(TEL-   |
|                     | LURICCORR_DATA_PATH/hitran/).  |
|                     | $Default = aer_v_3.8.1.2$  |
| LNFL_LINE_DB_FORMAT | Format of the line file: gives the length in terms of characters per |
|                     | line.  |
|                     | Default = 100.0  |
| LBLRTM_ICNTNM       | Continua and Rayleigh extinction [0,1,2,3,4,5].                      |
|                     | Default = 5  |
| LBLRTM_IAERSL       | Aerosols [0,1].  |
|                     | Default = 0  |
| LBLRTM_MPTS         | Number of optical depth values.                                      |
|                     | Default = 5  |
| LBLRTM_NPTS         | Number of values for each panel.                                     |
|                     | Default = 5  |
| LBLRTM_V1           | Beginning wavenumber value for the calculation.                      |
|                     | Default = 1.9  |
| LBLRTM_V2           | Ending wavenumber value for the calculation.                         |
|                     | Default = 2.4  |
| LBLRTM_SAMPLE       | Number of sample points per mean halfwidth [between 1 to 4, de-      |
|                     | fault=4].  |
|                     | Default = 4  |
| LBLRTM_ALFALO       | Average collision broadened halfwidth [cm-1/atm].                    |
|                     | Default = 0.0  |
| LBLRTM_AVMASS       | Average molecular mass [amu] for Doppler halfwidth.                  |
|                     | Default = 0.0  |
| LBLRTM_DPTMIN       | Minimum molecular optical depth below which lines will be rejected.  |
|                     | Default = 0.0002   |
| LBLRTM_DPTFAC       | Factor multiplying molecular continuum optical depth.                |
|                     | Default = 0.001  |
| LBLRTM_TBOUND       | Temperature of boundary [K].   |
|                     | Default = 0.0  |
| LBLRTM_SREMIS1      | Emissivity coefficient 1.  |
|                     | Default = 0.0  |
| LBLRTM_SREMIS2      | Emissivity coefficient 2.  |
|                     | Default = 0.0  |
| LBLRTM_SREMIS3      | Emissivity coefficient 3.  |
|                     | Default = 0.0  |



| Parameter      | Description                                   |
|----------------|---|
| LBLRTM_SRREFL1 | Reflectivity coefficient 1.                   |
|                | [0.0  |
| LBLRTM_SRREFL2 | Reflectivity coefficient 2.                   |
|                | Default = 0.0                                 |
| LBLRTM_SRREFL3 | Reflectivity coefficient 3.                   |
|                | Default = 0.0                                 |
| LBLRTM_MODEL   | Atmospheric profile [0,1,2,3,4,5,6].          |
|                | Default = 0                                   |
| LBLRTM_ITYPE   | Type of path [1,2,3].                         |
|                | Default = 3                                   |
| LBLRTM_NOZERO  | Zeroing of small amounts of absorbers [0,1].  |
|                | Default = 0                                   |
| LBLRTM_NOPRNT  | Do not print output? [0,1].                   |
|                | Default = 0                                   |
| LBLRTM_IPUNCH  | Write out layer data to TAPE7 [0,1].          |
|                | Default = 0                                   |
| LBLRTM_RE      | Radius of earth [km].                         |
|                | Default = 0.0                                 |
| LBLRTM_HSPACE  | Altitude definition for space [km].           |
|                | Default = 120.0                               |
| LBLRTM_H2      | Upper height limit [km].                      |
|                | Default = 0.0                                 |
| LBLRTM_RANGE   | Length of a straight path from H1 to H2 [km]. |
|                | Default = 0.0                                 |
| LBLRTM_BETA    | Earth centered angle from H1 to H2 [degrees]. |
|                | Default = 0.0                                 |
| LBLRTM_LEN     | Path length [0,1].                            |
|                | Default = 0                                   |



| Parameter           | Description   |
|---------------------|---|
| LBLRTM_HOBS         | Height of observer.   |
|                     | Default = 0.0   |
| LBLRTM_AVTRAT       | Maximum Voigt width ratio across a layer.                             |
|                     | Default = 2.0   |
| LBLRTM_TDIFF1       | Maximum layer temperature difference at ALTD1 [K].                    |
|                     | Default = 5.0   |
| LBLRTM_TDIFF2       | Maximum layer temperature difference at ALTD2 [K].                    |
|                     | Default = 8.0   |
| LBLRTM_ALTD1        | Altitude of TDIFF1 [km].  |
|                     | Default = 0.0   |
| LBLRTM_ALTD2        | Altitude of TDIFF2 [km].  |
|                     | Default = 0.0   |
| LBLRTM_DELV         | Number of wavenumbers [cm-1] per major division.                      |
|                     | Default = 1.0   |
| CHIP_EXTENSIONS     | If TRUE,  |
|                     | treat image extensions as a single observation to be fitted for as a  |
|                     | single combined spectrum.   |
|                     | If FALSE,   |
|                     | treat image extensions as independent science data to be fitted for   |
|                     | independently.  |
|                     | Default = FALSE   |
| MAP_REGIONS_TO_CHIP | Where there are multiple ranges and multiple chips, with the possi-   |
|                     | bility of wavelength overlaps, there's ambiguity as to which region   |
|                     | is assigned to which chip.  |
|                     | This parameter allows the user to explicitly specify this association |
|                     | as a string with a comma separated list of integers, where each       |
|                     | integer is a chip index.  |
|                     | The number of integers in the list can be either one, which then      |
|                     | applies to all ranges, or a value for each range can be individually  |
|                     | specified.  |
|                     | e.g. "1" and "1,1,1,1" are equivalent if there are 4 ranges.          |
|                     | If the value is "NULL", then it is assumed the values are read from   |
|                     | the TAG[WAVE_INCLUDE] FITS BINTABLE from a column named               |
|                     | MAPPED_TO_CHIP.   |
|                     | For further details see Appendix A2 of the molecfit user manual.      |
|                     | The default is for the ranges to be mapped to CHIP1, i.e.             |
|                     | Default = "1"   |



| Parameter   | Description   |
|-------------|---|
| EXPERT_MODE | If TRUE,  |
|             | read in starting values from the TAG[INIT_FIT_PARAMETERS] |
|             | FITS BINTABLE that is a similar format to the             |
|             | BEST_FIT_PARAMETERS.fits output file. See Appendix A5     |
|             | of the molecfit user manual for more details.             |
|             | If FALSE,   |
|             | Do not read in the starting values.                       |
|             | Default = FALSE   |

#### 6.3.2 Calctrans Recipe

Table 6.2 explains the individual calctrans recipe parameters in more detail and in the order they appear in the .rc file.

| Parameter                   | Description  |
|-----------------------------|--|
| USE_ONLY_INPUT_PRIMARY_DATA | Value=TRUE implies that only the primary data array contains the input science flux data.  |
|                             | Value=FALSE implies that the FITS extensions also contains input<br>science flux.<br>Default = FALSE   |
| USE_DATA_EXTENSION_AS_DFLUX | Only valid if USE_ONLY_INPUT_PRIMARY_DATA=TRUE.<br>This parameter specifies the FITS extension index that contains the<br>1-sigma error on the science flux (DFLUX).<br>A value of 0 implies that there is no DFLUX available.<br>Default = 0. |



| Parameter                  | Description   |
|----------------------------|---|
| USE_DATA_EXTENSION_AS_MASK | Only valid if USE_ONLY_INPUT_PRIMARY_DATA=TRUE.   |
|                            | This parameter specifies the FITS extension index that contains the                                 |
|                            | mask associated with the science flux data. A value of 0 implies                                    |
|                            | that there is no mask data.   |
|                            | Default = TRUE  |
| USE_INPUT_KERNEL           | If TRUE, use the kernel library if it is provided.  |
|                            | Default = TRUE  |
| CALCTRANS_MAPPING_KERNEL   | If USE_INPUT_KERNEL is TRUE,  |
|                            | List of extensions to map from the kernel library file (CALCTRANS_KERNEL_LIBRARY or KERNEL_LIBRARY) |
|                            | to the extensions of the input file (SCIENCE_CALCTRANS or SCI-<br>ENCE).                            |
|                            | [string with comma separated integers that represent the extension                                  |
|                            | numbers]  |
|                            | e.g. Assuming an input science file has extensions "1,2,3,4", then                                  |
|                            | CALCTRANS_MAPPING_KERNEL="2,2,2,2" implies a mapping exten-   |
|                            | sion 2 of the kernel library  |
|                            | for all the extensions in the input science file,   |
|                            | i.e. mapping 2 to 1, 2 to 2, 2 to 3, 2 to 4.  |
|                            | means that extension 2 of the kernel library is used for extensions 1, 2, 3 and 4.                  |
|                            | CALCTRANS_MAPPING_KERNEL="1,2,3,4" similarly maps 1 to 1, 2 to                                      |
|                            | 2, 3 to 3, 4 to 4.  |
|                            | If USE_INPUT_KERNEL is FALSE,   |
|                            | this parameter is unused  |
|                            | If set to NULL, check if the TAG[CALCTRANS_MAPPING_KERNEL]  |
|                            | FITS BINTABLE   |
|                            | values are provided.  |
|                            | The FITS BINTABLE must have one column [KER-  |
|                            | NEL_LIBRARY_EXT].   |
|                            | Default = NULL  |



| Parameter           | Description   |
|---------------------|---|
| MAPPING_ATMOSPHERIC | List of extensions to map from the associated file (ATM_PARAMETERS)   |
|                     | to the extensions of the input file (SCIENCE_CALCTRANS or SCI-  |
|                     | ENCE).  |
|                     | [string with comma separated integers that represent the extension numbers]                                       |
|                     | e.g. Assuming an input science file has extensions "1,2,3,4", then  |
|                     | MAPPING_ATMOSPHERIC="2,2,2,2" implies a mapping extension 2 of the associated file                                |
|                     | for all the extensions in the input science file,   |
|                     | i.e. mapping 2 to 1, 2 to 2, 2 to 3, 2 to 4.  |
|                     | MAPPING_ATMOSPHERIC="1,2,3,4" similarly maps 1 to 1, 2 to 2, 3  |
|                     | to 3, 4 to 4.   |
|                     | If set to NULL, check if the TAG[MAPPING_ATMOSPHERIC] FITS<br>BINTABLE value                                      |
|                     | is provided.  |
|                     | The FITS BINTABLE must have one column  |
|                     | [ATM_PARAMETERS_EXT].   |
|                     | Default = NULL  |
| MAPPING_CONVOLVE    | List of extensions to map from the associated file  |
|                     | (LBLRTM_RESULTS) to the extensions of the output (TELLURIC_CORR).   |
|                     | [string with comma separated integers that represent the extension  |
|                     | numbers]  |
|                     | e.g. Assuming an input science file has extensions "1,2,3,4", then  |
|                     | MAPPING_CONVOLVE="2,2,2,2" implies a mapping extension 2 of   |
|                     | the associated file   |
|                     | for all the extensions in the input science file,   |
|                     | i.e. mapping 2 to 1, 2 to 2, 2 to 3, 2 to 4.<br>MAPPING_CONVOLVE="1,2,3,4" similarly maps 1 to 1, 2 to 2, 3 to 3, |
|                     | 4 to 4.   |
|                     | If set to NULL, check if the TAG[MAPPING_CONVOLVE] FITS   |
|                     | BINTABLE value is   |
|                     | provided.   |
|                     | The FITS BINTABLE must have one column  |
|                     | [LBLRTM_RESULTS_EXT].   |
|                     | Default = NULL  |



| continuation of | f Table 6.2 |
|-----------------|-------------|
| oontinuation o  |             |

| Parameter       | Description  |
|-----------------|--|
| CHIP_EXTENSIONS | If TRUE, treat image extensions as a single observation to be fitted |
|                 | for as a single combined spectrum.                                   |
|                 | If FALSE, treat image extensions as independent science data to      |
|                 | be fitted for independently.   |
|                 | Default = FALSE  |
| SCALE_PWV       | Value read from science file to scale the precipitable water vapor   |
|                 | (PWV) of the telluric spectra. If SCALE_PWV='auto', the mean value   |
|                 | of TEL AMBI IWV START/END is read from the header of the input       |
|                 | science file. If SCALE_PWV= a user-provided numerical value, use     |
|                 | this as the scale factor. If SCALE_PWV= a FITS header keyword        |
|                 | name, read in the scale factor from the header of the input science  |
|                 | file using this keyword. If SCALE_PWV='none', perform no scaling.    |
|                 | Default = none. See Sec. A.7 for more details.                       |
| HDR_MJD         | FITS header keyword to read the Modified Julian Date at the          |
|                 | START of the exposure from telluric and science header. The          |
|                 | telluric header was copied to BEST_FIT_PARAMETERS by molec-          |
|                 | fit_model and read from there.                                       |
|                 | Default = MJD-OBS  |
| HDR_EXP         | FITS header keyword to read the exposure time from tel-              |
|                 | luric and science header. The telluric header was copied to          |
|                 | BEST_FIT_PARAMETERS by molecfit_model and read from there.           |
|                 | Default = ESO OBS EXECTIME   |
| HDR_AIR1        | FITS header keyword to read the airmass at the start of the ex-      |
|                 | posure from the telluric and science header. The telluric header     |
|                 | was copied to BEST_FIT_PARAMETERS by molecfit_model and read         |
|                 | from there.  |
|                 | Default = ESO TEL AIRM START   |
| HDR_AIR2        | FITS header keyword to read the airmass at the end of the ex-        |
|                 | posure from the telluric and science header. The telluric header     |
|                 | was copied to BEST_FIT_PARAMETERS by molecfit_model and read         |
|                 | from there.  |
|                 | Default = ESO TEL AIRM END   |
| SGWL            | Savitzky-Golay filter smoothing window length in pixels.             |
|                 | Default = 15. See Sec. A.8 for more details.                         |
| SGWL_ASMAX      | Treat the Savitzky-Golay filter smoothing window length as a maxi-   |
|                 | mum length.  |
|                 | Default = FALSE  |

### 6.3.3 Correct Recipe

Table 6.3 explains the individual correct recipe parameters in more detail and in the order they appear in the .rc file.



| Table 6.3: | Correct recipe parameters. |
|------------|----------------------------|
|------------|----------------------------|

| Parameter                   | Description  |
|-----------------------------|--|
| USE_ONLY_INPUT_PRIMARY_DATA | Value=TRUE implies that only the primary data array contains the     |
|                             | input science flux data.   |
|                             | Value=FALSE implies that the FITS extensions also contains input     |
|                             | science flux.  |
|                             | Default = FALSE  |
| USE_DATA_EXTENSION_AS_DFLUX | Only valid if USE_ONLY_INPUT_PRIMARY_DATA=TRUE.                      |
|                             | This parameter specifies the FITS extension index that contains the  |
|                             | 1-sigma error on the science flux (DFLUX). A value of 0 implies that |
|                             | there is no DFLUX.   |
|                             | Default = 0.   |
| USE_DATA_EXTENSION_AS_MASK  | Only valid if USE_ONLY_INPUT_PRIMARY_DATA=TRUE.                      |
|                             | This parameter specifies the FITS extension index that contains the  |
|                             | mask associated with the science flux data. A value of 0 implies     |
|                             | that there is no mask data.  |
|                             | Default = 0.   |
| SUPPRESS_EXTENSION          | Suppress arbitrary filename extension : TRUE (apply) or FALSE        |
|                             | (don't apply).   |
|                             | Default = FALSE  |



| Parameter       | Description  |
|-----------------|--|
| MAPPING_CORRECT | List of extensions to map from the telluric correction file (TEL-<br>LURIC_CORR)   |
|                 | to the extensions of the intput file (SCIENCE_CALCTRANS or SCI-<br>ENCE).  |
|                 | [string with comma separated integers that represent the extension numbers]  |
|                 | e.g. Assuming an input science file has extensions "1,2,3,4", then MAPPING_CORRECT="2,2,2,2" implies a mapping extension 2 of the telluric correction file |
|                 | for all the extensions in the input science file,  |
|                 |  |
|                 | i.e. mapping 2 to 1, 2 to 2, 2 to 3, 2 to 4.<br>means that extension 2 of the telluric correction is used for extensions 1, 2, 3 and 4                     |
|                 | of the input spectrum.   |
|                 | MAPPING_CORRECT="1,2,3,4" similarly maps 1 to 1, 2 to 2, 3 to 3, 4 to 4.   |
|                 | If set to NULL, check if the TAG[MAPPING_CORRECT] FITS   |
|                 | BINTABLE value is  |
|                 | provided.  |
|                 | The FITS BINTABLE must have one column [TEL-LURIC_CORR_EXT].   |
|                 | Default = NULL   |
| WLC_REF         | Indicates that the reference for the wavelength calibration :  |
|                 | - If it is set to "DATA", is the input data.   |
|                 | - If it is set to "MODEL", is the output model.  |
|                 | In the "MODEL" case, the wavelength given in the output spectrum has been corrected with the result of the fit.  |
|                 | If set to 'MODEL', the degree of the polynomial used   |
|                 | for the wavelength correction should be 0 or at most 1,  |
|                 | except if the inclusion regions cover a large  |
|                 | or well-sampled spectral range of the input spectrum.  |
|                 | Default = DATA   |
| CHIP_EXTENSIONS | If TRUE,   |
|                 | treat image extensions as a single observation   |
|                 | to be fitted for as a single combined spectrum.  |
|                 | If FALSE,  |
|                 | treat image extensions as independent science data   |
|                 | to be fitted for independently.  |
|                 | Default = FALSE  |



# 6.4 Input Spectrum

MOLECFIT 4.4.2 accepts the following formats:

- FITS binary table,
- 1D FITS image.

if USE\_ONLY\_INPUT\_PRIMARY\_DATA is TRUE, then the assumption is that the format is a 1D FITS image. Otherwise the input must be a FITS binary table.

The CDELT1, CDELT2 and CDELT3 FITS header keywords are now *deprecated* in favour of the corresponding FITS header keywords CD1\_1, CD2\_2 and CD3\_3. If your input spectrum is missing the latter keywords, a warning will be raised about the input spectrum. This warning can be resolved by adding these keywords to the FITS header.

### 6.5 Output Products

With the exception of the ASCII log file esorex.log, all MOLECFIT output files are in the same format as the input files as described above. A comprehensive list is given in Appendix B.7.



# 7 Guide for MOLECFIT 1.5.9 Users

# 7.1 Execution Format

MOLECFIT 1.5.9 consisted of three main standalone executables:

- molecfit Find best-fit parameters for molecules and continuum
- calctrans Calculate transmission model given best fit parameters
- corrfilelist Perform telluric correction for a set of spectra given the transmission model

These would run with a specified parameter file (.par) that would list all the non-default parameter values to use including the input and output filenames. A common practice would be to include the parameters for all three executables in a single .par file and execute as follows:

> molecfit VISIR\_HR.par

> calctrans VISIR\_HR.par

> corrfilelist VISIR\_HR.par

where input spectra files could be of either FITS or ASCII format.

For MOLECFIT versions 3.x and later, these applications are now represented as *EsoRex* recipes and interface in compliance of *EsoRex* pipeline standards. They are executed from the command line in the following manner:

> esorex --recipe-config=Pars2Use.rc recipe\_name SOF\_filename

Where Pars2Use.rc contains the list of "processing parameters", the SOF (set of files) is an ASCII list of the input and output filenames which are expected to be in FITS format. Note: ASCII data files are not supported.

It should be noted that the MOLECFIT 1.5.9 executables calctrans\_lblrtm and calctrans\_convolution are not represented as *EsoRex* recipes.

# 7.2 Parameter Format

In accordance with *EsoRex* standards, *EsoRex* calls will return an error if the .rc file specified contains a parameter that is not relevant to the recipe in use. Thus, unlike the original MOLECFIT, all the parameters to be used with the three molecfit steps — model, correct, and calctrans cannot be defined in a single parameter file. Instead, each step must be defined using an independent recipe configuration (.rc) file.

A recipe configuration file with all the default values can be generated in the working directory through *EsoRex* using the following command:

```
> esorex --create-config= filename recipe_name
```

For example:



```
> esorex --create-config=Default_model.rc molecfit_model
> esorex --create-config=Default_correct.rc molecfit_correct
> esorex --create-config=Default_calctrans.rc molecfit_calctrans
```

The above example will generate the default .rc files:

Default\_model.rc Default\_correct.rc Default\_calctrans.rc

respectively in the working directory. These .rc files can then be used as templates, thereby easing the burden of writing an .rc file from scratch.

The majority of the parameters declared in these .rc files should be familiar to the 1.5.9 user. In many cases there are some subtle differences in the parameter name and in the value format. For example the logical parameter to specify whether fitting for the continuum is to be performed has been changed from:

FIT\_CONT: 1
to
FIT\_CONTINUUM = TRUE

### 7.3 Input Output Filename Specifications

The names of all input and output files involved in a recipe execution are to be specified in the .sof file given in the command line execution. These are ASCII files that specify one file per line in the form:

filename tagname

where file pathnames may contain enviroment variables, and tagnames are specific keywords that declare the file's role in the process — e.g. input science spectra, input best-fit parameters, etc.

For example an SOF file for a MOLECFIT correct recipe execution may contain:

\$HOME/molecfit/raw/CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits SCIENCE
\$HOME/molecfit\_output/MODEL\_MOLECULES.fits MODEL\_MOLECULES
\$HOME/molecfit\_output/ATM\_PARAMETERS.fits ATM\_PARAMETERS

where

| \$HOME          | environment variable   |
|-----------------|--|
| SCIENCE         | tagname for input file that contains the science spectra to use                      |
| MODEL_MOLECULES | tagname for input file that contains a list of air molecules to include in the model |
| ATM_PARAMETERS  | tagname for the output file that contains the best fit parameters of the model       |

A full list of the SOF tag names are given in appendix A3.4



# 7.4 Plot Files

MOLECFIT 3.x and later versions do not support the 1.5.9 plot file commands, as *Reflex* offers other methods to graph results.

# 7.5 Extra LNFL and LBLRTM Control Parameters

From v3.x MOLECFIT supports extra optional parameters for additional control of the third party lnfnl and lblrtm executions. These are listed in Table 7.1: Description of the LBRTM parameters is available in following location in the kit distribution -

molecfit-kit-<version>/molecfit\_third\_party-<version>lblrtm /docs/html/ lblrtm\_instructions\_frame.html

| Parameter           | Description  |
|---------------------|--|
| LNFL_LINE_DB        | File name of the line list. This must be stored in the directory           |
|                     | TELLURICCORR_DATA_PATH/hitran/.  |
|                     | $Default = aer_v_{3.8.1.2}$  |
| LNFL_LINE_DB_FORMAT | Format of the line file: gives the length in terms of characters per line. |
|                     | Default = 100.0  |
| LBLRTM_ICNTNM       | Continua and Rayleigh extinction   |
|                     | Valid range: [0 – 5]   |
|                     | Default = 5  |
| LBLRTM_IAERSL       | Aerosols   |
|                     | Valid range: [0, 1]  |
|                     | Default = 0  |
| LBLRTM_MPTS         | Number of optical depth values.  |
|                     | Default = 5  |
| LBLRTM_NPTS         | Number of values for each panel.   |
|                     | Default = 5  |
| LBLRTM_V1           | Beginning wavenumber value for the calculation.                            |
|                     | Default = 1.9  |
| LBLRTM_V2           | Ending wavenumber value for the calculation.                               |
|                     | Default = 2.4  |
| LBLRTM_SAMPLE       | Number of sample points per mean halfwidth                                 |
|                     | Valid range: [1 – 4]   |
|                     | Default = 4  |
| LBLRTM_ALFALO       | Average collision broadened halfwidth ( $cm^{-1}/atm$ ).                   |
|                     | Default = 0.0  |
| LBLRTM_AVMASS       | Average molecular mass (amu) for Doppler halfwidth.                        |
|                     | Default = 0.0  |



| Parameter      | Description   |
|----------------|---|
| LBLRTM_DPTMIN  | Minimum molecular optical depth below which lines will be rejected. |
|                | Default = 0.0002  |
| LBLRTM_DPTFAC  | Factor multiplying molecular continuum optical depth.               |
|                | Default = 0.001   |
| LBLRTM_TBOUND  | Temperature of boundary (K).  |
|                | Default = 0.0   |
| LBLRTM_SREMIS1 | Emissivity coefficient 1.   |
|                | Default = 0.0   |
| LBLRTM_SREMIS2 | Emissivity coefficient 2.   |
|                | Default = 0.0   |
| LBLRTM_SREMIS3 | Emissivity coefficient 3.   |
|                | Default = 0.0   |
| LBLRTM_SRREFL1 | Reflectivity coefficient 1.   |
|                | Default = 0.0   |
| LBLRTM_SRREFL2 | Reflectivity coefficient 2.   |
|                | Default = 0.0   |
| LBLRTM_SRREFL3 | Reflectivity coefficient 3.   |
|                | Default = 0.0   |
| LBLRTM_MODEL   | Atmospheric profile   |
|                | Valid range: [0 – 6].   |
|                | Default = 0   |
| LBLRTM_ITYPE   | Type of path  |
|                | Valid range: [1 – 3].   |
|                | Default = 3   |
| LBLRTM_NOZERO  | Zeroing of small amounts of absorbers                               |
|                | Valid range: [0 – 1].   |
|                | Default = 0   |
| LBLRTM_NOPRNT  | Do not print output.  |
|                | Valid range: [0 – 1]  |
|                | Default = 0   |
| LBLRTM_IPUNCH  | Write out layer data to TAPE7                                       |
|                | Valid range: 0 – 1.   |
|                | Default = 0   |
| LBLRTM_RE      | Radius of earth [km].   |
|                | Default = 0.0   |
| LBLRTM_HSPACE  | Altitude definition for space [km].                                 |
|                | Default = 120.0   |
| LBLRTM_H2      | Upper height limit [km].  |
|                | Default = 0.0   |
| LBLRTM_RANGE   | Length of a straight path from H1 to H2 [km].                       |
|                | Default = 0.0   |
| LBLRTM_BETA    | Earth centered angle from H1 to H2 [degrees].                       |



| Parameter | Description   |
|-----------|---------------|
|           | Default = 0.0 |



| Parameter     | Description  |
|---------------|--|
| LBLRTM_LEN    | Path length  |
|               | Valid range = $[0, 1]$ .                           |
|               | Default = 0  |
| LBLRTM_HOBS   | Height of observer.                                |
|               | Default = 0.0                                      |
| LBLRTM_AVTRAT | Maximum Voigt width ratio across a layer.          |
|               | Default = 2.0                                      |
| LBLRTM_TDIFF1 | Maximum layer temperature difference at ALTD1 [K]. |
|               | Default = 5.0                                      |
| LBLRTM_TDIFF2 | Maximum layer temperature difference at ALTD2 [K]. |
|               | Default = 8.0                                      |
| LBLRTM_ALTD1  | Altitude of TDIFF1 [km].                           |
|               | Default = 0.0                                      |
| LBLRTM_ALTD2  | Altitude of TDIFF2 [km].                           |
|               | Default = 0.0                                      |
| LBLRTM_DELV   | Number of wavenumbers [cm-1] per major division.   |
|               | Default = 1.0                                      |

### 7.6 Simple Example

Assume that we want to process a science frame CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits which is located in the working directory and that all desired parameters have been defined in the associated .rc files

Model.rc Calctrans.rc Correct.rc

Step 1: invoke

> esorex --recipe-config=Model.rc molecfit\_model Model.sof

where Model.sof contains a single line specifying the location of the science frame:

./CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits SCIENCE

This will produce several FITS files, including:

| MODEL_MOLECULES.fits | Contains the list of molecules used in the model            |
|----------------------|---|
| ATM_PARAMETERS.fits  | Contains the atmospheric parameter values used in the model |
| BEST_FIT_PARAMETERS  | Contains the best fit values derived for the model          |

Step 2: run

> esorex --recipe-config=Calctrans.rc molecfit\_calctrans Calctrans.sof

where Calctrans.sof contains the location of the science frame and the parameter values as derived from the model recipe execution:



./CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits
./MODEL\_MOLECULES.fits
./ATM\_PARAMETERS.fits
./BEST\_FIT\_PARAMETERS.fits

SCIENCE MODEL\_MOLECULES ATM\_PARAMETERS BEST\_FIT\_PARAMETERS

This will produce several fits files including

TELLURIC\_CORR.fits

which contains the data to perform a convolution that will correct a spectrum of telluric contamination.

Step 3: invoke:

> esorex --recipe-config=Correct.rc molecfit\_correct Correct.sof

where Correct.sof contains the location of the science frame and the convolution telluric correction data derived from the calctrans recipe execution:

./CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits SCIENCE ./TELLURIC\_CORR.fits TELLURIC\_CORR

This will produce several fits files including:

SCIENCE\_TELLURIC\_CORR\_CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits

Which is the telluric corrected version of

CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits



### 7.7 Old Parameter Renames

We list the 1.5.9 MOLECFIT parameters in groups that users of the original MOLECFIT would be familiar with and tabulate them with their equivalence in the new MOLECFIT.

#### 7.7.1 Input Data

| MOLEC          | FIT 1.5.9         |         | MOLECFIT >3.x    |                   |          |                 |  |
|----------------|-------------------|---------|------------------|-------------------|----------|-----------------|--|
| Name           | Туре              | Default | Name             | Туре              | Default  | Comment         |  |
| filename       | String            | "none"  | -                | _                 | -        | Declared in SOF |  |
| listname       | String            | "none"  | -                | -                 | -        | Not supported   |  |
| trans          | Integer           | 1       | TRANSMISSION     | Logical           | TRUE     |                 |  |
| col_lam        | String            | "undef" | COLUMN_LAMBDA    | String            | "lambda" |                 |  |
| col_flux       | String            | "undef" | COLUMN_FLUX      | String            | "flux"   |                 |  |
| col_dflux      | String            | "undef" | COLUMN_DFLUX     | String            | "dflux"  |                 |  |
| default_error  | Double            | 0.01    | DEFAULT_ERROR    | Double            | 0.01     |                 |  |
| wlgtomicron    | Double            | 1.0     | WLG_TO_MICRON    | Double            | 1.0      |                 |  |
| vac_air        | String            | "vac"   | WAVELENGTH_FRAME | String            | "VAC"    |                 |  |
| wrange_include | Dbls <sup>3</sup> | "none"  | WAVE_INCLUDE     | Dbls <sup>4</sup> | "NULL"   |                 |  |
| wrange_exclude | Dbls <sup>1</sup> | "none"  | WAVE_EXCLUDE     | Dbls <sup>2</sup> | "NULL"   |                 |  |
| prange_exclude | Dbls <sup>1</sup> | "none"  | PIXEL_EXCLUDE    | Dbls <sup>2</sup> | "NULL"   |                 |  |

#### 7.7.2 Results

| MOLEC         | FIT 1.5.9 |          | MOLECFIT >3.x |      |         |               |
|---------------|-----------|----------|---------------|------|---------|---------------|
| Name          | Туре      | Default  | Name          | Туре | Default | Comment       |
| output_dir    | String    | "output" | -             | -    | _       | Not supported |
| output_name   | String    | "none"   | _             | -    | _       | Not supported |
| plot_creation | String    | XP       | _             | -    | _       | Not supported |
| plot_range    | Logical   | 0        | -             | -    | -       | Not supported |

#### 7.7.3 Fit Precision

| M    | IOLECFIT | 1.5.9               | MOLECFIT >3.x |         |                     |  |
|------|----------|---------------------|---------------|---------|---------------------|--|
| Name | Туре     | Default             | Name          | Comment |                     |  |
| ftol | Double   | $1 \times 10^{-10}$ | FTOL          | Double  | $1 \times 10^{-10}$ |  |
| xtol | Double   | $1 \times 10^{-10}$ | XTOL          | Double  | $1 \times 10^{-10}$ |  |

<sup>1</sup>String representation of a space-separated list of doubles – e.g. "4.230 5.238 6.239", or the string value "none" <sup>2</sup>String representation of a comma-separated list of doubles – e.g. "4.230,5.238,6.239", or the string value "NULL"



### 7.7.4 Molecular Columns

| MOLECFIT 1.5.9 |                   |       |            | MOLEC              | FIT >3.x |         |
|----------------|-------------------|-------|------------|--------------------|----------|---------|
| Name           | Type Default      |       | Name       | Туре               | Default  | Comment |
| list_molec     | Strs <sup>5</sup> | "H2O" | LIST_MOLEC | Strs <sup>6</sup>  | "NULL"   |         |
| fit_molec      | Ints <sup>7</sup> | "1"   | FIT_MOLEC  | Ints <sup>8</sup>  | "NULL"   |         |
| rel_col        | Dbls <sup>9</sup> | "1.0" | REL_COL    | Dbls <sup>10</sup> | "NULL"   |         |

#### 7.7.5 Background and Continuum

| MOLECFIT 1.5.9 |         |         | MOLECFIT >3.x              |         |         |         |  |
|----------------|---------|---------|----------------------------|---------|---------|---------|--|
| Name           | Туре    | Default | Name                       | Туре    | Default | Comment |  |
| flux_unit      | Integer | 0       | FLUX_UNIT                  | Integer | 0       |         |  |
| fit_back       | Integer | 1       | FIT_TELESCOPE_BACKGROUND   | Logical | TRUE    |         |  |
| telback        | Double  | 0.1     | TELESCOPE_BACKGROUND_CONST | Double  | 0.1     |         |  |
| fit_cont       | Integer | 1       | FIT_CONTINUUM              | Logical | TRUE    |         |  |
| cont_n         | Integer | 0       | FIT_CONTINUUM_N            | Integer | 0       |         |  |
| cont_const     | Double  | 1.0     | CONTINUUM_CONST            | Double  | 1.0     |         |  |

#### 7.7.6 Wavelength Solution

| MOLI      | ECFIT 1.5 | .9      |           | MOLEC   | FIT >3.x |         |
|-----------|-----------|---------|-----------|---------|----------|---------|
| Name      | Туре      | Default | Name      | Туре    | Default  | Comment |
| fit_wlc   | Integer   | 1       | FIT_WLC   | Logical | TRUE     |         |
| wlc_n     | Integer   | 1       | WLC_N     | Integer | 1        |         |
| wlc_const | Double    | 0.0     | WLC_CONST | Double  | 0.0      |         |

<sup>&</sup>lt;sup>3</sup>String representation of a space-separated list of strings, e.g. "H2O O2 O3", or the string value "none"

<sup>&</sup>lt;sup>4</sup>String representation of a comma-separated list of strings, e.g. "H2O,O2,O3", or the string value "NULL"

<sup>&</sup>lt;sup>5</sup>String representation of a space-separated list of integers, e.g. "1 0 1", or the string value "none"

<sup>&</sup>lt;sup>6</sup>String representation of a comma-separated list of integers, e.g. "1,0,1", or the string value "NULL"

<sup>&</sup>lt;sup>7</sup>String representation of a space-separated list of doubles, e.g. "4.230 5.238 6.239", or the string value "none"

<sup>&</sup>lt;sup>8</sup>String representation of a comma-separated list of doubles, e.g. "4.230,5.238,6.239", or the string value "NULL"


### 7.7.7 Line Spread Functions

| MOLECFIT 1.5.9 |         | MOLECFIT >3.x |                      |         |         |                 |
|----------------|---------|---------------|----------------------|---------|---------|-----------------|
| Name           | Туре    | Default       | Name                 | Туре    | Default | Comment         |
| fit_res_box    | Integer | 1             | FIT_RES_BOX          | Logical | TRUE    |                 |
| relres_box     | Double  | 1.0           | RES_BOX              | Double  | 1.0     |                 |
| fit_res_gauss  | Integer | 1             | FIT_RES_GAUSS        | Logical | TRUE    |                 |
| res_gauss      | Double  | 1.0           | RES_GAUSS            | Double  | 1.0     |                 |
| fit_res_       | Integer | 1             | FIT_RES_LORENTZ      | Logical | TRUE    |                 |
| lorentz        |         |               |                      |         |         |                 |
| res_lorentz    | Double  | 1.0           | RES_LORENTZ          | Double  | 1.0     |                 |
| kernmode       | Integer | 1             | KERNMODE             | Logical | FALSE   |                 |
| kernfac        | Double  | 3.0           | KERNFAC              | Double  | 3.0     |                 |
| varkern        | Integer | 0             | VARKERN              | Logical | FALSE   |                 |
| kernel_file    | String  | "none"        | USE_INPUT_KERNEL     | Logical | TRUE    | Flag has to be  |
|                |         |               |                      |         |         | true for kernel |
|                |         |               |                      |         |         | file to be used |
|                |         |               | MODEL_MAPPING_KERNEL | String  | "NULL"  | lf value is     |
|                |         |               |                      |         |         | "NULL" then can |
|                |         |               |                      |         |         | be specified in |
|                |         |               |                      |         |         | SOF file        |



### 7.7.8 Ambient Parameters

| MOLECFIT 1.5.9 |        |               | MOLECFIT >3.x       |        |               |
|----------------|--------|---------------|---------------------|--------|---------------|
| Name           | Туре   | Default       | Name                | Туре   | Default       |
| utc            | Double | -1.0          | UTC_VALUE           | Double | -1.0          |
| utc_key        | String | "UTC"         | UTC_KEYWORD         | String | "UTC"         |
| telalt         | Double | 90.0          | TELESCOPE_ANGLE_    | Double | 90.0          |
|                |        |               | VALUE               |        |               |
| telalt_key     | String | "ESO TEL ALT" | TELESCOPE_ANGLE_    | String | "ESO TEL ALT" |
|                |        |               | KEYWORD             |        |               |
| rhum           | Double | 15.0          | RELATIVE_HUMIDITY_  | Double | 15.0          |
|                |        |               | VALUE               |        |               |
| rhum_key       | String | "ESO TEL      | RELATIVE_HUMIDITY_  | String | "ESO TEL      |
|                |        | AMBI RHUM"    | KEYWORD             |        | AMBI RHUM"    |
| pres           | Double | 750.0         | PRESSURE_VALUE      | Double | 750.0         |
| pres_key       | String | "ESO TEL AMBI | PRESSURE_KEYWORD    | String | "ESO TEL AMBI |
|                |        | PRES START"   |                     |        | PRES START"   |
| temp           | Double | 15.0          | TEMPERATURE_VALUE   | Double | 15.0          |
| temp_key       | String | "ESO TEL      | TEMPERATURE_KEYWORD | String | "ESO TEL      |
|                |        | AMBI TEMP"    |                     |        | AMBI TEMP"    |
| m1temp         | Double | 15.0          | MIRROR_TEMPERATURE_ | Double | 15.0          |
|                |        |               | VALUE               |        |               |
| m1temp_key     | String | "ESO TEL TH   | MIRROR_TEMPERATURE_ | String | "ESO TEL TH   |
|                |        | M1 TEMP"      | KEYWORD             |        | M1 TEMP"      |
| geoelev        | Double | 2635.0        | ELEVATION_VALUE     | Double | 2635.0        |
| geoelev_key    | String | "ESO TEL      | ELEVATION_KEYWORD   | STRING | "ESO TEL      |
|                |        | GEOELEV"      |                     |        | GEOELEV"      |
| longitude      | Double | -70.4051      | LONGITUDE_VALUE     | Double | -70.4051      |
| longitude_     | String | "ESO TEL      | LONGITUDE_KEYWORD   | String | "ESO TEL      |
| key            |        | GEOLON"       |                     |        | GEOLON"       |
| latitude       | Double | -24.6276      | LATITUDE_VALUE      | Double | -24.6276      |
| latitude_key   | String | "ESO TEL      | LATITUDE_KEYWORD    | String | "ESO TEL      |
|                |        | GEOLAT"       |                     |        | GEOLAT"       |

### 7.7.9 Instrument Parameters

| MOLECFIT 1.5.9 |        | MOLECFIT >3.x |                    |        |                     |
|----------------|--------|---------------|--------------------|--------|---------------------|
| Name           | Туре   | Default       | Name               | Туре   | Default             |
| slitw          | Double | 0.4           | SLIT_WIDTH_VALUE   | Double | 0.4                 |
| slit_key       | String | "NONE"        | SLIT_WIDTH_KEYWORD | String | "ESO INS SLIT1 WID" |
| pixsc          | Double | 0.086         | PIX_SCALE_VALUE    | Double | 0.086               |
| pixsc_none     | String | "NONE"        | PIX_SCALE_KEYWORD  | String | "NONE"              |



#### 7.7.10 Atmostpheric Profiles

| MOLECFIT 1.5.9 |         |                      | MOLECFIT >3.x         |         |            |  |
|----------------|---------|----------------------|-----------------------|---------|------------|--|
| Name           | Туре    | Default              | Name                  | Туре    | Default    |  |
| ref_atm        | String  | "equ.atm"            | REFERENCE_ATMOSPHERIC | String  | "equ.fits" |  |
| gdas_dir       | String  | "data/profiles/grib" | -                     | -       | -          |  |
| gdas_prof      | String  | "auto"               | GDAS_PROFILE          | String  | "auto"     |  |
| layers         | Integer | 1                    | LAYERS                | Logical | TRUE       |  |
| emix           | Double  | 5.0                  | EMIX                  | Double  | 5.0        |  |
| pwv            | Double  | -1.0                 | PWV                   | Double  | -1.0       |  |



# 8 The Model

In this section, the atmospheric model used for MOLECFIT is described in more detail. First, the building and properties of the atmospheric profiles required for the calculation of emission and absorption spectra are discussed (Section 8.1). Then, we explain the properties of the radiative transfer code used (Section 8.2). The contribution of the different molecules to the resulting atmospheric spectra is discussed in Section 8.3. Moreover, we provide some information on the modelling of the telescope emission (Section 8.4). Finally, we describe how the resulting model is adapted to the input science spectrum (Section 8.5).

### 8.1 Atmospheric profiles and meteorological data

Information concerning the composition of the atmosphere is available at various levels. To the end of creating a uniform profile with the variables temperature, pressure, and density of various molecular species as a function of geoelevation, three sources of input are merged: standard profile (produced for MIPAS onboard the ENVISAT satellite), GDAS profile, and EMM data.

The largest amount of molecular density information is contained in the atmospheric standard profiles. However, they are only available for specific geographical latitudes and do not contain any time information whatsoever (see Section 8.1.1). To compensate the lack of time information, one can rely on the EMM (see Section 8.1.3). It provides the most frequent updates and is specific to the selected observing site. Unfortunately, it cannot provide molecular species information apart from water vapor (relative humidity measurements) and is restricted to a local on-site measurement, i.e. a single geoelevation data point only. To bridge the gap between these two data sources, GDAS provides a global grid of profile measurements (with approximate grid spacing of 110 km) to an altitude of  $\sim 26$  km with updates every three hours. GDAS does not contain molecular species apart from H<sub>2</sub>O, though (see Section 8.1.2).

These three data sources and the required processing for use with MOLECFIT are described in detail below (see also Noll et al. [RD06]).

### 8.1.1 MIPAS profiles

The atmospheric standard profiles provide the basis for the model atmosphere used in MOLECFIT (see parameter REF\_ATM in Section 6.3) including information on pressure, temperature, and molecular abundance as function of height (121 levels in the range 0-120 km). Up to now, the RFM homepage [RD24] provides standard profiles for mid-latitude (Lat =  $45^{\circ}$ , both, day and night), polar winter/summer (Lat =  $75^{\circ}$ ) and equatorial day-time conditions in such a configuration (J. Remedios 2001). So far, the following molecules are included in this standard profile: N<sub>2</sub>, O<sub>2</sub>, CO<sub>2</sub>, O<sub>3</sub>, H<sub>2</sub>O, CH<sub>4</sub>, N<sub>2</sub>O, HNO<sub>3</sub>, CO, NO<sub>2</sub>, N<sub>2</sub>O<sub>5</sub>, CIO, HOCI, CIONO<sub>2</sub>, NO, HNO<sub>4</sub>, HCN, NH<sub>3</sub>, F11, F12, F14, F22, CCl<sub>4</sub>, COF<sub>2</sub>, H<sub>2</sub>O<sub>2</sub>, C<sub>2</sub>H<sub>2</sub>, C<sub>2</sub>H<sub>6</sub>, OCS, SO<sub>2</sub>, and SF (see Table 8.3). Additional molecule profiles for F13 (CCIF3), F21 (CHCl<sub>2</sub>F), F113 (C<sub>2</sub>Cl<sub>3</sub>F<sub>3</sub>), F114 (C<sub>2</sub>Cl<sub>2</sub>F<sub>4</sub>), F115 (C<sub>2</sub>ClF<sub>5</sub>), and CH<sub>3</sub>Cl are available. Apart from these data, less resolved profiles (a tropical, sub-arctic summer/winter and a US standard profile) are available with 50 geoelevation layers including the molecules H<sub>2</sub>O, CO<sub>2</sub>, O<sub>3</sub>, N<sub>2</sub>O, CO, CH<sub>4</sub>, and O<sub>2</sub> only.



**Comparison between atmospheric standard profiles** In this section, an equatorial day-time equ.fits and a mid-latitude profile ngt.atm, corresponding to a latitude Lat =  $45^{\circ}$ , will be compared. The location of Paranal (Lat =  $24.6^{\circ}$ ) is between these two profiles. In Figure 8.1 both profiles are shown. Although the distribution of several molecules (e.g. N<sub>2</sub>, O<sub>2</sub>, CO<sub>2</sub>) does not vary, significant differences between the two profiles are visible. To investigate the impact of the input profile differences on the output spectra, LBLRTM was run with the same input parameters, but with varying standard profiles.

The resulting spectra are shown in Figures 8.2/8.3. These plots reveal output radiance spectra differing by at most  $\pm 10\%$ . The same is true for the transmission spectra, although somewhat less obvious due to numerical instabilities. Calculating the broad-band ratios in the main filter ranges  $UBVRI_cJHKLMN$  indicates deviations of less than 2% (see Table 8.1). Hence, one can conclude that the differences between the two standard atmospheric profiles are negligible at this stage. Anu Dudhia [RD08], the author of the RFM code, recommends the equatorial profile equ.fits to be used for typical applications at Cerro Paranal.

 Table 8.1: Broad-band comparison of the relative ratios between the equ.fits and the ngt.atm atmospheric standard profiles.

| Filter     | $\lambda_{min}$<br>[ $\mu$ m] | $\lambda_{\sf max}$<br>[ $\mu$ m] | Radiance<br>ratio [%] | Transmission<br>ratio [%] |
|------------|-------------------------------|-----------------------------------|-----------------------|---------------------------|
| U          | 0.33                          | 0.40                              | 0.02                  | 0.06                      |
| В          | 0.39                          | 0.50                              | 0.00                  | 0.02                      |
| V          | 0.50                          | 0.60                              | 0.14                  | 0.56                      |
| R          | 0.58                          | 0.82                              | 0.05                  | 0.28                      |
| <i>I</i> c | 0.73                          | 0.85                              | 0.01                  | 0.04                      |
| J          | 1.10                          | 1.34                              | -0.00                 | -0.00                     |
| Н          | 1.50                          | 1.80                              | -0.00                 | -0.01                     |
| K          | 2.00                          | 2.40                              | -0.05                 | -0.06                     |
| L          | 3.56                          | 4.12                              | -0.00                 | -0.05                     |
| М          | 4.52                          | 4.96                              | -0.69                 | 1.37                      |
| Ν          | 7.40                          | 13.60                             | -0.60                 | 1.39                      |





Figure 8.1: Comparison of the equatorial (equ.fits) and the mid-latitude night time atmospheric profile (ngt.atm). Red lines correspond to the equatorial, blue lines to the mid-latitude profile.



| Doc. Number:  | VLT-MAN-ESO- |  |  |
|---------------|--------------|--|--|
|               | 19550-5772   |  |  |
| Doc. Version: | 4.4.2        |  |  |
| Released on:  | 2025-02-11   |  |  |
| Page:         | 79 of 120    |  |  |



**Figure 8.2:** Direct comparison between the radiance spectra of equatorial day time (red line) and mid-latitude night time atmospheric standard profile (grey line) over the entire wavelength range  $\lambda = 0.3 - 30 \,\mu$ m (top panel). Blue lines mark the wavelength range ( $\lambda = 4 - 28 \,\mu$ m) plotted in the three panels below. *Second panel:* Same as in top panel, but for the limited wavelength range. *Third and bottom panel:* Residuals equ-ngt and ratio equ/ngt of the radiance spectra, respectively.



| Doc. Number:  | VLT-MAN-ESO- |  |  |
|---------------|--------------|--|--|
|               | 19550-5772   |  |  |
| Doc. Version: | 4.4.2        |  |  |
| Released on:  | 2025-02-11   |  |  |
| Page:         | 80 of 120    |  |  |







| P[hPa] | HGT[km] | T[K]  | RELHUM[%] |
|--------|---------|-------|-----------|
| 903    | 0.971   | 294.5 | 49.9      |
| 900    | 0.976   | 295.8 | 35.5      |
| 850    | 1.467   | 293.7 | 30.9      |
| 800    | 1.985   | 291.0 | 29.8      |
| 750    | 2.533   | 288.2 | 27.4      |
| 700    | 3.112   | 284.5 | 26.0      |
| 650    | 3.726   | 280.7 | 18.8      |
| 600    | 4.379   | 276.2 | 11.4      |
| 550    | 5.077   | 271.4 | 8.7       |
| 500    | 5.827   | 266.0 | 7.5       |
| 450    | 6.638   | 259.9 | 7.5       |
| 400    | 7.522   | 252.7 | 10.5      |
| 350    | 8.494   | 244.7 | 25.1      |
| 300    | 9.578   | 236.0 | 53.4      |
| 250    | 10.813  | 227.0 | 66.5      |
| 200    | 12.267  | 218.8 | 37.7      |
| 150    | 14.069  | 209.2 | 17.3      |
| 100    | 16.489  | 200.3 | 32.4      |
| 50     | 20.571  | 206.1 | 0.0       |
| 20     | 26.324  | 221.4 | 0.0       |
|        |         |       |           |

### 8.1.2 GDAS profiles

The GDAS data provided by NOAA are a model-based set of meteorological data dedicated to weather forecast studies. The models are archived by the ARL, as a global, 1 degree latitude/longitude data set based on pressure surfaces (starting from Dec. 2004). Apart from various meteorological parameters for the surface, vertical profiles for 23 pressure levels ranging from 0 to about 26 km are provided for the geopotential height, temperature, relative humidity, and wind components (not used in MOLECFIT) for three dimensions. An example is shown in Table 8.2.

The MOLECFIT software package provides the entire GDAS data for the location of Cerro Paranal from Dec. 2004 to Sep. 2013 on a 3 h basis taken from the NOAA archive<sup>11</sup>. Later dates (on a 6 h basis) or data for a different site are automatically downloaded from an online archive<sup>12</sup>. See also Section 8.1.4.

### 8.1.3 ESO Meteo Monitor

The EMM provides information on the local meteorological conditions at the ESO sites La Silla and Paranal. The data at Paranal are taken by a local meteo station mounted on a 30 m high mast installed in October 1984 [RD07]. This meteo station provides the following meteorological information on a 20 min average basis:

<sup>&</sup>lt;sup>11</sup>ftp://arlftp.arlhq.noaa.gov/pub/archives/gdas1/

<sup>&</sup>lt;sup>12</sup>http://nomad1.ncep.noaa.gov/pub/gdas/rotating/



Starting from January 1st 1985, currently  $\sim$  400 000 data points are measured with the following accuracy [RD07]:

| wind direction: | ~5.63deg                          |
|-----------------|-----------------------------------|
| wind speed:     | ~2% over 10m/s                    |
| temperature:    | ~0.1deg                           |
| humidity:       | linearity about 1%                |
| seeing:         | better than 10% above 0.25 arcsec |

The data can be retrieved online<sup>13</sup> on a daily basis, or as download provided by M. Sarazin<sup>14</sup> and are cumulatively shown in Figure 8.4. Thus, for any requested average time interval, like e.g. December and January, ample measurements are available. In compiling these data, care has to be taken to remove bad measurements before further processing.



Figure 8.4: Histograms of EMM data (from Jan. 1985 to Jan. 2008). *Left panel*: temperature; *middle panel*: relative humidity; *right panel*: pressure.

<sup>13</sup>http://archive.eso.org/asm/ambient-server

<sup>&</sup>lt;sup>14</sup>http://www.eso.org/gen-fac/pubs/astclim/paranal/database/



### 8.1.4 Processing of ESO Meteor Monitor data, GDAS, and MIPAS profiles

The main disadvantage of the GDAS profiles is that they do not represent the local atmospheric conditions of the geographical position and height of the observing site as accurately as provided by the EMM, and even more so for the MIPAS profiles. Therefore, one has to investigate how the three sources of information can be merged into a single profile.

**GDAS profile processing** The GDAS profiles originate from a server at NOAA<sup>15</sup>. They are retrieved via a dedicated software package GRIB (see Section 9). GRIB downloads a large data set containing the specific GDAS information for the requested point in time. As this data set contains a model for the complete globe, subsequently, the data points for the specified geolocation are extracted. Moreover, as the GDAS data are taken on a 3 hr basis only, two profiles need to be retrieved surrounding the requested point in time. The parameters OBSDATE, UTC, LONGITUDE, and LATITUDE are required for this task (see Section 6.3). They are usually provided by standard and ESO FITS keywords. In the following, we will describe how the resulting two profiles are combined to best match the date of the observations.

If the profiles exist locally, no download from the web-server is required. In addition, the MOLECFIT software distribution contains a compilation of all Cerro Paranal GDAS profiles for the dates from Dec. 01, 2004 to Sep. 30, 2013. Thus, before requesting the data from the web-server (in the case that they do not exist locally already), this database is checked for the existence of the appropriate profiles. For updates of this data set and the retrieval of data for other observing sites, see Section D.1.4.

Unfortunately, the web-server does not provide GDAS profiles for all dates or an internet connection may not always be available. Therefore, MOLECFIT incorporates a fall-back alternative to ensure availability of GDAS data in all occasions. To that end, the monthly averaged profiles from the Cerro Paranal sky model are included as well (for a detailed description see [RD09] and Noll et al. [RD06]). If after checking the local database or the web-server, a profile is still missing, the best-matching profile for the two-month bins Dec.-Jan., Feb.-Mar., ... are taken. The corresponding files, GDAS\_t0\_sX.fits with X=1 to 6, are located in "share/molecfit/data/profiles/lib/", with X=1 for December/January, X=2 for February/March, X=3 for April/May, X=4 for June/July, X=5 for August/September, X=6 for October/November. This is also done if the site is not Cerro Paranal.

The described procedure for the GDAS profile retrieval is performed if the parameter GDAS\_PROF (see Section 6.3) is set to "auto", which is the default. As an alternative, if GDAS\_PROF is set to "null" a specific GDAS-like profile (see Table 8.2) must be provided in the SOF. Finally, it is possible to avoid the use of GDAS profiles by setting GDAS\_PROF to "none".

**Time averaged profiles** Typically, the requested observation date does not fall exactly onto a single GDAS time slot. Instead of simply retrieving the closest dataset the two neighbouring profiles are obtained. In order to combine the two, a time-weighted average is calculated, i.e. performing a linear interpolation.

**Merging GDAS and MIPAS profiles** Next, the resulting GDAS profile is merged with the MIPAS standard profile. To that end, the MIPAS profile is regridded to a new irregular height grid with 50 levels (see Figure 8.5) spanning the whole geoelevation range from 2-120 km for Cerro Paranal. The GDAS profile is regridded to the

<sup>&</sup>lt;sup>15</sup>http://140.90.198.158/pub/gdas/rotating/



same grid in the range 2-26 km and then used to substitute the respective columns in the MIPAS profile. In addition, the four height levels from 20-26 km are not only a simple substitute of the MIPAS data, but a weighted mix of GDAS and MIPAS profile, in order to provide a smooth transition from one dataset to the other. The influence of the GDAS profile decreases with increasing height: 80%, 60%, 40%, 20% at 20 km, 22 km, 24 km, 26 km, respectively. Beyond 26 km, no GDAS information is available.

The discussed fixed grid of layers is used if the parameter LAYERS (see Section 6.3) is set to 1, which is the default. A value of 0 will cause the building of a natural grid consisting of all layers of the MIPAS and the GDAS profile. If local meteo data are used, the observer altitude GEOELEV is also added. The transition from GDAS to MIPAS is performed by means of a decrease of the relative difference of pressure, temperature, and water vapor concentration of both profiles at the height of the uppermost valid GDAS layer up to an altitude which is 1.2 times higher. The resulting grid of layers is (slightly) more accurate than the fixed grid, but also consists of a significantly higher number of levels.



**Figure 8.5:** *Composition of atmospheric profile:* relative importance of EMM data (red), GDAS data (green) and MIPAS data (blue) as function of geoelevation. Note that the interface boundary of GDAS and MIPAS data varies depending on the availability of the GDAS data.

**Combining GDAS/MIPAS profiles with EMM data** Observed data from ESO telescopes provide on-site measurements at the ground layer for pressure, temperature, and humidity originating from the ESO meteo monitor at Cerro Paranal. A detailed study of the GDAS data (see Figure 8.6), which represent the local troposphere, including information concerning the dominant wind direction as a function of altitude reveals a gradual reversal (rotation of 180°) at a geoelevation of 5 km, the so-called mixing altitude  $h_{mix}$  (see [RD09]). Beyond this altitude, the wind direction remains constant independent of the observation date. Thus, it can safely be assumed that at this altitude the influence of the local environment (as determined from the EMM data) has diminished.

In order to smoothly integrate the EMM data, all GDAS values for pressure, temperature, and humidity below the altitude of the observatory are set to the EMM value. Values above the aforementioned mixing altitude are left untouched. Intermediate values are linearly interpolated resulting in a smooth transition. To this end, first, a logarithmically interpolated value of the GDAS data corresponding to the observatory's altitude  $h_{tel}$  is calculated for pressure, temperature, and humidity. These values describe the reference point for the linear decrease of the relative difference between EMM and GDAS data and in the interval  $h_{tel} - h_{mix}$ .

The resulting profile is a smooth combination of all input data, i.e. MIPAS, GDAS, and EMM.

The default mixing altitude of 5 km can be manipulated by changing the parameter EMIX (see Section 6.3). This could be interesting for other observing sites. Setting EMIX to a value lower than the observer altitude GEOELEV



causes a profile building without local meteo data. In this case, the parameters PRES, TEMP, and RHUM are ignored.



**Figure 8.6:** *GDAS wind direction as function of geoelevation:* red, cyan, blue, and green curves represent summer, autumn, winter, and spring, respectively. The black symbols show the all year average with the corresponding scatter. The vertical dashed line marks the geoelevation of Paranal. At  $\sim$  5 km height, a constant plateau is reached.

**Scaling of the merged water vapour profile to a given PWV** The height profiles of the molecular abundances are defined by the atmospheric standard profile. The only exception is water vapor, where the profile is a combination of modelled and observed data as described above. The GDAS and EMM data used cannot be controlled by the user. There might be cases where this is not satisfying. For example, the user might be interested to use water vapor columns derived from independent measurements for a better start profile. For this purpose, MOLECFIT allows the user to enter a PWV value (parameter PWV, see Section 6.3), which is used to scale the merged water vapor profile. In this way, the input water vapor content of the atmosphere can be fixed. Only the shape of the profile is then ruled by the GDAS and EMM data. The profile scaling factor RELCOL, which is used in the context of the fitting procedure, refers to the modified profile. By default, the PWV option is switched off as indicated by a value of -1.

### 8.2 Radiative transfer code

MOLECFIT uses the radiative transfer code Line-By-Line Radiative Transfer Model (LNFL v3.2 / LBLRTM v12.11), which is widely used in atmospheric and climate research studies.



### 8.2.1 Line File / Line-By-Line Radiative Transfer Model (LNFL/LBLRTM)

LBLRTM is developed within the Radiative Transfer Working Group of the AER (see also Clough et al. [RD05], [RD19], and [RD17]). It is publicly available. LBLRTM can handle all molecules incorporated in the aer\_v\_<version> line parameter database [RD17] and offers a wide range of possibilities to adjust input parameters (see [RD25] for more details).

The AER code package used here consists of two programmes: (a) the "Line File" code LNFL, which extracts user selected spectral lines from the line parameter database, and provides these in appropriate form as input for (b) the radiative transfer code LBLRTM. Within MOLECFIT, the most recent versions LNFL v3.2 and LBLRTM v12.11 are used.

Some LBLRTM key features are (taken from [RD17]):

- the Voigt line shape is used at all atmospheric levels with an algorithm based on a linear combination of approximating functions;
- it has been and continues to be extensively validated against atmospheric radiance spectra from the ultra-violet to the sub-millimeter;
- it incorporates the self- and foreign-broadened water vapor continuum model, MT\_CKD as well as continua for carbon dioxide, and for the collision induced bands of oxygen at 1600 cm<sup>-1</sup> ( $\lambda$  = 6.25  $\mu$ m) and nitrogen at 2350 cm<sup>-1</sup> ( $\lambda$  = 4.255  $\mu$ m);
- all parameters of the line database are used including the pressure shift coefficient, the halfwidth temperature dependence, and the coefficient for the self-broadening of water vapor;
- a version of the Total Internal Partition Function (TIPS) programme is used for the temperature dependence of the line intensities;
- the effects of CO<sub>2</sub> line coupling are treated as first order with the coefficients for carbon dioxide generated from Niro et al. [RD10];
- temperature dependent cross section data such as those available with the aer\_v\_<version> database may be used to treat the absorption due to heavy molecules, e.g. the halocarbons;
- an algorithm is implemented for the treatment of the variation of the Planck function within a vertically inhomogeneous layer as discussed in Clough et al. [RD11];
- algorithmic accuracy of LBLRTM is approximately 0.5% and the errors associated with the computational procedures are of the order of five times less than those associated with the line parameters so that the limiting error is that attributable to the line parameters and the line shape;
- its computational efficiency mitigates the computational burden of the line-by-line flux and cooling rate calculation (Clough et al. [RD11]), for example linear algebraic operations are used extensively in the computationally intensive parts of LBLRTM so that vectorisation is particularly effective with a typical vectorised acceleration of 20;
- FFT instrument function with a choice of 9 apodisation functions;



- includes a realistic spectral sea surface emissivity model in the infrared (Masuda et al. [RD12]; Wu&Smith [RD13]);
- input atmospheric profiles in either altitude or pressure coordinates;
- interfaces with other radiative transfer models (like RRTM), and also the forward model for inversion algorithms (e.g. Tropospheric Emission Spectrometer (TES) and Infrared Atmospheric Sounding Interferometer (IASI));
- these attributes provide spectral radiance calculations with accuracies consistent with the measurements against which they are validated and with computational times that greatly facilitate the application of the line-by-line approach to current radiative transfer applications.

In principle, the user can change the setup of the LBLRTM. See the LBLRTM parameters in Table6.2. A change of the fixed LBLRTM input parameters is only recommendable for those users, which have a very good knowledge of the physics of atmospheric radiative transfer and/or LBLRTM.

### 8.2.2 aer line database

For calculating molecular spectra, the  $aer_v_<version>$  database [RD17] is used. It is built from HITRAN 2016 [RD20] and contains several updates. Version 3.8.1.2 covers the full spectral range from  $0 - 25, 232 \text{ cm}^{-1}$  (i.e. down to  $0.4 \mu \text{m}$ ) provides spectral information for 42 molecules. In total, more than 8,900,000 spectral lines are included. The majority is based on modelled data. However, only those 30 molecules are taken into account, which are present in the atmospheric standard profile. The remaining ones are minor trace gases and do not contribute significantly neither to radiance, nor transmission spectra (see Section 8.3). Table 8.2.2 provides an overview of all molecules based on the aer database (and known by LBLRTM) and those contained in the standard atmospheres (Column 4). See further details in Appendix D.1.3



Х

|                     | Index        | Molecule                      | Name                | Std Atmosphere | LBLRTM |
|---------------------|--------------|-------------------------------|---------------------|----------------|--------|
|                     | 1            | H <sub>2</sub> O              | Water               | Х              | Х      |
|                     | 2            | CO <sub>2</sub>               | Carbon dioxide      | Х              | Х      |
|                     | 3            | O <sub>3</sub>                | Ozone               | Х              | Х      |
|                     | 4            | N <sub>2</sub> O              | Nitrous oxide       | Х              | Х      |
|                     | 5            | CO                            | Carbon monoxide     | Х              | Х      |
|                     | 6            | $CH_4$                        | Methane             | Х              | Х      |
|                     | 7            | O <sub>2</sub>                | Oxygen              | Х              | Х      |
|                     | 8            | NO                            | Nitric oxide        | Х              | Х      |
|                     | 9            | SO <sub>2</sub>               | Sulfur dioxide      | Х              | Х      |
|                     | 10           | NO <sub>2</sub>               | Nitrogen dioxide    | Х              | Х      |
|                     | 11           | $NH_3$                        | Ammonia             | Х              | Х      |
|                     | 12           | $HNO_3$                       | Nitric acid         | Х              | Х      |
|                     | 13           | OH                            | Hydroxyl            |                | Х      |
|                     | 14           | HF                            | Hydrogen fluoride   |                | Х      |
|                     | 15           | HCI                           | Hydrogen chloride   |                | Х      |
|                     | 16           | HBr                           | Hydrobromic acid    |                | Х      |
|                     | 17           | HI                            | Hydrogen iodide     |                | Х      |
|                     | 18           | CIO                           | Chlorine monoxide   | Х              | Х      |
|                     | 19           | OCS                           | Carbonyl sulfide    | Х              | Х      |
|                     | 20           | H <sub>2</sub> CO             | Formaldehyde        |                | Х      |
|                     | 21           | HOCI                          | Hypochlorous acid   | Х              | Х      |
|                     | 22           | $N_2$                         | Nitrogen            | Х              | Х      |
|                     | 23           | HCN                           | Hydrogen cyanide    | Х              | Х      |
|                     | 24           | CH <sub>3</sub> Cl            | Chloromethane       |                | Х      |
|                     | 25           | $H_2O_2$                      | Hydrogen peroxide   | Х              | Х      |
|                     | 26           | $C_2H_2$                      | Acetylene           | Х              | Х      |
|                     | 27           | C <sub>2</sub> H <sub>6</sub> | Ethane              | Х              | Х      |
|                     | 28           | $PH_3$                        | Phosphine           |                | Х      |
|                     | 29           | COF <sub>2</sub>              | Carbonyl fluoride   | Х              | X      |
|                     | 30           | SF <sub>6</sub>               | Sulfur hexafluoride | Х              | Х      |
|                     | 31           | H <sub>2</sub> S              | Hydrogen sulfide    |                | Х      |
|                     | 32           | HCOOH                         | Formic acid         |                | Х      |
|                     | 33           | HO <sub>2</sub>               | Hydroperoxyl        |                | X      |
|                     | 34           | 0                             | Oxygen              |                | Х      |
|                     | 35           | CIONO <sub>2</sub>            | Chlorine Nitrate    | Х              | Х      |
|                     | 36           | NO+                           | Nitrosonium         |                | Х      |
|                     | 37           | HOBr                          | Hypobromous Acid    |                | Х      |
|                     | 38           | $C_2H_4$                      | Ethylene            |                | Х      |
|                     | 39           | CH₃OH                         | Methanol            |                | Х      |
|                     | 40           | CH₃Br                         | Methyl Bromide      |                | Х      |
|                     | 41           | C <sub>3</sub> H <sub>8</sub> | Propane             |                | Х      |
|                     | 42           | $C_2N_2$                      | Cyanogen            |                | X      |
|                     | 43           | $C_4H_2$                      | Diacetylene         |                | X      |
|                     | 44           | HC <sub>3</sub> N             | Cyanoacetylene      |                | Х      |
| Document Classifica | tion: P45lic | 2                             | Hydrogen            |                | Х      |
|                     | 46           | CS                            | Carbon Monosulfide  |                | Х      |
|                     |              | ~~                            |                     | 1              |        |

47

 $SO_3$ 

Sulfur trioxide

Table 8.3: List of molecules as provided by the aer line parameter database



### 8.3 Molecular spectra

The process of fitting molecular spectra is a complex task, which requires optimised input parameters. One of the key inputs for the fitting is the number of molecules that are included in the fitting. Fewer molecules in the fitting process result in the code finding a solution in significantly shorter amounts of time. Also the fitting process is a lot more robust. However, if too few molecules are included in the fit, the results may not provide a satisfying residual for model and observation. Optimally, one should include exactly those molecules in the fit, which will significantly contribute over the wavelength range of interest.

To the end of providing some insight into what molecules are required for the fit at a specific wavelength interval, in the following, we provide guidelines with a specific focus on the test data provided with this software package for the instruments CRIRES, VISIR, and XSHOOTER.

We have computed spectra with LBLRTM covering the complete wavelength range from  $0.3 - 30 \mu m$  using all molecules (i.e.  $C_2H_2$ ,  $C_2H_6$ ,  $CH_4$ ,  $CO_2$ ,  $COF_2$ , CO,  $CIONO_2$ , CIO, F14,  $H_2O_2$ ,  $H_2O$ , HCN, HNO<sub>3</sub>, HOCI,  $N_2O$ ,  $N_2$ ,  $NH_3$ ,  $NO_2$ , NO,  $O_2$ ,  $O_3$ , OCS,  $SF_6$ ,  $SO_2$ ), individually, i.e. one at a time. At fixed wavelength, we have then calculated the normalised total radiance/transmission (by summing up the contributions of all molecules and subtracting the continuum) and the resulting relative importance of individual molecules.

The Figures C.1-C.5 shown in the Appendix, give all molecules that, over the displayed wavelength range, exhibit at fixed wavelength  $\lambda$  a relative importance of at least 5%. The molecular data have been rebinned to 3000 data points for each individual wavelength range. This results in a varying resolution and in more molecules becoming important over smaller wavelength regimes. C<sub>2</sub>H<sub>6</sub>, e.g., has a few significant lines in some of the wavelength ranges shown (e.g. Figure C.2), but does not show up in the overview plot (Figure C.1). Typically, transmission (blue) and radiance (red) plots do not differ significantly. Hence, we do not show them separately. Note that these plots do not allow calculation of absolute fluxes.

These plots can be used to identify the important molecules over any wavelength range. For the range shown in Figure C.2, e.g., the user ought to include  $H_2O$ ,  $CH_4$ , and  $O_3$ . The fitting might mildly profit also from including  $C_2H_6$ .

Alternatively, a generic atlas for the molecules listed above is available at share/esopipes/datastatic/molecfit-4.4.2/molecule\_atlas\_UVB-MIR.fits, covering the wavelength range from the UVB to the MIR; alternatively, atlases of the relevant molecules for specific ESO instruments are available in share/esopipes/datastatic/molecfit-4.4.2, and are available in the GUI in the Reflex workflow.

### 8.4 Thermal emission by telescope

The telescope structure and the observing instrument cause unavoidable thermal emission in the IR. In particular, the telescope main mirror is a significant source of radiation. Hence, for radiance spectra, this background component has to be considered. A simple approximation is the calculation of a grey body spectrum, which equals a black body (BB) of temperature T times a wavelength-independent emissivity  $\epsilon$ . Since the emitting source, i.e. the main mirror, also absorbs a fraction of  $1 - \epsilon$  of the incoming sky radiation, the relative contribution of the telescope emission to the observed radiation is increased. Therefore, the apparent grey body radiation in flux-calibrated spectra can be derived by

$$F_{\text{tel}} = \frac{\epsilon}{(1-\epsilon)} BB(T).$$
 (1)



For *T*, MOLECFIT uses the temperature of the primary mirror (input parameter M1TEMP, see Section 6.3). Since the mirror temperature is close to the ambient temperature of about 280-290 K (Paranal), the grey body emission is expected to be important at wavelengths longwards of the *H*-band. The emissivity is a free fit parameter (see parameter TELBACK). An initial value can be provided in the parameter file. The default value is 0.1.

### 8.5 Adaptation of model to input spectrum

A good correspondence of the calculated model spectrum and the observed spectrum is usually prevented by the broadening of the spectral lines by the instrument, small errors in the wavelength calibration, uncertainties in the flux calibration in the case of emission spectra, or the non-flat standard star continuum in the case of transmission spectra. Hence, these unavoidable shortcomings of observed data have to be accounted for in the fitting procedure. For this reason, MOLECFIT modifies the model spectrum using a polynomial fit of the continuum and the wavelength grid. In addition, the model gets convolved with a kernel mimicking the instrumental profile. In the following, we discuss the fit parameters related to this adaptation process in detail.

### 8.5.1 The continuum

The model spectrum is scaled by a polynomial of degree  $n_c$  (CONT\_N in the parameter file; see Section 6.3)

$$F_{\text{out}}(\lambda) = F_{\text{in}}(\lambda) \sum_{i=0}^{n_{\text{c}}} a_i \lambda^i.$$
(2)

For deriving the  $n_c + 1$  coefficients  $a_i$ , the zero point of the wavelength grid is shifted to the center of the fit range. For  $a_0 = 1$  and all other  $a_i = 0$ , the model spectrum remains unchanged. This is the default configuration for the initial coefficients. In the parameter file, the initial value of the constant term of the polynomial (parameter CONT\_CONST) can be set manually. The continuum correction is carried out independently for each fit range listed in the RANGE\_INCLUDE file if such a file is provided. A fit range (or the full spectrum) is further split if it is distributed over more than one chip.

Before correcting the continuum, optionally a flux conversion is carried out. Details on the options selected by the parameter FLUX\_UNIT are given in Section 6.3. If the required data units are not included in FLUX\_UNIT, this factor must be incorporated into the  $a_0$  coefficient of the polynomial. As a general rule, it is advisable to choose  $a_0$  close to the mean flux (emission) or maximum flux (transmission) of the input spectrum (after consideration of FLUX\_UNIT) to optimise the performance of MOLECFIT.

The expert mode (see Appendix A.5) allows the user to specify the initial values for all the polynomial coefficients of the continuum.

### 8.5.2 The wavelength solution

The wavelength grid of the model spectrum is adapted to that of the observed spectrum by applying a Chebyshev polynomial of degree  $n_w$  (WLC\_N in the parameter file; see Section 6.3)

$$\lambda' = \sum_{i=0}^{n_{\rm w}} b_i t_i,\tag{3}$$



where

$$t_{i} = \begin{cases} 1 & \text{for } i = 0 \\ \lambda & \text{for } i = 1 \\ 2 \lambda t_{i-1} - t_{i-2} & \text{for } i \ge 2 \end{cases}$$
(4)

and  $\lambda$  ranging from -1 to 1. The temporary conversion of the wavelength grid to a fixed interval results in coefficients  $b_i$  independent of the wavelength range and step size of the input spectrum. For  $b_1 = 1$  and all other  $b_i = 0$ , the model spectrum remains unchanged. This is the default configuration for the initial coefficients. In the parameter file, the initial value of the constant term of the polynomial (parameter wLC\_CONST) can be set manually. This parameter corresponds to a wavelength shift relative to half the full wavelength range. For each chip or FITS extension, the wavelength fit is carried out independently.

For checks or improvements of the input wavelength grid, the model wavelengths rebinned to the input grid are provided in the results tables of MOLECFIT and calctrans (column "mlambda", see Section 6.5). The wavelengths are always given in  $\mu$ m and vacuum. Note that the reliability of this absolute wavelength calibration depends on the quality of the fit. Outside the selected fitting ranges, the wavelengths have to be interpolated or extrapolated by the Chebyshev polynomial. In particular, at optical and near-IR wavelengths, where strong absorption bands suitable for fitting are rare, the provided wavelengths have to be taken with care. For this reason, MOLECFIT does not provide an automatic wavelength solution correction by default.

However, it can be set by using WLC\_REF=MODEL in MOLECFIT\_CORRECT (see Table 6.3).

#### 8.5.3 The resolution

The model spectrum is convolved with up to three different line spread functions (LSFs) in order to get similar line shapes as in the observed spectrum. If a profile is not desired, it can be skipped by setting its width and fit flag in the parameter file to zero (see Section 6.3).

The first kernel is a simple boxcar

$$F_{\text{box}}(i) = \begin{cases} 1 & \text{for } -w_{\text{box}}/2 \le i \le w_{\text{box}}/2\\ 0 & \text{for } i < -w_{\text{box}}/2 \cap i > w_{\text{box}}/2 \end{cases}$$
(5)

which is adapted to the pixel scale (where the pixel number is represented by *i*) and normalised to an integral of 1. In the parameter file, the width  $w_{box}$  (parameter RES\_BOX) has to be given as fraction of the slit width, which is determined by the parameters SLITW, the slit width in arcsec, and PIXSC, the pixel scale in arcsec (see Section 6.3). By default, RES\_BOX is set to 1, i.e. the slit width. The fit parameter RES\_BOX can only vary between 0 and 2. A value larger than 1 is relevant if the slit width as given in the FITS header is not accurate, which sometimes happened with CRIRES before its upgrade.

The second convolution kernel is a Gaussian

$$F_{\text{gauss}}(i) = \frac{2}{\text{FWHM}} \sqrt{\frac{\ln 2}{\pi}} e^{-4\ln 2 i^2/\text{FWHM}^2}$$
(6)

centered on 0. The FWHM is given by the driver file parameter RES\_GAUSS in pixels (see Section 6.3). It is restricted to values below 100 pixels. The default value is 1 pixel. The number of pixels in the kernel amounts to KERNFAC (default: 3) times FWHM.



Finally, the third kernel is a Lorentzian

$$F_{\text{lorentz}}(i) = \frac{1}{\pi} \frac{W_{\text{lorentz}}}{i^2 + (W_{\text{lorentz}}/2)^2}$$
(7)

centered on 0, where  $w_{\text{lorentz}}$  is the FWHM. The width is adjusted by the driver file parameter RES\_LORENTZ, which has to be provided in pixels (see Section 6.3). It is restricted to values below 100 pixels. The default value is 1 pixel. The number of pixels in the kernel amounts to KERNFAC (default: 3) times  $w_{\text{lorentz}}$ . Compared to a Gaussian, the Lorentzian approaches the 0-level flux significantly slower, at much larger distances from the maximum.

Note that width zero components can occur in typical conditions, e.g. the line profile is very close to a pure Lorentzian shape. If this is not intended, the user should reduce the number of degrees of freedom by fixing individual fit components. A zero here identifies a unity convolution (i.e. no change of the input spectrum).

The combination of a Gaussian and a Lorentzian is called a Voigt profile. The flag KERNMODE (see Section 6.3) allows the user to apply only a single Voigt profile kernel, which is calculated by an approximate formula that takes the FWHM of Gaussian and Lorentzian as input. In this case (KERNMODE = 1), KERNFAC gives the kernel size in FWHM of the derived Voigt profile and not the FWHM of Gaussian and Lorentzian, as it is done for the default mode of two independent convolutions (KERNMODE = 0). For the fit results, the KERNMODE selection should be less important than the relative contributions of boxcar, Gaussian, and Lorentzian to the fitted line profile. Significant changes in the line profile can cause deviations in the water vapor column of more than 10% (cf. Section 10).

The parameter VARKERN allows the user to fit a kernel that linearly increases with wavelength (see Section 6.3). If the flag is set to 1, this option is selected. It is suitable for dispersion-dominated kernels and constant wavelength bins. In this case, the initial FWHM parameters are given for the central wavelength of the full wavelength range (considering the data of all chips). The default VARKERN = 0 assumes a constant kernel for the entire wavelength range. This option is suitable for narrow wavelength ranges and slit/object profile-dominated kernels.

Finally, the user can provide an optimised kernel via a FITS file – identified in the parameter kernel\_file (see Section 6.3). If this option is used (i.e. the default "none" is replaced by the corresponding file name), the fixed input kernel overrules the creation of a kernel based on boxcar, Gaussian, and Lorentzian components. Since there will be no fit of the kernel shape and width, the line profile has to be known well. Note that VARKERN = 1 will have no effect, since the pixel-based input kernel is wavelength independent.



| Doc. Number:  | VLT-MAN-ESO- |  |  |
|---------------|--------------|--|--|
|               | 19550-5772   |  |  |
| Doc. Version: | 4.4.2        |  |  |
| Released on:  | 2025-02-11   |  |  |
| Page:         | 93 of 120    |  |  |

# 9 Installation

ESO pipelines can be installed via several methods, depending on your OS, most of which facilitate easy installation, upgrade and removal. Please see the "ESO Data Reduction Pipelines and Workflow Systems" page (*https://www.eso.org/pipelines*).



# 10 Code Performance

Discussion of the scientific performance of MOLECFIT are given in A. Smette, H. Sana, S. Noll, H. Horst, W. Kausch, S. Kimeswenger, M.Barden, C. Szyszka, A. M. Jones, A. Gallene, J. Vinther, P. Ballester and J. Taylor (2015, A&A 576, A77), and W. Kausch, S. Noll, A. Smette, S. Kimeswenger, M. Barden, C. Szyszka, A. M. Jones, H. Sana, H. Horst, and F. Kerber, (2015, A&A 576, A78).



# 11 Tips and tricks

In the following, we provide a summary of rules that should be taken into account for a successful application of MOLECFIT:

- Pixels with possible defects that could affect the fit quality can be excluded from the fit in two ways. First, the critical pixels can be listed as a parameter or in a FITS file specified in the parameter prange\_exclude (see Section 6.3). Second, pixels can be skipped by adding a mask column to an input ASCII or FITS table, or adding an mask extension to a FITS image. In both cases, the name of the column/extension has to be given by the fourth COLUMNS parameter.
- The resulting best-fit parameters of MOLECFIT are written in the BEST\_FIT\_PARAMETERS.fits file. In the case of a complex fit, the more reliable fit parameters could be taken from this file and used as (fixed) input for another iteration of the fitting procedure.
- Changing the fit parameters FTOL and XTOL (see Section 6.3) can significantly affect the code run time and the quality of the fit. In the case of unsatisfying fit results, it may be an option to change the default values. However, the effect is often unpredictable, since more relaxed convergence criteria can lead to worse as well as better fit quality.
- To achieve an optimal performance of the code, one should fit only those molecules that significantly contribute to the wavelength range of the fit. We suggest to base the selection of relevant molecules on the information given in Section 8.3.
- In principle, PWV values can be measured by means of all kinds of spectra of bright standard stars which show telluric lines in absorption and atmospheric emission spectra in the thermal IR. However, a good fit requires significant H<sub>2</sub>O features. This criterion cannot be fulfilled if the water lines are very weak as in the optical. Moreover, too low resolution can smooth out the crucial lines, which can make the fit very unstable or even impossible.
- For MOLECFIT applications aiming at the derivation of the atmospheric water vapour content or the telluric absorption correction of astronomical spectra, it is often sufficient to set the abundances of other molecules to a fixed value. For the more frequent molecules in the atmosphere (see Section 8.3), the column density from the input standard profile, i.e. RELCOL = 1, is usually relatively close to the true value. It has to be taken into account that the MIPAS standard profiles were created in 2001 (see Section 8.1.1), which causes deviations for greenhouse gases that indicate a significant long-term increase in atmospheric abundance. For example, the global CO<sub>2</sub> concentration increased by about 6% in one decade, which suggests RELCOL = 1.06.
- The user can explicitly set the initial value of the constant term of the polynomials for the continuum correction and the wavelength solution. The higher-order coefficients are automatically set to reasonable start values if required (see Section 6.3). Setting WLC\_CONST is only recommended if a wavelength shift towards a certain direction is expected. The CONT\_CONST is more critical. If the continuum level of the input spectrum strongly deviates from 1, even after setting the scaling factor FLUX\_UNIT (see Section 6.3), it is prudent to adapt this term.
- For a correct wavelength fit especially at high resolution, it is important to have a correct setting of the VAC\_AIR parameter. The wavelength system depends on the instrument and the wavelength calibration



approach by the data reduction pipeline. IR data (e.g. CRIRES, VISIR) tend to be provided in vacuum wavelengths, whereas data at shorter wavelengths tend to be provided in air wavelengths (e.g. UVES, X-shooter). If the user does not know this necessary input, it can be easily derived by running MOLECFIT with both VAC\_AIR options.

- For the instrumental profile created by the convolution of boxcar, Gaussian, and Lorentzian kernels (see Section 8.5), a derivation from the fit can be difficult. If the  $\chi^2$  degeneration by the kernel parameters causes a bad fit, we recommend to modify the initial values of these parameters or to fix the width of a kernel element. Some knowledge on the true functional form of the instrumental profile can be quite helpful. For a first test, the width of the Lorentzian might be fixed. For CRIRES, a value of 0.75 plus a large kernel size could be reasonable (A. Smette 2012, priv. comm.). For our tests, we used 0.5 pixels for all instruments, although X-shooter spectra do not indicate a significant contribution of a Lorentzian. The boxcar kernel can probably be neglected if the slit width does not have a significant impact on the line FWHM. If the user has access to a suitable line profile kernel, it can be imported via the parameter KERNEL\_FILE (see Section 8.5.3). The three-component profile fitting is switched off in this case.
- The run time of the radiative transfer code LBLRTM depends on the width of the fitted wavelength ranges. For a better performance, it is, therefore, recommended to use fit ranges as narrow as possible. It also makes the polynomial continuum fit more reliable. For e.g. X-shooter, several representative ranges over the entire wavelength range covering lines of all critical molecules could be defined. A suitable bin size is 10 nm. CRIRES spectra are sufficiently narrow that the full wavelength range can be used for the fitting procedure.
- For spectra covering a wide wavelength range and narrow fit ranges, the degree of the polynomial for the wavelength solution WLC\_N should be set to 0 to avoid unpredictable wavelength corrections outside the fit ranges if a telluric absorption correction is required. For X-shooter, it is also strongly recommended to set VARKERN = 1 in order to consider the change of the instrumental profile with wavelength (see Section 6.3).
- For telluric absorption correction in the thermal IR, it could be advantageous to use the strong sky emission extracted as 1D spectrum as input for molecfit\_model, molecfit\_calctrans will then calculate the transmission function belonging to the best-fit sky radiance spectrum. Finally, the object spectrum can be corrected for telluric absorption via molecfit\_correct.



# A Expert Fitting

The spectral domain of the input data may be split into several ranges either by user declaration or by physical specifications of multi chip detectors. The expert mode offers greater access to the molecfit fitting parameters with regards these ranges and allows for more flexible iterative fitting. On the other hand, it increases the complexity of the fitting procedure and there is a chance of erroneous results if the masked files are changed and the range and chip related parameters are not carefully adapted. For this reason, it has not been implemented as standard mode for molecfit.

### A.1 Mapping Multiple Extension Data

It is possible to represent several science spectra to be processed in mutiple extensions of a single fits file. When the parameter "USE\_ONLY\_INPUT\_PRIMARY\_DATA" is set to false, molecfit will assume that all extensions in the input science data fit contain spectra data to be processed. When the "CHIP\_EXTENSIONS" flag is set to false it is assumed that they are to be processed independently of each other. In this format, the output files will have multiple extensions each corresponding to the extension index of the input science frame. For example consider the following science frame, CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits, that contains science spectra in extensions 1 and 2, but not in the primary:

| Index | Extension | Туре         | Dimension | Comment                             |
|-------|-----------|--------------|-----------|-------------------------------------|
| 0     | Primary   | Image        | 0         |                                     |
| 1     | SCIENCE1  | Binary table | 9x1024    | Contains the first science spectra  |
| 2     | SCIENCE2  | Binary table | 9x1024    | Contains the second science spectra |

 Table A.1: Table showing the summary for CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits

The molecfit\_model recipe would then generate derivative files with two extensions: SCIENCE1 and SCI-ENCE2. For example, the BEST\_FIT\_PARAMETER.fits file would consist of the following:

| Table A.2: Table showing the summary for BEST_ | FIT PARAMETERS.fits |
|--|---------------------|
| Tuble ALL Tuble one and a build of been        |                     |

| Index | Extension | Туре   | Dimension | Comment                                     |
|-------|-----------|--------|-----------|---|
| 0     | Primary   | Image  | 0         |   |
| 1     | SCIENCE1  | Binary | 3x30      | Contains the fitted parameters for SCIENCE1 |
| 2     | SCIENCE2  | Binary | 3x30      | Contains the fitted parameters for SCIENCE2 |

When dealing with multiple extension input and multiple extension parameters there may arise a need to specify which extension parameters are to be applied to which science extension input. In particular when applying a specific telluric correction that has already been generated for a set of science spectra. For this purpose the following mapping parameters are available:



| Table A.3:   | Table showing th | ne mapping | parameters |
|--------------|------------------|------------|------------|
| 14010 / 1101 | rabio onoming a  | io mapping | paramotoro |

| Parameter Name           | Maps from Extension of: | Maps to Extension of:                |
|--------------------------|-------------------------|--------------------------------------|
| MODEL_MAPPING_KERNEL     | MOLECFIT_SCIENCE        | MOLECFIT_MODEL_KERNEL_LIBRARY        |
| CALCTRANS_MAPPING_KERNEL | MOLECFIT_SCIENCE        | MOLECFIT_CALCTRANS_KERNEL_LIBRARY    |
| MAPPING_ATMOSPHERIC      | MOLECFIT_SCIENCE        | MOLECFIT_ATM_PARAMETERS              |
| MAPPING_CONVOLVE         | MOLECFIT_SCIENCE        | BEST_FIT_PARAMETERS                  |
|                          |                         | and                                  |
|                          |                         | ATM_PARAMETERS                       |
|                          |                         | (process will create an extension in |
|                          |                         | MOLECFIT_TELLURIC_CORR               |
|                          |                         | in line with the                     |
|                          |                         | MOLECFIT_SCIENCE extension)          |
| MAPPING_CORRECT          | MOLECFIT_SCIENCE        | MOLECFIT_TELLURIC_CORR               |

The mapping parameters are defined as a string of comma separated extension indices that lists, in order of the primary (index=0) and extensions (index>0) of the input science fits file, the extension number of the associated file they are to be mapped too. For example assume that the science frame contains data in extensions 1,2 and 3, then the identity mapping would be "0,1,2,3", that is Primary is mapped to Primary, extension 1 is mapped to extension 1, extension 2 is mapped to extension 2 and extension 3 is mapped to extension 4. Similarly, a mapping of "2,2,2,2" implies use extension 2 of the associated file for all the input science extensions.

It is further possible to define these extension mappings in separate fits files, by setting the above parameters to NULL and declaring the fits files in the SOF file with the associated tagnames which is the same as the parameter name. Those fits files must then contain the index list in a single column binary table. The name of the column must be the name of the parameter with the suffix "\_EXT". For example, if parameter MODEL\_MAPPING\_KERNEL is set to "NULL" then the fits file specified in the SOF file with tag MODEL\_MAPPING\_KERNEL must have a single column binary table with the column name as MODEL\_MAPPING\_KERNEL\_EXT.

### A.2 Mapping Ranges to Chips

With the definition of multiple ranges and multiple chips some with overlapping wavelength ranges, ambiguity may arise as to which chip a range is to be associated with. To prevent such ambiguity there is a mapping parameter MAP\_REGIONS\_TO\_CHIP which is a string of comma separated chip indices that lists, in order of the range indices. If this string is set as "NULL" it is then assumed that there is a fits table assigned to specify the wavelength inclusion ranges and that there is a column "MAPPED\_TO\_CHIP" from which the mappings are declared.

### A.3 Expert Range Specific Continuum Modelling

The default when fitting for continuum is to model the continuum for each region as an independent polynomial representation of an order which is common for all regions. The fitting options can be extended to specify a sub-



selection of regions to fit for only and for each region specify a unique polynomial order for model representation. The following parameters define these range specifc values:

### CONTINUUM\_N:

A list of integers specifying the polynomial order for each range. For example CONTIUUM\_N="0,3,0,1" implies that the polynomial order for ranges 1,2,3,4 are 0, 3, 0 and 1 respectively. If the string is set to "NULL" then it is assumed that there is a fits table assigned to specify the wavelength inclusion ranges and that there is a column "CONT\_POLY\_ORDER" from which the orders are declared.

### FIT\_CONTINUUM:

A list of integers (0 or 1) specifying whether a range is to be fitted for or not. For example FIT\_CONTIUUM="0,1,0,1" implies that the ranges 2 and 4 are to be fitted for but ranges 1 and 3 are not. If the string is set to "NULL" then it is assumed that there is a fits table assigned to specify the wavelength inclusion ranges and that there is a column "CONT\_FIT\_FLAG" from which the flags are declared.

### A.4 Expert Chip Specific Wavelength Correcting

The default when fitting for wavelength corrections is to model the pixel specific corrections for each chip as an independent polynomial of an order which is common for all chips. As with expert continuum modelling the fitting options can be extended to specify a sub-selection of the ranges associated with each chip to include in the fittings. Note, unlike expert continuum modelling there is no means to uniquely specify a polynomial order for each chip. Further, a chip is not explicitly declared as to be fitted for or not. Rather this is ascertained by whether at least one range associated with that chip has been flagged for fitting. The ranges are flagged with the following parameter.

### FIT\_WLC:

A list of integers (0 or 1) specifying whether a range is to be included in a wavelength correction of the chip that that range is assigned to. For example FIT\_WLC="0,1,0,1" implies that the ranges 2 and 4 are to be included in such fittings but ranges 1 and 3 are not. If the string is set to "NULL" then it is assumed that there is a fits table assigned to specify the wavelength inclusion ranges and that there is a column "WLC\_FIT\_FLAG" from which the flags are declared.

### A.5 Expert Mode Initial Fitting Values

As with any non-linear fitting algorithm, molecfit requires initial values (guess values) as a starting point for the fitting predure. These values can have a strong impact on both the run-time and the successful conclusion of the fitting. The closer the initial values are to the correct values the greater the chance of success and the shorter the run-time and vice versa. With regards the polynomial coefficients for continuum modelling and wavelength correction, the default parameters: CONTINUUM\_CONST and WLC\_CONST only allow the initial definition for these polynomial as a simple shift. The molecfit Expert Mode allows initial values for all the coefficients of all the polynomials being fitted for to be read from a binary table of fits a file. The accepted format of such a table includes the format as rendered in the BEST\_FITS\_PARAMETER. fits file allowing the user to select the results from a previous fitting as the starting point of a new fitting.



To use this expert mode the .rc file must include the optional parameter EXPERT\_MODE set to TRUE and the .sof file must include the pathname for an initial values fits file with tagname "INIT\_FIT\_PARAMETERS". The expected format of the fits file is to contain a binary table with at least two columns: "parameter" and "value". The expert mode will override default initial values from the values in this table associated with parameter names that match expected format (see table below).

| Parameter Name String Format:         | Case Sensitive: | Examples:               |
|---------------------------------------|-----------------|-------------------------|
| "boxfwhm"                             | YES             | boxfwhm                 |
| "gaussfwhm"                           | YES             | gaussfwhm               |
| "lorentzfwhm"                         | YES             | lorentzfwhm             |
| rel_mol_col_[MOLECULE NAME]           | YES             | rel_mol_col_H2O         |
|                                       |                 | rel_mol_col_O3          |
| "*RANGE [range idx]*COEF [coef idx]*" | NO              | Range 1, Coef 0         |
|                                       |                 | Range 1, chip 1, coef 0 |
|                                       |                 | range 1 coef 0          |
| "*CHIP [chip idx]*COEF [coef idx]*"   | NO              | Chip 2, Coef 1          |
|                                       |                 | Chip 2, Coef 1          |

The expert mode will only utilise coefficient values found that are associated with the list of coefficients that have been assigned to any of the previously declared chips or ranges. Any coefficient that is not included in the fits table will be given the value 0.0.

### A.6 Expert Analysis of the BEST\_FIT\_PARAMETERS.fits FILE

Whilst the BEST\_FIT\_PARAMETER.fits file is generated by molecfit\_model for use as input for the molecfit\_calctrans recipe, the expert user may look at the contents of this file to determine if the parameter fitting process of molecfit\_model has progressed as expected. For such individuals we highlight the structure of this file.

The file consists of a single binary table with three columns labelled: parameter; value; and uncertainty. The parameter column and the value column contains the name and value/final fitted value of each parameter involved within the process. This includes parameters that were chosen to be fitted, parameters that were chosen to be fixed and those parameters for which fitting is not applicable, eg the number of data points. The uncertainty column shows: a value of -1 for those parameters for which fitting is not applicable; 0.0 for those parameters set as a fixed value; and the estimated error of fit for those actually fitted.

For the most part the parameter names listed will make sense, but a few need further clarification.

### A.6.1 The value of the Status variable

Status is the variable that represents the integer output 'info' from the mpfit function. If the user has terminated execution, info is set to the (negative) value of iflag. see description of fcn. otherwise, info is set as follows.



info = 0 improper input parameters.

info = 1 both actual and predicted relative reductions in the sum of squares are at most ftol.

info = 2 relative error between two consecutive iterates is at most xtol.

info = 3 conditions for info = 1 and info = 2 both hold.

info = 4 the cosine of the angle between fvec and any column of the jacobian is at most gtol in absolute value.

info = 5 number of calls to fcn has reached or exceeded maxfev.

info = 6 ftol is too small. no further reduction in the sum of squares is possible.

info = 7 xtol is too small. no further improvement in the approximate solution x is possible.

info = 8 gtol is too small. fvec is orthogonal to the columns of the jacobian to machine precision.

#### A.6.2 Wavelength correction

Coefficients for the polynomial fittings for range specific continuum models and the chip specific wavelength corrections:

The wavelength correction is defined as a Chebyshev polynomial correction to the given wavelength calibrations of the chip pixels. For each coefficient of each chip, the parameter name is given in the format: Chip [chip idx], coeff [coef idx]

where *coef idx* is the coefficient associated with the Chebyshev function of that order. For example, if the wavelength correction function for chip 2 is a quadratic, then the three associated coefficient parameters would be labeled:

Chip 2, coef 0 Chip 2, coef 1 Chip 2, coef 2

and x represents the wavelength normalized to the whole spectral range of the chip.

If we symbolise these coefficient parameters as  $c_0$ ,  $c_1$  and  $c_2$  respectively, then the correction function is defined as

$$c_0 T_0(x) + c_1 T_1(x) + c_2 T_2(x)$$
(8)

where  $T_0$ ,  $T_1$  and  $T_2$  are the first three Chebyshev functions

$$T_o(x) = 1$$
  
 $T_1(x) = x$  (9)  
 $T_2(x) = 2x^2 - 1$ 

It is important to note that the identity correction (i.e. no correction) is defined as  $c_0 = 0$ ,  $c_1 = 1$  and  $c_i = 0$  for all  $i \neq 1$ .



Thus to model the correction function as a shift to be fitted would be to fit a value for  $c_0$  and set  $c_1$  as fixed with the value 1. This maybe confusing as such a fit is declared by specifying that the polynomial order be zero when in fact it will be of order one by only one coefficient is actually fitted.

### A.6.3 Continuum

The situation with the continuum ranges is similar except that standard polynomials are used instead of Chebyshev, and that the parameter name is given by Range [range idx], chip [chip idx] coef [coef idx]

and that the polynomial order of fit can be unique per range. The storage of the range coefficients is that of a rectangular table of size the number of ranges by the maximum polynomial order specified. Coefficients that are not used are fixed at 0.0 and will be displayed in the parameter column with value 0.0 and uncertainty 0.0. Note that the spectral range of the chip is normalized to [-1,1]. For a constant shift (i.e.,  $c_1 = 1$ ,  $c_i = 0$  for i > 1), the shift of wavelength is given by [ $c_0 (\lambda_{max} - \lambda_{min}) + (\lambda_{max} + \lambda_{min})$ ]/2, where  $\lambda_{min}$  and  $\lambda_{max}$  are the minimum and maximum wavelengths of the spectral range, respectively.

### A.7 Scaling the transmission function to the PWV of the science spectrum

In some cases, it is advantageous to execute molecfit\_model on a telluric star spectrum and molecfit\_calctrans on a science spectrum. It may happen that the amount of precipitable water vapour (PWV) has changed significantly between the observing time of the two spectra.

Therefore, since version 4.3, the molecfit\_calctrans recipe includes the capability to scale the transmission function to the PWV of the science spectrum as provided as input to molecfit\_calctrans. The PWV of the science spectrum may be specified via a numerical value or via values located in header keywords of the science product. This is controlled via the SCALE\_PWV parameter which must be set to enable the scaling (see Tab. 6.2). The H2O column of the ATM\_PARAMETERS table is then scaled by the ratio of the PWV values. The FITS headers of the molecfit\_calctrans output products will then contain the additional keywords 'ESO DRS PWV SCI' (Value for the science spectrum), 'ESO DRS PWV TELL' (Best fit from telluric spectrum) and their ratio 'ESO DRS PWV RATIO' (the value of 'ESO DRS PWV SCI' divided by 'ESO DRS PWV TELL').

In addition to these header keywords, the molecfit\_calctrans output products will also contain the header keywords 'ESO DRS PWV DELTA MJD' and 'ESO DRS PWV DELTA AIRM', providing the difference in Modified Julian Date and Airmass between the science spectrum and telluric spectrum. These are helpful in order to keep track of the time and airmass that has elapsed between the spectra.

In order for these keywords to be meaningful, the appropriate header keywords must be specified via the molecfit\_calctrans parameters HDR\_MJD, HDR\_EXP, HDR\_AIR1 and HDR\_AIR2 (see Tab. 6.2). These header keywords are read directly from the science spectrum, but for the telluric spectrum they are read from a copy of the telluric header that is copied to BEST\_FIT\_PARAMETERS by molecfit\_model. For these values to be successfully read, the latest version of molecfit\_model must have produced the BEST\_FIT\_PARAMETERS table. If SCALE\_PWV is not the default ('none'), then an error will be raised if any of these HDR\_\* keywords are missing. An error does not occur in the default case, but a warning may instead be given.

The success of the PWV scaling depends on the fact that the exposure time of the telluric used by molecfit\_model is short, so that the airmass does not change much during the exposure. Hence the airmass



mean value is negligibly different from its median value.

*Caveat:* Currently the PWV scaling works in the simplest case for single extension datasets. Users trying to process more complex multiple extension datasets (e.g. KMOS) may encounter difficulties, in which case it is recommended to leave SCALE\_PWV as the default 'none'. The next molecfit release will include a fix for this issue.

### A.8 Quality of the correction parameter QC MEAN\_ABS\_DEV MAX

In previous versions of molecfit\_calctrans it was not possible to determine how well the airmass-scaled transmission matches the science spectrum. Since version 4.3, an algorithm has now been incorporated into molecfit\_calctrans that determines the quality of the correction. The corrected spectrum is smoothed by running a Savitzky-Golay filter of width SGWL pixels (see Tab. 6.2). The corrected spectrum is divided by the smoothed spectrum in an attempt to remove slopes and other intrinsic features in the spectrum. The normalized result is then subtracted from 1.0 and a mean absolute deviation of the resulting spectrum is calculated (excluding NaN values over the included regions). The value of this quantity is reported in the keyword 'ESO QC MEAN\_ABS\_DEV MAX' of the products of molecfit\_calctrans. The value of this keyword may be compared between different output products of molecfit\_calctrans in order to gauge the quality of the fit as determined by various other parameters.

The optional parameter SGWL\_ASMAX (see Tab. 6.2) instructs the above algorithm to be run multiple times for a variety of window lengths, namely from SGWL down to a minimum of 5 at decrements of 2 pixels. In this case the value of 'ESO QC MEAN\_ABS\_DEV MAX' will represent the maximum over this interval. Additional header keywords include 'ESO QC MEAN\_ABS\_DEV MAX WIDTH', representing the width of the Savitzky-Golay filter at the maximum which the 'ESO QC MEAN\_ABS\_DEV MAX' value was determined, and 'ESO QC MEAN\_ABS\_DEV MAX' value of SGWL.



## **B** Parameters

### **B.1 Molecfit Parameter Name Changes:** MOLECFIT\_MODEL

**Table B.1:** Table showing how the parameter names for MOLECFIT\_MODEL have changed between the original MOLECFIT (4.2.x) and the current version (4.4.2)

| MOLECFIT 4.4.2+             | Туре               | MOLECFIT 4.2.x |
|-----------------------------|--------------------|----------------|
| USE_ONLY_INPUT_PRIMARY_DATA | BOOL               | -              |
| USE_DATA_EXTENSION_AS_DFLUX | INT (0 implies no) | _              |
| USE_DATA_EXTENSION_AS_MASK  | INT (0 implies no) | -              |
| USE_INPUT_KERNEL            | BOOL               | -              |
| MODEL_MAPPING_KERNEL        | STRING             | KERNEL_FILE    |
| LIST_MOLEC                  | STRING             | LIST_MOLEC     |
| FIT_MOLEC                   | STRING             | FIT_MOLEC      |
| REL_COL                     | STRING             | RELCOL         |
| WAVE_INCLUDE                | STRING             | WRANGE_INCLUDE |
| WAVE_EXCLUDE                | STRING             | WRANGE_EXCLUDE |
| PIXEL_EXCLUDE               | STRING             | PRANGE_EXCLUDE |
| TELLURICCORR_PATH           | STRING             | -              |
| TELLURICCORR_DATA_PATH      | STRING             | -              |
| TMP_PATH                    | STRING             | -              |
| OPENMP_THREADS              | INT                | -              |
| SILENT_EXTERNAL_BINS        | BOOL               | -              |
| TRANSMISSION                | BOOL               | TRANS          |
| COLUMN_LAMBDA               | STRING             | COL_LAM        |
| COLUMN_FLUX                 | STRING             | COL_FLUX       |
| COLUMN_DFLUX                | STRING             | COL_DFLUX      |
| COLUMN_MASK                 | STRING             | COL_MASK       |
| MASK_BINARY                 | BOOL               | -              |
| DEFAULT_ERROR               | DOUBLE             | DEFAULT_ERROR  |
| WLG_TO_MICRON               | DOUBLE             | WLGTOMICRON    |
| WAVELENGTH_FRAME            | STRING             | VAC_AIR        |
| ( has value VAC or AIR)     |                    |                |
| CLEAN_MODEL_FLUX            | BOOL               | CLEAN_MFLUX    |
| FTOL                        | DOUBLE             | FTOL           |
| XTOL                        | DOUBLE             | XTOL           |
| FLUX_UNIT                   | DOUBLE             | FLUX_UNIT      |
| FIT_TELESCOPE_BACKGROUND    | BOOL               | TELBACK        |
| TELESCOPE_BACKGROUND_CONST  | DOUBLE             | TELBACK        |
| FIT_CONTINUUM               | BOOL               | FIT_CONT       |
| CONTINUUM_N                 | INT                | CONT_N         |
| CONTINUUM_CONST             | DOUBLE             | CONT_CONST     |
| FIT_WLC                     | BOOL               | FIT_WLC        |



#### continuation of Table B.1

| MOLECFIT 4.4.2+            | Туре   | MOLECFIT 4.2.x  |
|----------------------------|--------|-----------------|
| WLC_N                      | INT    | FIT_WLC         |
| WLC_CONST                  | DOUBLE | WLC_CONST       |
| FIT_RES_BOX                | BOOL   | FIT_RES_BOX     |
| RES_BOX                    | DOUBLE | RELRES_BOX      |
| FIT_RES_GAUSS              | BOOL   | FIT_RES_GAUSS   |
| RES_GAUSS                  | DOUBLE | RES_GAUSS       |
| FIT_RES_LORENTZ            | BOOL   | FIT_RES_LORENTZ |
| RES_LORENTZ                | DOUBLE | RES_LORENTZ     |
| KERNMODE                   | BOOL   | KERNMODE        |
| KERNFAC                    | DOUBLE | KERNFAC         |
| VARKERN                    | BOOL   | VARKERN         |
| OBSERVING_DATE_KEYWORD     | STRING | OBSDATE_KEY     |
| OBSERVING_DATE_VALUE       | INT    | OBSDATE         |
| UTC_KEYWORD                | STRING | UTC_KEY         |
| UTC_VALUE                  | DOUBLE | UTC             |
| TELESCOPE_ANGLE_KEYWORD    | STRING | TELALT_KEY      |
| TELESCOPE_ANGLE_VALUE      | DOUBLE | TELALT          |
| RELATIVE_HUMIDITY_KEYWORD  | STRING | RHUM_KEY        |
| RELATIVE_HUMIDITY_VALUE    | DOUBLE | RHUM            |
| PRESSURE_KEYWORD           | STRING | PRES_KEY        |
| PRESSURE_VALUE             | DOUBLE | PRES            |
| TEMPERATURE_KEYWORD        | STRING | TEMP_KEY        |
| TEMPERATURE_VALUE          | DOUBLE | TEMP            |
| MIRROR_TEMPERATURE_KEYWORD | STRING | M1TEMP_KEY      |
| MIRROR_TEMPERATURE_VALUE   | DOUBLE | M1TEMP          |
| ELEVATION_KEYWORD          | STRING | GEOELEV_KEY     |
| ELEVATION_VALUE            | DOUBLE | GEOELEV         |
| LONGITUDE_KEYWORD          | STRING | LONGITUDE_KEY   |
| LONGITUDE_VALUE            | DOUBLE | LONGITUDE_      |
| LATITUDE_KEYWORD           | STRING | LATITUDE_KEY    |
| LATITUDE_VALUE             | DOUBLE | LATITUDE        |
| SLIT_WIDTH_KEYWORD         | STRING | SLITW_KEY       |
| SLIT_WIDTH_VALUE           | DOUBLE | SLITW           |
| PIX_SCALE_KEYWORD          | STRING | PIXSC_KEY       |
| PIX_SCALE_VALUE            | DOUBLE | PIXSC           |
| REFERENCE_ATMOSPHERIC      | STRING | REF_ATM         |
| GDAS_PROFILE               | STRING | GDAS_PROF       |
| LAYERS                     | BOOL   | LAYERS          |
| EMIX                       | DOUBLE | EMIX            |
| PWV                        | DOUBLE | PWV             |
| LNFL_LINE_DB               |        | -               |
| LNFL_LINE_DB_FORMAT        |        | -               |



#### continuation of Table B.1

| MOLECFIT 4.4.2+ | Туре | MOLECFIT 4.2.x |
|-----------------|------|----------------|
| LBLRTM_ICNTNM   |      | -              |
| LBLRTM_IAERSL   |      | -              |
| LBLRTM_MPTS     |      | -              |
| LBLRTM_NPTS     |      | -              |
| LBLRTM_V1       |      | -              |
| LBLRTM_V2       |      | -              |
| LBLRTM_SAMPLE   |      | -              |
| LBLRTM_ALFALO   |      | -              |
| LBLRTM_AVMASS   |      | -              |
| LBLRTM_DPTMIN   |      | -              |
| LBLRTM_DPTFAC   |      | -              |
| LBLRTM_TBOUND   |      | -              |
| LBLRTM_SREMIS1  |      | -              |
| LBLRTM_SREMIS2  |      | -              |
| LBLRTM_SREMIS3  |      | -              |
| LBLRTM_SRREFL1  |      | _              |
| LBLRTM_SRREFL2  |      | _              |
| LBLRTM_SRREFL3  |      | _              |
| LBLRTM_MODEL    |      | _              |
| LBLRTM_ITYPE    |      | _              |
| LBLRTM_NOZERO   |      | _              |
| LBLRTM_NOPRNT   |      | _              |
| LBLRTM_IPUNCH   |      | _              |
| LBLRTM_RE       |      | _              |
| LBLRTM_HSPACE   |      | _              |
| LBLRTM_REF_LAT  |      | _              |
| LBLRTM_H1       |      | _              |
| LBLRTM_H2       |      | _              |
| LBLRTM_RANGE    |      | _              |
| LBLRTM_BETA     |      | _              |
| LBLRTM_LEN      |      | _              |
| LBLRTM_HOBS     |      | _              |
| LBLRTM_AVTRAT   |      | _              |
| LBLRTM_TDIFF1   |      | _              |
| LBLRTM_TDIFF2   |      |                |
| LBLRTM_ALTD1    |      | _              |
| LBLRTM_ALTD2    |      |                |
| LBLRTM_DELV     |      |                |
| CHIP_EXTENSIONS | BOOL | CHIP           |



### B.2 Molecfit Parameter Name Changes: MOLECFIT\_CALCTRANS

**Table B.2:** Table showing how the parameter names for MOLECFIT\_CALCTRANS have changed between the original MOLECFIT (4.2.x) and the current version (4.4.2)

| MOLECFIT 4.4.2+             | Туре               | MOLECFIT 4.2.x |
|-----------------------------|--------------------|----------------|
| USE_ONLY_INPUT_PRIMARY_DATA | BOOL               | -              |
| USE_DATA_EXTENSION_AS_DFLUX | INT (0 implies no) | -              |
| USE_DATA_EXTENSION_AS_MASK  | INT (0 implies no) | -              |
| USE_INPUT_KERNEL            | BOOL               | -              |
| CALCTRANS_MAPPING_KERNEL    | STRING             | KERNEL_FILE    |
| MAPPING_ATMOSPHERIC         | STRING             | -              |
| MAPPING_CONVOLVE            | STRING             | -              |
| CHIP_EXTENSIONS             | BOOL               | CHIP           |

### B.3 Molecfit Parameter Name Changes: MOLECFIT\_CORRECT

**Table B.3:** Table showing how the parameter names for MOLECFIT\_CORRECT have changed between the original MOLECFIT (4.2.x) and the current version (4.4.2)

| MOLECFIT 4.4.2+             | Туре               | MOLECFIT 4.2.x |
|-----------------------------|--------------------|----------------|
| USE_ONLY_INPUT_PRIMARY_DATA | BOOL               | -              |
| USE_DATA_EXTENSION_AS_DFLUX | INT (0 implies no) | -              |
| USE_DATA_EXTENSION_AS_MASK  | INT (0 implies no) | -              |
| SUPPRESS_EXTENSION          | BOOL               | -              |
| MAPPING_CORRECT             | STRING             | -              |
| CHIP_EXTENSIONS=FALSE       | BOOL               | CHIP           |



#### **B.4** molecfit\_model SOF **Tag names**

 Table B.4: Table showing the SOF Tagnames and their purpose as used in the MOLECFIT\_MODEL recipe.

| Tag Name             | Tag Purpose   |
|----------------------|---|
| SCIENCE              | INPUT (mandatory)   |
|                      | Contains the spectral data affected by telluric contamination.                                  |
| STD_MODEL            | INPUT   |
|                      | Alternative tag name for SCIENCE  |
| MOLECULES            | INPUT   |
|                      | Only used if recipe parameter, LIST_MOLEC, is set to NULL, in                                   |
|                      | which case this file must be provided.  |
|                      | Declares the LIST_MOLEC parameters in a binary table extension                                  |
|                      | with three columns: LIST_MOLEC, FIT_MOLEC, and REL_COL.   |
| WAVE_INCLUDE         | INPUT   |
|                      | Only used if recipe parameter, WAVE_INCLUDE, is set to NULL, but                                |
|                      | does not have to be provided.   |
|                      | Declares the WAVE_INCLUDE parameters in a binary table exten-                                   |
|                      | sion with two columns: LOWER_LIMIT, and UPPER_LIMIT   |
| WAVE_EXCLUDE         | INPUT   |
|                      | Only used if recipe parameter, WAVE_EXCLUDE, is set to NULL, but                                |
|                      | does not have to be provided.   |
|                      | Declares the WAVE_EXCLUDE parameters in a binary table exten-                                   |
|                      | sion with two columns: LOWER_LIMIT, and UPPER_LIMIT.  |
| PIXEL_EXCLUDE        |   |
|                      | Only used if recipe parameter, PIXEL_EXCLUDE, is set to NULL, but                               |
|                      | does not have to be provided.<br>Declares the PIXEL_EXCLUDE parameters in a binary table exten- |
|                      | sion with two columns: LOWER_LIMIT, and UPPER_LIMIT   |
| GDAS                 | INPUT & OUTPUT (optional)   |
| GDAD                 | User-supplied GDAS profile FITS file  |
| ATM_PROFILE_STANDARD | INPUT (optional)  |
|                      | User defined atmospheric standard profile.  |
| ATM_PROFILE_COMBINED | INPUT (optional).   |
|                      | User specified file that contains comprehensive grid data of mete-                              |
|                      | orological and molecular abundance information necessary to con-                                |
|                      | struct an atmospheric parameter set to use in the LBLRTM tools.                                 |
| KERNEL_LIBRARY       | Library of user provided kernels.   |
| MAPPING_KERNEL       | FITS binary table mapping the kernel in the library   |
|                      | KERNEL_LIBRARY with the SCIENCE spectrum.   |
| MODEL_KERNEL_LIBRARY | Alternative tag name for KERNEL_LIBRARY   |
| MODEL_MAPPING_KERNEL | Alternative tag name for MAPPING_KERNEL   |


#### B.5 molecfit\_calctrans SOF Tag Names

Table B.5: Table showing the SOF Tagnames and their puprose as used in the MOLECFIT\_CALCTRANS recipe

| Tag Name                 | Tag Purpose   |
|--------------------------|---|
| SCIENCE                  | INPUT (mandatory)   |
|                          | Contains the spectral data of interest with telluric contamination. |
| SCIENCE_CALCTRANS        | INPUT   |
|                          | Alternative tag name for SCIENCE                                    |
| MODEL_MOLECULES          | INPUT (mandatory)   |
|                          | Local copy of a FITS file that contains the LIST_MOLEC parameters   |
|                          | as used by the model recipe execution.                              |
| KERNEL_LIBRARY           | Library of user provided kernels.                                   |
| MAPPING_KERNEL           | FITS binary table mapping the kernel in the library KER-            |
|                          | NEL_LIBRARY with the SCIENCE spectrum.                              |
| CALCTRANS_KERNEL_LIBRARY | Alternative tag name for KERNEL_LIBRARY                             |
| CALCTRANS_MAPPING_KERNEL | Alternative tag name for MAPPING_KERNEL                             |
| MAPPING_ATMOSPHERIC      | FITS binary table mapping the parameters in MOLEC-                  |
|                          | FIT_ATM_PARAMETERS with the SCIENCE spectrum.                       |
| MAPPING_CONVOLVE         | FITS binary table mapping the parameters in both                    |
|                          | BEST_FIT_PARAMETERS and ATM_PARAMETERS with the                     |
|                          | SCIENCE spectrum.   |
| ATM_PARAMETERS           | INPUT (mandatory)   |
|                          | Atmospheric model file containing the necessary parameters to be    |
|                          | used in the LBLRTM tools.   |
| BEST_FIT_PARAMETERS      | INPUT (mandatory)   |
|                          | Parameter file containing the values (assumed derived by the        |
|                          | model recipe) fitted for.   |

#### B.6 molecfit\_correct SOF Tag Names

Table B.6: Table showing the SOF Tagnames and their puproses as used in the MOLECFIT\_CORRECT recipe

| Tag Name        | Tag Purpose   |              |          |           |        |         |      |        |
|-----------------|---|--------------|----------|-----------|--------|---------|------|--------|
| SCIENCE         | INPUT   | (mandat      | ory)     |           |        |         |      |        |
|                 | Contains the spectral data of interest with telluric contamination. |              |          |           |        |         |      |        |
| MAPPING_CORRECT | FITS  | binary       | table    | mapping   | the    | kernel  | in   | MOLEC- |
|                 | FIT_TELLURIC_CORR with the SCIENCE spectrum.                        |              |          |           |        |         |      |        |
| TELLURIC_CORR   | Telluric correction data.   |              |          |           |        |         |      |        |
| TELLURIC_DATA   | Tellurio  | c correction | on data, | mandatory | if WLC | C_REF=M | IODE | L.     |



## **B.7 Molecfit Output Files**

#### B.7.1 molecfit\_model Output Files

| File name                | File purpose   |
|--------------------------|--|
| MODEL_MOLECULES.fits     | The list of molecules used in the model execution.                             |
|                          | This is either constructed from the LIST_MOLEC parameter sup-                  |
|                          | plied in the .rc file or is a local copy of the file specified by the          |
|                          | MOLECULES tag in the SOF file.   |
|                          | To be used as input for calctrans execution when specified in the              |
|                          | MODEL_MOLECULES tag of the calctrans related SOF file.                         |
| ATM_PARAMETERS.fits      | The list of atmospheric parameters used in the model execution.                |
|                          | To be used as input for calctrans execution when specified by                  |
|                          | the ATM_PARAMETERS tag of the calctrans related SOF file.                      |
| BEST_FIT_PARAMETERS.fits | The list of fitted parameter used derived in the model execution.              |
|                          | To be used as input for calctrans execution when specified by                  |
|                          | the BEST_FIT_PARAMETERS tag of the calctrans related SOF file.                 |
| BEST_FIT_MODEL.fits      | Spectra of the fitted model as derived in the model execution.                 |
| GDAS.fits                | The GDAS meteorological profile data used with MIPAS and EMM                   |
|                          | data to construct an atmospheric model.  |
|                          | Contains grid data for pressure height temperature and relative hu-            |
|                          | midity at a physical range around the required location at the time            |
|                          | of observation.  |
|                          | Note this is either an interpolation from GDAS_BEFORE.fits and                 |
|                          | GDAS_AFTER.fits or is a local copy of the profile file specified in            |
|                          | the relevant SOF file with the GDAS tagname.                                   |
| GDAS_BEFORE.fits         | GDAS meteorological profile data from a time prior to the science              |
|                          | frame capture time. Used with GDAS_AFTER.fits to interpolate                   |
|                          | GDAS.fits  |
|                          | Note this is either a copy extracted from the NOAA packaged data <sup>14</sup> |
|                          | or is a local copy of the profile file specified in the relevant SOF file      |
|                          | with the GDAS_BEFORE tagname.  |
| GDAS_AFTER.fits          | GDAS meteorological profile data from a time after the science                 |
|                          | frame capture time. Used with GDAS_BEFORE.fits to interpolate                  |
|                          | GDAS.fits  |
|                          | Note this is either a copy extracted from the NOAA packaged data <sup>14</sup> |
|                          | or is a local copy of the profile file specified in the relevant SOF file      |
|                          | with the GDAS_AFTER tagname.   |



#### continuation of Table B.7

| File name                 | File purpose  |
|---------------------------|---|
| ATM_PROFILE_STANDARD.fits | MIPAS atmospheric composition data used with GDAS and EMM                     |
|                           | data to construct an atmospheric model.                                       |
|                           | Note this is either a copy extracted from the ESA packaged data <sup>15</sup> |
|                           | or is a local copy of the profile file specified in the relevant SOF file     |
|                           | with the ATM_PROFILE_STANDARD tagname.  |
| ATM_PROFILE_COMBINED.fits | The result of combining the GDAS, MIPAS and EMM data and                      |
|                           | represents the gridded atmospheric profile data to be used in the             |
|                           | LNFL and LBLRTM third-party tools. Required as input for the                  |
|                           | calctrans recipe.   |

#### **B.7.2** molecfit\_calctrans **Output Files**

 Table B.8: Table showing output files of molecfit\_calctrans and their purpose.

| File name           | File purpose   |
|---------------------|--|
| LBLRTM_RESULTS.fits |  |
| TELLURIC_DATA.fits  | Input spectrum in BINTABLE format  |
| TELLURIC_CORR.fits  | Contains the telluric correction transformation data to be used for convolution with the uncorrected spectrum. |

#### B.7.3 molecfit\_correct Output Files

Table B.9: Table showing output files of molecfit\_correct and their purpose

| File name                      | File purpose  |
|--------------------------------|---|
| SCIENCE_TELLURIC_CORR_[RFNAME] | The input science frame corrected for telluric contami- |
|                                | nation.   |

Where [RFNAME] is the name of the raw input science file - e.g. if the input science file was

CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits

Then the above output files would be

SCIENCE\_TELLURIC\_CORR\_CRIRES\_HighSNR\_Telluric\_input\_AM1p472.fits

 $<sup>^{14}</sup> Located in the installed file / \texttt{share/molecfit/data/profiles/gdas/gdas_profiles_C-70.4-24.6.tar.gz$ 

 $<sup>^{15}{\</sup>tt Located}$  within the installed subdirectory /share/molecfit/data/profiles/mipas



| Doc. Number:  | VLT-MAN-ESO- |  |  |
|---------------|--------------|--|--|
|               | 19550-5772   |  |  |
| Doc. Version: | 4.4.2        |  |  |
| Released on:  | 2025-02-11   |  |  |
| Page:         | 112 of 120   |  |  |

# C Molecular Spectra

The following figures provide the transmission spectra for specific molecules. Digital version of these atlases are available as FITS binary tables under the share/esopipes/datastatic/molecfit-<version>/



| Doc. Number:  | VLT-MAN-ESO- |  |  |
|---------------|--------------|--|--|
|               | 19550-5772   |  |  |
| Doc. Version: | 4.4.2        |  |  |
| Released on:  | 2025-02-11   |  |  |
| Page:         | 113 of 120   |  |  |



**Figure C.1:** *Influence of individual molecules as a function of wavelength:* this figure shows the relative importance of all molecules at a given wavelength, which exceed more than 5% of the total radiance (in red) or transmission (in blue).



| Doc. Number:  | VLT-MAN-ESO- |  |  |
|---------------|--------------|--|--|
|               | 19550-5772   |  |  |
| Doc. Version: | 4.4.2        |  |  |
| Released on:  | 2025-02-11   |  |  |
| Page:         | 114 of 120   |  |  |







| Doc. Number:  | VLT-MAN-ESO- |  |  |
|---------------|--------------|--|--|
|               | 19550-5772   |  |  |
| Doc. Version: | 4.4.2        |  |  |
| Released on:  | 2025-02-11   |  |  |
| Page:         | 115 of 120   |  |  |



Figure C.3: Influence of individual molecules as a function of wavelength: same as Figure C.1, but for the VISIR test data wavelength regime.



| Doc. Number:  | VLT-MAN-ESO- |  |  |
|---------------|--------------|--|--|
|               | 19550-5772   |  |  |
| Doc. Version: | 4.4.2        |  |  |
| Released on:  | 2025-02-11   |  |  |
| Page:         | 116 of 120   |  |  |







| Doc. Number:  | VLT-MAN-ESO- |
|---------------|--------------|
|               | 19550-5772   |
| Doc. Version: | 4.4.2        |
| Released on:  | 2025-02-11   |
| Page:         | 117 of 120   |



**Figure C.5:** *Influence of individual molecules as a function of wavelength:* same as Figure C.1, but for the X-Shooter VIS-arm test data wavelength regime.



## D Maintenance and License Issues

#### D.1 Maintenance

## D.1.1 Introduction

The package MOLECFIT relies on some external code and data, which might undergo a change due to development after the current release. Although it is not possible to foresee all possible modifications, some hints can be given to use MOLECFIT even with updates from the external sources at a later stage. This particularly applies to the radiative transfer code LNFL/LBLRTM, and the GDAS data. In this appendix, we summarise the issues to be taken into account for a successful usage of MOLECFIT with later versions of LNFL/LBLRTM, the line database, and the GDAS data.

## D.1.2 Radiative transfer code LNFL/LBLRTM

The radiative transfer code package LNFL/LBLRTM is developed by AER [RD25] and can directly be obtained from there. MOLECFIT is delivered with LNFL v3.2, LBLRTM v12.11, and the modified AER line database v3.8.1.2.

In principle, MOLECFIT should also work with later versions as its principle usage has been unchanged for several years. This means that MOLECFIT is expected to provide all functionality as long as it is retained. In particular, the principle usage of the TAPE<XX> files [RD28], and the format of the LBLRTM input file TAPE5 MUST remain unchanged. However, as LBLRTM is widely used in atmospheric research, we do not expect major changes in the near future. In the course of the former DR-, and later SM-projects, we used several versions without problems.

What might change is the installation procedure of the LNFL/LBLRTM codes. If a future release of MOLECFIT is provided with a newer version of the radiative transfer code, the install script might need to be modified accordingly. For that purpose, the developers from AER provide a README file in the tree of the sources. In order to maximise flexibility, MOLECFIT only searches for executables labelled lnfl and lblrtm in the <INST\_DIR>/bin/ directory. Therefore, a soft link to the actual binaries of LNFL and LBLRTM has to be provided by the installation script. To facilitate the modification in the script, the corresponding section is marked there.

We recommend the following procedure for the update of MOLECFIT to later versions of LNFL/LBLRTM:

- Updates of the LNFL/LBLRTM are usually once or twice per year (see Section "What's new" at [RD25]). Therefore, a quarterly check for new versions is sufficient.
- Download the latest version of LNFL/LBLRTM from [RD25]
- Make sure that the principle usage has not changed. Check the documents in the docs directory in the tree of the LBLRTM sources. There, you should find a FAQ document, the release notes, and a detailed HTML description of the TAPE5 input file for LBLRTM.
- Read the README files whether the installation procedure has changed. These files should be located in the src/build directory of LNFL and LBLRTM.



- Check whether there is also a line list update and follow the steps described in Section D.1.3.
- Adapt the installation script of MOLECFIT.
- Evaluate by means of the delivered examples.

## D.1.3 AER Line parameter list

Usually the line parameter list delivered with the LNFL/LBLRTM package is updated at the same time as the code. We recommend to use the version which is delivered together with the corresponding code version.

To integrate the new line list, simply unpack the tarball and copy the new line parameter file (usually labelled aer\_v\_<version> to the directory <INST\_DIR>/data/hitran/ of your MOLECFIT installation. This can be done also be done by modifying the install script, as the corresponding section is marked. The parameter LNFL\_LINE\_DB should be updated accordingly.

Note that it has turned out that at long wavelengths relevant for VISIR, MOLECFIT can be significantly slower if the recommended aer line list is used instead of the basic HITRAN 2008 database. Unfortunately, this line list is no more available at the HITRAN website [RD20], which now provides the 2016 version. The new version has not been tested in terms of performance in MOLECFIT. Hence, it cannot be guaranteed that it works at all. If you want to test an original HITRAN line list, you have to put it in the data/hitran/ folder and modify \_LINE\_DB in the lblrtm\_setup file. In addition, the parameter \_LINE\_DB\_FMT has to be changed from 100 to 160 because of a different file format.

The version of the database provided with version 4.3 of molecfit was built from version aer\_v\_3.8.1 (https://github.com/AER-RC/AER\_Line\_File/wiki/What's-New#aer-line-file) but modified by removing duplicated lines. In addition, as suggested by Romain Allart, the H2O lines at wavenumber = 16984.8471 cm<sup>-1</sup> (588.7608 nm) and wavenumber = 15437.8563 cm<sup>-1</sup> (647.7583 nm) have also been removed.

## D.1.4 GDAS data

In contrast to the radiative transfer code, which is updated once/twice per year, the GDAS profiles are updated permanently. MOLECFIT initially searches in a local tarball for the corresponding profiles (script <INST\_DIR>/bin/get\_gdas\_profiles.sh), and tries to download it from the web in the case they are not found locally. The latter is done by the script <INST\_DIR>/bin/extract\_grib.sh with the help of the GRIB software.

However, the incorporated download server<sup>16</sup> does not seem to be very stable and organised in a predictable way. It might happen, that the profiles are not found. Therefore data tarballs updated daily are available from <a href="https://ftp.eso.org/pub/dfs/pipelines/skytools/molecfit/gdas">https://ftp.eso.org/pub/dfs/pipelines/skytools/molecfit/gdas</a>. They can be downloaded and placed into the <INST\_DIR>/data/profiles/gdas folder in case the automatic update mechanism fails. Note that this directory needs to be write enabled to have the new version permanently available.

Additionally MOLECFIT contains the script used to produce the updated files for the website. It can also be used to update the local GDAS database. The script downloads the GDAS data archive from a different server<sup>17</sup>.

<sup>&</sup>lt;sup>16</sup>http://nomad3.ncep.noaa.gov/pub/gdas/rotating/

<sup>&</sup>lt;sup>17</sup>ftp://ftp.arl.noaa.gov/archives/gdas1/



The data there is organised on weekly basis, each stored in a single file labelled gdas1.<mmm><yy>.w#, being <mmm> the month (e.g. jan, feb, mar,...) and <yy> the year. The extension "w#" describes the number of the week of this month:

#=1 - days 1-7 of the month #=2 - days 8-14 #=3 - days 15-21 #=4 - days 22-28 #=5 - days 29 - rest of the month

For example, the file gdas1.apr07.w3 contains data of the third week of April 2007 (15th to 21st of April 2007). More information on the file structure is given here<sup>18</sup>.

These files provide the GDAS profiles of the world wide grid. Therefore, it is necessary to extract the profiles in the same way as described in Section 8.1. This can be achieved with the shell script update\_gdas\_db.sh. It downloads the data on monthly basis, and extracts the profiles as required by MOLECFIT, and adds them to the local database. However, this may take a while as the amount of data to be downloaded is large (up to 580 MB per week) and the server is fairly slow.

The shell script invokes a C-programme extract\_gdas\_profiles. Although, both tools are part of the molecfit package, they can be used independently of MOLECFIT to provide the possibility to update the GDAS archive on different machines. This might be useful as some disc space is required temporarily. If this is intended, both tools have to be installed in the following way on the target machine:

- create subfolders bin/ and data/downloads/paranal/ in the target directory on the target machine.
- Copy the shell script update\_gdas\_db.sh and the C source file extract\_gdas\_profiles.c to the bin/ folder in target directory on the target machine.
- There compile the C programme by invoking

gcc extract\_gdas\_profiles.c -lm -o extract\_gdas\_profiles

• Copy the tarball file gdas\_profiles\_C-70.4-24.6.tar.gz delivered with the MOLECFIT package to the directory <targetdir@targetmachine>/data/downloads/paranal/. This tarball is being updated.

To update the local GDAS database please follow these steps:

• The final release of MOLECFIT contains GDAS profiles up to March 2014. So, if this is your first update, invoke the update script with the parameter oct13:

cd <INST\_DIR>/

OR

<sup>&</sup>lt;sup>18</sup>http://www.ready.noaa.gov/gdas1.php



cd <targetdir@targetmachine>

(depending on your installation (see above)

• invoke the update script

bin/update\_gdas\_database.sh oct13 P

and proceed until the end of the last fully missing month, if necessary. Note: You only can download full months. The file data/paranal/last\_month.txt shows a history of past updates.

- Copy the resulting tarball data/downloads/paranal/gdas\_profiles\_C-70.4-24.6.tar.gz to the folder <INST\_DIR>/data/profiles/gdas/ in your MOLECFIT installation path.
- Optionally: delete the downloaded files in data/downloads/gdas/.

Some dates are missing. Please check the website<sup>19</sup> for more details. There are no GDAS data available before 1st of December 2004.

#### D.2 License issues

#### D.2.1 LNFL/LBLRTM

URL: http://rtweb.aer.com/lblrtm\_frame.html

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<sup>&</sup>lt;sup>19</sup>http://ready.arl.noaa.gov/archives.php



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**Principal References** Clough, S. A., M. W. Shephard, E. J. Mlawer, J. S. Delamere, M. J. Iacono, K. Cady-Pereira, S. Boukabara, and P. D. Brown, Atmospheric radiative transfer modeling: a summary of the AER codes, Short Communication, J. Quant. Spectrosc. Radiat. Transfer, 91, 233-244, 2005.

Clough, S.A., M.J. Iacono, and J.-L. Moncet, Line-by-line calculation of atmospheric fluxes and cooling rates: Application to water vapor. J. Geophys. Res., 97, 15761-15785, 1992.

#### D.2.2 cmpfit library

URL: http://www.physics.wisc.edu/ craigm/idl/cmpfit.html URL: http://cow.physics.wisc.edu/ craigm/idl/idl.html

DISCLAIMER MPFIT: A MINPACK-1 Least Squares Fitting Library in C

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#### D.2.3 HITRAN

**URL:** http://www.cfa.harvard.edu/hitran/ **URL**: http://www.cfa.harvard.edu/hitran/Updated/ref-table.pdf

**Principal references** Rothman et al., "The HITRAN 2008 molecular spectroscopic database", Journal of Quantitative Spectroscopy and Radiative Transfer, vol. 110, pp. 533-572 (2009)

#### D.2.4 GDAS

URL: http://ready.arl.noaa.gov/gdas1.php



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