KSPECTRUM version 1.0.5 User Manual

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Contents

1	\mathbf{Des}	scription of the code 3				
	1.1	Production of high-resolution spectrum, full control of the un-				
		certainty				
	1.2	Range of applications	6			
	1.3	Physical hypothesis	7			
		1.3.1 Sub-lorentzian profiles and CIA	7			
		1.3.2 Broadening by the rest of the gas	9			
		1.3.3 High-temperature transitions	9			
	1.4	A better respect of spectroscopy	10			
		1.4.1 (no) line selection \ldots \ldots \ldots \ldots \ldots \ldots	10			
		1.4.2 (no) truncation of lines and "continuums"	10			
	1.5	Parallel computing	11			
	1.6	The multi-pass approach	11			
	1.7	Estimation of remaining computation time	12			
	1.8	Not starting from scratch after a crash	13			
	1.9	Automatic tabulation of Lorentz and Voigt functions	13			

2	Installation and prerequisites	15					
	2.1 First thing to do	15					
	2.2 Tweaking the "Makefile"	16					
	2.3 Setting up array sizes	16					
	2.4 Compilation	16					
	2.5 Installation of MPICH	17					
	2.6 Installation of LBL databases	18					
3	Using the code 21						
	3.1 User input files	21					
	$3.1.1$ The "data.in" file \ldots \ldots \ldots \ldots	21					
	3.1.2 The "options.in" file	22					
	3.2 Other input files	25					
	3.2.1 Atmospheric composition	25					
	3.2.2 Special molecules	26					
	3.2.3 Narrowband intervals	26					
	3.3 Running KSPECTRUM using MPICH	26					
4	Results	28					
	4.1 Results files	28					
	4.2 More output files	28					
	4.3 What's next ?	29					
5	The fun stuff	30					
	5.1 Code license	30					
	5.2 Questions and Answers	30					
\mathbf{A}	Appendix - Validation	33					
	A.1 Low pressures / temperatures	33					
	A.2 High pressures / temperatures	33					

1 Description of the code

1.1 Production of high-resolution spectrum, full control of the uncertainty

KSPECTRUM has been designed to produce high-resolution spectrum of any gas mixture, in any thermodynamical conditions, from line-by-line databases. In practice, it is far from being the case: see sections 1.2 and 1.3, mostly because we have no idea of sub-lorentzian profiles and collision-induced absorption (CIA) in the general case. This code will be improved gradually when our knowledge of spectroscopy evolves. Anyway, the purpose of the present code is clearly NOT to produce spectrum that will be used in engineering applications (i.e. spatial missions). Rather, it is intended at producing reliable spectrum that may be used for subsequent radiative transfer analysis (production of radiative transfer parameterizations into planetary GCMs).

By "high-resolution" spectrum, we mean that the spectral resolution is high enough so that individual lines are well resolved. This point is discussed below. The main purpose of producing high-resolution spectrum, is the possibility to compute k-distribution data sets.

The main idea that lead to the development of KSPECTRUM is to produce a code that can compute high-resolution data for virtually any conditions, with a full control of the accuracy:

- Any value of $k_a(\nu)$ has to be computed with a specified relative error ϵ_1 .
- The maximum relative error made when considering a linear profile $k_a^*(\nu)$ between two computed values $k_a(\nu_i)$ and $k_a(\nu_{i+1})$ may also be specified: ϵ_2 (see Fig. 1).

Ideally, when computing the absorption coefficient $k_a(\nu)$ at a given wavenumber ν , the contribution of every line should be summed up:

$$k_{a}(\nu) = \sum_{l=1}^{N} k_{a,l}(\nu)$$
(1)

where $k_{a,l}(\nu)$ is the contribution of the N lines index l to $k_a(\nu)$. In practice, it is impossible to use such a method to compute $k_a(\nu)$ at high



Figure 1: Schematic representation of the computed $k_a(\nu_i)$ spectrum, and of the maximal error made when considering a linear variation $k_a^*(\nu)$ between two consecutive computed result points.

spectral resolution, because the order of magnitude of N is typically several hundreds of thousands, when not millions.

The first constraint of accuracy will be used to set up a faster algorithm: we start by discretizing the infrared spectrum in a given number N_b of "narrowband" intervals, whose spectral limits are known. Let us take the example of any spectral interval $[\nu_{min}, \nu_{max}]$. Each one of the N lines will be examined for this interval. For a great number of lines, $[\nu_{min}, \nu_{max}]$ will be located in the far wings, and therefore the values of $k_a(\nu_{min})$ and $k_a(\nu_{max})$ are not very different. If the relative error $\epsilon = \frac{|k_a(\nu_{max})-k_a(\nu_{min})|}{\min(k_a(\nu_{min}),k_a(\nu_{max}))}$ is lower than the specified value ϵ_1 , then a constant value (for instance $\frac{k_a(\nu_{min})+k_a(\nu_{max})}{2}$) can be taken as $k_{a,l}(\nu)$, for every value $\nu \in [\nu_{min}, \nu_{max}]$, with a total relative error over $k_a(\nu)$ that will be lower than ϵ_1 .

The second accuracy constraint (ϵ_2) will be used for optimizing the spectral discretization: schematically, narrowband intervals $[\nu_{min}, \nu_{max}]$ will be discretized in such a way that line centers are well described. Discretization steps will take greater values in line wings. The computation of the spectral grid uses a tabulation of the Lorentz and Voigt functions. The method is

explained below for a Lorentz line profile.

For an isolated Lorentz line, we have:

$$f_L = \frac{\gamma_L}{\gamma_L^2 + (\nu - \nu_c)^2}$$
(2)

with γ_L the Lorentz line width, and $\nu_c = \nu_0 + \delta P$ the (corrected) wavenumber at line center.

This function can be written as:

$$f_L = \frac{1}{\gamma_L} \frac{1}{1 + x^2}$$
(3)

with $x = \frac{\nu - \nu_c}{\gamma_L}$. Next, function $f(x) = \frac{1}{1+x^2}$ has been tabulated in order to determine a series of triplets $[x_1, x_2, x_3]$ such that for any value $x_0 \in [x_1, x_2]$, the maximum relative difference between f(x) and $f^*(x) = f(x_0) + \frac{f(x_3) - f(x_0)}{x_3 - x_0}(x - x_0)$ (defined for $x \in [x_0, x_3]$) never exceeds ϵ_2 . Figure 2 may help.



Figure 2: Purpose of tabulating function f(x): for any $x_0 \in [x_1, x_2]$, the maximum relative difference between f and f^* is lower than the imposed value ϵ_2 .

In practice, for the computation of the spectral grid: starting from a wavenumber value ν_i , the algorithm will have to determine the next grid

point ν_{i+1} (see Fig. 1). For every line l whose center wavenumber is within $[\nu_{min}, \nu_{max}]$, the code will have to identify the triplet $[x_{1,l}, x_{2,l}, x_{3,l}]$ so that $x_{i,l} = \frac{\nu_i - \nu_{c,l}}{\gamma_{L,l}} \in [x_{1,l}, x_{2,l}]$. We know that the position of the next grid point could safely be taken at $x_{3,l}$, if line l was alone. Taking the minimum value $\nu_{i+1} = \min(\nu_{i+1,l} = \gamma_{L,l}x_{3,l} + \nu_{c,l})$ (over all lines present in the interval) ensures that the relative difference between k_a and k_a^* never exceeds ϵ_2 over $[\nu_i, \nu_{i+1}]$.

The principle is exactly the same for the Voigt function $f_V(\nu) = \sqrt{\frac{\ln(2)}{\pi}} \frac{1}{\gamma_D} \frac{y}{\pi} \int_{-\infty}^{+\infty} \frac{e^{-t^2}}{y^2 + (x-t)^2} dt$, with $x = \sqrt{\ln(2)} \frac{\nu - \nu_c}{\gamma_D}$, $y = \sqrt{\ln(2)} \frac{\gamma_L}{\gamma_D}$, and γ_D the Doppler linewidth. This function depends on y, the ratio of Lorentz and Doppler linewidths. This function will therefore have to be tabulated as a function of x, but also as a function of y.

I really don't see the point in giving any more details. If anyone ever reach this point without falling asleep and reads this, please send me an email at: v.eymet@gmail.com; I have been writing codes and manuals for the last 8 years, and no one ever told me if it has been useful in any way.

A special mention should be made tough to the issue encountered for meshing the spectral grid between close strong lines. In the case ν_i is located between two strong lines, the value of ν_{i+1} computed by the above-described algorithm may be too far from ν_i to satisfy accuracy criterion over ϵ_2 , because of the strong overlap between the two lines. In this case, a special treatment should be applied for a correct discretization of wavenumbers.

1.2 Range of applications

KSPECTRUM is currently using the following LBL databases: HITRAN (2004 and 2008) [1, 2], HITEMP (2004) [3] (H_2O , CO_2 , CO and OH only) and CDSD [4] (CO_2 only). The code uses the HITRAN database for temperatures lower than a user-defined temperature "T_hitemp" (defined in the "data.in" file, see section /refpara:userfiles). For temperatures greater than this value, it lets the user chose between HITEMP or CDSD for CO_2 lines, and will automatically use HITEMP for H_2O , CO and OH transitions. The temperature upper limit of validity of the present code is therefore the same as the HITEMP database (about 1000K), except when computing the spectrum of a pure CO_2 atmosphere: the CDSD database is supposed to be accurate up to 3000K.

No physical limit is imposed for pressures. However, the upper limit

in terms of pressure is probably what is found at ground level on Venus: ground pressure is 92 bars, and ground temperature is 735K. At such levels of pressure and temperature, the atmosphere is in super-critical thermodynamic state, and can no longer be considered as a mixture of gas. Therefore KSPECTRUM should probably not be used for a pressure greater than 100atm. The lower pressure limit will be imposed by the code itself, and more specifically by the spectral discretization algorithm. It may crash for values of the pressure lower than 10^{-7} atm, because spectral lines are so narrow that wavenumber steps are really small ($\approx 10^{-7}cm^{-1}$). Besides the fact that the computation of the spectral grid may take a lifetime in these conditions, such small wavenumber steps may not be properly taken into account by the algorithm.

In terms of molecules, the HITRAN database takes into account the transitions of 42 (for the 2008 version) molecular species, along with their isotopes. The list of molecules can be found in the HITRAN documentation [1], or directly in the "data/molparam.txt" file.

Other limits may be imposed for specific gas mixtures. See section 1.3 for more details.

1.3 Physical hypothesis

This section presents how we manage the fact that there is a lack of available spectroscopic data for various aspects of the theory.

1.3.1 Sub-lorentzian profiles and CIA

The main issue is that we have very little knowledge about:

- The real line shape, for each molecule, at each wavenumber. We know that CO_2 lines have a sub-lorentzian behavior in line wings, which is why thermal infrared signals can be transmitted from ground level up to space in very specific near-IR spectral windows through the otherwise very thick atmosphere of Venus.
- Collision-induced absorption (CIA), that are mainly due to the fact that in high pressure and/or high temperature conditions, collisions between molecules temporarily create new molecular species, with their own energetic transitions.

Sub-lorentzian profiles have been studied since 1969: in [5], the Lorentz profile $f_L(\nu - \nu_0)$ is corrected by a function $\chi(\nu - \nu_0)$ that accounts for the sub-lorentzian nature of the lineshape. Function $\chi(\nu - \nu_0)$ is given for pure CO_2 and for mixtures of CO_2 and other species (N_2, He, Ar, O_2, H_2) in very specific spectral ranges, for various values of the temperature. In [6] and [7], function χ is given respectively for pure CO_2 and for CO_2 broadened by O_2 and N_2 , in the 4.3 μ m region, for different temperatures. It was then shown in [8] that function χ is asymmetric (with respect to ν_0) for CO_2 in the 4.3 μ m region, at 296K.

Finally, Perrin and Hartmann [9] and Tonkov et al. [10] provide χ functions for pure CO_2 , respectively in the 4.3 μ m region, for $T \in [190 - 800]K$ and in the 2.3 μ m region, at 296K. These results are used in KSPECTRUM in order to compute χ profiles. Various options (see section 3) let the user chose whether χ functions from the literature must be used in their strict range of validity, or if they have to be used in different spectral ranges / for other molecules than CO_2 .

Collision-induced transitions are even more difficult to take into account: there are some measurements from Tonkov at al. [10] in the 2.3 μ m region, at room temperature. Central wavenumbers and intensities are given for 8 transitions, but the authors clearly state that the data they provide should be considered with caution, because of the large uncertainties of their measurements/computations. Other measurements of CO_2 CIA have been reported in the literature: [11, 12, 13, 14]. Data (lines central wavenumber and intensity) are either not provided, or given with poor accuracies, or provided in a very limited range of validity (in very specific spectral regions, for only one value of pressure and temperature, etc). Practically, it is not possible to use literature results to take into account collision-induced (or pressure-induced) absorption lines properly.

The only results that could practically be used for computing CO_2 CIA is the work from Gruszka and Borysow [15]. In addition to the paper, the authors provide a fortran computer code [16] that will compute CIA for CO_2 , in the 0-250 cm⁻¹ wavenumber range, in the 200-800K temperature range (this was motivated by studies of Venus' atmosphere, since CIA is a dominant source of opacity in the far-infrared (between 0 and $550 cm^{-1}$) on Venus' atmosphere [17] where temperature reach 735K at ground level). This code has been modified (for computing CIA coefficient for only one value of ν) and integrated into KSPECTRUM¹. CIA of CO_2 will therefore accurately be taken into account within these validity ranges. Options (see section 3) allow the use of CIA computation outside these validity ranges. In particular, the 2.3 and 4.7 μ m spectral bands of CO_2 are not taken into consideration.

1.3.2 Broadening by the rest of the gas

Another limitation of the code is the hypothesis that is done concerning γ_{other} : lines of a given species *i* can be broadened by collisions with the same species *i*, or by collisions with other species *j*. The self-broadened half-width γ_{self} given by LBL databases, and that account for collisions between molecules of the same species, can be considered as reliable. However, LBL databases also give us parameter γ_{air} , the air-broadened half-width, that corresponds to the broadening of lines by collisions between species *i* and... the air on the terrestrial atmosphere !

Of course, when computing a spectrum for a non-terrestrial atmosphere, parameter γ_{other} should not take the value of γ_{air} indicated in LBL databases, because the rest of the gas is no longer Earth' air. However, we have no other choice than taking $\gamma_{other} = \gamma_{air}$, because there are no other data available. This choice can be justified by the fact that γ_{other} will probably never be very different from γ_{air} anyway.

1.3.3 High-temperature transitions

We know that (a large number of) new transitions are activated at high temperatures. The CDSD database offers a solution for CO_2 lines: it is valid up to 3000K [4]. A new HITEMP database is in preparation, that should cover a huge number of high-temperature transitions for H_2O . More generally, the accuracy and range of application of public LBL databases is clearly increasing. However, the accuracy of spectra computations always depend on the available LBL parameters. The current level of knowledge will always impose a limitation on the accuracy of numerical simulations.

WARNING: there are some inconsistencies within the current HITEMP database: some values of the γ_{self} parameter are null ! This is an issue because, in the case you want to produce a high-resolution spectra for a single molecular species, only this parameter will be taken into account in

 $^{^1 \}rm Note that M.$ Gruszka's code is not supposed to be redistributed. However, the authors mention that the code can be used in academic applications, such as KSPECTRUM .

the computation of the Lorentz line width γ_L . A null γ_{self} will result in a null value for γ_L , and this will end in a crash of the code during spectral discretization. The spectral discretization algorithm has been improved in order to detect null values of γ_{self} ; in this case, instead of using a null value, the γ_{self} parameter will be set to the closest non-null value found among the LBL database.

1.4 A better respect of spectroscopy

The general idea of KSPECTRUM is to ensure the simple accuracy criterion that are discussed in section 1.1. In order to achieve this for any situation (gas mixture composition, values of temperature and pressure), it is absolutely necessary that the coding is well separated from the physical problem. In particular, the available computing power should not impose limitations on the versatility of the code. We believe this is possible with to-days masscomputing power.

1.4.1 (no) line selection

Existing computation codes that can produce high-resolution spectrum always involve line selection at some point. The purpose of selecting lines is to reduce the total computation time, by reducing the number of lines N whose contribution has to be computed at each wavenumber point ν (see Eq. 1).

However, KSPECTRUM does not need to select lines that have to be taken into account: the contribution of all transitions are effectively computed. See section 1.1 for details (accuracy parameter ϵ_1).

1.4.2 (no) truncation of lines and "continuums"

In the same manner, line profiles are not truncated as they probably would in a classical approach: the contribution of each line at each wavenumber ν is computed, no matter how far from the line center ν is. This is also a direct consequence of the way accuracy parameter ϵ_1 is taken into consideration.

KSPECTRUM does therefore not need to use a "continuum" that is classically added to the computed spectrum to correct line truncation side-effects. Values of $k_a(\nu)$ are computed with a specified relative accuracy ϵ_1 .

2008/12/09 update: this is no longer completely true, since I have enabled a line profile truncation possibility in the code (see section 3.1, end of

"options.in" description). This option should be used with great care, since it is most probable that very intense and distant lines will not be taken into account when truncating line profiles.

1.5 Parallel computing

Because of accuracy requirements that are imposed, computation times could be prohibitive on a single-processor machine. KSPECTRUM has been designed for running on multi-processor machines / clusters. It uses MPICH instructions so that multiple processes can be run at the same time. The four time-consuming loops of the code have been parallelized. As communication times between processes are negligible (compared to computation times), the total computation time will effectively be divided by the number of processors KSPECTRUM is running on (provided that all processors / machines in the cluster run at the same speed and are free of other time-consuming processes.)

Of course, this means MPICH has to be installed before using the code. See section 2.5 for installing MPICH / creating a cluster of machines.

1.6 The multi-pass approach

We are now talking about the limitations of fortran 77, which has been used for coding KSPECTRUM. The source code is constantly checked against array overflows, unused or uninitialized variables, etc (classical programming errors), and there should be no problem of bad coding. However, one limitation of fortran 77 is that the size of each array has to be declared. And in the present case, the most obvious limitation is the size of the arrays that hold the LBL data. For a typical atmosphere (Earth, Venus), less than 10 molecular species have to be taken into account, which gives a total number of lines of approximately 2.10^5 when using the HITRAN LBL database. However, when using the CDSD database for CO_2 lines, the number of lines that have to be taken into account can exceed 10^7 . And there is no way 9 arrays of size 10^7 can be declared on a classical machine that holds 1GB RAM (KSPECTRUM actually needs to read 9 LBL parameters from databases, which imposes the declaration of 9 data arrays). The default size of theses arrays is 2.10^6 , and can be reduced for low-memory configurations (see section 2.3).

With such a limitation, the only way several millions of lines can be taken into account is to perform a multi-pass computation: for instance, when the user specifies the use of the CDSD database for CO_2 transitions, 10 millions lines have to be taken into account (for CO_2 only). With an array size of 10⁶, KSPECTRUM will read the first 10⁶ lines (first pass), and perform the computation of the $k_a(\nu)$ spectrum with these first set of lines (it will actually compute the contribution of this first million of lines to the final result). Then it will perform a second pass, using the second million of lines in the database: it will compute the contribution of the second million of lines to the final result. And so on, until all lines have been taken into account.

An attentive reader could tell me this approach is incorrect if applied to the spectral discretization. Indeed, in order to compute a correct spectral grid for any $[\nu_{min}, \nu_{max}]$ narrowband spectral interval, the algorithm needs to take into account ALL the lines whose central wavenumber is within $[\nu_{min}, \nu_{max}]$. When reading chunks of LBL database, it is obvious that line parameters are missing for many $[\nu_{min}, \nu_{max}]$ intervals. This is the reason why the spectral discretization algorithm does not use the multi-pass approach: it automatically looks into every LBL database file to identify relevant line parameters for each $[\nu_{min}, \nu_{max}]$ interval (in practice, a complete parsing of LBL data files is necessary in order to accelerate the process of line identification). This is also why LBL data files must be readable by any KSPECTRUM process (see section 2.6).

Finally, we should mention that the total computation time for a multipass computation is not higher than for a 1-pass computation, since all lines have to be taken into account anyway. The spectral discretization is performed during the first pass and is known for subsequent passes, which does not increase the total computation time. The main advantage of this multipass approach is the possibility to run the code on low-memory configurations.

1.7 Estimation of remaining computation time

When using the multi-pass approach, the code will be able to guess the remaining time before the computation of a spectra is finished for the current atmospheric level. This is only possible for a multi-pass computation, because in order to have an estimation of the remaining computation time, the code needs to know the spectral grid (that is computed during the first pass of each atmospheric level), and how many points of the grid have already been computed. The "status.txt" file will report the estimated time (in real-life date/hour format) when the computation for the next atmospheric level is going to start.

1.8 Not starting from scratch after a crash

Imagine you want to compute the spectrum of a given atmosphere mainly composed of CO_2 , using the CDSD database, over 80 atmospheric levels, for the whole infrared range. This should typically require several weeks of CPU time, even for a multi-processor machine or a small-size cluster. If you run KSPECTRUM on a dedicated multi-processor machine, if no one else has access to this machine, and if there is no power cut during all the computation, there should be no problem. Now imagine you run it on a PC cluster, with many other users that daily access these machines. Someone will eventually reboot one of these machines, crashing your whole computation. I would say that you can not decently expect more than 48 hours between two consecutive reboots. In these conditions, the computation will never end.

Fortunately, KSPECTRUM can resume a computation if a crash has been detected. It will start over from the last backup point, which occurs every time the computation of the spectrum has been achieved on a given narrowband interval. Multi-pass computations will be resumed as well.

1.9 Automatic tabulation of Lorentz and Voigt functions

Section 1.1 presents the principle of the spectral discretization algorithm used in KSPECTRUM. It needs a tabulation of the Lorentz and Voigt functions. These tabulations depend on the value of ϵ_2 , the user-defined accurately that is required over the spectral grid. Tabulation files reside within the "data" directory, and are indexed according to the value of ϵ_2 : for instance, "data/tabulation_lorentz0.01.txt" is the file where $[x_1, x_2, x_3]$ triplets have been recorded for the Lorentz function, and for an accuracy $\epsilon_2 = 0.01$ (a 1% uncertainty).

Values of ϵ_1 and ϵ_2 may be changed by the user (see section 3.1). If the user specifies a value of ϵ_2 for which there is no known file, KSPECTRUM will have to perform the tabulation prior to start the spectrum computation. Of course, the code will record tabulation results in the appropriate file, so that

the tabulation for this value of ϵ_2 will never have to be performed again. This tabulation is performed by a fully parallel loop, so that it can benefit from the number of processors KSPECTRUM has been launched on.

2 Installation and prerequisites

2.1 First thing to do

is to declare your fortran 77 compiler in the F77 environment variable. In order to do so, you should export this environment variable in your .bashrc, .cshrc or .profile file; for instance, add the following line into your .bashrc file:

>export F77="g77"

in case you are using the GNU fortran 77 compiler (g77). Replace "g77" by your local fortran 77 compiler if needed.

Then you need to type:

>source .bashr

in order to take the modifications into account. Replace ".bashrc" by whatever you are using. The purpose of defining the F77 environment variable is to provide the correct compiler to compilation commands that will be used later by the installation script. If this variable is not defined, the name of your fortran 77 compiler will be asked every time a compilation will occur.

Then you need to use script "install_kspectrum.bash" that is provided with the archive. This script should be placed in the same directory than the provided zipped archive. Use command "./install_kspectrum.bash" in order to run it. This script will:

- Untar the archive, if the corresponding version of KSPECTRUM is not already installed
- Make appropriate links in the newly created directory
- Go into the "data" directory, compile program "make_data.exe" and run in, in order to generate test files "composition.in" and "narrow-bands.in" (see section 3.2).
- Remind you to install LBL databases into the "data" directory (see section 2.6).
- Erase the zipped archive, and erase itself.

Then you can go into the newly created directory (named "kspectrumX.X.X" with "X.X.X" the version number) to continue the installation.

2.2 Tweaking the "Makefile"

Before compiling, you will have to find out what compilation options are right for your compiler, and your machine. Open the "Makefile" file, and look at variables "FOR", "ARCH" and "OPTI". Variable "FOR" is used to specify your fortran 77 compiler. As KSPECTRUM uses MPICH, you will most likely use the "mpif77" compilation command, that has been installed along with MPICH.

Variable "ARCH" is used to specify machine architecture. "-m486" is probably a good choice for a PC running a 32bits linux. Use the documentation of your fortran compiler to find out what architecture option you can use.

Variable "OPTI" is used to specify code optimization options. The default options should be enough. Please note that you definitely must use option "-Wno-globals" for compiling parallel code.

You might also want to set variable "DEBUG" (look for its definition in the file). You can expect faster execution times if you leave it empty.

2.3 Setting up array sizes

As explained in the previous section (see 1.6), one limitation of fortran 77 code is that you must define array sizes before compilation. Arrays sizes used by the present code are defined within the "includes/max.inc" file. You should at least look at it before compiling, and more precisely at the value of variable "Nline_mx". Its default value is 2000000 (2.10⁶), because most machines will be able to compile KSPECTRUM with this value (you need at least 1GB RAM). You can enable a lower value if you do not have enough memory. The code will then perform a multi-pass computation (see section 1.6), but total computation time will not be increased.

2.4 Compilation

Once you checked compilation options and array size definitions, you can use the following command in order to compile the executable file:

> make all

If compilation fails, use the compiler error message to determine what went wrong. The most probable error causes are: a bad definition of architecture compilation option, or an inappropriate value in code optimization options.

If you ever need to modify the source files (in directory "source"), you can quickly recompile the code using "make all" again. This will only recompile the modified source files, and link objects files in order to produce the new executable file.

If you have to modify the value of any variable defined in includes files (directory "includes"), you will have to recompile the whole code from scratch. Use the following command to erase all objects files, and then recompile them properly:

> make clean all

Odd errors may happen if you change an include file and then recompile using only the "make all" command (old value of the modified variable will remain in the unchanged object files).

2.5 Installation of MPICH

If you do not already have MPICH installed on the machine / group of machines you want to run KSPECTRUM on, you will first have to download mpich2 from http://www.mcs.anl.gov/research/projects/mpich2; make sure you download version 1.0.7. Next, untar the downloaded archive, and install it on every system that will be part of your cluster:

> ./configure - prefix = /path/to/installation/directory

> make

> make install

Before running the MPD daemon, you must create a ".mpd.conf" file in your home folder:

> echo secretword=[secretword] » /.mpd.conf

> chmod 600 .mpd.conf

using any "secretword".

Next, you will need to be able to connect via ssh to every other machine of your cluster, with no password request. For this, you must first create a DSA key:

> ssh-keygen -t dsa

leaving all fields blank (use the "enter" key to answer each question). Then you will have to add this DSA key to the list of authorized keys:

> cd .ssh

> cat id_dsa.pub > authorized_keys

Finally, create the list of machines that belong to your cluster. This list must reside within the "mpd.host" file on your home folder. Each line must contain the name of the machine, by order of availability:

[host1].[domain] [host2].[domain] [host3].[domain] etc You can then try to run the MPD daemon: > mpdboot -n [#]

with [#] the number of hosts you want to run MPD on (typically, the number of machines in your cluster). If you encounter no error, you can use command "mpdtrace" to check the number of hosts the MPD daemon is running on. This should give you the list of machines in your cluster.

2.6 Installation of LBL databases

KSPECTRUM needs to access LBL databases in order to run. LBL databases are not provided with the code, you will have to find and download them. The code expects to find LBL databases in the "data" directory. Of course, you have to install KSPECTRUM on a directory that is common to all machines the code will run on (shared disk), so that each process will be able to access the LBL databases through the "data" directory. As LBL databases can use a lot of disk space, it is recommended to install LBL database files on the local disk of each machine, and then redirect a link from the (shared) "data" directory to each LBL file directory.

LBL data file directories must be the following:

- "data/HITRAN2004": must contain the uncompressed by-molecule HI-TRAN data files in its 2004 version. If you create a link, it must point to the "HITRAN2004/By-Molecule/Uncompressed-files" directory on the local disk of each machine. When the code is set to use the HI-TRAN2004 database (see "options.in" in the 3.1.2 section), this folder must exist within the "data" folder.
- "data/HITRAN2008": must contain the uncompressed by-molecule HI-TRAN data files in its 2008 version. If you create a link, it must point to the "HITRAN2008/By-Molecule/Uncompressed-files" directory on

the local disk of each machine. When the code is set to use the HI-TRAN2008 database (see "options.in" in the 3.1.2 section), this folder must exist within the "data" folder.

- "data/HITEMP": must contain the HITEMP uncompressed data files. If a link, it must point to the "HITEMP/uncompressed_data" local directory.
- "data/CDSD": must contain CDSD uncompressed data files. If a link, it must point to the "CDSD_1000_UPDATED/uncompressed_data" local directory.
- "CDSD_Venus": if you want to use the CDSD version that was specially compiled for the atmosphere of Venus. This directory must contain the uncompressed files of the Venus version of CDSD data base. If a link, it must point to the "Venus/uncompressed data" local directory.

In practice, let's say you have downloaded and unzipped the 2008 version of the HITRAN database somewhere on the local disk of the machine you want to use for your computation. KSPECTRUM will need the "by-molecule" data files that are provided with HITRAN. For instance, you have these files in the following folder:

 $HOME/LBL_databases/HITRAN/HITRAN2008/By-Molecule/Uncompressed-files/$

On the other hand, KSPECTRUM is installed in the following folder: \$HOME/great_codes/kspectrum1.0.5/

In order to let KSPECTRUM know where the by-molecule data files of the HITRAN2008 database are located, go to the "data" folder:

cd \$HOME/great_codes/kspectrum1.0.5/data

and create a symbolic link called "HITRAN2008" to the folder that contains the data files:

ln -s \$HOME/LBL_databases/HITRAN/HITRAN2008/By-Molecule/Uncompressed-files/ ./HITRAN2008

Here you go. You have just installed the HITRAN2008 database in KSPECTRUM. You can do the same for the 2004 version of HITRAN if you want to use it (you can select which database you want to use by setting the right option in the "options.in" file, see section 3.1.2) and for the HITEMP and CDSD databases.

Frequent updates are published about HITRAN data. If you want to use these new data files, follow these directions: let's say you want to download the update for SO₂ from the HITRAN website (section "HITRAN updates"). Download the updated file "09_hit09.par" within the folder that contains your by-molecule data files. KSPECTRUM will still look for files that are labelled "08", so you will have to link the new file to the old one: within the by-molecule data folder, rename the old SO₂ file:

mv 09_hit08.par 09_hit08.par.ori

Then create a link from the new file to the old one:

ln -s 09_hit09.par 09_hit08.par

Repeat this step for every update file.

Finally, there is also supplemental data, that is not directly provided among the "by-molecule" data. These files reside within the "HITRAN2008/Supplemental" folder. There are currently three data files within this "supplemental" folder: "30_hit08.par", "35_hit08.par" and "42_hit08.par". In order to add then to your "by-molecule" folder, you can just copy/paste them, or once again create symbolic links (does not take more disk space); within your "by-molecule" data folder, type the following commands:

ln -s ../../Supplemental/30_hit08.par . ln -s ../../Supplemental/35_hit08.par . ln -s ../../Supplemental/42_hit08.par .

3 Using the code

3.1 User input files

There are two files in the main KSPECTRUM folder: "data.in" and "options.in". The "options.in" file is where the user can chose between several options, for each algorithm. The "data.in" file is where the user can specify the numerical values of a number of variables the code will use. Below is a description of these two files.

3.1.1 The "data.in" file

- Section "Production of k spectrum": the user must specify the values of ϵ_1 and ϵ_2 (see section 1.1). Default values are 1%.
- Section "Narrowband discretization": the user can specify the minimum and maximum values of the wavenumber range the spectrum will be computed for, and the maximum error percentage over the Planck intensity for discretization of the wavenumber range into narrowband intervals. These data should be specified only if the user chose to compute the narrowband discretization instead of using pre-defined narrowband intervals (via an option in the "options.in" file). Refer to the description of the corresponding option for more details.
- "Parallel computing": the user should specify the number of chunks the code will use for the computation of the k spectrum. Once the spectral grid is computed, the code knows the number of k values that will have to be computed. The computational load that will be sent to each process is the number of k values divided by the number of chunks. If this number of chunks is large, each process will have to compute a small number of k values. A chunk will be sent each time a process is found idle. The advantage is that if processors run at different speeds / have to run other processes, processes that run on slow processors will be sent a lower computational load (fewer chunks), and processes that run of fast processors will be sent a greater number of chunks.
- Section "Levels and narrowbands limits": first, the high-temperature level has to be indicated; this value is used to switch between HITRAN (at low temperatures) and HITEMP or CDSD (at high temperatures).

By using a high value (higher than the highest value of temperature you want to compute spectrum for) you can be sure only HITRAN is used. The user can then specify the indexes of the first and last atmospheric level / narrowband interval if only a limited number of atmospheric levels / narrowband intervals should be treated instead of every atmospheric level / narrowband interval. The choice itself is done via the corresponding option.

• Section "Special options for degraded mode": the user should provide here values the code will use when running into degraded mode. Three values are used for the degraded spectral discretization algorithm (number of line widths to consider in the definition of a line's central zone, the number of points used to discretize lines central zones and the number of points used to discretize regions between lines) and one is used for line profiles truncation (distance to line centers). One last value is the percentage of lines that will be rejected when required (see "options.in").

3.1.2 The "options.in" file

- Section "Narrowband discretization": the user can chose between computing automatically the narrowband discretization or use pre-defined narrowband intervals. If the narrowband discretization has to be performed by the code, it will use the three corresponding values in the "data.in" file (section "Narrowband discretization"): the specified wavenumber range will then be discretized into a number of narrowband intervals, and the Planck intensity relative variation between the limits of each narrowband interval will not exceed the specified maximum error percentage over Plank intensity. If the user decides to use a pre-defined narrowband discretization, a file called "narrowbands.in" must be found in the "data" directory. This file should contain the number of narrowband intervals, and the wavenumber limits of each interval. See the provided "data/narrowbands.in" example file.
- Section "Choice of spectroscopic database": the user has to decide whether the default HITRAN database to use is the 2004 or the 2008 version; the corresponding link must be established within the "data" folder (see section 2.6). The user can then choose between the HITEMP or the CDSD database for a temperature greater than "T_hitemp"

(whose value was set in the "data.in" file). In the case of the atmosphere of Venus, the user should decide whether the code will use the standard LBL database (HITRAN 2004 or 2008) or the CDSD version that was specially compiled for Venus. Finally, the user should decide whether LBL data reorganization should occur on-the-fly (when reading LBL data) or once all LBL data has been read. In practice, the second choice is better for this last option, since data sorting will be much faster.

- Section "Composition": the user should specify whether the code will use only the main isotope or all isotopes when the concentration of isotopes are not specified for a given molecule.
- Section "Sub-Lorentzian profiles": the user has to chose whether or not the sub-lorentzian nature of lines has to be taken into consideration. See section 1.3 for more information about how the code will manage sub-lorentzian profiles. In particular, function χ is given for pure CO_2 only, in specific spectral ranges, and for a limited range of temperature. The user can chose to use literature data outside its range of validity (over the whole IR spectrum), and for other molecules than CO_2 (use with great care !). Finally, the asymmetric nature of function χ can be taken into account, but results will not be very different from using a symmetric function.
- Section "Collision-Induced absorption": the use can chose whether or not collision-induced absorption (CIA) should be simulated for CO_2 . See section 1.3. The user can subsequently choose CIA computation from the work of Gruszka [15] or Baranov [18] ² or both, in which case both sources of opacity are added. The user can also chose to compute CIA coefficients outside the wavenumber validity range (0-250 cm^{-1}) and the temperature validity range (200-800 K).
- Section "Code behavior": the user can chose whether the code will erase results files at start time or not. It is preferable to erase these files, but in any case, previous computation results should have been recorded

²When using CIA from the work of Baranov 2004, the code needs an additional file. This file is named "CO2_dimer_data", and is located within the "data" folder. However, since this file is not free for distribution, it is not provided with the code. Please contact the author for more information.

before starting a new computation. Even if erasing has been required, files will not be erased in the case the previous computation was not finished (crashed), and computation resuming has been required. The user can chose to display or not computation times on screen. Finally, it should be specified whether the code has to resume an interrupted computation, or start again from the beginning.

- Section "Sensitivities": the user can chose to compute sensitivities of results (cross-section and absorption coefficient) to total pressure, temperature and species concentrations. When computation of sensitivities are enabled, results are recorded in files "results/dk_dXXX".
- Section "Levels and narrowbands limits": the user must chose whether the code must compute spectrum for every atmospheric level / narrowband interval, or only for a limited number of levels / intervals. If a limitation is imposed, the code will use the corresponding values found in the "data.in" file (section "Levels and narrowbands limits").
- Section "Spectral discretization algorithm": the user should choose between the reference spectral discretization algorithm or a degraded spectral resolution. Degraded resolution will compute the spectral grid much faster, and will also make the computation of the kspectrum faster since less points will have to be computed. However, the accuracy criterion ϵ_2 will not be satisfied by the degraded resolution algorithm. This degraded resolution algorithm takes input values (see "data.in"). When using the reference spectral discretization algorithm, the user should also provide the accuracy level that will use the special spectral grid algorithm for wavenumbers located between close strong lines. A value of 0 disables this special algorithm: wavenumbers ν_i will be computed using the standard meshing algorithm based on Lorentz and Voigt functions tabulation, described in 1, but remember that in this case, the spectral grid may be too coarse between strong lines. A value between 1 and 4 will enable the special meshing algorithm. The highest the value, the more accurate this algorithm will be (there will be more values of ν_i , therefore making the k-spectra computation longer). A value of 3 may be enough in most cases.
- Section "line truncation": the user has the ability to truncate line profiles at a fixed distance from line centers (see "data.in"). When this

option is used, the accuracy criterion ϵ_1 is no longer satisfied. Line truncation will make kspectrum computations much faster (around 20 times faster). However, there may be cases where the resulting spectra is completely wrong (typically, when negelecting the contribution of distant, yet very intense transitions). Be very carefull when line truncation is used ! When a truncation is required, the user has also the ability to set the contribution of constant lines to zero, so that absorption is null between very distant lines. A last option enables weak lines rejection: in order to make the code faster, the computation can be performed using only the stronger lines (those who have the highest intensities). When line rejection is enabled, the code will reject the x% weaker lines, with x the percentage provided in the "data.in" file.

3.2 Other input files

3.2.1 Atmospheric composition

is provided via the "data/composition.in". This file must contain a label for the atmospheric composition (could be used in future version for planetspecific physics, such as sub-lorentzian profiles or CIA), the number of atmospheric levels, the number of molecular species, the label of each molecular species, and, for each atmospheric level: altitude in km (unused at present date), pressure in atm, temperature in Kelvin, and the molar fraction (or partial pressure) of each molecular species. See the example file.

Molecule labels have to be identical to labels used in the HITRAN nomenclature. Labels of molecules defined in the HITRAN database can be found in the "data/molparam.txt" file. They are obvious for most species: "H2O" is used for water vapor, "CO2" for carbon dioxide, etc. Please use capital letters when providing molecule labels in the "data/composition.in" file, and note that molecule labels must be provided between brackets (example: " $[H_2O]$ ") so that the code can identify molecular species.

You can also use molecule labels that do not belong to the HITRAN nomenclature, when specifying the molar fraction of an isotope. For instance, you can specify the molar fraction of water vapor (using the "H2O" label), and you might want to specify the molar fraction of deuterium separately. In this case, you have to define a new molecule label, for instance "HDO". This label should also be used in the "data/special_molecules.in" file. See description of this input file at section 3.2.2.

One last remark about isotopes: when no isotopic abundance has been explicitly provided via the "data/composition.in" file for a given molecular species, the algorithm can chose to take into account only the main isotope of this species, or use all isotopes whose line parameters are provided in LBL databases. The user can chose between both behaviors in the "options.in" file (see section 3.1.2).

3.2.2 Special molecules

When special molecules have been provided in the "data/composition.in" file (i.e. molecules whose label is unknown from the HITRAN nomenclature), information has to be provided about these molecules in the "data/special_molecules.in" file. See the provided example file. This file must contain, for each special molecule: its label (for instance "HDO"), the label of its "mother-molecule" (for instance "H2O" for deuterium), and its molar mass, in g/mol. If the special molecule can count multiple isotopes, the molar mass of each isotope must be provided. In the example file, there are three lines for "HDO", that correspond to isotopes $HH^{17}O$, $HH^{18}O$ and $H^2H^{18}O$.

When an isotope (such as "HDO") is specified within the composition file "data/composition.in", and multiple definitions of this isotope are found in the special molecules files "data/special_molecules.in", the algorithm in charge of defining which lines have to be read in the LBL files will affect the provided total partial pressure (that is specified for "HDO" in the composition file) to each isotope, prorata to their abundance in the Earth atmosphere.

3.2.3 Narrowband intervals

have to be provided in the "data/narrowbands.in" file, provided that the user chose to use a pre-defined narrowband discretization (see section 3.1). This file must contain the number of narrowband intervals, and, for each interval, its lower and upper limits, in terms of wavenumber (in cm^{-1}).

3.3 Running KSPECTRUM using MPICH

Once everything is installed and the executable file "kspectrum.exe" file has been compiled, you can try to run a computation. I would recommend that, for the first time, you run KSPECTRUM over a single atmospheric level, for a single narrowband interval, using a simple atmospheric composition (the provided example composition for instance).

Use the following command to run the code:

> mpirun -np [#] kspectrum.exe

with [#] the number of processes that have to run.

Because communication times are small compared to computation times in KSPECTRUM, it is a good idea to chose a number of processes equal to the number of (physical) processors of your cluster, plus one. One process, the master process, is dispatching computational loads to every other processes (slave processes), and gathering results from them. It does not require any significant CPU time, therefore it is OK to have a number of slave processes equal to the number of processors, so that each slave process can use a processor (or each processor will have only one slave process running on it).

In practice, if your cluster is composed of n processors, you can use:

> mpirun -np n+1 kspectrum.exe

4 Results

4.1 Results files

KSPECTRUM will produce results files in the "results" directory. There is one file per atmospheric level. Files are named "k***" with "***" the index of the atmospheric level. For instance, "results/k001" is the result file produced for the first atmospheric level (with the highest pressure level).

Each file contains 3 columns. The fist column gives values of wavenumbers, in cm^{-1} . The second column is the total cross-section of the gas mixture (in $cm^2/molecule$), at the corresponding wavenumber. The third column is the total absorption coefficient (in m^{-1}) at the corresponding wavenumber.

4.2 More output files

Additionally, the user can check a number of log files produced by KSPEC-TRUM during its execution:

- "results/composition_info.txt": information returned by the algorithm that reads the atmospheric composition file and sets abundances for each molecular species. This file shows what molecules have been identified, the number of isotopes and their respective abundances. For each molecular species, the sum of abundances of all the isotopes should be equal to 1.
- "results/calculation_info.txt": various output information: values of accuracy criterion ϵ_1 and ϵ_2 are given, along with the gain from line classification: the total number of lines is printed, as well as the number of lines whose contribution has to be explicitly computed at each value of ν . These are cumulated values (over each narrowband interval, over each atmospheric level). The difference is an estimation of the computation time gained by line classification (using criterion ϵ_1).

These two files should be saved together with spectrum result files in order to keep a record. In particular, file "results/calculation_info.txt" will be needed when producing k-distribution data sets.

• "optimizations/LBL_files": shows which LBL data files have been read.

• "status.txt": is continuously updated during computation. Some lines are appended to this file each time a backup is performed, i.e. when the computation of the spectra is over for a given narrowband spectral interval. It shows the time the backup took place at, for what atmospheric level / narrowband interval, and how many values of k have been computed for this interval. When running a multi-pass computation, it can also display the predicted time/date when computation will be over for the current atmospheric level.

4.3 What's next ?

You can use KSPECTRUM for the sole reason of producing high-resolution $k_a(\nu)$ spectrum. However, you might want to use the results of KSPEC-TRUM in order to produce k-distribution data sets that can be used for actual radiative transfer simulations. See the documentation of code "k-dist" for the production of k-distribution data from high-resolution spectrum.

5 The fun stuff

5.1 Code license

KSPECTRUM is distributed under the General Public License. See the COPYING file within the main directory. This means that you can use / redistribute part of all the source code in your own source code, provided that your source code is also distributed under the GPL. You can not produce commercial code from the provided sources. I will eventually know it and kill you.

5.2 Questions and Answers

I am in a big hurry, I don't want to read all that boring stuff, I just want my results, immediately.

The easy way is to ask me to use my code to produce your results. And as I am fed up of working for free, that means you will have to pay. Please send me an email: v.eymet@gmail.com

There is an alternative though: you can hire a grad student / post-doc to do the work. It will still cost you money, but it will take longer.

I can't manage to compile the source code / I don't understand compilation error messages.

There should be not problems using the gnu fortran 77 compiler (g77) and mpich2. Send me an email with a description of the tools you are using, on what hardware/system, and what error messages you get: v.eymet@gmail.com

Compilation was successful, but the code crashes.

I know that when a code has been extensively used for years and all bugs have been patiently tracked, identified and corrected, there still remains a huge ugly bug for every 1000 lines of code. My code counts nearly 14000 lines of code, and it has not yet been extensively used. So yes, there may be a bug. Send me an email with the description of what you want to do, attach input files when possible. I will correct the bug (and do it for free !). **Error "Nkmx has been reached"** This is one of the most likely errors. You are probably trying to compute a spectra for a very low value of pressure, right? There are a lot of very thin and isolated lines, the code takes ages to compute the spectral discretization, and then crashes with this error. This is because the number of wavelength the spectra should be computed for is too high. There are a number of ways to get around this issue:

- First, try to edit the "includes/max.inc" file, and to increase the value of variable "Nkmx", in order to allow more memory to spectral arrays. You will have to recompile the code completely to take the modification into account: use "make clean" and then "make all" (or directly "make clean all"). Be very careful when increasing the value of "Nkmx": if you do not increase it enough, the same error will happen again (not enough memory for spectral discretization). If you increase it too much, you will not be able to compile the code (not enough available memory on the system). The default value for "Nkmx" is 10⁵, you should first try to set it at a value of 5.10⁵. Then increase it if the same error occurs again, until you reach the limit at which compilation and / or execution is impossible.
- If you are low in memory and you can not increase further the value of "Nkmx", you can try to use a coarser spectral resolution, by setting the appropriate options in the "options.in" file (within the "Spectral discretization algorithm" section).
- You can also try using smaller narrowband spectral intervals: Nkmx is the number of values of ν a given narrowband interval will be discretized into, so using smaller narrowbands can make the discretization possible.
- If all the above suggestion failed, or you have the feeling there is another problem, feel free to contact me. When possible, send me your "composition.in", "narrowbands.in", "data.in" and "options.in" files with a description of the problem you encounter.

What about all this unknown spectroscopic data...?

As mentioned in section 1.3, we know very little about: sub-lorentzian profiles, collision-induced absorption (CIA) lines, and broadening by the rest

of the gas mixture. Hypothesis have been made: sub-lorentzian profiles are given in the case of pure CO_2 , and in specific (spectral, temperature) ranges of validity. These limitations can be overridden by the user (see section 3.1.2). Broadening by the rest of the gas has been considered as if the rest of the gas mixture was terrestrial air, and CIA can be accurately simulated for CO_2 only, in the 0-250 cm^{-1} spectral range only.

I am aware this situation is not acceptable, since KSPECTRUM is supposed to be used for any gas mixture, in any spectral region, and in a wide range of thermodynamic conditions. However, such a code would require a good knowledge of all the missing data (sub-lorentzian profiles, CIA transitions and broadening by the rest of the gas mixture) for every molecular species, broadened by every other molecular species, for any spectral region, for every temperature and pressure level. The best I can propose is to improve the code for very specific applications, when the corresponding spectroscopic data is known. Please let me know your needs: v.eymet@gmail.com

A (not fully satisfying) solution can be found in the case you know well the CIA continuum you want to take into account, i.e. you know how to compute CIA opacities for your configuration, but it has not been included into KSPECTRUM : in this case, you should first compute your hi-resolution spectrum without any CIA calculations. Once hi-resolution spectrum without CIA have been computed, you can compute and add CIA opacities by yourself at each ν point.

A Appendix - Validation

KSPECTRUM results have been checked in various situations against other results.

A.1 Low pressures / temperatures

Results from KSPECTRUM have been compared to line-by-line spectrum produced by P. Dubuisson (Laboratoire d'Optique Atmosphérique, Lille) [19, 20, 21, 22, 23] for Earth's atmosphere. Various compositions / spectral intervals have been considered:

- H_2O , CO_2 and O_3 , in the [748 752] cm^{-1} range.
- H_2O , CO_2 and O_3 , in the [4900 5040] cm^{-1} range.
- H_2O , CO_2 , O_3 and CH_4 , in the $[4200 4240] \ cm^{-1}$ range.

In each case, spectrum have been computed for 49 atmospheric levels, from ground level (1 atm, 292K) to $2.878.10^{-6}$ atm (0.29 Pa, 345K) at the top of the atmosphere.

Figures 3 to 6 represent both spectrum in several spectral ranges, at different pressure and temperature levels.

A.2 High pressures / temperatures

Absorption spectrum computed by KSPECTRUM in the case of Venus' atmosphere have been used to compute absorption coefficient average $\bar{k_a}$ over 68 narrowband spectral intervals covering the [40-5825] cm^{-1} range. These average absorption coefficients have been compared to average absorption coefficients computed from k-distribution data used for radiative transfer simulation in Venus' atmosphere [24, 25].

Comparisons have been performed at ground level (90.9atm, 735K), at an altitude of 5km (65.8 atm, 697K) and at an altitude of 55km (0.52atm, 302K).

In each case, orders of magnitude of average absorption coefficients computed from KSPECTRUM high-resolution spectrum are in good agreement with average absorption coefficients from k-distribution data, especially in the 1.7 and 2.3 μm "transparency windows" (this validates sub-lorentzian profiles representation) and in the far infrared ($\nu < 300 \ cm^{-1}$) (which validates the collision-induced absorption continuum).

Table 1 provides a comparison between average absorption coefficients computed at ground level.



Figure 3: Absorption spectra for a H_2O , CO_2 and O_3 mixture at pressure of 0,298 atm ($x_{H_2O} = 3, 34.10^{-4}, x_{CO_2} = 3, 30.10^{-4}, x_{O_3} = 1, 20.10^{-7}$) and a temperature of 238,6K, in the [748 - 752] cm^{-1} spectral range.



Figure 4: Absorption spectra for a H_2O , CO_2 and O_3 mixture at pressure of 6,37.10⁻² atm ($x_{H_2O} = 3,25.10^{-6}, x_{CO_2} = 3,30.10^{-4}, x_{O_3} = 1,74.10^{-6}$) and a temperature of 218,6K, in the [750, 2 - 740, 6] cm^{-1} spectral range.



Figure 5: Absorption spectra for a H_2O , CO_2 and O_3 mixture at pressure of 0,583 atm ($x_{H_2O} = 3, 04.10^{-3}, x_{CO_2} = 3, 30.10^{-4}, x_{O_3} = 5, 16.10^{-8}$) and a temperature of 270,3K, in the [4940 - 4980] cm^{-1} spectral range.



Figure 6: Absorption spectra for a H_2O , CO_2 , O_3 and CH_4 mixture at pressure of 0,259 atm ($x_{H_2O} = 1,74.10^{-4}, x_{CO_2} = 3,30.10^{-4}, x_{O_3} = 1,54.10^{-7}, x_{CH_4} = 1,56.10^{-6}$) and a temperature of 232,1K, in the [4214 - 4220] cm^{-1} spectral range.

Table 1: Average absorption coefficients $\bar{k_a}$ computed from KSPEC-TRUM high-resolution output spectra and from k-distribution data used for radiative transfer simulations ([24]) for some narrowband spectral intervals, at ground level (90,1 atm, 735K). Values of $\bar{k_a}$ extracted from the kdistribution data set are not a reference, but are rather used as a comparison point.

$ u_{inf}$	ν_{sup}	k_a k-distributions	k_a KSPECTRUM
520	570	60,5	48,8
570	615	574,1	529,8
615	645	$2,36.10^3$	$2,21.10^3$
645	680	$7,56.10^3$	$8,02.10^3$
680	720	$3,71.10^3$	$3,74.10^3$
720	760	899,3	913,1
760	785	205,3	179,9
785	815	58,4	42,2
815	845	14,0	8,5
845	900	8,9	5,1
900	935	15,7	13,9
2100	2155	14,5	5,9
2155	2280	$3,22.10^3$	$2,56.10^3$
2280	2390	$5,28.10^4$	$5,42.10^4$
2390	2450	$1,48.10^3$	$1,06.10^3$
2450	2540	15,9	10,6
2540	2670	$2, 8.10^{-2}$	$3, 1.10^{-2}$
2670	2745	$1, 4.10^{-2}$	$1, 1.10^{-2}$
2745	2790	$2,0.10^{-2}$	$4, 0.10^{-3}$
2790	2925	$7,9.10^{-3}$	$2, 5.10^{-2}$
2925	3040	$2, 2.10^{-1}$	$1, 1.10^{-2}$
3040	3225	$2, 2.10^{-2}$	$9, 9.10^{-3}$
3225	3450	2,2	0,18
3450	3760	604, 6	592,2
3760	3875	28,2	19,4
3875	4030	0,17	$9, 4.10^{-2}$
4030	4135	$4, 1.10^{-2}$	$1, 9.10^{-2}$
4135	4350	$3, 2.10^{-3}$	$1, 1.10^{-3}$
4350	4550	$6, 8.10^{-3}$	$3, 3.10^{-4}$
4550	4950	4,8	3,9
4950	5200	10,2	9,4
5200	5700	$2, 7.10^{-2}$	$1, 6.10^{-2}$
5700	5825	$3, 4.10^{-3}$	$1, 6.10^{-3}$

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