The Cosmic Linear Anisotropy Solving System (CLASS) I: Overview

Julien Lesgourgues\textsuperscript{a,b,c}

\textsuperscript{a} Institut de Théorie des Phénomènes Physiques, 
École Polytechnique Fédérale de Lausanne, 
CH-1015, Lausanne, Switzerland.

\textsuperscript{b} CERN, Theory Division, 
CH-1211 Geneva 23, Switzerland.

\textsuperscript{c} LAPTh (CNRS - Université de Savoie), BP 110, 
F-74941 Annecy-le-Vieux Cedex, France.

Abstract: The Cosmic Linear Anisotropy Solving System (CLASS) is a new accurate Boltzmann code, designed to offer a more user-friendly and flexible coding environment to cosmologists. CLASS is very structured, easy to modify, and offers a rigorous way to control the accuracy of output quantities. It is also incidentally a bit faster than other codes. In this overview, we present the general principles of CLASS and its basic structure. We insist on the friendliness and flexibility aspects, while accuracy, physical approximations and performances are discussed in a series of companion papers.
1. Why a new code?

There are several excellent Boltzmann codes publicly available on the market, in particular CMBFAST [1], CAMB [2] and CMBEASY [3]. The second one is still regularly maintained and upgraded, and offers lots of functionality. What could we expect from a new Boltzmann code? Our answer would be: user-friendliness, flexibility, accuracy and speed.

By friendliness, we mean: a code easy to understand, easy to compile and to run on any platform, offering a convenient input interface, in which input parameters could be passed in several ways.

Flexibility refers to the possibility to generalize the code to more complicated cosmological scenarios, or to interface it with various other programs (for parameter extraction, or for computing different observables).

All accuracy parameters should be grouped in a unique place in order to facilitate precision tuning. It should be possible to take the decision to run with various well-identified and calibrated precision levels, without needing to tune all accuracy parameters each time.

Since parameter extraction codes (e.g. CosmoMC [4], MultiNest [5], CosmoPMC [6]) typically require $10^4$ to $10^6$ calls to a Boltzmann code (depending on the complexity of the model), any improvement in the running time for a fixed precision setting would be appreciable.
Existing codes all contain several nice features, and brought decisive progress with respect to their predecessors. **CMBFAST** reduced drastically the running time thanks to the line-of-sight integration method, and included CMB lensing calculations for the first time. **CAMB** and **CMBFAST** incorporate several refinements in terms of structure of the code, lensing accuracy, approximation schemes, speed, etc. Criticizing these codes would be totally out of place. But who would claim that they are friendly and flexible? Anybody can download them and learn how to run in one hour. But modifying these codes is another story. Identifying the places where the modifications should be performed is a very involved task. If the goal is to introduce minimal modifications to the cosmological evolution, to interface the code with another one, to compute another observable (e.g. lensing spectra), a lot of work is already necessary. For implementing more complicated changes (like e.g. adding dark matter species with a non-trivial distribution or an interaction term), one should be – or become – a dedicated expert.

Concerning accuracy, all current Boltzmann codes have been claimed to be accurate at the 0.1% level [7]. This is marginally sufficient for a code supposed to analyze Planck data, for which the error bars on temperature are really small for intermediate angular scales (those with a very small cosmic variance and no significant foreground contamination). With post-Planck CMB experiments, cosmic shear surveys or 21cm data, even more accuracy might be needed. But how can we be sure that existing Boltzmann code are really accurate even at the 0.1% level? **CMBFAST** and **CMBEASY** are not maintained anymore, their recombination algorithm is already outdated. In ref. [8], these aspects were fixed for the purpose of a **CMBFAST** vs. **CAMB** comparison, which revealed indeed some very good agreement. But since the two codes have not been developed in an independent way, their relative agreement does not prove their absolute accuracy. Having an independent code would be the only way to check this absolute accuracy, and to try to push the precision further. **CAMB** offers an option to tune some “accuracy boost parameters”, but the user does not know exactly to which precision level a given setting corresponds. Without an independent and up-to-date code, **CAMB** ’s accuracy can only be calibrated “with respect to itself”, i.e. with respect to its infinite-accuracy-parameter limit.

These issues of friendliness, flexibility and control of accuracy were the main three motivations for developing a new code, the Comic Linear Anisotropy Solving System (**CLASS**)\(^1\). The fact that this code is also slightly faster than its competitors is less important, but does not hurt.

2. The **CLASS** way

2.1 User friendliness

**CLASS** is written in C, a language familiar to most scientific programmers, not very different from Fortran 90 but more diffuse. Compiling **CLASS** requires no specific version of the compiler, no special package or library. In the **Makefile**, the compiling command is set by default to **gcc -04**, i.e. to the GNU C compiler (installed on most computers) with

\(^1\)available at [http://class-code.net](http://class-code.net)
optimization level 4. We checked that the code is compatible with other compilers. After a
make class command, the code should be ready to run. On a multi-core PC, one should
try to run the code in parallel. This simply requires the code to be compiled with an
OpenMP option, e.g. gcc -O4 -fopenmp (with the version 4.2 or higher of gcc). The
OpenMP flag usually has a different name for other compilers.

The code can be executed with a maximum of two input files, e.g.

./class explanatory.ini chi2pl1.pre

The file with a ini extension is the cosmological parameter input file, and the one with a
pre extension is the precision file. Both files are optional: all parameters are set to
default values corresponding to the “most usual choices”, and are eventually replaced by
the parameters passed in the two input files. For instance, if one is happy with default
accuracy settings, it is enough to run with ./class explanatory.ini. Input files do not
necessarily contain a line for each parameter, since many of them can be left to default
value. The example file explanatory.ini is very long and somewhat indigestible, since
it contains all possible parameters, together with lengthy explanations. We recommend to
keep this file unchanged for reference, and to copy it in e.g. test.ini. In the latter file,
the user can erase all sections in which he/she is absolutely not interested (e.g., all the part
on isocurvature modes, or on tensors, or on non-cold species, etc.). Another option is to
create an input file from scratch, copying just the relevant lines from explanatory.ini.
For the simplest applications, the user will just need a few lines for basic cosmological
parameters, one line for the output entry (where one can specifying which power spectra
must be computed), and one line for the root entry (specifying the prefix of all output
files).

The syntax of the input files is explained at the beginning of explanatory.ini. Typ-
ically, lines in those files look like:

parameter1 = value1
free comments
parameter2 = value2 # further comments
# commented_parameter = commented_value

and parameters can be entered in arbitrary order. This is rather intuitive. The user should
just be careful not to put an “=” sign not preceded by a “#” sign inside a comment: the
code would then think that one is trying to pass some unidentified input parameter.

The syntax for the cosmological and precision parameters is the same. It is clearer
to split these parameters in the two files .ini and .pre, but there is no strict rule about
which parameter goes into which file: in principle, precision parameters could be passed
in the .ini, and vice-versa. The only important thing is not to pass the same parameter
twice: the code would then complain and not run.

The CLASS input files are also user-friendly in the sense that many different cosmological
parameter bases can be used. This is made possible by the fact that the code does not
only read parameters, it “interprets them” with the level of logic which has been coded in
the input.c module. For instance, the Hubble parameter, the photon density, the baryon
density and the ultra-relativistic neutrino density can be entered as:

\[ h = 0.7 \]
\[ T_{\text{cmb}} = 2.726 \quad \# \text{Kelvin units} \]
\[ \omega_{\text{b}} = 0.02 \]
\[ N_{\text{eff}} = 3.04 \]

(in arbitrary order), or as

\[ H_0 = 70 \]
\[ \omega_{\text{g}} = 2.5\text{e}^{-5} \quad \# g \text{ is the label for photons} \]
\[ \Omega_{\text{b}} = 0.04 \]
\[ \omega_{\text{ur}} = 1.7\text{e}^{-5} \quad \# \text{ur is the label for ultra-relativistic species} \]

or any combination of the two. The code knows that for the photon density, one should
pass one (but not more than one) parameter out of \( T_{\text{cmb}}, \omega_{\text{g}}, \Omega_{\text{g}} \) (where
small \( \omega \)‘s refer to \( \omega_i \equiv \Omega_i h^2 \)). It searches for one of these values, and if needed, it
converts it into one of the other two parameters, using also other input parameters. For
instance, \( \omega_{\text{g}} \) will be converted into \( \Omega_{\text{g}} \) even if \( h \) is written later in the file than
\( \omega_{\text{g}} \): the order makes no difference. Lots of alternatives have been defined. If the
code finds that not enough parameters have been passed for making consistent deductions,
it will complete the missing information with in-built default values. On the contrary, if
it finds that there is too much information and no unique solution, it will complain and
return an error. The code also writes a root_params.ini file in output, in which the user
can check which non-default values have been used. Like in CAMB, this file can be used as
an input file for another run.

In summary, the input syntax has been defined in such way that the user does not
need to think too much, and can pass his preferred set of parameters in a nearly informal
way.

Apart from easy compilation and easy input, user-friendliness is achieved by sticking to
well-known notations for all background and perturbation quantities (in the synchronous
and Newtonian gauge): namely, those from Ma & Bertschinger [1]. Most equations are
taken literally from this paper. The units are inverse mega-parsecs for wavenumbers \( k \) and
for the Hubble rate, and mega-parsecs for conformal time \( \tau \). For background densities \( \rho_i \)
and pressure \( p_i \), we use more unusual units. The goal is to be able to write the Friedmann
equation like

\[ H = \left( \sum_{i=0}^{N} \rho_{i} \right)^{1/2}. \]

Hence, everywhere in the code, the density \( \rho_{i} \) stands for \( [8\pi G \rho_i / 3] \), in units of squared
mega-parsecs (same for the pressure \( p_{i} \)). The output CMB multipoles \( C_l \)’s are dimen-
sionless (unlike those of CAMB which are in squared micro-Kelvins: hence, the $C_l^{TT}$'s from CLASS should be equal to those from CAMB divided by $[2.726 \times 10^6]^2$, if $T_{cmb}$ has been set to 2.726 K. Unless otherwise specified in the input file, the units and the definition of output quantities are written in comment lines at the top of each output file.

### 2.2 Flexibility

It is difficult to summarize in a few pages why we believe that CLASS is much easier to modify than other Boltzmann codes. Broadly speaking, we tried to achieve this goal by respecting three golden rules:

- **no hard coding.** Any feature or equation which could be true in one cosmology and not in another one should not be written explicitly in the code, and should not be taken as granted in several other places. Discretization and integration steps should be defined automatically by the code, instead of being set to something which might be optimal for minimal models, and not sufficient for other ones. Physical relations which are likely to change in different cosmological scenarios should be localized in a small number of well-identified places in the code, that the user can access easily. Any step which depends of these relations should be performed automatically rather than in a model-dependent way. We provide several concrete examples below.

- **clear structure.** The tasks of the code must be clearly separated: one module for the background evolution, another one for the thermodynamics evolution, another one for the perturbation evolution, etc. There should be no duplicate equations: a given physical assumption should be formulated in a single place.

- **dynamical allocation of all indices.** One might be tempted to decide that in a given array, matrix or vector, a given quantity is associated with an explicit index value. However, when modifying the code, extra entries will be needed and will mess up the initial scheme; the user will need to study which index is associated to which quantity, and possibly make an error. All this can be avoided by using systematically a dynamical index allocation. This means that all indices remain under a symbolic form, and in each, run the code attributes automatically a value to each index. The user *never* needs to know this value.

As a result of these rules, the CLASS files contain essentially no numbers, apart from:

- coefficient of physical equations, which can be transformed into input parameters if there is any reason that these coefficients could depend on the cosmology,

- unit conversion factors and constants, defined in the include/*.h files,

- default values of input parameters, defined in the input module.

Let us illustrate the flexibility of CLASS through a few examples.
Sampling steps. We need to integrate over time (or redshift) a set of background, thermodynamical and perturbation equations. For background equations, we use an adaptive Runge-Kutta integrator. Such integrators need to be called several times over small time steps. The edge of these time steps define the discrete values of time at which background quantities will be tabulated and stored, in view of being interpolated in other modules. CLASS infers the step sizes automatically from the rate at which background equations change. So, in a non-standard cosmological model where something special would happen at a given time, the step size would automatically decrease around that time. The same occurs with the perturbation equations when the user uses the same Runge-Kutta integrator, which is only an option (the default integrator for perturbations, called ndf15, is described in [10]). For thermodynamical quantities, the equations are solved by the RECFAST module [12, 13], which we did not change much (apart from small modifications leading to smoother results, as mentioned in [14]). RECFAST chooses some step sizes for integrating and sampling thermodynamical quantities during recombination. But at low redshift, the evolution is modified according to some theoretical ansatz for the enhanced free electron fraction $\Delta x_e(z)$ during reionization. This function is chosen by the user. For whatever function $\Delta x_e(z)$, CLASS finds automatically the step sizes in redshift space which are sufficient for capturing the evolution $x_e(z)$, given a dimensionless tolerance parameter reionization_sampling (each step size is given by $(d \ln x_e/dz)^{-1}$ times this parameter).

A similar logic is used by the code for defining the discrete times at which the source functions $S(k, \tau)$ (needed to compute the observable spectra) are sampled; or for defining the momentum bins in which the phase-space distribution of each massive neutrinos and other non-cold relics are calculated [14]. As a result, the user is free to change most physical equations (e.g. Friedmann and Einstein equations, reionization function, phase-space distribution of neutrinos) without ever needing to think about sampling issues. The only step sizes which are not found automatically in CLASS v1.0 are those of wavenumbers $k$ (for source functions $S(k, \tau)$ and transfer functions $\Delta l(k)$) and of multipoles $l$ (for transfer functions $\Delta l(k)$ and harmonic spectra $C_l$'s), which are either linear, logarithmic or a combination of the two. Those step sizes are controlled by dimensionless parameters which can all be tuned in the precision parameter file.

Reionization. After computing the electron fraction $x_e(z)$ imposed by Helium/Hydrogen recombination using RECFAST, the code must assume a given enhancement $\Delta x_e(z)$ of this function at low redshift, accounting for reionization when the first stars form. By default, this function is often assumed to be a double step described by two hyperbolic tangents, corresponding to Helium and Hydrogen reionization. For each of these two steps, the user should pass input parameters describing the mean redshift of the transition, its width, its amplitude, and the redshift $z_{\text{max}}$ above which we decide to neglect reionization. In CLASS, this function is not hard-coded, in the sense that the user should just write inside the function thermodynamics_reionization_function() his favorite ansatz for $\Delta x_e(z)$, depending on an arbitrary number of input parameters. The code will do the rest automatically: finding appropriate sampling steps, ensuring the continuity of the total $x_e(z)$ around $z_{\text{max}}$, finding the relation between the reionization redshift and the optical depth.
(as in CAMB, any of the two can be passed in input, the other one will be inferred). Hence, studying alternative reionization models is trivial with CLASS.

**Non-standard massive neutrinos and non-cold dark matter relics.** Other publicly available Boltzmann codes include one or several massive neutrino eigenstates, with one or several masses, but sharing a unique phase space distribution $f(q)$ taken to be an exact Fermi-Dirac. Changing this assumption (for studying non-standard massive neutrinos with chemical potentials or non-thermal distortions, light or heavy sterile neutrinos, warm dark matter candidates, etc.) requires a large number of modifications to these codes, in which the Fermi-Dirac distribution is hard-coded. The user interested in above models should redefine $f(q)$ and its derivative $[d \ln f/d \ln q]$ in several places, eventually switch to a finer sampling of background and perturbed quantities in $q$ space, redefine the relation between mass and density, redefine some approximation schemes in the relativistic/non-relativistic limits, etc. In CLASS, for each non-cold species $i$, the user should only modify in a single place the function $f_i(q)$. This function can depend on an arbitrary number of parameters, which are easy to pass from the input file. All the rest is done automatically: finding the sampling in $q$-space using some dimensionless tolerance parameters; finding the derivative $[d \ln f/d \ln q]$; finding the mass-to-density relation; using approximation schemes, etc. The user is even free to specify in the input file that for one or several of these species, the distribution function should be read in a file rather than in a function. This is useful for warm dark matter candidates, which phase-space distribution function can be non-trivial and computable only numerically. All these features are described in more detail in [14].

**Adding new species in the code.** CLASS v1.0 includes the following species with the following labels: photons ($g$, compulsory); baryons ($b$, compulsory); cold dark matter ($cdm$, compulsory only in synchronous gauge); massless neutrinos and other ultra-relativistic species ($ur$, optional); arbitrary number of massive neutrinos and other non-cold dark matter relics ($ncdm$, optional); a non-perfect fluid with constant linear equation of state and sound speed ($fl$, optional); and a cosmological constant ($lambda$, optional). Including more species is trivial thanks to the dynamical index allocation. The indices of the above quantities are never hard-coded; for instance, in the vector of all background quantities at a given time, the photon density reads $\text{background}[\text{ba.index}_\text{bg}_\text{rho}_g]$, where “index bg” means “one index in the background vector”, “rho_g” means “photon density”, and “ba.” is necessary because this index is defined inside the background structure, usually abbreviated as ba in the code. The user willing to code more species should simply try to duplicate any reference to another species, and adapt the names and the equations of motion. For instance, let us assume that one wants to introduce another imperfect fluid with different properties from $fl$, and call it e.g. new. The easiest would be to search for all occurrences of the letters $fl$ in the modules input, background, perturbations, to duplicate them and to replace $fl$ by new. Then, the user can simply change the equations of motion for new, and define new input parameters specifying the properties of the fluid in the same place where the $fl$ parameters $\Omega_{fl}, w_{fl}$ and $cs^2_{fl}$ are defined. By doing so, the user will be guided to define new variables for the indices, e.g. $index_{bg}_\text{rho}_{new}$.
for the background density, or `index_pt_delta_new` for the density perturbations. All equations will be written in terms of these index names, without needing to every worry about changes in the number of equations, about explicit values of the indices, or about the order in which quantities are arranged inside vectors and lists. The user will also be guided to define a flag `has_new` which will be set to “true” inside the `input` module only when the input value of the `new` density is non-zero. All references to the new fluid in the code will be inside a condition “if (has_new == _TRUE_) {...}”. This offers two advantages. First, when a species is not present, the code ignores completely the lines referring to it. So, it does not slow down the code to implement hundreds of new species and to run with only a few of them being non-zero. Second, if the density of e.g. the `new` component is null, its index values are never assigned, but also never needed by the code.

Adding new approximation schemes in the perturbation module. All Boltzmann codes use approximation schemes in order to reduce the time spent in the perturbation integration. The most famous one is the unavoidable tight-coupling approximation. In CLASS, the notion of approximation is formalized in such way that defining other approximation schemes or skipping some of them is straightforward. Any approximation scheme has a nickname: examples are the Tight-Coupling Approximation `tca`, the Ultra-relativistic Fluid Approximation `ufa`, the Radiation Streaming Approximation `rsa` (all described in [10]) or the Non-Cold Dark Matter Fluid Approximation `ncdmfa` (described in [13]). Each of these approximation are associated with “status flags” (e.g. `tca_on`, `tca_off`) and “method flags” (e.g. `tca_class`, `tca_camb`, or just `tca_none` if this approximation should never be used). One of the “method flags” is passed in input as a precision parameter, to state which equations should be used when the approximation is turned on. In the routine `subroutine_approximations()`, one defines the condition under which an approximation should be “on” or “off”. For instance, the Ultra-relativistic Fluid Approximation should be “on” only if there are non-relativistic species in the problem (this depends on the flag `has_ur`), and if the ratio $k/aH$ exceeds a threshold value defined as a precision parameters. Before integrating perturbations for a given wavenumber $k$, the code will automatically check how many approximations need to be switched on or off for this wavenumber, and will find each switching time $t_i$. The integration is then performed inside each interval $[t_i, t_{i+1}]$ over which the approximation scheme does not change. It means that at each switching time $t_i$, the system of differential equations is redefined with a new size and new initial values. With this logic, it is very simple to define new approximations, or to modify the conditions under which an approximation should be switched on or off, or finally to define a new method corresponding to different equations when the approximation is switched on. Again, if one needs to code such a new approximation, the easiest way is to search for all occurrence of one other nickname (e.g. `ufa`), to duplicate these occurrences, adapt the name, and adapt the conditions and physical equations defining this approximation.

Adding new source functions, new transfer functions, etc... In the jargon of the code, a source function is just a function $S(k, \tau)$ of wavenumber and time which can be inferred from background, thermodynamical and perturbed quantities, and should be
stored during the execution of the perturbation module in order to be used by other modules. In fact, what remains in memory after the integration of perturbed quantities is only a list of tabulated source function. For CMB spectra, we need a temperature and a polarization source function. For CMB lensing and cosmic shear, we need another source, equal to the Newtonian gravitational potential: \( S(k, \tau) = \phi(k, \tau) \). For the matter power spectrum, we must store one more source function equal to the matter density perturbation: \( S(k, \tau) = \delta_m(k, \tau) \). For outputting “density transfer functions”, i.e. the density of each species \( i \) at time \( \tau \) relative to the one at initial time, one should store several extra source functions equal to \( S(k, \tau) = \delta_i(k, \tau)/\delta_i(k, \tau_{ini}) \) for each \( i \). In the code, all source functions are defined with a dynamical index and associated to a flag. For instance, if the CMB spectra are needed (which is not always the case), the flag for the temperature source \texttt{has_source_t} will be set to true, and any statement regarding this particular source will be inside a condition “if \( \texttt{has_source_t == \_TRUE\_} \) {...}”. Inside these regions, the temperature source, which is one element in the vector of all sources, will be referred as \texttt{source[pt.index_tp_t]} where \texttt{index_tp_} means “index for the type of source function” and \texttt{t} means “temperature”; this index is defined inside the perturbation structure, usually called \texttt{pt}. Once more, following this logic, defining any other source function would be easy. The user would need to track all occurrences of e.g. \texttt{has_source_t} or \texttt{index_tp_t}, duplicate the corresponding lines, and replace them with the name and definition of the new source functions.

**Other examples.** The very same logic based on flags and dynamical indices is found everywhere in the code, in the declaration of modes (scalars, tensors, etc.), of initial conditions (adiabatic, different isocurvature types), of transfer functions \( \Delta_l(k) \), of observables (e.g. \( C_l^{XX'} \) where \( X \) stands for temperature, polarization, lensing potential, etc.)... This means that there are infinite ways to extend the code easily, using only the search and the copy/paste commands, plus the many explanatory comments written throughout the C files, and some basic theoretical knowledge of cosmological perturbations.

### 2.3 Control of accuracy

The number of accuracy parameters in a Boltzmann code is surprisingly large. In the \texttt{CAMB} input file, only a few “accuracy boost” parameters and accuracy flags are visible; but they control many other quantities. In \texttt{CLASS}, we grouped all accuracy parameters within a single structure, the “precision structure” declared in the \texttt{include/common.h} file. As far as we remember, the user will not find a single accuracy parameter declared and hard-coded locally in any function or module. All these values are initialized in the \texttt{input} module and eventually overwritten by declarations in the input file <...>.pre. This is convenient, since the user willing to tune the accuracy of \texttt{CLASS} does not need to search for precision variables dispersed throughout the code. The code is released together with a few precision files which have been calibrated using a rigorous process (see [1] for more details).
2.4 Speed

Speaking of the speed of the code only makes sense by referring to a given precision level. The speed of CLASS compared to that of CAMB is presented in [11] for a few accuracy settings and in the minimal ΛCDM context. In this case, we found that CLASS is faster by a factor 2.5. This difference seems to result from the sum of several improvements, rather than from a single trick. Progress on the side of approximation schemes and integrators is described in [10] for minimal ΛCDM, and in [14] for models with massive neutrinos. Other improvements related to numerical strategies and algorithms are impossible to summarize in a paper, since the code was really written independently: inevitably, the differences with respect to CAMB are plethoric, and it is a difficult task to identify which one are the most decisive for speeding up the code.

The user should always try to compile with the most aggressive optimization flags and with the OpenMP option, in order to be able to run in parallel. We checked that if the number of OpenMP threads $N$ is not too large, the running time scales almost linearly with $1/N$. If several runs are performed with the same precision parameters, the code will find automatically that it can read spherical Bessel functions in a file where they have been written previously, in order to save a bit of time (a similar feature was implemented in CMBFAST). For usual precision settings, the computation of Bessel functions is anyway very fast. When running many occurrences of CLASS in a loop or in a parameter extraction code, one needs to compute the Bessel function only once, and keep them in memory (this is easy to achieve, as explained in the next section).

3. Structure of the code

3.1 Directories

After downloading CLASS, one can see that files are split between the following directories:

- **source/** contains the C files for each CLASS module, i.e. each block containing some part of the physical equations and logic of the Boltzmann code.

- **tools/** contains purely numerical algorithms, applicable in any context: integrators, simple manipulation of arrays (derivation, integration, interpolation), Bessel function calculation, quadrature algorithms, parser, etc.

- **main/** contains the main module class.c with the main routine class(...), to be used in interactive runs (but not necessarily when the code is interfaced with other ones).

- **test/** contains alternative main routines which can be used to run only some part of the code, to test its accuracy, to illustrate how it can be interfaced with other codes, etc.

- **include/** contains all the include files with a .h suffix.

- the root directory contains the Makefile and some example of input files.
3.2 Logic of each module

All important modules are inside the source/ directory. The structure of CLASS is formalized in a rigorous way. It relies on 11 modules that we will call generically module_i, where i is supposed to be a label between 1 and 11. A module coincides with a C file source/module_i.c and an include file include/module_i.h. It is also associated with a structure that we will call generically structure_i. Finally, the module module_i.c contains at least two functions module_i_init.c(...) and module_i_free.c(...) which are the essential ones.

The overall logic of CLASS is summarized by the following two principles:

1. before the function module_i_init() has been executed, structure_i contains some input parameters relevant for this module. These input parameters have been written inside the structure when the first module (the input.c module) has been executed. For instance, in the case of the perturbation.c module, the structure perturbation contains information about which modes (scalar, tensors, ...) or which initial conditions (adiabatic, isocurvature...) are present.

2. after the module module_i_init() has been executed, structure_i contains everything that other modules need to know. Intermediate quantities only computed internally in the module and not stored in structure_i are lost. For instance, after the execution of the perturbation.c module, the structure perturbation contains a table of source functions $S(k, \tau)$.

3. when the various information stored in structure_i is not useful anymore, a call to module_i_free.c(...) frees the memory dedicated to structure_i.

The arguments of module_i_init() should be: the precision structure precision which contains all accuracy parameters for the whole code; the structures of the previous modules structure_1, ..., structure_(i-1) which contain relevant information; and the structure structure_i, which contains a few input parameter before the execution of this function, and is entirely filled afterward. So, formally, executing the whole code amounts in calling:

```
module_1_init(precision, structure_1)
module_2_init(precision, structure_1, structure_2)
...
module_11_init(precision, structure_1, structure_2, ..., structure_11)
/* done, now free everything */
module_11_free(structure_11)
...
module_1_free(structure_1)
```

The main routine in class.c is therefore extremely compact (it looks a bit longer than above only because of error management, as explained below).
Each module contains more functions than just \texttt{module\_i\_init()} and \texttt{module\_i\_free()}. These other functions are always named \texttt{module\_i\_<...>\(\)} (i.e., they always start with the name of the module). They can be divided in two categories:

1. functions used only internally by each module. In the \texttt{module\_i\.c} file, these functions are always written after \texttt{module\_i\_init()} and \texttt{module\_i\_free()}.

2. functions which can be called by other modules. Typically, when a module \( j \) needs quantities from an earlier module \( i < j \), it can either read the information directly inside \texttt{structure\_i}, or call a function inside \texttt{module\_i\.c} which knows how to read quantities in \texttt{structure\_i} and returns a pointer to the needed information. For instance, after the \texttt{background\.c} module has been executed, the \texttt{background} structure contains interpolation tables for all background quantities. If another module needs background quantities at a given conformal time \( \tau \), instead of entering into the structure \texttt{background}, this module can call the function \texttt{background\_at\_tau(tau, ...)}), which will return the desired quantities. Functions called by other modules are always written before \texttt{module\_i\_init()} in the \texttt{module\_i\.c} file, since they are those deserving more visibility.

### 3.3 The CLASS backbone

Executing CLASS amounts to execute eleven functions of the type \texttt{module\_i\_init()}, for the following eleven modules:

1: \texttt{input\.c}

2: \texttt{background\.c}

3: \texttt{thermodynamics\.c}

4: \texttt{perturbations\.c}

5: \texttt{bessel\.c}

6: \texttt{transfer\.c}

7: \texttt{primordial\.c}

8: \texttt{spectra\.c}

9: \texttt{nonlinear\.c}

10: \texttt{lensing\.c}

11: \texttt{output\.c}
So, the main() function is identical to the scheme described in the previous section, with the above module names, called in the above order. This order is defined by the fact that a function module_i_init() needs the structures structure_j for j < i to be already filled. There is actually a little bit of freedom. For instance, the only argument of the bessel_init() function is the precision structure, initialized in the input.c module and passed to all module_i_init() functions; as well as the bessel structure, for which input parameter are initialize as usual in the input.c module, and the rest is filled within the bessel_init() function. The bessel structure is used only by the transfer.c module. This means that bessel_init() could actually be called at any time between input_init() and transfer_init(). But there is not as much flexibility for other modules, since most of them really use the structure of the previous module (i − 1), while their own structure is requested by the next module (i + 1).

This structure implements a very clear separation of the physical tasks. The user can easily know where is the region that he eventually needs to modify. Moreover, somebody can only be interested in computing the background evolution, or the thermodynamical evolution, or the transfer functions. In this case, instead of using the full sequence of modules, it is possible to execute only the first few modules. The alternative main() functions contained in the test/ directory offer such examples.

3.4 Quick glance at each module

- input.c, the first module, is a bit different from the other ones. It is the only module not associated with a structure (an input structure would have been useless), and it is also the only module which has two _ini() functions. The first one, input_init_from_arguments(), is the first function called by CLASS when the code is run interactively using the main routine in main/class.c. It reads a maximum of two arguments, the <...>.ini input file and the <...>.pre precision file. It sets all input and precision parameters to default values, and then eventually replace some of them with the values indicated in the file(s) (but the same parameter cannot be reset twice in the files, otherwise the code will complain). As we said earlier, the fact that input parameters go into <...>.ini and precision parameters into <...>.pre is not a strict rule: in practise the code just considers the sum of the two files, and everything could be passed in a single file. The second function, input_init(), is the first function to be called when the code is interfaced with another one. In this case, input parameters are not read from the arguments of the function, but from a structure called file_content which contains the same information as the files, in roughly the same format. So, if CLASS is to be run e.g. within a parameter extraction code, this code should write input/precision parameter inside a file_content structure, and then call input_init(). Actually, the role of the function input_init_from_arguments() is to convert the content of the input files into a structure file_content, and then to call input_init(). The list of arguments for these two functions is long: indeed, they need a pointer towards each of the structures of all other modules, in order to initialize them. Since input.c is the
only module not having its own structure, it is also the only one without a \_free() function.

- \texttt{background.c} simply solves the background equations (in particular, the Friedmann equation), and stores into its structure \texttt{background} an interpolation table for all background quantities as a function of time. Other modules often call the function \texttt{background\_at\_tau()} which returns these quantities interpolated at some conformal time $\tau$, and the function \texttt{background\_tau\_of\_z()} which converts a value of redshift into a value of conformal time. This module also stores in the \texttt{background} structure useful background-related quantities like the age of the universe.

- \texttt{thermodynamics.c} solves for the thermodynamical evolution with RECFAST, corrects it for reionization, and stores into its structure \texttt{thermo} an interpolation table for all thermodynamical quantities as a function of redshift. Other modules often call the function \texttt{thermodynamics\_at\_z()} which returns these quantities interpolated at some redshift $z$. This module also stores in the \texttt{thermo} structure useful quantities like the recombination time, reionization time, reionization optical depth if it was not passed in input, reionization redshift if it was not passed in input, sound horizon at recombination, etc.

- \texttt{perturbation.c} solves the evolution of all perturbations, and stores the source functions $S(k, \tau)$ in a table inside its structure \texttt{perturbs}. Note that when perturbations are integrated for a given wavenumber, background and thermodynamical quantities are not evolved another time: they are interpolated using the functions \texttt{background\_at\_tau()} and \texttt{thermodynamics\_at\_z()}, in order to enforce a clear separation of the tasks. As a result, the Friedmann equation only appears in a single line in the \texttt{background.c} module.

- \texttt{bessel.c} computes spherical Bessel functions and stores them in its structure \texttt{bessels}. We already mentioned that this module could also be called earlier, just after the input module. Actually, when \texttt{CLASS} is to be run several time within e.g. a parameter extraction code, always with the same accuracy parameters, one should keep the bessel functions in memory and not call this module each time. In practise, this can be done in the following way: call once \texttt{input\_init()} and \texttt{bessel\_init()}; then, for each new model, call again \texttt{input\_init()} and all other \_init() functions in each module, but not \texttt{bessel\_init()} again. Free all structures by calling in reverse order all \_free() functions, but not \texttt{bessel\_free()}. Finally, when everything is finished, do a final call of \texttt{bessel\_free()}.

- \texttt{transfer.c} computes transfer functions $\Delta_l(k)$ (by convolving source functions and Bessel functions), and stores them in its structure \texttt{transfers}.

- \texttt{primordial.c} computes the primordial power spectra (for each mode and initial condition) and stores them in its structure \texttt{primordial}. In \texttt{CLASS v1.0}, only simple analytical formulas are implemented in this module, but in the future, it will be able to call an inflation simulation module if requested.
• **spectra.c** computes observable power spectra out of source functions, transfer functions and primordial spectra, and stores them in its structure **spectra**.

• **nonlinear.c** gives an estimate of the non-linear version of the previous spectra, according to some scheme chosen by the user, and stores them in the structure **nonlinear**. Very soon, a renormalization scheme will be available, as well as other methods.

• **lensing.c** computes lensed temperature and polarization CMB spectra, using the unlensed spectra and the CMB lensing potential spectrum, and stores them in the structure **lensing**. **CLASS** computes the lensed CMB spectra from all-sky correlation functions \([\mathcal{E}]\), i.e. with the same method as **CAMB**, but with a different numerical implementation written by S. Prunet, based on quadrature weights.

• **output.c** just writes the output in some files. This module does not need to be called when **CLASS** is used inside another code, e.g. a parameter extraction code. Otherwise, the function **output_init()** is the last one called in the main routine, before freeing structures. Actually, all fields in the **output** structure can be freed at the end of **output_init()**, so there is no need to call **output_free()**, which is not even defined.

### 3.5 Error management

Error management is based on the fact that all functions are defined as integers returning either **_SUCCESS_** or **_FAILURE_**. Before returning **_FAILURE_**, they write an error message in the structure of the module to which they belong. The calling function will read this message, append it to its own error message, and return a **_FAILURE_**; and so on and so forth, until the main routine is reached. This error management allows the user to see the whole nested structure of error messages when an error has been found. The structure associated to each module contains a field for writing error messages, called **structure_i.error_message**. So, when a function from a module \(i\) is called within module \(j\) and returns an error, the goal is to write in **structure_j.error_message** a local error message, and to append to it the error message in **structure_i.error_message**. These steps are implemented in a macro **class_call()**, used for calling whatever function:

```
class_call(module_i_function(...,structure_i),
           structure_i.error_message,
           structure_j.error_message)
```

So, the first argument of **call_call()** is the function we want to call; the second argument is the location of the error message returned by this function; and the third one is the location of the error message which should be returned to the higher level. The user will find in **include/common.h** a list of additional macros, all starting by **class_...()**, which are all based on this logic (for instance, the macro **class_test()** offers a generic way to
return an error in a standard format if a condition is not fulfilled. A typical error message from CLASS looks like:

```
Error in module_j_function1
=> module_j_function1 (L:340) : error in module_i_function2(...) => module_i_function2 (L:275) : error in module_k_function3(...) ...
=> module_x_functionN (L:735) : your choice of input parameter blabla=30 is not consistent with the constraint blabla<1
```

where the L’s refer to line numbers in each file. These error messages are very informative, and are built almost entirely automatically by the macros. For instance, in the above example, it was only necessary to write inside a `class_test()` macro the words `your choice of input parameter blabla = %g is not consistent with the constraint blabla < %g`, `blabla, blabla_max`. All the rest was added step by step by the various `class_call()` macros.

4. What is already there, and what is next?

In terms of cosmological species, as already mentioned previously, CLASS v1.0 includes photons (g, compulsory); baryons (b, compulsory); cold dark matter (cdm, compulsory only in synchronous gauge); massless neutrinos and other ultra-relativistic species (ur, optional); arbitrary number of massive neutrinos and other non-cold dark matter relics (ncdm, optional); a non-perfect fluid with constant linear equation of state and sound speed (fl, optional); and a cosmological constant (lambda, optional). Initial conditions can be an arbitrary mixture of an arbitrary number of arbitrarily correlated adiabatic and isocurvature modes. For each auto-correlation and cross-correlation spectra, the user can enter an amplitude, a tilt and a running. Non-cold dark matter relics can be tuned with a variety of options described in the `explanatory.ini` input file and illustrated in [14]: free masses, temperatures, chemical potentials for flavor species, mixing angles (used to go from the flavor basis to the mass basis), degeneracy parameters, and as mentioned above, possibility to change the phase-space distribution, using an analytical expression or tabulated values in a file.

The code includes tensor modes, which have not been tested thoroughly in v1.0 like the scalar ones (progress is still expected in this direction). It computes the CMB anisotropy spectra for the products $TT$, $TE$, $EE$, $BB$, as well as the auto-correlation and cross-correlation spectra involving the CMB lensing potential $\phi$. It computes also the lensed anisotropy spectra. It can output the matter power spectrum, today or for an arbitrary list of redshifts passed by the user; same for the density transfer functions of each species.

Coding the Newtonian gauge is almost finished, but only the synchronous gauge is operational in version 1.0. The next steps will consist in improving the way to deal with tensors, in making the Newtonian gauge operational, and in coding open/closed models. We wish to release
interfaces between CLASS and the parameter extraction codes CosmoMC, CosmoPMC and MultiNest. It would be easy to add more general parametrizations for the reionization history, and to adapt to CLASS our CAMB inflationary module [16, 17]. We will release very soon a renormalization algorithm embedded in the nonlinear.c module, in which we will also implement non-linear approximations like HALOFIT [18]. It will finally be easy to add more observables, like e.g. correlation functions and power spectra for cosmic shear survey.

Acknowledgments

We wish to thank François Bouchet for inspiring this project, as well as Martin Bucher, Damien Girard, Jan Hamann and Alain Riazuelo for useful input in its earliest stages. Throughout the realization of CLASS, Karim Benabed and Simon Prunet provided essential help and support. Benjamin Audren and Simon Prunet coded respectively the renormalization and lensing algorithms. In the past ten months, Thomas Tram boosted the last coding steps without sparing time and efforts. Enthusiastic moral support from several friends and colleagues was very much appreciated.

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