Neutrino decoupling including flavour oscillations and primordial nucleosynthesis

Julien Froustey, Cyril Pitrou, and Maria Cristina Volpe

Abstract. We revisit the decoupling of neutrinos in the early universe with flavour oscillations. We rederive the quantum kinetic equations which determine the neutrino evolution based on a BBGKY-like hierarchy, and include the full collision term. We focus on the case of zero chemical potential and solve these equations numerically. We also develop an approximate scheme based on the adiabatic evolution in the matter basis. In fact, the large difference between the oscillations and cosmological time scales allows to consider averaged flavour oscillations which can speed up the numerical integration by two orders of magnitude, when combined with a direct computation of the differential system Jacobian. The approximate numerical scheme is also useful to gain more insight into the physics of neutrino decoupling. Including the most recent results on plasma thermodynamics QED corrections, we update the effective number of neutrinos to $N_{\text{eff}} = 3.0440$. Finally we study the impact of flavour oscillations during neutrino decoupling on the subsequent primordial nucleosynthesis.
1 Introduction

The hot Big Bang model predicts that several physical phenomena take place when the Universe temperature reaches the MeV scale. Long before the temperature reaches this threshold, the Universe consists in a plasma of coupled photons, electrons, positrons, neutrinos and antineutrinos at equilibrium. But when the temperature drops below $\sim 2\,\text{MeV}$, weak interactions become too weak to keep (anti)neutrinos in thermal contact with the electromagnetic plasma: neutrinos decouple and form the cosmic neutrino background, a key prediction of the standard cosmological model. Soon after, the temperature decreases below the electron mass and $e^\pm$ pairs annihilate into photons, reheating the electromagnetic plasma compared to the bath of neutrinos. If one considers those two events to be well-separated in time, entropy conservation leads to the standard ratio for the temperatures of neutrino and photon backgrounds, $T_\gamma / T_\nu = (11/4)^{1/3}$.
However, the overlap between neutrino decoupling and \( e^\pm \) annihilations, known as *incomplete neutrino decoupling*, leads to slightly non thermal neutrino spectra, and to an increased neutrino energy density (both typically of order 1%), which is usually described by an effective number of thermalised neutrinos \( N_{\text{eff}} \) departing from 3 [1–5]. An accurate prediction of the neutrino spectra requires to take into account multiple physical effects, including QED radiative corrections to the plasma equation of state [3, 6, 7]. Furthermore, the small but non-vanishing masses of neutrinos and their mixings are the cause of the famous neutrino oscillations, given that mass eigenstates differ from flavour eigenstates [8]. Neutrino mixings can thus influence the process of neutrino decoupling, in particular the flavour dependence of spectral distortions. Flavour oscillations have already been included in calculations of neutrino decoupling [9–12], yet approximating some collision terms for computational purposes, either neglecting off-diagonal components or replacing them by damping approximations.

A robust and precise prediction of the consequences of incomplete neutrino decoupling is crucial since neutrinos impact many cosmological stages:

1. During Big-Bang Nucleosynthesis (BBN), neutrinos control neutron/proton conversions as they participate to weak interactions, and the frozen neutron abundance subsequently affects nuclear reactions and light element relics [5, 13].

2. During the Cosmic Microwave Background (CMB) formation, the free streaming of neutrinos is crucial to predict the CMB angular spectrum. Most famously, the neutrino shear is directly responsible for a difference between the two Bardeen potentials [14], which then disappears in the matter era. Also, the value of \( N_{\text{eff}} \) affects the cosmological expansion, and thus also the radiative transfer of CMB. From these effects, CMB alone can be used to place constraints on \( N_{\text{eff}} \) (\( N_{\text{eff}} = 2.99 \pm 0.17 \) at 68% confidence [15]) or in combination with BBN constraints on primordial light elements [13].

3. In the late universe, neutrino free streaming also affects structure formation, via its effect on the growth of perturbations. This is used to place the constraint \( \sum_\nu m_\nu < 0.12 \text{ eV} \) (see e.g. [15]) on the sum of neutrino masses.

It is striking that neutrino masses play a key role in both the earliest stage 1 and the latest stage 3 for very different reasons. In stage 1, neutrino oscillations, which are due to small neutrino mass-squared differences and mixing angles, affect the non-thermal part of the spectra, as they lead to less distortion in electron-type neutrinos and more distortion in other types than if there were no oscillations at all. Also oscillations lead to a mild modification of \( N_{\text{eff}} \). In stage 3, and due to cosmological redshifting, all neutrinos undergo at some point a transition from being very relativistic (they behave gravitationally like decoupled photons) to being non-relativistic (they then behave like cold dark matter). This transition depends only on neutrino masses and not on mixing angles, since frozen neutrino spectra inherited from stage 1 are generated incoherently in the mass basis. Finally, stage 2 would also be affected beyond the standard cosmological model, if we were to consider exotic physics with increased neutrino self-interactions, so that they would still behave effectively as a perfect fluid around CMB formation [16, 17].

This interplay between the various cosmological eras implies that it is crucial to understand neutrino decoupling as precisely as possible, in order to use these predictions as initial conditions for the subsequent eras. For instance, current constraints from CMB on cosmological parameters [15] were placed using \( N_{\text{eff}} = 3.046 \) when solving numerically for the linear evolution of cosmological perturbations.
For stage 1, the inclusion of neutrino masses and mixings requires the numerical solution of the full neutrino quantum kinetic equations (QKEs). Various approaches were used to derive them, e.g., a perturbative expansion of the density matrix [18], or the Closed-Time-Path (CTP) formalism for the two-point function [19, 20]. A hierarchy can be built for the neutrino density matrix, corresponding to a relativistic generalization of the Bogoliubov-Born-Green-Kirkwood-Yvon (BBGKY) equations. This formalism has been applied to derive the most general mean-field equations for astrophysical neutrinos [21, 22], introducing notably neutrino-antineutrino pairing correlations and wrong helicity contributions due to the neutrino mass.

The main goal of this work is to reevaluate the standard value of $N_{\text{eff}}$ and the distorted neutrino spectra, including all relevant effects to reach a $10^{-4}$ precision, also including the effect of neutrino masses. To this aim we first derive the neutrino QKEs, extending the work of [21] for astrophysical environments, and implement two-body collisions in an isotropic and homogeneous environment, including neutrino self-interactions. Then we numerically solve these QKEs, but also present an approximate solution where an adiabatic evolution is considered, exploiting the different timescales of collisions, mean-field and mixing terms, nearby neutrino decoupling. This procedure allows to maintain the required precision while decreasing substantially the computation speed, gaining some physical insight on the role of flavours oscillations in neutrino decoupling. The numerical results we present correspond to the case of zero chemical potential. Finally we investigate the impact of neutrino masses and mixings on BBN predictions, implementing the contribution of the numerical solution of the full QKEs, going beyond works available in the literature [9, 24, 25].

The manuscript is structured as follows. The formalism used to determine the neutrino evolution in the early universe is described in section 2, several technical details being gathered in appendices. In section 3, the approximate scheme used in computations is presented; whereas results for the key observables (neutrino spectra and $N_{\text{eff}}$) are given in section 4. Finally section 5 is devoted to the effect that incomplete neutrino decoupling has on the nucleosynthesis, and to the comparison with previous results [5] obtained without taking into account neutrino masses and mixings. Natural units ($\hbar = c = k_B = 1$) are used throughout the manuscript.

2 Derivation of quantum kinetic equations

In this section, we present a derivation from first principles of the neutrino quantum kinetic equations, which generalize the Boltzmann kinetic equation for distribution functions to account for mixing. We present the BBGKY hierarchy that was historically derived for a non-relativistic $N$–body system and heavily used in nuclear physics [26–30], but that can also be applied to a relativistic system such as neutrinos and antineutrinos in the early universe. We extend the work done in [21], where the BBGKY formalism was applied to derive extended mean-field equations for astrophysical applications, and include the collision term. Neutrino QKEs were previously derived using different approaches (see e.g. [18, 20]).

\footnote{Note that helicity, or spin coherence, [19, 23], that requires anisotropy, is not considered in the present work.}
2.1 BBGKY formalism

The exact evolution of a \(N\)-body system under the Hamiltonian \(\hat{H}\) is given by the Liouville-von Neumann equation for the many-body density

\[
\frac{i}{\hbar} \frac{d \hat{D}}{dt} = [\hat{H}, \hat{D}] ,
\]

where \(\hat{D} = |\Psi \rangle \langle \Psi|\), with \(|\Psi\rangle \rangle \) the quantum state, from which we define the \(s\)-body reduced density matrices,

\[
\hat{\rho}^{(1\cdots s)} \equiv \frac{N!}{(N-s)!} \text{Tr}_{s+1\cdots N}\hat{D} ,
\]

with components (we drop the superscript \((1\cdots s)\), redundant with the number of indices):

\[
\hat{\rho}_{ij_1\cdots j_s}^{(1\cdots s)} \equiv \langle \hat{a}_{i}^{\dagger} \hat{a}_{j_1} \cdots \hat{a}_{j_s} \rangle ,
\]

where the indices \(i, j\) label a set of quantum numbers (species \(\phi\), momentum \(\vec{p}\), helicity \(h\)) which describe a one-particle quantum state. For instance,

\[
\sum_i \hat{a}_i^\dagger = \sum_{\phi_i} \sum_{h_i} \int [d^3 \vec{p}_i] \hat{a}_i^\dagger (\vec{p}_i, h_i) \quad \text{with} \quad [d^3 \vec{p}_i] \equiv \frac{d^3 \vec{p}_i}{(2\pi)^32E_i} .
\]

The central object is the one-body reduced density matrix [18],

\[
\hat{\rho}_i^j \equiv \langle \hat{a}_i^\dagger \hat{a}_j \rangle ,
\]

whose diagonal entries correspond to the standard occupation numbers.

The Hamiltonian for this system is given by the sum of the kinetic and the two-body interaction terms,

\[
\hat{H} = \hat{H}_0 + \hat{H}_{\text{int}} = \sum_{i,j} \hat{t}_j^i \hat{a}_i^\dagger \hat{a}_j + \frac{1}{4} \sum_{i,j,k,l} \hat{v}_{ijl}^{ik} \hat{a}_{i}^\dagger \hat{a}_{k}^\dagger \hat{a}_l \hat{a}_j .
\]

The interaction matrix elements are fully anti-symmetrized by construction:

\[
\langle ik | \hat{H}_{\text{int}} | jl \rangle \equiv \hat{v}_{ijl}^{ik} = -\hat{v}_{jil}^{ki} = \hat{v}_{jli}^{ki} .
\]

This set of definitions ensures proper transformation laws under a unitary transformation \(\psi' = U_t \psi\): all lower indices are covariant while upper indices are contravariant, namely,

\[
\hat{g}^{\alpha}_b^d = U_i^a \hat{g}^{\alpha}_i^j U_d^j , \quad \hat{t}^{\alpha}_b^d = U_i^a \hat{t}^{\alpha}_i^j U_d^j , \quad \hat{v}_{bd}^{\alpha\beta} = U_i^a U_\nu^c \hat{v}_{ijl}^{\alpha\beta} U_\nu^l U_d^j U^c_i .
\]

The evolution equation for \(\hat{g}\) can be obtained directly via the Ehrenfest theorem. One can also apply partial traces to (2.1), which leads to the well-known BBGKY hierarchy [21, 31–34], whose first two equations read explicitly \(^2\) (Einstein summation convention implied):

\[
\begin{align*}
\frac{d \hat{g}^{\alpha}_b^d}{dt} &= \left( \hat{t}^{\alpha}_i^j \hat{g}^{\alpha}_b^d - \hat{g}^{\alpha}_b^d \hat{t}^{\alpha}_i^j \right) + \frac{1}{2} \left( \hat{v}_{ml}^{ik} \hat{g}^{\alpha}_{i}^{ml} - \hat{g}^{\alpha}_{i}^{mk} \hat{v}_{ijl}^{ik} \right) , \\
\frac{d \hat{v}_{ijl}^{ik}}{dt} &= \left( \hat{t}^{\alpha}_i^j \hat{v}_{ijl}^{ik} + \hat{t}^{\alpha}_r \hat{v}_{rlj}^{ikp} + \frac{1}{2} \hat{v}_{rml}^{ik} \hat{v}_{ijl}^{mpr} - \hat{v}_{ml}^{ik} \hat{v}_{ijl}^{rmp} - \frac{1}{2} \hat{v}_{rml}^{ik} \right) \\
&+ \frac{1}{2} \left( \hat{v}_{rml}^{ik} \hat{v}_{ijl}^{rkm} \hat{v}_{ijl}^{rjm} + \hat{v}_{rml}^{ik} \hat{v}_{ijl}^{rkm} \hat{v}_{ijl}^{rjm} - \hat{v}_{rml}^{ik} \hat{v}_{ijl}^{rkm} \hat{v}_{ijl}^{rjm} \right) .
\end{align*}
\]

\(^2\)We made explicit the components of the tensors compared to the expressions found in [21] or [28, 29].
More than simply recasting in a less compact form the very complicated problem (2.1), this hierarchy furnishes a set of evolution equations which depend on higher-order reduced density matrices, and lead to natural truncation schemes for practical applications. The simplest non-trivial closure is the so-called Hartree-Fock or mean-field approximation, which corresponds physically to the propagation of particles in a potential due to the interactions with the particles of the background. Mathematically, it consists in neglecting the correlated parts in the two-body density matrix and the higher order density matrices. Separating the uncorrelated and the correlated contributions, the two-body density matrix reads [28, 30]

\[ \rho^{ik}_{jl} \equiv 2\rho^i_j \rho^k_l + C^{ik}_{jl} \equiv \rho^i_j \rho^k_l - \rho^i_j \rho^k_l + C^{ik}_{jl}. \]  

(2.10)

Inserting this decomposition into (2.9), we get:

\[ i \frac{d\rho^i_j}{dt} = \left( [t^i_j + \Gamma^i_j] \rho^k_l - \rho^k_l \left[ t^i_j + \Gamma^i_j \right] \right) + \frac{1}{2} \left( \bar{\tau}^{ik}_{ml} C_{ml}^{jk} - C_{ml}^{ik} C_{ml}^{jk} \right) = \left[ \hat{t} + \hat{\Gamma}, \rho \right]_j^i + i \hat{C}^i_j, \]

(2.11)

where the mean-field potential \( \hat{\Gamma} \) is defined as (for once, we make explicit the summation)

\[ \Gamma^i_j = \sum_{k,l} \bar{\tau}^{ik}_{jl} \rho^k_l . \]  

(2.12)

The mean-field approximation then consists in neglecting \( C^{ik}_{jl} \simeq 0 \) and keeping only the commutator part in (2.11). However, in the context of neutrino decoupling in the early universe, one seeks a generalization of the Boltzmann equation for neutrino distribution functions [1–5], which describes the evolution of densities under two-body collisions. In other words, we need to truncate the hierarchy (2.9) assuming the molecular chaos ansatz: correlations between the one-body density matrices arise from two-body interactions between uncorrelated matrices. This prescribes the form of \( C^{ik}_{jl}(t) \), leading to the following formal expression for the collision term (see Appendix A for details):

\[ C^{ik}_{jl} = \frac{1}{4} \left( \bar{v}_{i1}^{j1} \bar{v}_{j1}^{i1} \bar{v}_{j2}^{j2} \bar{v}_{i2}^{i2} \left( \hat{1} - \hat{\rho} \right)^{j1} \left( \hat{1} - \hat{\rho} \right)^{i1} - \bar{v}_{i1}^{j1} \bar{v}_{j1}^{i1} \bar{v}_{j2}^{j2} \bar{v}_{i2}^{i2} \left( \hat{1} - \hat{\rho} \right)^{j1} \left( \hat{1} - \hat{\rho} \right)^{i1} \right) \]  

(2.13)

The collision term has the standard structure gain \( \text{loss} + \text{h.c.} \), which will be made more explicit when we give the full expressions for a system of neutrinos and antineutrinos interacting with standard model weak interactions. In (2.13), the indices \( (i_k, j_k) \) will correspond to a definite momentum \( \vec{p}_k \).

We will now focus on the case of the early universe and consider three active species of neutrinos in a background of electrons, positrons (and photons). The influence of baryons can be discarded given their negligible density compared to relativistic species (the baryon-to-photon ratio is \( \eta \equiv n_b/n_\gamma \simeq 6.1 \times 10^{-9} \) from the most recent measurement of the baryon density [15]).
2.2 The case of neutrinos in the early universe

Assuming the universe to be homogeneous and isotropic in the period of interest, the density matrices read,

\[
\langle \hat{a}^\dagger_{\nu_a}(\vec{p}', h') \hat{a}_{\nu_a}(\vec{p}, h) \rangle = (2\pi)^3 2E_p \delta^{(3)}(\vec{p} - \vec{p}') \delta_{hh'} \delta_{\alpha\beta}(p, t) \delta_{h-} ,
\]

\[
\langle \hat{b}^\dagger_{\nu_a}(\vec{p}, h) \hat{b}_{\nu_a}(\vec{p}', h') \rangle = (2\pi)^3 2E_p \delta^{(3)}(\vec{p} - \vec{p}') \delta_{hh'} \delta_{\alpha\beta}^\dagger(p, t) \delta_{h+} .
\]

The Kronecker delta ensures that only left-handed neutrinos and right-handed antineutrinos are included, whereas wrong helicity contributions can be present in anisotropic environments [22]. The energy function is \( E_p = p \) for neutrinos (while it would be \( E_p = \sqrt{\vec{p}^2 + m_e^2} \) for electrons and positrons). Moreover, in the subspace of charged leptons, the density matrices are diagonal and correspond to the distribution functions \( f_e(p, t) \) and \( f_e(p, t) \).

In the following, we will apply the BBGKY formalism to a system of neutrinos, leaving the inclusion of antineutrinos\(^4\) to Appendix E. Note that, for a relativistic system, the hierarchy is given by an infinite set of equations. Therefore, the one-body density matrix will be the neutrino one, with the notation \( \varrho^a_{\beta\gamma} \) (instead of \( \varrho^{\alpha\beta}_{\nu\nu} \)) for clarity. Furthermore, all quantities being diagonal in momentum space, we only deal with the diagonal values \( A(p) \) of operators \( A^{\vec{p}'}_{\vec{p}} = A(p)\delta_{\vec{p}\vec{p}'} \), where the Kronecker symbol” in momentum space is \( \delta_{\vec{p}\vec{p}'} = (2\pi)^3 2E_p \delta^{(3)}(\vec{p} - \vec{p}') \).

We now calculate the relevant expressions of the vacuum, the mean-field (2.12) and collision (2.13) terms for neutrino evolution.

2.2.1 Vacuum term

The neutrino kinetic term is easily calculated in the mass basis, where it is diagonal by definition (the basis elements being the eigenstates of the vacuum Hamiltonian \( \hat{H}_0 \)):

\[
t^i_b(p)\big|_{\text{mass basis}} \simeq p \delta^a_b + \frac{m_e^2}{2p} \delta^a_b \equiv p \delta^a_b + \frac{M^2}{2p} |^a_b .
\]

Since terms proportional to the identity do not contribute to flavour evolution, \( M^2 \) can be taken as the matrix of mass squared differences. In the flavour basis, the vacuum term is obtained following the transformation laws (2.8):

\[
t^i_j = \left( U \frac{M^2}{2p} U^\dagger \right)^i_j ,
\]

with \( U \) the Pontecorvo-Maki-Nakagawa-Sakata (PMNS) mixing matrix [8].

2.2.2 Weak interactions

Neutrinos and antineutrinos in the early universe interact with each others and with the electrons and positrons composing the homogeneous and isotropic plasma. The interaction

\(^3\)The annihilation and creation operators satisfy the non-zero equal time anticommutation rules

\[
\{ \hat{a}_{\nu_a}(\vec{p}, h), \hat{a}^\dagger_{\nu_a}(\vec{p}', h') \} = (2\pi)^3 2E_p \delta^{(3)}(\vec{p} - \vec{p}') \delta_{hh'} \delta_{\alpha\beta} \quad \{ \hat{a}^\dagger_{\nu_a}(\vec{p}, h), \hat{a}^\dagger_{\nu_a}(\vec{p}', h') \} = \{ \hat{a}_{\nu_a}(\vec{p}, h), \hat{a}_{\nu_a}(\vec{p}', h') \} = 0
\]

Similar relations hold for the antiparticle operators.

\(^4\)Note that the antineutrino density matrix \( \varrho^\dagger_{\beta\gamma} \equiv \langle \hat{b}_{\nu_a}^\dagger \hat{b}_{\nu_a} \rangle \) is defined with a transposed convention, compared to the neutrino density matrix, to have similar evolution equations and transformation properties.
Hamiltonian is thus given by the charged- and neutral-current terms from the standard model of weak interactions, expanded at low energies compared to the gauge boson masses. The different expressions and subsequent interaction matrix elements (2.7) are displayed in the appendix B.

Mean-field potential With the set of all relevant $\tilde{v}^{ik}_{jl}$, one can compute the mean-field potential from (2.12). This procedure is outlined in [21], and we just quote here the result:

$$\Gamma^\alpha_\beta = \sqrt{2} G_F (n_e - n_\bar{e}) \delta_e^\alpha \delta_\beta^e + \sqrt{2} G_F (n_\nu - n_\bar{\nu}) \delta_\nu^\alpha \delta_\beta^\nu - \frac{2\sqrt{2} G_F P}{m_W^2} (\rho_e + P_e + \rho_\bar{e} + P_\bar{e}) \delta_e^\alpha \delta_\beta^e - \frac{8\sqrt{2} G_F P}{m_Z^2} (\rho_\nu + P_\nu + \rho_\bar{\nu}) \delta_\nu^\alpha \delta_\beta^\nu.$$  \hspace{1cm} (2.18)

The first two terms are the particle/antiparticle asymmetric mean-field potentials arising from the $V-A$ Hamiltonian. Expanding the gauge boson propagators to next-to-leading order leads to the symmetric terms proportional to the neutrino momentum $p$. This expression is derived in the flavour basis in which $\delta_e^\alpha$ is the Kronecker symbol. However it can be directly read in any basis, through the contravariant (covariant) transformation of upper (lower) indices (2.8).

The various thermodynamic quantities involved are

$$n_e = 2 \int \frac{d^3 p}{(2\pi)^3} f_e(p), \quad \rho_e + P_e = 2 \int \frac{d^3 p}{(2\pi)^3} \left( E_p + \frac{p^2}{3E_p} \right) f_e(p),$$

and the corresponding quantities for antiparticles are obtained by replacing $f_e \to f_\bar{e}$ and $\varrho^\alpha_\beta \to \bar{\varrho}^\alpha_\beta$.

The mean-field potentials up to first order in $1/m_{W,Z}^2$ do not usually take into account the non-relativistic nature of electrons and positrons [9–12, 18]. Instead, our expression involves both the energy density and the pressure of charged leptons, as mentioned for instance in [35]. As expected, we recover the more common expression in the ultra-relativistic limit $\rho_e + P_e \to (4/3)\rho_e$.

Collision integral The collision term is derived by inserting all possible matrix elements in the general expression (2.13). This leads to collision integrals previously derived in [18, 20], and progressively included in numerical computations, except for the self-interactions, whose off-diagonal components were approximated by damping terms or discarded [9–11]. In Appendix C, we illustrate how our formalism applies by carrying out an explicit derivation for neutrino-neutrino scattering, displaying the full matrix structure of the statistical factor.

2.2.3 Quantum Kinetic Equations

We present here the QKE for $\varrho(p,t)$, obtained from (2.11) after dividing each term by the momentum-conserving function $\delta(p')$ from (2.14). Moreover, the time derivative $d/dt$ becomes $\partial/\partial t - H p \partial/\partial p$ to account for the expansion of the universe, $H \equiv \dot{a}/a$ being the Hubble

\footnotetext[5]{The absence of extra complex conjugation on $\bar{\varrho}$ compared to [21] is due to the transposed definition of the antineutrino density matrix.}
rate, given by Friedmann’s equation \( H^2 = (8\pi G/3)\rho \). The QKEs read:

\[
 i \left[ \frac{\partial}{\partial t} - H_p \frac{\partial}{\partial p} \right] \varrho = \left[ U \frac{M^2}{2p} U^\dagger, \varrho \right] + \sqrt{2} G_F \left[ N_e + N_\nu, \varrho \right] - 2\sqrt{2} G_F p \left[ \frac{E_e + P_e}{m_W^2} + \frac{4}{3} \frac{E_\nu}{m_Z^2}, \varrho \right] + i \mathcal{I} \tag{2.20}
\]

with the matrices defined in flavour space \( N_e \equiv \text{diag}(n_e - n_\bar{e}, 0, 0) \), \( N_\nu \equiv n_\nu - n_\bar{\nu} \), \( E_e \equiv \text{diag}(\rho_e + \bar{\rho}_e, 0, 0) \) (likewise for \( P_e \)), and \( E_\nu \equiv \rho_\nu + \bar{\rho}_\nu \). Similarly, the QKEs for the antineutrino density matrix read (cf. Appendix E):

\[
 i \left[ \frac{\partial}{\partial t} - H_p \frac{\partial}{\partial p} \right] \varrho^- = - \left[ U \frac{M^2}{2p} U^\dagger, \varrho^- \right] + \sqrt{2} G_F \left[ N_e + N_\nu, \varrho^- \right] + 2\sqrt{2} G_F p \left[ \frac{E_e + P_e}{m_W^2} + \frac{4}{3} \frac{E_\nu}{m_Z^2}, \varrho^- \right] + i \mathcal{I} \tag{2.21}
\]

Note that only eq. (2.20) will be solved numerically, since we will be focussing on the case of zero chemical potential for which antineutrinos evolve as neutrinos.

The collision term is the sum of the contributions from different physical processes: scattering with charged leptons \((\nu e^\pm \leftrightarrow \nu e^\pm)\), annihilation \((\nu \bar{\nu} \leftrightarrow e^+ e^-)\) and self-interactions (involving only \( \nu \) and \( \bar{\nu} \)). The expressions for the processes involving charged leptons are exactly the same as the ones quoted in [10] [eqs. (2.19)–(2.21)], and we do not report them here for brevity. This reference, however, does not contain the full expressions for neutrino self-interactions, derived for instance in [20]. Our expression for the self-interactions contribution to the collision integral reads:

\[
 i \mathcal{I}^{[\nu]} = \frac{1}{2} \frac{25 G_F^2}{2 \pi} \int \frac{d^3 \vec{p}_1}{(2\pi)^3} \left[ (1 - \varrho_1) \varrho_2 + \text{Tr}(\cdots) \right] \varrho_3 (1 - \varrho_4) \varrho_1 (1 - \varrho_2) \varrho_4 + \text{Tr}(\cdots) \right] \tag{2.22}
\]

with the statistical factors for scattering and annihilation processes:

\[
 F_{sc}(\nu^{(1)}, \nu^{(2)}, \nu^{(3)}, \nu^{(4)}) = [\varrho_4 (1 - \varrho_2) + \text{Tr}(\cdots)] \varrho_3 (1 - \varrho_1) + (1 - \varrho_1) \varrho_3 (1 - \varrho_2) \varrho_4 + \text{Tr}(\cdots)]
\]

\[
 - (1 - \varrho_1) \varrho_2 + \text{Tr}(\cdots) \right] (1 - \varrho_3) \varrho_1 (1 - \varrho_4) \varrho_3 (1 - \varrho_2) + \text{Tr}(\cdots) \right] , \tag{2.23}
\]

\[
 F_{sc}(\bar{\nu}^{(1)}, \bar{\nu}^{(2)}, \bar{\nu}^{(3)}, \bar{\nu}^{(4)}) = [(1 - \bar{\varrho}_2) \bar{\varrho}_4 + \text{Tr}(\cdots)] \varrho_3 (1 - \varrho_1) + (1 - \varrho_1) \varrho_3 (1 - \bar{\varrho}_2) + \text{Tr}(\cdots)]
\]

\[
 - (1 - \bar{\varrho}_2) \bar{\varrho}_4 + \text{Tr}(\cdots) \right] (1 - \varrho_3) \varrho_1 (1 - \varrho_4) (1 - \bar{\varrho}_2) + \text{Tr}(\cdots) \right] , \tag{2.24}
\]

\[
 F_{\text{ann}}(\nu^{(1)}, \bar{\nu}^{(2)}, \nu^{(3)}, \bar{\nu}^{(4)}) = [(1 - \varrho_3) (1 - \varrho_4) + \text{Tr}(\cdots)] (1 - \bar{\varrho}_2) \bar{\varrho}_4 + (1 - \varrho_1) (1 - \bar{\varrho}_2) [\bar{\varrho}_4 \varrho_3 + \text{Tr}(\cdots)]
\]

\[
 - [(1 - \varrho_3) (1 - \varrho_4) + \text{Tr}(\cdots)] \varrho_2 \varrho_1 (1 - \varrho_4) (1 - \bar{\varrho}_2) + \text{Tr}(\cdots) \right] , \tag{2.25}
\]

where we chose the more compact notation \( \varrho_k = \varrho(p_k) \), and \( \text{Tr}(\cdots) \) means the trace of the term in front of it.

\footnote{It is equivalent with eq. (96) of ref. [20] (one only needs to swap the variables \( \vec{p}_3 \leftrightarrow \vec{p}_4 \) in the second and fourth terms of (2.23)). Our expression makes more explicit the gain – loss + h.c.” structure of this collision term.}
2.3 Reduced set of equations

The full QKE (2.20) can be recast in a form more suitable for a numerical resolution. Though neutrino density matrices will deviate from kinetic and chemical equilibrium, electrons and positrons undergo very efficient electromagnetic interactions with the photon background, ensuring that their distribution function remains a Fermi-Dirac one at the photon temperature $T_\gamma$ [36]. Due to a very low baryon-to-photon ratio $\eta$, the difference between the electron and positron number densities is very small compared to the number density of relativistic species (e.g. photons or neutrinos). When electrons and positrons are still relativistic, this implies that their chemical potentials can be safely ignored as they are of the same order as $\eta$. When they annihilate at temperatures lower than the electron mass, the number density difference remains constant leading to a complete asymmetry when positrons have disappeared, and thus to a sizeable chemical potential for electrons, see e.g. figure 30 of ref. [13]. However the relic number density of electrons is of the order of $\eta$ and their effect on neutrino decoupling can be completely ignored. We will thus neglect the chemical potential of $e^\pm$ since its effect on neutrino decoupling would be of the order of $\eta$. In addition, we also neglect any CP violating phase in the PMNS matrix (4.6) or CP breaking reactions, implying that the equality $\bar{\varrho} = \varrho$ will be ensured at all times [24, 25]. Therefore, we will only solve the equation (2.20), in which the antisymmetric mean-field

\[ N_e + N_\nu \] vanishes. Moreover, since deviations from the equilibrium distribution $\varrho \propto I$ are small (cf. numerical results below), the mean-field term proportional to $E_\nu$ will be very close to the identity (because we are at zero chemical potential), so it will give a negligible contribution within the commutator. We thus discard this term in the numerical resolution.

The most time consuming part of the QKE is the computation of the collision term. Following [1], the nine-dimensional collision integrals are reduced to two-dimensional ones. Finally, we define the comoving temperature $T_{cm} \propto a^{-1}$ [4], which corresponds to the physical temperature of all species when they are strongly coupled, i.e. $T_\nu = T_\gamma = T_{cm}$ when $T_{cm} \gg 1$ MeV. From this proxy for the scale factor, we define the comoving variables [2, 9]

\[ x = m_e/T_{cm}, \quad y = p/T_{cm}, \quad z = \frac{T_\gamma}{T_{cm}}, \] (2.26)

which are respectively the reduced scale factor, the comoving momentum, and the dimensionless photon temperature, such that $\varrho(p,t)$ is now expressed $\varrho(x,y)$. We also introduce the dimensionless thermodynamic quantities $\tilde{\varrho} = \varrho(x/m_e)^4$ and $\tilde{P} = P(x/m_e)^4$.

Therefore, the QKE is rewritten:

\[
\frac{\partial \varrho(x,y_1)}{\partial x} = \frac{i}{xH} \frac{m_e}{x} \left[ U \frac{M^2}{2y_1} U^\dagger, \varrho \right] + i \frac{2 \sqrt{2} G_F}{xH} y_1 \left( \frac{m_e}{x} \right)^{\frac{5}{2}} \left[ \frac{\tilde{\varrho}_e + \tilde{\varrho}_\nu}{m^2_{\nu}}, \varrho \right] + \frac{1}{xH} \mathcal{I},
\] (2.27)

with the two-dimensional collision integral\(^7\) (recall that we assume $f_e = f_{\bar{e}}$, which regroups some terms):

\(^7\)We integrated out the energy delta-function via $\int p_4 dp_4 \delta(E_1 + E_2 - E_3 - E_4) = E_1 + E_2 - E_3$, since $p_4 dp_4 = E_4 dE_4$. In (2.28) $E_4$ stands for $E_1 + E_2 - E_3$. 

- 9 -
\[
I = \frac{G_F^2}{2\pi^3 y_1} \left( \frac{m_e}{x} \right)^5 \int y_2 dy_2 y_3 dy_3 \bar{E}_4 \times \frac{1}{2} \times 
\left[ 4 \left[ 2d_1 + 2d_3 + d_2(1, 2) + d_2(3, 4) - d_2(1, 4) - d_2(2, 3) \right] \times \left( F_{sc}^{LL}(\nu^{(1)}, e^{(2)}, \nu^{(3)}, e^{(4)}) + F_{sc}^{RR}(\nu^{(1)}, e^{(2)}, \nu^{(3)}, e^{(4)}) \right) - 4x^2 \left[ d_1 - d_2(1, 3) \right] / \bar{E}_3 \bar{E}_4 \times \left( F_{sc}^{LR}(\nu^{(1)}, e^{(2)}, \nu^{(3)}, e^{(4)}) + F_{sc}^{RL}(\nu^{(1)}, e^{(2)}, \nu^{(3)}, e^{(4)}) \right) + 4 \left[ d_1 + d_3 - d_2(1, 4) - d_2(2, 3) \right] \times \left( F_{sc}^{LL}(\nu^{(1)}, \bar{\nu}^{(2)}, e^{(3)}, e^{(4)}) + F_{sc}^{RR}(\nu^{(1)}, \bar{\nu}^{(2)}, e^{(3)}, e^{(4)}) \right) + 2x^2 \left[ d_1 + d_2(1, 2) \right] / \bar{E}_3 \bar{E}_4 \times \left( F_{sc}^{LR}(\nu^{(1)}, \bar{\nu}^{(2)}, e^{(3)}, e^{(4)}) + F_{sc}^{RL}(\nu^{(1)}, \bar{\nu}^{(2)}, e^{(3)}, e^{(4)}) \right) + \left[ d_1 + d_3 + d_2(1, 2) + d_2(3, 4) \right] \times F_{sc}(\nu^{(1)}, \nu^{(2)}, \nu^{(3)}, \nu^{(4)}) + \left[ d_1 + d_3 - d_2(1, 4) - d_2(2, 3) \right] \times \left( F_{sc}(\nu^{(1)}, \bar{\nu}^{(2)}, \nu^{(3)}, \bar{\nu}^{(4)}) + F_{sc}(\nu^{(1)}, \bar{\nu}^{(2)}, \nu^{(3)}, \bar{\nu}^{(4)}) \right) \right]
\]

The \(d\)-functions are \(d_i = (x/m_e)d_i^{DHS}\), with \(d_i^{DHS}\) defined in [1] as functions of the momenta \(p\), hence the prefactor \(x/m_e\). Note that [10] use a different convention (4 times greater \(D\)-functions and opposite sign for \(d_2\)).

In addition to the QKEs, the remaining dynamical equation is the energy conservation equation \(\dot{\rho} = -3\H (\rho + P)\), rewritten as an equation on \(\z(x)\) [3, 7]. See Appendix D for the complete expression including QED corrections to the plasma equation of state.

### 3 Adiabatic transfer of averaged oscillations

Solving the full QKE (2.27) is a priori a considerable numerical challenge because of the need to resolve numerically both the effect of the mean-field terms and of computationally expensive collision integrals. However, previous numerical results [9, 10] seem to indicate that the expected oscillations are somehow “averaged” while there is a comparatively slow evolution due to collisions.

We thus expect a clear separation of time-scales to hold, allowing for an effective description which correctly captures the salient features of the dynamical evolution. For convenience, let us rewrite the QKE (2.27) in the compact form:

\[
\frac{\partial \rho}{\partial x} = -i[H, \rho] + \mathcal{K},
\]

with

\[
H \equiv \frac{1}{x\mathcal{H}} \left[ \frac{x}{m_e} U \bar{M}_e^2 U^\dagger - 2\sqrt{2} G_F y \left( \frac{m_e}{x} \right)^5 \bar{E}_e + \bar{P}_e \right]
\]

and \(\mathcal{K} \equiv \frac{1}{x\mathcal{H}} I\). We treat the \(y\) dependence of \(H\) implicitly, as the following procedure must be applied for each \(y\). Since the mean-field Hamiltonian \(H\) is Hermitian, it can be diagonalized by the unitary transformation

\[
\mathcal{H} = U_m H U_m^\dagger \quad \text{with} \quad (H_m)^j_k = (H_m)^j_k \delta^j_k.
\]

The density matrix in the matter basis reads \(\rho_m = U_m^\dagger \rho U_m\), and evolves according to

\[
\frac{\partial \rho_m}{\partial x} = -i[H_m, \rho_m] - \left[ U_m^\dagger \frac{\partial U_m}{\partial x}, \rho_m \right] + U_m^\dagger \mathcal{K} U_m.
\]
The first approximation that we consider is the adiabatic approximation \[8, 37\] which consists in neglecting the time evolution of the matter PMNS matrix compared to the inverse effective oscillation frequency:

\[
\text{Adiabatic approximation} \quad \left| U_m^\dagger \frac{\partial U_m}{\partial x} \right| \ll |H_m|. \tag{3.5}
\]

This condition means that the effective mixing matrix varies very slowly compared to the effective oscillation frequency, so that the matter basis evolves adiabatically. Such adiabaticity condition is particularly important in presence of Mikheev-Smirnov-Wolfenstein (MSW) resonances [38, 39]. Note that the sign of the mean-field contribution to \(H\) (3.2) is opposite to the one encountered due to charged-current neutrino-electron scattering at lowest order, important for astrophysical environments (Sun, supernovae, binary neutron star mergers).

We numerically checked (figure 1) that the condition (3.5) is indeed satisfied throughout the range of temperatures of interest.

If we now assume that many oscillations take place before the collision term varies substantially and write the collision term in matter basis

\[
K_m \equiv U_m^\dagger K U_m,
\]

its variation frequency \(\sim K_m^{-1}(\partial K_m/\partial x)\) must be small compared to the effective oscillation frequency \(H_m\), namely

\[
\text{Averaged oscillations} \quad \left| K_m^{-1} \frac{\partial K_m}{\partial x} \right| \ll |H_m|. \tag{3.6}
\]

If this new separation of time-scales holds, and the variation of the collision term is negligible during an oscillation period (see figure 2), we can average the evolution over many oscillations (the collision term produces at constant rate neutrinos with random initial phases, which average out). More precisely, we can write

\[
(\bar{\rho}_m)^j_k (x, y) = e^{-i(H_m)^j_k x} R_j^k (x, y) e^{i(H_m)^k_j x} \quad \Rightarrow \quad \frac{\partial R_j^k}{\partial x} = e^{i(H_m)^j_k x} (K_m)^{-1} e^{-i(H_m)^k_j x}, \tag{3.7}
\]

where we also assumed a slow variation of \(H_m\), as a consequence of the adiabatic approximation. If (3.6) holds, \(\partial R_j^k/\partial x\) is integrated over many oscillations and the non-diagonal parts vanish.\(^8\) This leaves us with the effective equation in matter basis:

\[
\text{Adiabatic Transfer of Averaged Oscillations} \quad \begin{cases} \\
\frac{\partial \tilde{\rho}_m}{\partial x} = U_m^\dagger K U_m, \\
\tilde{\rho}_m = \tilde{\rho}_m 
\end{cases}, \tag{3.8}
\]

where the tilde means that we only keep the diagonal terms of \(\tilde{\rho}_m\), then convert it to the flavour basis to compute the collision term \(K\) and only keep the diagonal part of the collision term \(U_m^\dagger K U_m\) when transforming back to the matter basis.

In the flavour basis, the density matrix \(\varrho = U_m \tilde{\varrho}_m U_m^\dagger\) has non-diagonal components, while \(\tilde{\varrho}_m\) is diagonal. Therefore the collision term destroys the coherence between these components (since it aims at a diagonal \(\varrho\) in flavour space, with equilibrium distributions), which modifies the diagonal values of \(\varrho_m\) (whose non-diagonal terms average out). For clarity, we refer to this approximate numerical scheme to determine the neutrino evolution “Adiabatic Transfer of Averaged Oscillations” (ATAO) and we solve (3.8) instead of (3.1).

In the following section, we will numerically solve the QKEs in both the full case and the ATAO approximation and discuss the validity of the approximate numerical solution.

\(^8\)As it is sometimes stated, one could phrase it by saying that the off-diagonal terms are washed out by going to a comoving frame.
Figure 1. Evolution of the different quantities appearing in (3.4) in the normal hierarchy of masses. The condition (3.5) is satisfied throughout the evolution.

Figure 2. Comparison of the evolution of the collision term and the effective oscillation frequencies in the normal hierarchy of masses. The condition (3.6) is satisfied throughout the evolution.

4 Numerical results

4.1 Numerical implementation

We integrate numerically the QKE for neutrinos (2.27), or (3.8) in the ATAO approximation, along with the energy conservation equation (D.1). We use our own code NEVO (Neutrino EVOlver) written in Python with the scipy and numpy libraries.\textsuperscript{9}

\textsuperscript{9}Time consuming functions are compiled with the just-in-time compiler numba.
Solver and initial conditions  The collision term consists most of the time in nearly compensating gain and loss terms, and for energies larger than 0.1 MeV, the system is very stiff. Hence, one must rely on an implicit method. We chose the LSODA method which consists in a BDF method (with adaptative order and adaptative step) when the system is stiff, which switches to an explicit method when not stiff (the Adams method). It was first distributed within the ODEPACK [40] Fortran library, but we used the Python wrapper solve_ivp distributed with the Python scipy module. We remarked that when setting the absolute and relative error tolerances to $10^{-n}$, the spectra are typically obtained with precision better than $10^{-n+2}$, in agreement with section B.5 of [11]. Hence we fixed these error tolerances to $10^{-7}$ so as to obtain results with numerical errors below $10^{-5}$.

The initial common temperature of all species, that is all types of neutrinos and the electromagnetic plasma, is inferred from the conservation of total entropy. Choosing the initial comoving temperature $T_{\text{cm}, \text{in}} = 20$ MeV, the initial common temperature of all species is slightly larger because of early $e^\pm$ annihilations, and given by

$$ T_{\text{in}} = z_{\text{in}} T_{\text{cm}, \text{in}} , $$

with $z_{\text{in}}^{-1} = 7.42 \times 10^{-6}$. Had we chosen to start at $T_{\text{cm}, \text{in}} = 10$ MeV, the initial comoving temperature would be $z_{\text{in}}^{-1} = 2.98 \times 10^{-5}$, in agreement with Refs. [9, 41]. As initial condition for the density matrix we take

$$ \rho(x_{\text{in}}, y) = \begin{pmatrix} f^{(\text{in})}_{\nu} (y) & 0 & 0 \\ 0 & f^{(\text{in})}_{\nu} (y) & 0 \\ 0 & 0 & f^{(\text{in})}_{\nu} (y) \end{pmatrix} , \quad \text{with} \quad f^{(\text{in})}_{\nu} (y) \equiv \frac{1}{e^{y/z_{\text{in}}} + 1} . (4.1) $$

Momentum grid  The neutrino spectra are sampled with $N$ points on a grid in the reduced momentum $y$. When choosing a linear grid, we use the range $0.01 \leq y \leq 16 + [N/20]$, and integrals are evaluated with the Simpson method. However, for functions which decay exponentially for large $y$, it is motivated to use the Gauss-Laguerre quadrature which was already proposed in [11], and we confirm that this method typically requires half of the grid points to reach the same precision as the one obtained with a linear spacing. In practice, when choosing the nodes and weights of the quadrature, we restrict to $y \leq 20 + [N/5]$. When using $N = 80$, we have thus restricted nodes to $y \leq 36$, and we used Laguerre polynomials of order 439 to compute the weights with eq. (B.14) of [11]. Since the tools provided in numpy are restricted to much lower polynomial orders, we used Mathematica to precompute once for all in a few hours the nodes and weights. The results reported in this paper were performed with $N = 80$ and the Gauss-Laguerre quadrature, checking that with $N = 100$ the differences are smaller than the desired precision.

For each momentum $y_i$ of the grid, and with $N_{\nu}$ flavours, each density matrix has $N_{\nu}^2$ independent degrees of freedom ($N_{\nu}(N_{\nu} + 1)$ real parts and $N_{\nu}(N_{\nu} - 1)$ imaginary parts). In practice we reorganize these independent matrix entries into a vector $A^j(y_i)$ with $j = 1, \ldots, N_{\nu}^2$ and we concatenate them with the $y_i$ spanning the momentum grid. We thus solve for serialized variables, that is a giant vector of length $N N_{\nu}^2$. When using the ATAO approximation, one needs only to keep the diagonal part in the matter basis, and the giant vector is of size $N N_{\nu}$.\footnote{Results are then only converted at the very end in the flavour basis if desired.} Note that we do not store the binned density matrix components $\rho_{\alpha\beta}(y_i)$, which would be sub-optimal. Indeed, if neutrinos decoupled instantaneously, their distribution function would then be

$$ f^{(\text{eq})}_{\nu} (x, y) \equiv \frac{1}{e^y + 1} . (4.2) $$
Therefore, we can parametrize the density matrix $\rho^3_{\beta}(x, y) = \left[ \delta^\alpha_{\beta} + a^3_{\beta}(x, y) \right] \times f^{(eq)}_\nu(x, y)$, and we store the values of $a^3_{\beta}$, which encapsulate the deviation from instantaneous decoupling.

**Numerical optimization via Jacobian computation** The implicit method requires to solve algebraic equations and thus to obtain the Jacobian of the differential system. For the sake of this discussion, and to alleviate the notation, we ignore the different flavours and consider that we have only one neutrino flavour with spectrum $f(y)$. Noting the grid points $y_i$ and the values of the spectra $f_i = f(y_i)$ on the grid, the differential system is of the type $\partial_x f_i = C_i(x, f_j)$. The implicit method requires the Jacobian $J_{ij} = \partial C_i/\partial f_j$. If no expression is provided, it is evaluated by finite differences in the $\{f_i\}$ at a given $x$. Since the collision term involves a two-dimensional integral for each point of the grid, its computation on the whole grid is of order $O(N^3)$. Hence the computation of the Jacobian with finite differences is of order $O(N^4)$. It is however possible to reduce this time by providing a numerical method to compute the Jacobian with an $O(N^3)$ complexity. To use a simple example, let us only consider the contribution from the loss part of the neutrino self-interactions, without including Pauli-blocking factors. This component of the collision term, once computed numerically with a quadrature, is of the form

$$C_i(x, f_j) = - \sum_{j,k} w_j w_k g(y_i, y_j, y_k) f_i f_j.$$  \hfill (4.3)

In this expression $\sum_j w_j$ (resp. $\sum_k w_k$) accounts for the integration on $y_2$ (resp. $y_3$) in (2.28) using a quadrature, and the function $g$ takes into account the specific form of the factor multiplying the statistical function (which is for the contribution considered $f_i f_j$). Noting then that

$$\partial f_i/\partial f_j = \delta_{ij},$$ \hfill (4.4)

the Jacobian associated with the contribution (4.3) is

$$J_{im} = \partial C_i/\partial f_m = - \delta_{im} \sum_{j,k} w_j w_k g(y_i, y_j, y_k) f_i - \sum_k w_k g(y_i, y_m, y_k) f_i.$$ \hfill (4.5)

The complexity of the second sum is of order $O(N)$, and since the Jacobian has $N^2$ entries, it leads to a complexity of order $O(N^3)$. The first term is not worse even though the double sum is of order $O(N^2)$, because it concerns only the diagonal entries of the Jacobian due to the prefactor $\delta_{im}$. More generally for all contributions to the collision term, the complexity when computing the associated Jacobian is always of order $O(N^3)$. When restoring the fact that we do not have a single flavour but density matrices, the discussion is similar when using the serialized variables described above, and again the complexity is of order $O(N^3)$. When considering the various contributions, we need more integrals to compute the Jacobian than for the collision term, and mostly due to the fact that with Pauli-blocking factors the collision term for self-interactions is cubic or quartic in the density matrices. In practice, we found that it takes roughly five times more time to compute a Jacobian than a collision term. Hence, when compared with the finite difference method, providing a numerical method for the Jacobian leads to a factor $N/5$ speed-up. Note that we must also integrate $z$ with eq. (D.1) jointly with the density matrices, so that we must pad the Jacobian obtained with the previous description with one extra line and one extra column. Again, the corresponding entries can be deduced using (4.4) and their computation is also of order $O(N^3)$. It is worth mentioning that providing a method for the Jacobian is not specific to the ATAO
approximation. Indeed, when solving the full QKE one can also compute the Jacobian of the collision term, and one only needs to add the contribution from the vacuum and mean field commutators whose complexity is simply of order $O(N^2)$.

When compared with the full QKE method, the ATAO numerical resolution allows to gain at least a factor 5 in time. Hence when using both a method for the Jacobian and the ATAO approximation, we gain typically a factor $N$ and computations that would otherwise last days on CPU clusters, are reduced to just few hours on a single CPU. Moreover, nothing prevents the computation of collision terms and Jacobians to be parallelized on the momentum grid, as we checked on the 4 or 8 CPUs of desktop machines, reducing even further the computation time.

4.2 Oscillation parameters

For the numerical calculations, we employ the standard parametrization of the PMNS matrix which reads

$$U = \begin{pmatrix} c_{12}c_{13} & s_{12}c_{13} & s_{13} \\ -s_{12}c_{23} - c_{12}s_{23}s_{13} & c_{12}c_{23} - s_{12}s_{23}s_{13} & s_{23}s_{13} \\ s_{12}s_{23} - c_{12}c_{23}s_{13} & -c_{12}s_{23} - s_{12}c_{23}s_{13} & c_{23}s_{13} \end{pmatrix},$$

(4.6)

with $c_{ij} = \cos \theta_{ij}$, $s_{ij} = \sin \theta_{ij}$ and $\theta_{ij}$ the mixing angles. We use the most recent values from the Particle Data Group:

$$\begin{pmatrix} \Delta m_{21}^2/10^{-5}\text{eV}^2, \Delta m_{31}^2/10^{-3}\text{eV}^2, s_{12}^2, s_{23}^2, s_{13}^2 \end{pmatrix}_{\text{NH}} = (7.53, 2.53, 0.307, 0.545, 0.0218).$$

(4.7)

For completeness, we also give the most recent values of the physical constants used: the Fermi constant $G_F = 1.1663787 \times 10^{-5}\text{GeV}^{-2}$ and the gravitational constant $G = 6.70883 \times 10^{-39}\text{GeV}^{-2}$.

4.3 Neutrino temperature and spectra

A convenient parametrization of neutrino spectral distortions consists in separating effective temperatures and residual distortions [5], namely,

$$\varrho_\alpha(x, y) \equiv \frac{1}{e^{y/z_{\nu_\alpha}} + 1} \left[ 1 + \delta g_{\nu_\alpha}(x, y) \right],$$

(4.8)

where the reduced effective temperature $z_{\nu_\alpha} \equiv T_{\nu_\alpha}/T_{\text{cm}}$ is the reduced temperature of the Fermi-Dirac spectrum with zero chemical potential which has the same energy density as the real distribution:

$$\bar{\rho}_{\nu_\alpha} \equiv \frac{7 \pi^2}{90} \nu_\alpha^4.$$

(4.9)

We plot in figure 3 the evolution of the neutrino effective temperatures, with and without flavour oscillations. The higher values for the electronic flavour are due to the charged-current processes (that do not exist for muon and tau neutrinos), which increase the transfer of entropy from electrons and positrons. Likewise, the non-thermal residual distortions are more important for $\varrho_e$ (see figure 4). This increased energy density of neutrino species has historically been parametrized through the effective number of neutrino species $N_{\text{eff}}$, i.e.,

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11. [https://pdg.lbl.gov/2020/tables/rpp2020-sum-leptons.pdf](https://pdg.lbl.gov/2020/tables/rpp2020-sum-leptons.pdf)
12. [https://pdg.lbl.gov/2020/reviews/rpp2020-rev-phys-constants.pdf](https://pdg.lbl.gov/2020/reviews/rpp2020-rev-phys-constants.pdf)
the number of instantaneously decoupled neutrino species that would give the same energy density. Long after decoupling, this reads:

$$\rho_\nu = \left[ 1 + \frac{7}{8} \left( \frac{4}{11} \right)^{4/3} N_{\text{eff}} \right] \rho_\gamma \iff N_{\text{eff}} \equiv \left[ \frac{(11/4)^{1/3}}{z} \right]^4 \times \left( z_{\nu_e}^4 + z_{\nu_\mu}^4 + z_{\nu_\tau}^4 \right).$$

\[(4.10)\]

---

**Figure 3.** Evolution of the effective neutrino temperatures, with and without oscillations. Long before decoupling, they remain equal to the photon temperature \(z\), before freezing-out at different values depending on the interaction with the electromagnetic plasma. Without mixing, the distribution function (and thus, the effective temperatures) are identical for \(\nu_\mu\) and \(\nu_\tau\).

The final values of the comoving temperatures and \(N_{\text{eff}}\) are given in Table 1. The inclusion of QED corrections up to \(O(e^3)\) order reduces \(N_{\text{eff}}\) by \(\sim 0.001\), as predicted in ref. [7], and already observed in [12]. Indeed, without these corrections, but keeping the ones at order \(O(e^2)\), we get \(N_{\text{eff}} \simeq 3.0444\) (no oscillations), compared to \(N_{\text{eff}} \simeq 3.0434\) with the corrections up to \(O(e^3)\).

Flavour oscillations reduce the discrepancy between the different flavours, thus \(z_{\nu_e}\) is reduced while \(z_{\nu_\mu}\) and \(z_{\nu_\tau}\) are increased, with a very slightly higher value for \(z_{\nu_\mu}\). This enhanced entropy transfer towards \(\nu_\mu\) compared to \(\nu_\tau\) is due to the more important \(\nu_e - \nu_\mu\) mixing (cf. figure 5 and the corresponding discussion).

The deviation of the dimensionless temperatures with respect to 1 can be expressed as a relative change in the energy density, \(\delta \bar{\rho}_\nu = 4(z_\nu - 1)\). Our values for the increase in the neutrino energy density are \(\delta \bar{\rho}_{\nu_e} \simeq 0.70\%\), \(\delta \bar{\rho}_{\nu_\mu} \simeq 0.53\%\) and \(\delta \bar{\rho}_{\nu_\tau} \simeq 0.52\%\). This is in agreement with the results of ref. [10] (Table 1) or ref. [12] (Table 2), except for the relative variation of muon and tau flavours: these works obtain a higher reheating of \(\nu_\tau\) compared to \(\nu_\mu\), while we find the opposite. This is due to a difference in the values of the mixing angles.\(^\text{13}\) Nevertheless, if we use the mixing angles from [10], we obtain \(\delta \bar{\rho}_{\nu_e} \simeq 0.694\%\).

\(^{13}\)For instance, the older values used in [9] lead to higher distortions for \(\nu_\mu\) than for \(\nu_\tau\).
Figure 4. Frozen-out effective spectral distortions, with and without oscillations, for \( x_f \simeq 51 \) (corresponding to \( T_{cm,f} = 0.01 \text{MeV} \)). The full QKE results are indistinguishable from the ATAO approximate ones.

Table 1. Frozen-out values of the dimensionless photon and neutrino temperatures, and the effective number of neutrino species. The values without oscillations differ from [5] because of the inclusion of QED corrections at order \( \mathcal{O}(e^3) \) in this work (cf. Appendix D). \( N_{\text{eff}} \) is different between the ATAO approximation and full QKE calculations at order \( 10^{-6} \), which we attribute mainly to numerical errors. The implementations in the third and fourth lines are discussed in section 4.4. The post-averaging result corresponds to eq. (4.13).

\[
\delta \bar{\rho}_{\nu_e} \simeq 0.525 \% \quad \text{and} \quad \delta \bar{\rho}_{\nu_\mu} \simeq 0.530 \%.
\]

Furthermore, if \( \mathcal{O}(e^3) \) QED corrections are not included and only the diagonal components of the self-interaction collision term are kept, the spectra reach less flavour equilibration and the results of [10] are recovered (at the level of a few \( 10^{-5} \)): \( \delta \bar{\rho}_{\nu_e} \simeq 0.706 \% \), \( \delta \bar{\rho}_{\nu_\mu} \simeq 0.515 \% \) and \( \delta \bar{\rho}_{\nu_\tau} \simeq 0.522 \% \).

\[
\text{Finally, the results in Table 1 show the striking accuracy of the ATAO approximation, as expected since the conditions (3.5) and (3.6) are satisfied by several orders of magnitude (Figs. 1 and 2). The frozen-out values of the comoving temperatures and of } N_{\text{eff}} \text{ differ by } 10^{-6}, \text{ which is beyond our desired accuracy, and beyond the expected effect of neglected QED.}
\]
contributions.\textsuperscript{14}

The numerical solution of the QKE shows a larger $N_{\text{eff}}$ value (Table 1) compared to the no-oscillation case. To understand this slight increase of the total energy density of neutrinos, one should keep in mind that electron-positron annihilations, which is the dominant process during decoupling, are more efficient in producing electronic type neutrinos (because of the existence of charged-current processes). Now the mixing and mean-field terms tend to depopulate $\nu_e$ and populate the other flavors, which frees some phase space for the reactions which create $\nu_e$, while augmenting the effect of Pauli-blocking factors for reactions creating $\nu_{\mu,\tau}$. Since the former are the dominant reactions, the net effect is a larger entropy transfer from $e^\pm$, hence the larger value of $N_{\text{eff}}$. In the next section, we further clarify the effect of mixing and mean-field terms in the light of the ATAO approximation.

To conclude, we find that the value of $N_{\text{eff}}$ predicted by the Standard model of cosmology, including flavour oscillations and QED radiative corrections, is $N_{\text{eff}} = 3.0440$ with at least $10^{-4}$ precision.

4.4 ATAO transfer functions

The ATAO approximation allows to get some insight on the impact of the mixings and mean-field terms on the spectral modifications and on $N_{\text{eff}}$. To this purpose, let us define the ATAO transfer function

$$T(\alpha \rightarrow \beta, x \rightarrow x', y) = \left[ U_m(x', y) \left( U_m^\dagger(x, y) D(\alpha) U_m(x, y) \right) U_m^\dagger(x', y) \right]_{\beta}^\beta,$$  \hspace{1cm} (4.11)

where $D(\alpha)$ is a diagonal matrix with a non-vanishing (unit) component, that is $[D(\alpha)]_{\beta}^\gamma = \delta_{\beta}^{\gamma} \delta_{\alpha}^{\gamma}$ (no summation). Equation (4.11) corresponds to the probability for a state of flavour $\alpha$ and momentum $y$ generated at a scale factor $x$, “averaged” according to the ATAO approximation, to re-emerge as a flavour $\beta$ at later $x'$, if it is not affected by collisions in the meantime. When evaluated at $x' \rightarrow \infty$, the asymptotic $T(\alpha \rightarrow \beta, x, y) \equiv T(\alpha \rightarrow \beta, x \rightarrow \infty, y)$ provide information on neutrino flavour conversion from their last scattering with other species, until all neutrino spectra are frozen since mean-field and collisions are then negligible (figure 5).

If mean-field effects can be ignored, the asymptotic ATAO transfer function converges to the following expression

$$T^{\text{vac}}(\alpha \rightarrow \beta) \equiv \left[ U \left( U^\dagger D(\alpha) U \right) U^\dagger \right]_{\beta}^\beta,$$  \hspace{1cm} (4.12)

which is independent of $y$ and where the PMNS matter matrix is replaced by the vacuum one.

To gather further insight on the impact of the mixing and mean-field terms, we have performed two schematic calculations, including either the neutrino probabilities at the end of the evolution, i.e. $T_{\text{cm,f}} = 0.01$ MeV (“NO, post-aver.”), or keeping only the mixing and collision terms during the evolution (“without mean-field”). The corresponding results are shown in Table 1.

\textsuperscript{14}Higher order QED corrections or subdominant log-dependent contributions are not expected to modify $N_{\text{eff}}$ above order $10^{-5}$ [7].
Figure 5. Asymptotic ATAO transfer function $T(\alpha \to \beta, x, y)$ for $y = 5$. Left: Normal hierarchy. Right: Inverted hierarchy. The asymptotic values for large $x$ correspond to the vacuum oscillation averages (4.12).

In the first schematic calculation, we have introduced a post-averaging of the no-oscillation results as

\[
(g^{\text{post}})_{\beta} = \sum_{\alpha} (g^{\text{NO}})_{\alpha} T^{\text{vac}}(\alpha \to \beta).
\]  

(4.13)

From Table 1 one can see that the electronic spectra are suppressed and other neutrino types spectra are enhanced by the vacuum averaging procedure. One can nearly recover the oscillation case results by averaging the final results found without oscillations, thus showing that the different values of the effective neutrino temperatures between the no-oscillation case and the full oscillation case are likely to be due to the effect of the mixings. However, the post averaging of the no-oscillation case (which by construction preserves $N_{\text{eff}}$) does not capture the enhancement of $N_{\text{eff}}$ of the full oscillation case, discussed at the end of section 4.3.

In the second schematic calculation we have solved the QKEs (2.27) without the mean-field term, i.e., keeping only the vacuum and collision terms\(^{15}\) (Table 1). This is somehow an improvement of the “post averaging” procedure, since it neglects the variation of the transfer functions (which always have their asymptotic vacuum values), but accounts correctly for the effect of collisions. The accuracy of the results compared to the full treatment shows once more that the effect of the mean-field is very mild in this case. Indeed, the mean-field contribution becomes effective when $\varrho$ deviates from a matrix proportional to the identity, which only happens when $x \sim 3 \times 10^{-1}$: however at this point the mean-field contribution is becoming negligible compared to the vacuum one (cf. figure 5). Note that this would not hold if we introduced chemical potentials [24, 37, 42–45]. The higher value obtained for $N_{\text{eff}}$ in this case can be qualitatively understood. Since $T^{\text{vac}}(e \to e) < T(x \ll 1, e \to e)$, $\nu_e$ produced by collisions will be more converted into other flavours (in particular $\nu_\tau$) at early times compared to the full calculation. This frees some phase space for the reheating of $\nu_e$,

\(^{15}\)We thus have $U_m = U$ and the matter basis is the mass basis.
which is the dominant process. More entropy is transferred from $e^\pm$ annihilations, which increases slightly $N_{\text{eff}}$.

These transfer functions also shed some light on the importance of the precise value of the mixing angles, which explain some discrepancy with previous results (see section 4.3). Indeed, varying $\theta_{ij}$ within their uncertainty ranges slightly modify the $T(\alpha \rightarrow \beta)$ curves, which can cross each other. For instance, with the set of parameters used in [10], the asymptotic value $T^\text{vac}(e \rightarrow \tau)$ is higher than $T^\text{vac}(e \rightarrow \mu)$, contrary to figure 5. This higher conversion of electron neutrinos into tau neutrinos explains why their final temperatures are $z_{\nu_e} \gtrsim z_{\nu_\mu}$ (the values remaining very close).

**Sensitivity to the mass hierarchy** In the inverted hierarchy, for which $\Delta m^2_{31} < 0$, $N_{\text{eff}}$ is increased by $5 \times 10^{-6}$. In this case, $\nu_e$ can be generated above an MSW resonance (e.g. at about 4 MeV for $y = 5$), and are converted nearly entirely as $\nu_\mu$ and $\nu_\tau$ (figure 5). Again, this impacts subsequent collisions because it frees some phase space for $\nu_e$, which is beneficial for the total production of neutrinos. However, since neutrino decoupling occurs mainly at temperatures which are below the MSW resonance, the differences between normal and inverted hierarchies are extremely small.

To summarise, neutrino decoupling is mostly sensitive to the neutrino mixings, whereas it has little sensitivity to the mass-squared differences and therefore to the neutrino mass hierarchy.

5 **Flavour oscillations and Big Bang Nucleosynthesis**

Predicting a precise value of $N_{\text{eff}}$ in the standard cosmological model is timely since forthcoming generations of CMB experiments aim at measuring a possible contribution of light relic particles predicted by extensions of the standard model [46]. Yet CMB is not the only cosmological stage impacted by neutrinos, and $N_{\text{eff}}$ can be further constrained using the predicted abundances of light elements produced during BBN.

Indeed, incomplete neutrino decoupling, by giving rise to slightly non-thermal spectral distortions in neutrino spectra and modifying the photon to neutrino temperature ratio, affects BBN in various ways (see ref. [13] for a review).

1. The neutron-to-proton ratio freezes out from equilibrium when the rates of $n \leftrightarrow p$ interconversion reactions ($n + \nu_e \leftrightarrow p + e^-$, $n + e^+ \leftrightarrow p + \bar{\nu}_e$, $n \leftrightarrow p + e^- + \bar{\nu}_e$) drop below the Hubble expansion rate. The neutron fraction $X_n \equiv n_n/n_b$, with $n_b$ the baryon density, thus depends on when freeze-out occurs, and on deviations from standard nuclear statistical equilibrium that all depend on $z$, $z_{\nu_e}$ and $\delta g_{\nu_e}$ [5].

2. After the freeze-out, the neutron fraction decreases since neutrons continue to undergo beta decay until the onset of nucleosynthesis at $T_\gamma = T_{\text{Nuc}}$. The higher energy density of neutrinos for a given photon temperature, parametrized by $N_{\text{eff}}$, increases the Hubble rate compared to the instantaneous decoupling case, thus diminishing the number of neutrons that decayed. This is the so-called clock effect [47, 48], which tends to increase the fraction of neutrons at the beginning of nucleosynthesis $X_n(T_{\text{Nuc}})$, and consequently the helium fraction $Y_p \equiv 4n_{4\text{He}}/n_b \simeq 2X_n(T_{\text{Nuc}})$ since these neutrons are almost fully converted into $^4\text{He}$.

---

\[16\] This is not the case for very large $y$ but they are subdominant in the total energy density budget.
3. The production of other light elements from the remaining traces of neutrons is also controlled by the clock effect [4, 5]. For instance, deuterium is mainly destroyed from its equilibrium value to its frozen-out abundance at the end of BBN [49]. The higher expansion rate leaves less time for this destruction to happen, which leads to a net increase of the deuterium abundance.

There was some discrepancy in the literature about the sign of variation of the different abundances due to these effects, see for instance Table 3 in [9] and Table V in [4]. The extensive analysis of ref. [5] favoured the latter results. Though it did not include flavour oscillations, it predicted that the main conclusions would hold since the final neutrino spectra are qualitatively similar to the no-oscillations case, only “averaged”.

We aim at filling this gap and therefore introduce the results from section 4.3 in the BBN code PRIMAT [13]. This section is meant as an extension of the work [5], from which we will borrow the notation. We implement neutrino-induced corrections following the three levels of refinements introduced in [5].

\( i \) assuming that the three neutrino species have thermal spectra at the average temperature \( \hat{T}_\nu \equiv \frac{1}{3} (T_{\nu_e}^4 + T_{\nu_\mu}^4 + T_{\nu_\tau}^4)^{1/4} \) (“\( \hat{T}_\nu \)”),

\( ii \) using the proper effective temperature for \( \nu_e \), but without non-thermal distortions (“\( T_{\nu_e} \), no distortions”), and

\( iii \) using the real spectra from NEVO (“\( T_{\nu_e} \), with distortions”). Note that the total neutrino energy density, so \( N_{\text{eff}} \), is identical in all three implementations, therefore the clock effect contributions will be identical. We report the obtained values for the abundances of helium-4, deuterium, helium-3 and lithium-7 in Table 2, with the associated relative variations compared to the instantaneous decoupling case\(^{17} \) in Table 3.

| BBN framework | \( Y_P \) | \( D/H \times 10^5 \) | \( ^3\text{He}/H \times 10^5 \) | \( ^7\text{Li}/H \times 10^{10} \) |
|---------------|--------|-----------------|-----------------|-----------------|
| Inst. decoupling | 0.24705 | 2.4503 | 1.0728 | 5.6937 |
| \( \hat{T}_\nu \) (NO) | 0.24710 | 2.4594 | 1.0741 | 5.6702 |
| \( T_{\nu_e} \), with distortions (NO) | 0.24710 | 2.4594 | 1.0741 | 5.6701 |
| \( \hat{T}_\nu \) | 0.24710 | 2.4595 | 1.0741 | 5.6697 |
| \( T_{\nu_e} \), no distortions | 0.24707 | 2.4593 | 1.0741 | 5.6693 |
| \( T_{\nu_e} \), with distortions | 0.24716 | 2.4598 | 1.0742 | 5.6705 |

Table 2. Light element abundances, including all weak rate corrections [13] and QED corrections up to \( O(e^3) \) to plasma thermodynamics, for various implementations of neutrino-induced corrections. \(^3\text{He} \) stands for \(^4\text{He} + \text{T} \) and \(^7\text{Li} \) stands for \(^7\text{Li} + \text{Be} \) to account for slow radioactive decays. We compare the inclusion of results from neutrino decoupling with and without (NO) oscillations.

The variation of the \(^4\text{He} \) abundance due to incomplete neutrino decoupling is estimated by

\[
\delta Y_P = \delta X_n^{[\text{Nuc}]_n} = \delta X_n^{[\text{FO}]} + \delta X_n^{[\Delta t]},
\]

where the first equality comes from the almost total conversion of free neutrons into \(^4\text{He} \). \( \delta X_n^{[\text{FO}]_n} \) is the variation of the neutron fraction at freeze-out (point 1 above), and \( \delta X_n^{[\Delta t]} \) is the variation due to the different duration of beta decay (clock effect, point 2 above).

\(^{17}\)The instantaneous decoupling baseline is the same with or without flavour oscillations, since in this limit all three neutrino species have FD spectra at the comoving temperature \( T_{\text{cm}} \).
variation of the other abundances relative to the proton fraction $i/H \equiv n_i/n_H$ is given by \[ \delta(i/H) \simeq \delta X_i^{[\Delta t]} + \delta Y_P, \] where $\delta X_i^{[\Delta t]}$ is the variation of the final abundance due to the clock effect, and the $\delta Y_P$ contribution is actually $-\delta X_H$.

**Comparison of implementations** An a priori surprising conclusion of ref. [5] was the quasi-equivalence of the "\(\hat{T}_\nu\)" and full implementations. We recover this feature in the no-oscillation case (cf. for instance the first two lines of Table 3), while there is a sizeable difference when using the neutrino spectra with oscillations. Let us focus on the helium fraction $Y_P$. We plot the variation of the neutron fraction at freeze-out $\delta X_n^{[\text{FO}]}$ on figure 6. First note that the average temperature implementations give quasi identical results with and without oscillations, which is a direct consequence of the small difference of $N_{\text{eff}}$ in Table 1. Then, including the true $T_{\nu_e}$ reduces $\delta X_n^{[\text{FO}]}$: since $T_{\nu_e} > \hat{T}_\nu$, the weak rates increase and freeze-out is delayed, thus $X_n$ tracks its equilibrium value longer. This reduction of $X_n^{[\text{FO}]}$ is more important without oscillations because the effective $\nu_e$ temperature is much higher than the average temperature in this case (cf. figure 3). Finally, the spectral distortions alter the detailed balance relation which sets the neutron-to-proton ratio [5], shifting $X_n^{[\text{FO}]}$ in the opposite direction. Once again, this re-increase of the neutron fraction is more important in the no-oscillation case, since $|\delta g_{\nu_e}^{(\text{NO})}| > |\delta g_{\nu_e}|$ (cf. figure 4).

$N_{\text{eff}}$, having the same value in all three implementations, the difference between the last three lines of Tables 2 and 3 lies in the variation of $\delta X_n^{[\text{FO}]}$. This is somehow hidden for $^3\text{He}$ and $^7\text{Li}$ because they are the aggregated results of $(^3\text{He} + T)$ and $(^7\text{Li} + ^7\text{Be})$ respectively.

All in all, the final value of $\delta X_n^{[\text{FO}]}$ is higher with oscillations, and exceeds the average temperature value (i.e., the solid green curve is above the solid blue one, while the dash-dotted green and blue curves almost coincide). This could be surprising, since $z_{\nu_e}$ and $\delta g_{\nu_e}$ are both reduced by about 25% with mixing (Figs. 3 and 4), so we would expect the solid curves to be in homothetic ratio with the dash-dotted ones. However, $z_{\nu_e}$ is reduced by 25% \textit{compared to} $z = 1$, but is much more reduced, by $\sim 68\% \text{ compared to } \hat{z}$. That is why the gap between the solid blue and orange curves is 68% smaller than the gap between the dash-dotted blue and orange curves. Since the up-shifting of $X_n$ due to distortions is just reduced by $\sim 10\%$,$^{18}$ the "\(T_{\nu_e}, \text{ with distortions}\)" value in the oscillation case is higher.

$^{18}$It is not a 25% reduction since the relation between $\delta g_{\nu_e}$ and the modification of detailed balance is not exactly linear.

| BBN framework                  | $\delta Y_P$ (%) | $\delta (\text{D}/\text{H})$ (%) | $\delta (^3\text{He}/\text{H})$ (%) | $\delta (^7\text{Li}/\text{H})$ (%) |
|-------------------------------|-----------------|----------------------------------|------------------------------------|-----------------------------------|
| $\hat{T}_\nu$ (NO)            | 0.020           | 0.368                            | 0.121                              | -0.413                            |
| $T_{\nu_e}$, with distortions (NO) | 0.020           | 0.368                            | 0.121                              | -0.414                            |
| $\hat{T}_\nu$                 | 0.021           | 0.374                            | 0.123                              | -0.420                            |
| $T_{\nu_e}$, no distortions    | 0.007           | 0.367                            | 0.120                              | -0.429                            |
| $T_{\nu_e}$, with distortions  | 0.042           | 0.386                            | 0.127                              | -0.406                            |

**Table 3.** Relative variations of the light element abundances compared to the instantaneous decoupling limit, in the same frameworks as Table 2.
Figure 6. Neutron fraction variation around freeze-out, for different implementations of neutrino-induced corrections, with and without flavour oscillations.

Note however that, although the average temperature implementation is less accurate in the oscillation case than in the no-oscillation case, it is sufficient to provide the various abundances at a relative precision of a few $10^{-4}$, which is well beyond experimental uncertainties. Moreover, this method is particularly simple, since all information is contained in one parameter, the average effective temperature $\hat{T}_\nu(T_\gamma)$. It can be used in a BBN code from a table of its values, or be deduced from the dimensionless heating function $N$ [50, 51] that parametrizes the heat transfer from $e^\pm$ annihilations and which can be fitted to the desired precision.

Overall effect of flavour oscillations Let us now discuss the global difference in the final abundances due to the inclusion of oscillations. To keep the discussion simple, we will discuss the average temperature implementation, where all the information about neutrino spectra is encoded in $\hat{T}_\nu$ (the differences between the three implementations for a given (no-)oscillation case being explained above). We see from Table 1 that $N_{\text{eff}}$ is slightly higher when including oscillations, thus increasing the clock effect. For instance, there will be less time for the destruction of deuterium to take place, and we expect a higher D abundance. The same argument goes for $^3$He and T, causing an increase of $^3$He/H. Last, the abundance $^7$Li/H is dominated by primary $^7$Be, that is produced during nucleosynthesis: a faster expansion diminishes the $^7$Be yield, and thus the value of $^7$Li/H. The results corresponding to the cases $\hat{T}_\nu$ (NO) and $\hat{T}_\nu$ in Tables 2 and 3 can be understood using these simple heuristic arguments.

6 Conclusions

We derived the QKEs governing neutrino evolution at the epoch of weak decoupling using a BBGKY-like formalism, obtaining the mean-field terms up to $O(1/m_W^2)$ order and the collision terms with their full matrix structure. We solved the QKEs and presented the ATAO approximation which allows to increase the computation speed. This approximation is based
on the assumptions that there is a clear separation of time-scales between the oscillation frequencies and the collision rate, the off-diagonal terms of the density matrix in the matter basis are averaged out and the matter basis evolves adiabatically.

Results on $N_{\text{eff}}$ and the neutrino final spectra were presented with a numerical precision better than $10^{-4}$. A better precision would require the inclusion of several corrections. First, one would need to consider QED effects at order $e^4$ and sub-leading logarithmic-dependent terms at order $e^2$ [7]. But more importantly it would not be possible to consider a homogeneous cosmology since fluctuations inherited from the inflationary phase, and imprinted in the CMB, are of order $10^{-5}$. One would then need to consider fluctuations in the QKE as was done to estimate fluctuations in the CMB.\footnote{Furthermore, the physics of decoupling also depends on the Fermi and Newton constants, and the latter is only known with a $4 \times 10^{-5}$ precision.}

The obtained value of $N_{\text{eff}} \simeq 3.0440$ and the associated spectral distortions were used in the BBN code PRIMAT to investigate the consequences of incomplete neutrino decoupling with flavour oscillations on the primordial production of light elements, solving the discrepancy between [9] and [4], with results in agreement with [5]. Even though the subsequent variations occur at precisions well beyond experimental uncertainties, we were able to understand the physical processes at play, thus checking the validity of our results. The nuclear abundances, with all weak rates corrections included (as in [13]), and taking completely into account neutrino distorted spectra, are reported in the last line of Table 2. The next update of PRIMAT will include these results.

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A Derivation of the formal collision term

Compared to the Boltzmann treatment of neutrino evolution, which neglects flavour mixing, the QKE contains mean-field terms, and the collision term has a richer matrix structure with non-zero off-diagonal components. To derive this collision term, i.e., the contribution to the evolution of the one-body density matrix from two-body correlations, one needs an expression for the correlated part $C$ in (2.11). It is obtained from the evolution equation for $\varrho^{(12)}$, where we separate correlated and uncorrelated parts [28].

To do so, we need a splitting similar to (2.10) for the three-body density matrix,

$$\varrho^{ikm}_{jln} = 6 \varrho^{i}_{j} \varrho^{k}_{m} \varrho^{l}_{n} + 9 \varrho^{i}_{j} \varrho^{k}_{m} \varrho^{l}_{n} + C^{ikm}_{jln}.$$ (A.1)

This allows (2.9) to be rewritten as an equation for the two-body correlation function [21]. In the \textit{molecular chaos} ansatz, correlations are built through a collision between uncorrelated particles. These correlations then evolve “freely”, i.e., we do not take into account a mean-field background for $C$. The evolution equation is thus greatly simplified, retaining only the vacuum and Born terms:

$$\frac{dC^{ik}_{jl}}{dt} = \left[ t^{i}_{r} C^{rj}_{kl} + i^{k}_{p} C^{ip}_{jl} - C^{ik}_{rl} t^{r}_{j} - C^{ik}_{jl} t^{r}_{p} \right] + (\bar{1} - \varrho)^{i}_{j} (1 - \varrho)^{k}_{p} \varrho^{l}_{q} \varrho^{r}_{s} (\bar{1} - \varrho)^{r}_{t} (1 - \varrho)^{t}_{u},$$ (A.2)
where the second line will be labelled $B_{jkl}^{ik}$. We can actually solve this equation, starting from $C(t = 0) = 0$,  

$$
C_{jkl}^{ik}(t) = -i \int_0^t ds T_{mp}^{lk}(t, s) B_{nlq}^{mq}(s) T_{jpl}^{tnq}(t, s),
$$  

(A.3)

with the evolution operator  

$$
T_{jkl}^{ik}(s, s') = \exp \left( -i \int_{s}^{s'} d\tau \hat{\mathcal{H}}(\tau) \right) \left( \begin{array}{c}
\hat{\mathcal{T}}_{ik}^{jl} \\
\hat{\mathcal{T}}_{jkl}^{ik}
\end{array} \right).
$$  

(A.4)

Now we consider that there is a clear separation of scales [18], hence the duration of one collision is very small compared to the variation timescale of the density matrices (i.e., compared to the duration between two collisions, and the typical inverse oscillation frequency). Therefore, the argument inside the integral of (A.3) is only non-zero for $s \approx 0$: we can extend the integration domain to $+\infty$, while the operators keep their $t = 0$ value. Finally we symmetrize the integration domain\(^{20}\) with respect to 0 (with an extra factor of 1/2), which leads to the equation with collision term:  

$$
i \frac{d\hat{\varrho}_j^i}{dt} = \left[ \hat{I} + \hat{\mathcal{H}}, \hat{\varrho} \right] + \frac{i}{4} \int_{-\infty}^{+\infty} dt \left[ \hat{\mathcal{V}}, T(t, 0)B(0)T^\dagger(t, 0) \right]_{jk}^{ik}
$$  

(A.5)

\begin{align*}
&= \left[ (\hat{t}_k^i + \Gamma_k^i)\hat{\varrho}_j^k - \hat{\varrho}_j^i (\hat{t}_k^k + \Gamma_k^k) \right] \\
&- \frac{i}{4} \int_{-\infty}^{+\infty} dt e^{-i(E_m + E_l - E_j - E_k)t} \left[ \hat{\mathcal{V}}_{il} B_{jk}^{il} - B_{il}^{jk} \hat{\mathcal{V}}_{il} \right], \\
&\equiv \left[ \hat{I} + \hat{\mathcal{H}}, \hat{\varrho} \right] + i \hat{\mathcal{C}}_j^i
\end{align*}

(A.6)

$$
\equiv \left[ \hat{I} + \hat{\mathcal{H}}, \hat{\varrho} \right] + i \hat{\mathcal{C}}_j^i
$$  

(A.7)

The exponential of energies comes from the $T$ terms, using that the density matrix for a given momentum $\varrho(p)$ satisfies $\hat{\mathcal{H}}(p) = p \varrho(p)$.  

\section*{B Interaction potential matrix elements}

The relevant two-body interactions correspond to standard model interactions involving neutrinos and antineutrinos. In the early universe, they interact through weak processes with electrons, positrons and other (anti)neutrinos. Therefore, we must take as interaction Hamiltonian (2.6) the useful part of the standard model Hamiltonian of weak interactions, that is given by  

$$
\hat{H}_{\text{int}} = \hat{H}_{\text{CC}} + \hat{H}_{\text{NC}}^\text{mat} + \hat{H}_{\text{NC}}^\nu,
$$  

(B.1)

where we separated three contributions:  

- the charged current hamiltonian,  

$$
\hat{H}_{\text{CC}} = 2\sqrt{2} G F m_W^2 \int \left[ d^3 p_1 \right] \left[ d^3 p_2 \right] \left[ d^3 p_3 \right] \left[ d^3 p_4 \right] (2\pi)^3 \delta^{(3)}(\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4) \times \left[ \bar{\psi}_\nu(\vec{p}_1) \gamma_\mu P_L \psi_e(\vec{p}_4) \right] W^{\mu\nu}(\Delta) \left[ \bar{\psi}_e(\vec{p}_2) \gamma_\nu P_L \psi_\nu(\vec{p}_3) \right],
$$  

(B.2)

---

\(^{20}\)See section 6.1 in ref. [52] for a detailed discussion of this procedure.
with \( \psi(p) = \sum_h \left[ \hat{a}(p, h)u^h(p) + \hat{b}^\dagger(-p, h)v^h(-p) \right] \) the Fourier transform of the quantum fields, \( P_L = (1 - \gamma_5) / 2 \) the left-handed projection operator, and the gauge boson propagator

\[
W^{\mu\nu}(\Delta) = \frac{g^{\mu\nu} - \Delta^{\mu} \Delta^{\nu}}{m_W^2 - \Delta^2} \approx \frac{g^{\mu\nu}}{m_W^2} + \frac{1}{m_W^2} \left( \frac{\Delta^2 g^{\mu\nu}}{m_W^2} - \frac{\Delta^{\mu} \Delta^{\nu}}{m_W^2} \right). \tag{B.3}
\]

The lowest order in this expansion is the usual 4-Fermi effective theory. The momentum transfer is \( \Delta = p_1 - p_4 \) for a t-channel (\( \nu_e - e^- \) scattering), and \( \Delta = p_1 + p_2 \) for the s-channel (\( \nu_e - e^+ \)). At Fermi order, we get for instance (after a Fierz transformation):

\[
\tilde{v}_{\nu_e(3)e(4)}^{(1)a(2)} = 2\sqrt{2}G_F (2\pi)^3 \delta^{(3)}(\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4) \times [\tilde{v}_{\nu_e}^{\mu}(\vec{p}_1)\gamma^{\mu}P_L u_{\nu_e}^{h_3}(\vec{p}_3)] [\tilde{u}_{\nu_e}^{\mu}(\vec{p}_2)\gamma^{\nu}P_L u_{\nu_e}^{h_1}(\vec{p}_4)]. \tag{B.4}
\]

- the neutral current interactions with the matter background (electrons and positrons),

\[
\hat{H}_{NC}^{\mu\nu} = 2\sqrt{2}G_F m_Z^2 \sum_\alpha \int [d^3\vec{p}_1][d^3\vec{p}_2][d^3\vec{p}_3][d^3\vec{p}_4] (2\pi)^3 \delta^{(3)}(\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4)
\]

\[
\times [\tilde{v}_{\nu_e}^{\mu}(\vec{p}_1)\gamma^{\mu}P_L \psi_{\nu_e}(\vec{p}_3)] Z^{\nu\mu}(\Delta) [\tilde{\psi}_{\nu_e}(\vec{p}_2)\gamma^{\nu}P_L \psi_{\nu_e}(\vec{p}_4)]. \tag{B.5}
\]

where \( Z^{\mu\nu} \) is identical to \( W^{\mu\nu} \) with the replacement \( m_W \to m_Z \). The neutral-current couplings are \( g_L = -1/2 + \sin^2 \theta_W \) and \( g_R = \sin^2 \theta_W \), where \( \sin^2 \theta_W \approx 0.231 \) is the weak-mixing angle.

- the self-interactions of neutrinos,\(^{21}\)

\[
\hat{H}_{NC}^{\nu\nu} = \frac{G_F}{\sqrt{2}} m^2 \sum_{\alpha, \beta} \int [d^3\vec{p}_1][d^3\vec{p}_2][d^3\vec{p}_3][d^3\vec{p}_4] (2\pi)^3 \delta^{(3)}(\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4)
\]

\[
\times [\tilde{v}_{\nu_e}^{\mu}(\vec{p}_1)\gamma^{\mu}P_L \psi_{\nu_e}(\vec{p}_3)] Z^{\nu\mu}(\Delta) [\tilde{\psi}_{\nu_e}(\vec{p}_2)\gamma^{\nu}P_L \psi_{\nu_e}(\vec{p}_4)]. \tag{B.6}
\]

We show in Table 4 the set of interaction matrix elements derived from these Hamiltonians, which are needed for the neutrino collision term. To compute the mean-field potentials at order \( 1/m_{W,Z}^2 \), one needs the matrix elements from the expansion of the propagator (B.3), which are obtained similarly and not reproduced here for the sake of brevity.

At leading order, the charged-current processes are written as neutral-current ones thanks to Fierz rearrangement identities. Therefore one can write the global expression:

\[
\tilde{v}_{\nu_e(3)e(4)}^{(1)a(2)} = 2\sqrt{2}G_F (2\pi)^3 \delta^{(3)}(\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4)
\]

\[
\times [\tilde{v}_{\nu_e}^{\mu}(\vec{p}_1)\gamma^{\mu}P_L u_{\nu_e}^{h_3}(\vec{p}_3)] [\tilde{u}_{\nu_e}^{\mu}(\vec{p}_2)\gamma^{\nu}P_L u_{\nu_e}^{h_1}(\vec{p}_4)]. \tag{B.7}
\]

with, in the Standard model,

\[
G_L = \text{diag}(g_L + 1, g_L, g_L) \quad , \quad G_R = \text{diag}(g_R, g_R, g_R). \tag{B.8}
\]

One can also introduce non-standard interactions which promote the couplings to non-diagonal matrices [10].

\(^{21}\)To understand the different prefactor from \( \hat{H}_{NC}^{\text{mat}} \), start from the general neutral-current Hamiltonian:

\[
\hat{H}_{NC} = 2\sqrt{2}G_F m_Z^2 \sum_{f,f'} \int \cdots [\tilde{v}_{f f'}^{\gamma_{\nu}(g_L^f P_L + g_R^f P_R)\psi_f}] Z^{\gamma_{\nu}}(\Delta) [\tilde{\psi}_{f f'}^{\gamma_{\nu}(g_L^{f'} P_L + g_R^{f'} P_R)\psi_f}].
\]

Now the multiplicity of each term and the use of \( g_L^f = 1/2, g_R^f = 0 \) lead to the Hamiltonians above.
| Interaction process | $\bar{v}_{\nu_3}^{2(1)} \left[ \sqrt{2} G_F (2\pi)^3 \delta^{(3)}(\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4) \right] $ |
|---------------------|----------------------------------------------------------------------------------------------------------------------------------|
| $\nu_\alpha(1)\nu_\beta(2)\nu_\gamma(3)\nu_\delta(4)$ | $2 \times [\bar{u}_{\nu_\alpha}^h(\vec{p}_1)\gamma^\mu P_L u_{\nu_\beta}^h(\vec{p}_2)][\bar{u}_{\nu_\gamma}^h(\vec{p}_3)\gamma^\mu P_L u_{\nu_\delta}^h(\vec{p}_4)]$ |

Table 4. Interaction matrix elements at lowest order in the expansion of the gauge boson propagators (Fermi effective theory of weak interactions).

### C Neutrino self-interactions collision term

As an illustration of the use of the BBGKY formalism to derive the collision integrals, we detail the steps to obtain the neutrino-neutrino scattering contribution to (2.22).

Neutrino-neutrino scattering processes correspond to the terms in (2.13) for which the inner matrix elements are scattering ones $\bar{v}_{\nu_i}^0 v_{\nu_i}^0$. For simplicity, we focus here on the first term in the expression of $C_{ij}^{1}$ (2.13). Here, the index $i_1$ will refer to $\nu_\alpha(\vec{p}_1)$ and $i_2'$ to $\nu_\beta(\vec{p}_2')$.

There are two possible contributions to this collision matrix (note that we impose $\vec{p}_k = \vec{p}_k'$ for all $k$, which is enforced by the assumption of homogeneity (2.14)):

- when 1 and 3 have the same flavour, the scattering amplitude is:

$$
\bar{v}_{\nu_3}^{v_{\nu_1}(1)} v_{\nu_2}^{v_{\nu_2}(1)} v_{\nu_4}^{v_{\nu_3}(2)} v_{\nu_4}^{v_{\nu_4}(4)} \times \bar{v}_{\nu_3}^{v_{\nu_3}(3)} v_{\nu_4}^{v_{\nu_4}(4)} = 2G_F^2 \times (2\pi)^6 \delta^{(3)}(\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4) \delta^{(3)}(\vec{p}_1 - \vec{p}_2') \\
\times [\bar{u}_{\nu_\alpha}^h(\vec{p}_1)\gamma^\mu P_L u_{\nu_\beta}^h(\vec{p}_2)][\bar{u}_{\nu_\gamma}^h(3)\gamma^\mu P_L u_{\nu_\delta}^h(1)] \times [\bar{u}_{\nu_\alpha}(2)\gamma^\mu P_L u_{\nu_\beta}(4)][\bar{u}_{\nu_\gamma}(4)\gamma^\mu P_L u_{\nu_\delta}(2)] \\

= 2G_F^2 \times (2\pi)^6 \delta^{(3)}(\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4) \delta^{(3)}(\vec{p}_1 - \vec{p}_2') \\
\times p_{3\mu} p_{1\rho} p_{2\sigma} \gamma^\rho P_L \gamma^\sigma P_L \gamma^\mu P_L \gamma^\mu P_L \gamma^\mu P_L \\
\times \bar{p}_1^2 \Delta^2 \gamma^\mu P_L \gamma^\mu P_L \gamma^\mu P_L \\
= 2^5 G_F^2 \times (2\pi)^6 \delta^{(3)}(\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4) \delta^{(3)}(\vec{p}_1 - \vec{p}_2') \times (p_1 \cdot p_2)(p_3 \cdot p_4) \\ 

With this term, we have the matrix product

$$
\bar{v}_{\nu_3}^{v_{\nu_1}(3)} v_{\nu_2}^{v_{\nu_2}(4)} (1 - \theta)^{\delta^{(1)}(1 - \theta)^{\gamma^{(1)}}} (1 - \theta)^{\delta^{(2)}(1 - \theta)^{\gamma^{(2)}}} = [\text{Tr}[\theta_4 \cdot (1 - \theta_2)] \cdot \theta_3 \cdot (1 - \theta_1)]_{\beta_1}^{\alpha}.
$$
• when 1 and 4 have the same flavour, the scattering amplitude is:

\[
\tilde{\nu}_\gamma(1)\nu_e(2) \times \tilde{\nu}_\gamma(3)\nu_e(4) \times \tilde{\nu}_\gamma(3)\nu_e(4) \times \tilde{\nu}_\gamma(1)\nu_e(2)
\]

\[
= -2G_F^2 \times (2\pi)^6 \delta(3) (\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4) \delta(3) (\vec{p}_1 - \vec{p}_2)
\times [\tilde{\nu}_\gamma(1) (1) \gamma^\mu P_L u_{\nu_e}(4)] [\tilde{\nu}_\gamma(2) (2) \gamma^\mu P_L u_{\nu_e}(2)] [\tilde{\nu}_\gamma(3) (3) \gamma^\nu P_L u_{\nu_e}(3)] [\tilde{\nu}_\gamma(4) (4) \gamma^\nu P_L u_{\nu_e}(1)]
\]

\[
= -2G_F^2 \times (2\pi)^6 \delta(3) (\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4) \delta(3) (\vec{p}_1 - \vec{p}_2)
\times \rho_{3\lambda p_1 p_4 P_2 p_2} \times \gamma^\mu P_L \gamma^\nu P_L \gamma^\tau P_L \gamma^\lambda P_L \rho^\rho
\]

\[
= 2^5 G_F^2 \times (2\pi)^6 \delta(3) (\vec{p}_1 + \vec{p}_2 - \vec{p}_3 - \vec{p}_4) \delta(3) (\vec{p}_1 - \vec{p}_2) \times (p_1 \cdot p_2)(p_3 \cdot p_4)
\]

With this term, we have the matrix product

\[
\rho_{\delta(3)}(\rho_{\sigma(4)}) (1 - \rho_{\beta(1)}) (1 - \rho_{\gamma(2)}) = \left[ \rho_{4} \cdot (1 - \rho_{2}) \cdot \rho_{3} \cdot (1 - \rho_{1}) \right]_{\beta}.
\]

We chose the compact notation \( \rho_k \equiv \rho(p_k) \) for brevity, and used \( \rho_1 = \rho_\perp \) thanks to the momentum-conserving function \( \delta(3)(\vec{p}_1 - \vec{p}_2) \).

Considering all terms in (2.13), the scattering amplitude is always identical, and the matrix products arrange such that the final result has the expected gain – loss + h.c.” structure. Note that we considered here a particular ordering of the indices, while the full expression is symmetric through the exchange \((3, 4, 3', 4') \leftrightarrow (4, 3', 3')\). In other words, one must take twice the previous result to account for all non-zero combinations. Therefore,

\[
C^{(\nu \nu \leftrightarrow \nu \nu)} = (2\pi)^3 \delta(3)(\vec{p}_1 - \vec{p}_2) \frac{2^5 G_F^2}{2} \int [d^3 \vec{p}_2][d^3 \vec{p}_3][d^3 \vec{p}_4](2\pi)^4 \delta(4)(p_1 + p_2 - p_3 - p_4)
\times (p_1 \cdot p_2)(p_3 \cdot p_4) \times F_{sc}(\nu^{(1)}, \nu^{(2)}, \nu^{(3)}, \nu^{(4)})
\]

with the statistical factor:

\[
F_{sc}(\nu^{(1)}, \nu^{(2)}, \nu^{(3)}, \nu^{(4)}) = \left[ \rho_{4} \cdot (1 - \rho_{2}) + \text{Tr}(\cdots) \right] \rho_{3}(1 - \rho_{1}) + (1 - \rho_{1}) \rho_{3} \left[ (1 - \rho_{2}) \rho_{4} + \text{Tr}(\cdots) \right]
\]

\[
- \left[ (1 - \rho_{4}) \rho_{2} + \text{Tr}(\cdots) \right] \left[ (1 - \rho_{3}) \rho_{1} - \rho_{3} \left[ (1 - \rho_{3}) \rho_{4} + \text{Tr}(\cdots) \right] \right]
\]

where \( \text{Tr}(\cdots) \) means the trace of the term in front of it.

Finally, the collision integral \( \mathcal{I} \) which appears in the equation for \( \rho(p_1) \) is \( \mathcal{C} \) without the momentum-conserving delta-function \( C[\rho] = (2\pi)^3 2E_{1} \delta(3)(\vec{p}_1 - \vec{p}_2)\mathcal{I}[\rho] \).

### D Energy conservation and QED equation of state

The transfer of entropy from electron/positron annihilations into the photon and neutrino baths is governed by the continuity equation \( \dot{\rho} = -3H(\rho + P) \), which we rewrite as an equation on the dimensionless photon temperature \( z(x) [3, 7] \):

\[
\frac{dz}{dx} = \frac{x z J(x/z) - \frac{1}{2\pi^2 z^3} \frac{1}{x H} \int_0^\infty dy y^3 \text{Tr} \mathcal{I} \mathcal{G}_1(x/z) + G_2(x/z)}{x^2 J(x/z) + Y(x/z) + \frac{2\pi^2}{15} + G_2(x/z)}
\]

\[22\] This symmetry vanishes if 3 and 4 have the same flavour. However, this is precisely compensated by the extra factor of 2 in the matrix elements for identical flavour, cf. Table 4.
with

\[
J(\tau) \equiv \frac{1}{\pi^2} \int_0^\infty d\omega \omega^2 \frac{\exp(\sqrt{\omega^2 + \tau^2})}{(\exp(\sqrt{\omega^2 + \tau^2}) + 1)^2},
\]

\[
Y(\tau) \equiv \frac{1}{\pi^2} \int_0^\infty d\omega \omega^4 \frac{\exp(\sqrt{\omega^2 + \tau^2})}{(\exp(\sqrt{\omega^2 + \tau^2}) + 1)^2}.
\]

The $G_1$ and $G_2$ functions account for the modifications of the plasma equation of state due to finite-temperature QED corrections [3, 6, 7]. They can be calculated order by order in an expansion in powers of $\alpha = e^2/4\pi$. We use:

\[
G^{(2)}_1(\tau) = 2\pi\alpha \left[ \frac{K'(\tau)}{3} + \frac{J'(\tau)}{6} + J'(\tau)K(\tau) + J(\tau)K'(\tau) \right],
\]

\[
G^{(2)}_2(\tau) = -8\pi\alpha \left[ \frac{K(\tau)}{6} + \frac{J(\tau)}{6} - \frac{1}{2} K(\tau)^2 + K(\tau)J(\tau) \right]
+ 2\pi\alpha \tau \left[ \frac{K'(\tau)}{6} - K(\tau)K'(\tau) + \frac{J'(\tau)}{6} + J'(\tau)K(\tau) + J(\tau)K'(\tau) \right],
\]

\[
G^{(3)}_1(\tau) = -\sqrt{2}\pi\alpha^{3/2}\sqrt{J(\tau)} \times \tau \left[ 2j(\tau) - \tau j'(\tau) + \frac{\tau^2 j(\tau)^2}{2J(\tau)} \right],
\]

\[
G^{(3)}_2(\tau) = \sqrt{2}\pi\alpha^{3/2}\sqrt{J(\tau)} \left[ \frac{2J(\tau) + \tau^2 j(\tau)^2}{2J(\tau)} + 6J(\tau) + \tau^2 (3j(\tau) - \tau j'(\tau)) \right],
\]

where $(\cdots)' = d(\cdots)/d\tau$, and with the additional functions

\[
j(\tau) \equiv \frac{1}{\pi^2} \int_0^\infty d\omega \frac{\exp(\sqrt{\omega^2 + \tau^2})}{(\exp(\sqrt{\omega^2 + \tau^2}) + 1)^2},
\]

\[
K(\tau) \equiv \frac{1}{\pi^2} \int_0^\infty d\omega \frac{\omega^2}{\sqrt{\omega^2 + \tau^2} \exp(\sqrt{\omega^2 + \tau^2}) + 1},
\]

\[
k(\tau) \equiv \frac{1}{\pi^2} \int_0^\infty d\omega \frac{1}{\sqrt{\omega^2 + \tau^2} \exp(\sqrt{\omega^2 + \tau^2}) + 1}.
\]

Note that we discarded a logarithmic contribution to $G^{(2)}_{1,2}$ that is subdominant compared to $G^{(3)}_{1,2}$ [7]. Note that our expressions are formally different from those of previous literature. For instance (D.4) is formally different from the one in [3, 7], while (D.5) matches formally with [3], but not with [7]. Finally, (D.6) and (D.7) slightly differ from expressions reported in [7]. Actually, all expressions are identical, since one can prove (after integrations by parts and rearrangements) the following identities:

\[
J'(\tau) = -\tau j(\tau) , \quad K'(\tau) = -\tau k(\tau) , \quad Y'(\tau) = -3\tau J(\tau) , \quad 2K(\tau) + \tau^2 k(\tau) = J(\tau).
\]

### E Quantum Kinetic Equations with antiparticles

We present in this appendix the inclusion of antiparticles to the BBGKY formalism.
**Generalized definitions** One must adapt the definitions (2.3) and (2.6) to include the annihilation and creation operators $\hat{b}, \hat{b}^\dagger$. Throughout this appendix, we will emphasize the indices which are associated to antiparticles with a barred notation ($\bar{i}, \bar{\jmath}$). Therefore, with capital indices $I$ being either $i$ or $\bar{i}$, we have:

$$\rho^{I_1 \ldots I_s}_{J_1 \ldots J_s} \equiv \langle \hat{c}^\dagger_{J_s} \cdots \hat{c}^\dagger_{J_1} \hat{c}_{I_s} \cdots \hat{c}_{I_1} \rangle,$$

(E.1)

$$\hat{H}_0 = \sum_{I,J} t^I_J \hat{c}^\dagger_I \hat{c}_J,$$

(E.2)

$$\hat{H}_{\text{int}} = \frac{1}{4} \sum_{I,J,K,L} \tilde{v}^{IK}_{JL} \hat{c}^\dagger_I \hat{c}^\dagger_J \hat{c}_K \hat{c}_L,$$

(E.3)

where $\hat{c}_I = \hat{a}_i$ or $\hat{b}_{\bar{i}}$ depending on the index $I$ labelling a particle or an antiparticle.

The evolution equations (2.9) and (2.11) are naturally extended to the antiparticle case thanks to the global indices. The downside of this strategy is that the transformation law of tensors is now implicit: since $\hat{a}$ transforms like $\hat{b}^\dagger$ under a unitary transformation $\psi^a = U^a_i \psi_i$, the behaviour of upper and lower indices is inverted whenever they label an antiparticle degree of freedom, for instance:

$$t^i_j = U^a_i t^a_b U^b_j ; \quad t^{\bar{i}}_{\bar{j}} = U^a_{\bar{i}} t^a_{\bar{\jmath}} U^\dagger_{\bar{\jmath} b}.$$

(E.4)

Since we assume an isotropic medium, there are no “abnormal” or “pairing” densities [21–23] such as $\langle \hat{b} \hat{a} \rangle$, which ensures the separation of the two-body density matrix between the neutrino density matrix (for which we keep the notation $\rho$) and the antineutrino one $\bar{\rho}$. In order for $\bar{\rho}$ to have the same transformation properties as $\rho$, we need to take a transposed convention for its components:

$$\bar{\rho}^{\bar{i}}_{\bar{j}} = \rho^{\{J=j\}}_{\{I=i\}} = \langle \hat{c}^\dagger_{\bar{i}} \hat{c}_{\bar{j}} \rangle = \langle \hat{b}^{\dagger}_{\bar{i}} \hat{b}_j \rangle.$$

(E.5)

One could further take transposed conventions for the antiparticle indices in $t$ and $\tilde{v}$, which would ensure a clear correspondence between index position and transformation law–contrary to (E.4). For instance, $\tilde{t}^{I}_J \equiv t^I_J$ transforms as $t^I_J$. However, in order to keep a unique expression for the mean-field potential or the collision term, we stick to the general definitions above.

For instance, we have:

$$\Gamma^I_J = \sum_{K,L} \hat{v}^{IK}_{JL} \theta^L_K = \sum_{k,l} \hat{v}^{ik}_{jl} \theta^l_k + \sum_{k,l} \hat{v}^{ik}_{jl} \theta^l_k.$$

(E.6)

Since the annihilation and creation operators do not appear naturally in normal order in the Hamiltonian (B.1), recasting it in the form (E.3) leads to extra minus signs in $\tilde{v}$ involving antiparticles (cf. Table 4).

These conventions being settled, we can include the full set of interaction matrix elements and compute all relevant contributions to the neutrino QKEs (2.20). In the following, we derive the QKE for $\bar{\rho}$, which is not solved in this paper since we consider a zero asymmetry.\[23\]

\[23\]We just used the QKE for $\tilde{\rho}$ to check the numerical stability of the code.
QKE for antineutrinos Thanks to our conventions, the evolution equation for the antineutrino density matrix \( \bar{\rho} \) is similarly obtained within the BBGKY formalism, with some differences compared to the neutrino case. First and foremost, the evolution equation for \( \bar{\rho} \) correspond in the general formalism to the equation for \( \rho \):

\[
\frac{i}{\hbar} \frac{d\bar{\rho}^i_j}{dt} = \left( [t^j_i, \Gamma^j_i] \right) \rho^K_l - \bar{\rho}^j_i \left( t^K_l + \Gamma^K_l \right) + i \hat{C}^j_i, \tag{E.7}
\]

showing that taking the commutator with a transposed convention leads to a minus sign. Moreover,

- we express the kinetic terms \( t^j_i \), starting from the mass basis:

\[
t^j_i = U^a_j \frac{M^2}{2p} U^b_i \quad U^b_i = U^i_j \frac{M^2}{2p} \quad U^a_j = t^j_i; \tag{E.8}
\]

- \( \bar{v}^{jk} \) is the coefficient in front of \( \hat{b}^b_j \hat{a}^b_k \hat{a}^b_l \), so it will have the same expression (apart from the interchange of \( u \) and \( v \) spinors for neutrinos, which leaves the result identical) as the coefficient in front of \( \hat{a}^a_j \hat{a}^a_k \hat{a}^a_l \), that is \( -\bar{v}^{jk} \). Therefore, \( \Gamma^j_i = -\Gamma^j_i \).

Including these two results in (E.7) show that, compared to the neutrino case, the vacuum term gets a minus sign (from the reversed commutator), but not the mean-field. Formally,

\[
\frac{i}{\hbar} \frac{d\bar{\rho}^i_j}{dt} = \left[ -\hat{t} + \hat{\Gamma}, \bar{\rho} \right]^i_j + i \hat{C}^j_i. \tag{E.9}
\]

Two additional remarks:

- \( s \) and \( t \) channels are inverted when the particle 1 is an antineutrino (2 and 4 left unchanged). For instance, the scattering between \( \bar{\nu}_e \) and \( e^- \) is a \( s \)-channel (exchanged momentum \( \Delta = p_1 + p_2 \)), contrary to the scattering between \( \nu_e \) and \( e^- \) (\( \Delta = p_1 - p_2 \)). This changes the sign of \( \Delta^2 \), leading to another minus sign for \( \Gamma \) at order \( 1/m^2_{W,Z} \);

- the collision integral \( \bar{I} \) is obtained from \( I \) through the replacements \( \rho \leftrightarrow \bar{\rho} \) and \( g_L \leftrightarrow g_R \).

Considering all these remarks, we obtained the QKE for \( \bar{\rho} \) (2.21).

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