Radiative transfer effects during primordial helium recombination

Jens Chluba1*, Jeffrey Fung2† and Eric R. Switzer1,3‡

1 Canadian Institute for Theoretical Astrophysics, 60 St. George Street, Toronto, ON M5S 3H8, Canada
2 Department of Astronomy and Astrophysics, University of Toronto, 50 St. George Street, Toronto, ON M5S 3H8, Canada
3 Kavli Institute for Cosmological Physics, University of Chicago, 933 East 56th Street, Chicago, IL 60637, USA

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ABSTRACT

In this paper we refine the calculation of primordial helium recombination, accounting for several additional effects that were neglected or treated more approximately in previous studies. These include consideration of (i) time-dependent radiative transfer interaction between the $2^1P_1 - 1^1S_0$ and $2^3P_1 - 1^1S_0$ resonances; (ii) time-dependent radiative transfer for the partially overlapping $n^1P_1 - 1^1S_0$, $n^1D_2 - 1^1S_0$ and $n^3P_1 - 1^1S_0$ series with $3 \leq n \leq 10$; (iii) electron scattering within a kernel approach. We also briefly discuss the effect of electron scattering and H$_1$ quadrupole lines on the recombination of hydrogen. Although the physics of all considered processes is interesting and subtle, for the standard cosmology, with helium abundance $Y_p \approx 0.24$, the overall correction to the ionization history during helium recombination with respect to the previous implementation of CosmoRec remains smaller than $|\Delta N_e/N_e| \approx 0.05\%$. The dominant improvement is caused by consistent inclusion of resonance scattering for the $2^1P_1 - 1^1S_0$. For cosmologies with a large helium fraction, $Y_p \approx 0.4$, the difference reaches $|\Delta N_e/N_e| \approx 0.22\%$ at $z \approx 1800$, however, the overall correction to the CMB power spectra is small, exceeding $|\Delta C_l/C_l| \approx 0.05\%$ only at $l \gtrsim 3000$. In comparison to the current version of RecFast the difference reaches $|\Delta C_l/C_l| \approx 0.4\%$ at $l \approx 3000$ for $Y_p \approx 0.4$, and also for the standard value $Y_p \approx 0.24$ we find differences $|\Delta C_l/C_l| \gtrsim 0.1\%$ at $l \approx 2500$. The new processes are now included by the cosmological recombination code CosmoRec and can be activated as needed for most settings without affecting its runtime significantly.

Key words: Cosmic Microwave Background: cosmological recombination, temperature anisotropies, radiative transfer

1 INTRODUCTION

Over the past few years, several independent groups have considered (e.g. see Dubrovich & Grachev 2005; Chluba & Sunyaev 2006; Kholupenko & Ivanikh 2008; Wong & Scott 2007; Rubiño-Martín et al. 2008; Karshenboim & Ivanov 2008; Hirata 2008; Chluba & Sunyaev 2008; Jentschura 2009; Labzowsky et al. 2009; Grin & Hirata 2010; Ali-Haïmoud & Hirata 2010) the cosmological recombination problem of helium and hydrogen, with the premise of assessing how uncertainties in the theoretical treatment of the recombination process could affect the science return of PLANCK and future CMB experiments. Neglecting these recently-derived corrections to the standard recombination model (Seager et al. 1999, 2000) biases cosmological parameter estimates (Rubiño-Martín et al. 2010; Shaw & Chluba 2011). In particular, the inference of the slope of primordial scalar perturbations and baryon content. Especially when trying to distinguish between different competing inflationary models (see Komatsu et al. 2011, for recent constraints from WMAP), such inaccuracies should be avoided.

These efforts are now coming to an end, and it appears that all important corrections to the standard recombination scenario have been identified (e.g. see Pened et al. 2009; Rubiño-Martín et al. 2010, for overview). This has lead to the development of two new, independent recombination codes, CosmoRec (Chluba & Thomas 2011) and H$
u$Rec (Ali-Haïmoud & Hirata 2011), both of which include all important modifications to the recombination problem, superseding the physical model of RecFast. These new codes are fast, accurate, and self-contained in the sense that they do not need externally-calibrated effective rate modulation (“fudge”) factors from more complete recombination codes. A detailed code-comparison is currently in preparation, however, there are a few subtle radiative transfer effects during helium recombination that have not been discussed in detail thus far.

The current helium recombination model of CosmoRec is mainly based on the approach presented in (Chluba & Sunyaev 2010a, b), which we briefly recap in Sect. 2. Here we extend this model by explicitly solving the radiative transfer problem for the evolution of the high frequency distortion of the CMB introduced by neutral helium resonances. This extension allows us to include
redistribution of photons by resonance and electron scattering, and the full time-dependence of photon production and absorption (see Sect. 2.1) in an efficient and accurate manner.

The refinements to the helium recombination model discussed here are especially interesting since current constraints on the helium mass fraction, \( Y_p \), derived using ACT (Dunkley et al. 2011) and SPT (Keisler et al. 2011) data indicate a slightly larger helium abundance of \( Y_p = 0.313 \pm 0.044 \) and \( Y_p = 0.296 \pm 0.03 \), respectively. In that case it is expected that, e.g., the effect of resonance scattering on the \( ^2P_1 - ^1S_0 \) line is modified as compared to the standard cosmological model, with \( Y_p = 0.24 \). As this effect is currently only taken into account approximately (i.e., by using a correction function for the escape probability which was calibrated at fixed \( Y_p = 0.24 \) [Rubino-Martín et al. 2008], one might worry about the precision of the helium recombination calculation. Further, a detailed radiative transfer computation treats the feedback and interaction of overlapping excited levels (see Sect. 2.2). Our calculations show that for the analysis of future data from Planck, the considered processes are only marginally important (see Sect. 3.5).

Nevertheless, the new version of CosmoRec (version 2.0) now includes a helium radiative transfer module that can be activated as needed. This module might be useful for recalibration of the R- and Fast fudge-factors (see Sect. 3.5.2), once the final recombination code comparison is completed.

Finally, we briefly consider the effect of \( ^1S_0 \) quadrupole lines during hydrogen recombination. So far, only the \( nd \)-\( ls \) transitions were included using an effective transition rate between the \( nd \) and \( np \) state [Grim & Hirata 2010]. Here we confirm this approximation using a full radiative transfer computation. Furthermore, we include the quadrupole transitions among the excited levels and show that these actually dominate the correction. However, the final effect is negligible (see Sect. 3.7).

### 2 HELIUM RECOMBINATION PHYSICS

In this section we briefly outline how the radiative transfer calculations during helium recombination are carried out. The approach strongly relies on the treatment presented in Chluba & Thomas (2011) for the case of hydrogen recombination with several important differences, as we explain here.

#### 2.1 Time-dependent radiative transfer: main aspects

In simple words the helium and hydrogen recombination problem can be described as follows: free electrons are captured into some excited atomic state, forming a neutral atom under the emission of a photon in the free-bound continuum. From the excited levels, electrons can cascade towards energetically lower states, releasing one or more resonant photons. While most of the possible sequences of transitions are compensated by their inverse process (mediated by interactions with photons drawn from the ambient CMB), as recombination proceeds, more electrons reach the ground state of the atoms and distortions to the CMB blackbody spectrum are created. These distortions are very small in the Rayleigh-Jeans tail and close to the maximum of the CMB blackbody, however, in the Wien-tail they exceed the thermal radiation field (e.g. see Sunyaev & Chluba 2009, for more details about spectral distortions from the recombination epoch). In particular, non-thermal photons emitted at high frequencies (for hydrogen by the Lyman-series and for helium the \( n^1P_1 - ^1S_0 \) and \( n^3P_1 - ^1S_0 \) resonances) can lead to re-excitation of electrons from the ground state to higher levels and the continuum, and hence inhibit the recombination process. Therefore the recombination problem requires simultaneously solving the evolution of the populations of electrons in the excited states as well as the high frequency spectral distortions introduced by the recombination process. In particular it is important to compute at which rate photons can leave the cores of the resonances, thereby escaping from further (strong) interactions with the atoms.

Without additional approximations the recombination problem described above is prohibitively complicated. However, in the case of hydrogen it has now become possible to treat the recombination process with sufficient detail, achieving a precision \( \lesssim 0.1\% \) in the ionization history of the Universe close to \( z \approx 1100 \) [Chluba & Thomas 2011, Ali-Haimoud & Hirata 2011]. Here we extend the approach of Chluba & Thomas (2011) to account for the detailed physics of the helium recombination process. The main addition is that we explicitly solve the radiative transfer problem for the \( He^+ \) resonances, instead of relying on simple (occasionally even rough) approximations.

For the line emission and absorption process we use the 1 + 1 photon description (see Sect. 3.2 in Chluba & Thomas 2011). We neglect two-photon corrections to the shapes of the line profiles, as these are expected to lead to small changes in the ionization history (Switzer & Hirata 2008b). Nevertheless, the 1 + 1 photon treatment allows accounting for the full time-dependent evolution of the photon distribution, respecting detailed balance in the distant wings of the line profiles. Both aspects have been shown to be very important for hydrogen recombination (e.g., see Chluba & Sunyaev 2009b, 2010b), but thus far were only partially treated for helium.

We also include the redistribution of photons over frequency. Close to the Doppler cores of the optically thick lines, resonance

| Level | \( A_{ls} \) [s\(^{-1}\)] | \( v_{ls} \) [MHz] | \( \sigma_i \) [cm\(^2\)] | \( |\Delta z|/z \) | \( \Delta \text{thick} \) |
|-------|----------------|-----------------|----------------|----------------|----------------|
| \( ^2P_1 \) | 177.58 \times 10\(^3\) | 5.0691 | 5.1 \times 10\(^{-21}\) | – | 0.63 |
| \( ^2P_1 \) | 1.799 \times 10\(^3\) | 5.1305 | 5.0 \times 10\(^{-14}\) | 1.2 \times 10\(^{-2}\) | 120 |
| \( ^3P_1 \) | 56.1 \times 10\(^3\) | 5.5631 | 1.2 \times 10\(^{-21}\) | 7.8 \times 10\(^{-2}\) | – |
| \( ^3D_2 \) | 1.298 \times 10\(^3\) | 5.5793 | 4.7 \times 10\(^{-20}\) | 2.9 \times 10\(^{-3}\) | 2 |
| \( ^3P_1 \) | 5.663 \times 10\(^3\) | 5.5824 | 1.2 \times 10\(^{-14}\) | 5.6 \times 10\(^{-4}\) | 33 |
| \( ^3P_1 \) | 23.7 | 5.7325 | 4.8 \times 10\(^{-22}\) | 2.6 \times 10\(^{-2}\) | – |
| \( ^3D_2 \) | 748.5 | 5.7394 | 2.5 \times 10\(^{-20}\) | 1.2 \times 10\(^{-3}\) | 1.4 |
| \( ^3P_1 \) | 2.436 \times 10\(^3\) | 5.7408 | 4.9 \times 10\(^{-15}\) | 2.4 \times 10\(^{-4}\) | 13 |
| \( ^5P_1 \) | 12.1 | 5.8100 | 2.3 \times 10\(^{-22}\) | 1.2 \times 10\(^{-2}\) | – |
| \( ^5D_2 \) | 431.4 | 5.8135 | 1.4 \times 10\(^{-20}\) | 6.0 \times 10\(^{-4}\) | 1.2 |
| \( ^5P_1 \) | 1.258 \times 10\(^3\) | 5.8143 | 2.4 \times 10\(^{-15}\) | 1.4 \times 10\(^{-4}\) | 7 |
scattering is most important. Here we implement a Fokker-Planck approximation for the redistribution function, however, in addition we compare the results with those from a Monte Carlo approach (Switzer & Hirata 2008a). In the distant wings of the resonances and also in the cores of the optically thin lines, redistribution of photons by electron scattering becomes significant. We explain the treatment of this process in more detail below (see Sect. 2.3).

Finally, for helium recombination there is one very important difference relative to the hydrogen recombination problem: since already at $z \approx 2000$ a small fraction of neutral hydrogen atoms is present, high frequency photons emitted in the main helium resonances are able to additionally ionize hydrogen atoms. This removes helium photons from the cores of the lines and consequently strongly accelerates helium recombination (Kholupenko et al. 2007; Switzer & Hirata 2008b). Using the variables $x = \nu / \nu_\text{rest}$ and $\Delta n_i = \nu_3 / \Delta \nu_i$, where $\nu_3 = 5.1305 \times 10^{13}$ Hz is the $2^1S_1 - 1^1S_0$ transition frequency, and $\Delta \nu_i$ denotes the deviation of the photon occupation number from the CMB blackbody, the corresponding term in the transfer equation reads

$$ \frac{d \Delta n_i}{dz} \bigg|_{\nu = \text{abs}} = \frac{\sigma_{\text{abs}}(\nu) N_i c}{H [1 + z]} \Delta n_i. \quad (1) $$

Here $\sigma_{\text{abs}}(\nu)$ is the photoionization cross section from the ground state of hydrogen and $N_i$ denotes the $H_i$ ground state population. This expression has to be added to the radiative transfer equation describing resonance scattering, and line emission/absorption for the individual helium resonances (see Table 1 for transition rates and frequencies for some of the neutral helium lines) given in Chluba & Thomas (2011).

In the work presented here we do not explicitly discuss corrections to the $2^1S_1 - 1^1S_0$ two-photon channel. For example, stimulated $2^1S_0 - 1^1S_0$ two-photon decay should be included, accelerating helium recombination; at the same time one expects that feedback of photons from the main helium resonances can re-excite atoms to the $2^1S_0$ by $1^1S_0 - 2^1S_0$ two-photon absorption. Similar processes are very important for the $2s-1s$ two-photon channel of hydrogen (Chluba & Sunyaev 2004; Kholupenko & Ivanchik 2006), but for He I recombination the $1^1S_0 - 2^1S_0$ two-photon channel is much less important, allowing only $\lesssim 8\%$ of the helium atoms to become neutral (Rubino-Martín et al. 2008). We implemented the corresponding problem, and found that a detailed treatment leads to a correction $\lesssim 0.01\%$ in the ionization history of helium, which is in agreement with the analysis of Switzer & Hirata (2008a).

### 2.2 Overlapping resonances in the upper shells

In the previous treatments of He I recombination, the feedback of photons from higher to lower lines is modelled by the approximation that all the resonances are isolated (see Fig. 1), in the sense that within the proximity of any individual line, only the line itself contributes significant opacity (Chluba & Sunyaev 2004; Chluba & Thomas 2011). In this approach the standard Sobolev escape probability, $P_S(\tau_S) = [1 - e^{-\tau_S}] / \tau_S$, where $\tau_S$ is the Sobolev optical depth of the $i^{th}$ resonance, is replaced by the effective escape probability (see Chluba & Sunyaev 2010a, for more details)

$$ P_{\text{eff},S} \equiv P_{i,S} \left[ 1 - \frac{\Delta n_i}{\Delta n_i} e^{-\tau_{j,m}} \right], \quad (2a) $$

$$ P_{i,a} \equiv \frac{P_{i,S} P_{j,m}^e}{1 - P_{i,m}^e P_j}. \quad (2b) $$

Here, $P_{i,a} \equiv P_S(p_i, \tau_S)$, where $P_{i,m}^e = 1 - P_{i,m}^e$ is the death probability or line albedo of the $i^{th}$ resonance; $\Delta n_i(z)$ is the deviation of the line occupation number of the $i^{th}$ resonance from a blackbody; $\Delta n_i(z) e^{-\tau_{j,m}}$ is the distortion of the photon occupation number caused by the energetically higher line $j$ at the line center of $i$ after redshifting and absorption by neutral hydrogen; finally, $\tau_{j,m}$ is the H I continuum opacity between the upper and lower line.

In the case of hydrogen ($\tau_{j,m} \equiv 0$), the isolated line approximation works very well for the Lyman-series up to $n \approx 10 - 20$, since the resonances are sufficiently separated in frequency, while for larger $n$, overlap of the optically thick resonances becomes important (Ali-Ha¨ımoud et al. 2010). As pointed out earlier (Chluba & Sunyaev 2010a), for the resonances in the upper shells of helium, the isolated line approximation should already break down for $n \gtrsim 3$. This is because in every shell with $n \gtrsim 3$ photons encounter a triplet of $n^2P_1 - 1^1S_0$, $n^2D_1 - 1^1S_0$ and $n^2P_1 - 1^1S_0$ resonances (see Fig. 1). In particular, the $n^2D_1 - 1^1S_0$ quadrupole and $n^2P_1 - 1^1S_0$ dipole lines are separated by $\lesssim 30$ Doppler widths at $z \approx 2000$, while the $n^2P_1 - 1^1S_0$ dipole lines are optically thick up to comparable distances from the line center (see Table 1 for examples). Such proximity between the lines allows for a cross-communication in the damping wings, which becomes increasingly important towards the end of helium recombination; therefore, to correctly evaluate the escape probability of each line, the photon fields of nearby resonances must be taken into account. Furthermore, redistribution of photons by electron and line scattering may lead to additional subtle changes in the recombination dynamics that are not captured by the simple isolated line approximation. Explicitly solving the full radiative transfer equation, as described in the previous section, accounts for the physics of this problem self-consistently. We explain the different effects in more detail below.
2.3 Electron scattering

To account for the effect of electron scattering on the cosmological recombination process, we start with the Boltzmann collision term describing the redistribution of photons over frequency in an isotropic medium, e.g., see Sazonov & Sunyaev (2000):

$$\frac{\partial \Delta n_\nu}{\partial \tau_e} \bigg|_{z=0} = \int_0^\infty P_\nu \nu' \left[ \Delta n_\nu e^{\Delta \nu' + \nu' \Delta \tau_e} - \Delta n_{\nu'} \right] d\nu'. \quad (3)$$

Here $\Delta n_\nu$ is the deviation of the photon occupation number from the CMB blackbody; $\tau_e = \int \sigma_T N_e dr$ is the optical depth with respect to Thomson scattering; $\sigma_T \approx 6.65 \times 10^{-25} \text{cm}^2$ is the Thomson cross sections; $N_e$ is the free electron number density; and $P_\nu \nu'$ denotes the electron scattering kernel for thermal electron at temperature $T_e$. For the computations presented here, we use the Kompaneets equation kernel, $P_K$, as defined by Eq. (23) of Sazonov & Sunyaev (2004). It accounts for Doppler broadening, Doppler boosting, and electron recoil. However, both electron recoil and Doppler boosting remain rather small, and the redistribution of photons by electron scattering is dominated by Doppler broadening.

When the variations in the solution for the spectral distortion are very broad in comparison to the characteristic width of the scattering kernel, it is possible to resort to a Fokker-Planck approximation of the Boltzmann collision integral, Eq. (3). In this approach, $\Delta n_\nu$ is approximated using the first and second derivative of the photon occupation number, and the integral over frequency can be carried out analytically, leading to a second order parabolic partial differential equation (see, e.g., Sazonov & Sunyaev 2000). Such types of problems can be efficiently solved numerically, and the cosmological recombination code, CosmoRec, is readily extended to take this into account.

However, as explained earlier (Hirata & Sunyaev 2008), the characteristic width of the electron scattering kernel during helium recombination is given by

$$\frac{\Delta \nu}{\nu} \approx \sqrt{\frac{2 k T_e}{m_e c^2}} \approx 1.4 \times 10^{-3} \sqrt{\frac{1 + z}{2000}} \approx \frac{\sqrt{m_\nu \Delta \nu_\nu}{m_e} \nu}, \quad (4)$$

where $\Delta \nu_\nu$ is the Doppler width of the helium resonances, and $\sqrt{m_\nu \Delta \nu_\nu}{m_e} \approx 86$ arises from the helium to electron mass ratio. This turns out to be much wider than the typical variation of the spectral distortions found on the blue side of the high frequency $\text{He}^+ \nu$ resonances, and hence the simple Fokker-Planck approximation is expected to break down in regions close to the resonances. In this case, one faces solving the numerically expensive dense matrix equation $\mathbf{M} u = b$ for the evolution of the photon occupation number during helium recombination.

Fortunately, it turns out that in places where the variation of the phase space density is large (i.e. in the vicinity of the Doppler cores of the main $\text{He}^+ \nu$ resonances), electron scattering is always much less important than any of the line processes. This can be seen by comparing the mean free path to Thomson scattering, $\ell_T \approx 1/N_e \sigma_T$, with the one for resonance interactions (scattering + absorption) at the line center, $\ell_i \approx 1/N_{\text{He}^+} \sigma_i$. Here $N_{\text{He}^+}$ is the neutral helium number density and $\sigma_i$ the line center resonance scattering cross sections, which at $\approx 2000$ is given in Table 1 for the first few lines. At $z \approx 2000$ one has $N_{\text{He}^+}/N_e \approx 0.04$, so that even for the $5\text{P} \rightarrow 1\text{S}_0$ resonance (the one with the longest mean free path among the lines given in Table 1) we find $\ell_i/\ell_T \approx 14$. For the singlet $\text{P} \rightarrow 1\text{S}_0$ resonances one finds $\ell_i/\ell_T \gg 1$ even at very early stages of neutral helium recombination. Similarly, $\ell_i/\ell_T \gtrsim 1$ for the shown quadrupole lines, with a comfortable margin.

One can therefore resort to an iterative approach when solving the dense matrix equation at fixed redshift $z$: given the matrix equation $\mathbf{M} u = b$ describing the solution for the photon occupation number without accounting for the effect of electron scattering, one can compute the solution $u_0$, benefiting from the extreme sparseness of the problem. This solution is then used to compute the correction vector $\delta b_0 = -\mathbf{M} u_0$, which can be derived from the Boltzmann integral for electron scattering. In the next iteration, one now has to compute the correction to the solution, $\delta u_0$, from the matrix equation $\mathbf{M} \delta u_0 = \delta b_0$, which again can be calculated efficiently using banded matrix routines. This procedure can be repeated until convergence is reached, however, we find that usually one iteration per time-step is sufficient.

The above procedure is practically equivalent to the approach explained by Ali-Haëmoud et al. (2010) in the case of hydrogen recombination. However, during hydrogen recombination electron scattering is much less important for three reasons: (i) the width of the electron scattering kernel is only about $86 \sqrt{m_\nu \Delta \nu_\nu}{m_e} \approx 43$ times larger than the Doppler widths of the $\text{H}^+ \nu$ resonances; (ii) all the $\text{H}^+ \nu$ Lyman-series lines are extremely optically thick, so that in the vicinity of the resonances line scattering always dominates over electron scattering (Chluba & Sunyaev 2009a); (iii) the number of free electrons rapidly decreases as hydrogen recombination proceeds. Overall, electron scattering never becomes very important during hydrogen recombination (Chluba & Sunyaev 2009a). However, even in the case of hydrogen recombination, the effect of electron scattering on the recombination dynamics requires a treatment using the precise redistribution kernel (Ali-Haëmoud et al. 2010). We briefly discuss this in Sect. 3.3.

We emphasize that in most astrophysical situations the redistribution of photons by electron scattering can be safely described using a Fokker-Planck approximation (e.g., see Pozdniakov et al. 1979). The cosmological recombination epoch is one example for which this approximation fails, as we show below. It is the interplay between line emission/absorption processes, causing narrow features in the photon distribution, and broadband electron scattering which render it important to use the full redistribution kernel.

3 MODIFICATIONS TO THE COSMOLOGICAL IONIZATION HISTORY

In this section, we describe the corrections to the cosmological recombination history of helium arising from the additional processes treated in this paper. We proceed in a step-by-step manner, first comparing the results of our treatment with those obtained in earlier works. As a fiducial model we use the cosmological parameters $\Omega_b = 0.044$, $\Omega_{\text{dm}} = 0.216$, $T_0 = 2.725 \text{ K}$, $H_0 = 71.0 \text{ km s}^{-1} \text{ Mpc}^{-1}$, $Y_p = 0.24$, and $N_e = 3.046$. In addition, we assume flatness and an equation of state parameter $\omega = -1$ for dark energy.

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4 We neglect stimulated electron scattering and only follow the spectral distortion with respect to the CMB blackbody (see Chluba & Thomas 2011 for further justification).

5 In essence, one has to insert the solution $u_0$ into Eq. (3) and carry out the integrals to determine $b_0$. Since the integral has to be computed for every grid point, this computation still remains rather time-consuming.
3.1 Transport in the $^2\Pi_1 - ^1\Sigma_0$ and $^2\Pi_1 - ^1\Sigma_0$ resonances

In this section, we consider physical processes in the $^2\Pi_1 - ^1\Sigma_0$ and $^2\Pi_1 - ^1\Sigma_0$ transitions which most strongly regulate the formation of the neutral helium. One way to represent the corrections to the recombination dynamics of each line is by computing the effective escape probabilities (see Chluba & Sunyaev 2010a, Sect. 4.1; or Switzer & Hirata 2008a, Sect. V). This can be achieved in a multi-step process: (i) calculate the solution to the recombination problem based on an approximate analytic prescription for the escape probabilities of each line; (ii) compute the escape probability with a radiative transfer code using the obtained solutions for the level populations; (iii) iterate until convergence is achieved. Starting with the prescription implemented in CosmoRec v1.3 we found that usually one iteration with the new helium radiative transfer module was sufficient. Below we present the results for several cases, comparing with the standard Sobolev approximation and different intermediate approximations.

3.1.1 Escape from the $^2\Pi_1 - ^1\Sigma_0$ resonance

Figure 2 presents the effective escape probability of the $^2\Pi_1 - ^1\Sigma_0$ resonance for different settings, regarding feedback. For comparison, we show the standard Sobolev escape probability.

The $^2\Pi_1 - ^1\Sigma_0$ resonance no longer reach the $^2\Pi_1 - ^1\Sigma_0$ intercombination line because of absorption in the H I Lyman continuum. This feedback model was already incorporated into CosmoRec v1.3, however, the escape probability is slightly smaller when using the results from the radiative transfer code of CosmoRec v2.0 (curve labeled 'radiative transfer (with feedback)'). At early times differences at the level of $\approx 10\%$ are visible, while at $z \lesssim 2000$ the effective escape probability again becomes identical to the case without feedback. This result for the escape probability is consistent with the simpler radiative transfer computation presented in Fig. 6 of Chluba & Sunyaev 2010a. Physically, the main addition of the new radiative transfer module discussed here is line scattering, which can be neglected for $^2\Pi_1 - ^1\Sigma_0$. However, the feedback model of CosmoRec v1.3 in addition neglected part of the time-dependence, since the more detailed computation of Chluba & Sunyaev 2010a was too time-consuming to be solved quickly. CosmoRec v2.0 now includes all these effects simultaneously without compromising the performance too much.

3.1.2 Escape from the $^2\Pi_1 - ^1\Sigma_0$ resonance

Figure 3 shows the isolated line computations for the $^2\Pi_1 - ^1\Sigma_0$ resonance. We ran two cases with the radiative transfer code of CosmoRec v2.0, one with (curve labeled ‘radiative transfer (no feedback)’) and another without line scattering (‘radiative transfer (no feedback, no diffusion)). Again we compare with the standard Sobolev approximation, as well as the escape probability used in CosmoRec v1.3 (curve labeled ‘RM2008’), which is based on the work of Rubiño-Martin et al. 2008.

At low redshifts, the effect of H I continuum absorption becomes very large, leading to a strong increase in the escape probability of the $^2\Pi_1 - ^1\Sigma_0$ line (compare with Switzer & Hirata 2008a, Rubiño-Martin et al. 2008). Like for the $^2\Pi_1 - ^1\Sigma_0$ intercombination line, this is accounted for using a 1D integral approximation (Appendix B, Eq. (B.3) of Rubiño-Martin et al. 2008), but in addition with a redshift-dependent correction function to account for the effect of line-diffusion on the escape probability.

\footnote{To compute the escape probabilities in all cases, we use the solution for the populations of the levels that did not include the feedback correction. This allows us to directly compare all of them in one figure.}
In Rubiño-Martín et al. (2008) this effect was calibrated for fixed $Y_e = 0.24$, using a simple radiative transfer calculation, while here we consistently treat the transport problem case-by-case. As Fig. 4 indicates, for $Y_e = 0.24$ at both high and low redshifts, good agreement with Rubiño-Martín et al. (2008) is found. The main differences are visible close to $z \approx 2300$, reaching the level of 1%–5%. At this redshift the optical depth of the resonance is largest, so that similar to the escape problem of H i Lyman-$\alpha$ photons (see Chluba & Sunyaev 2009a, for more detailed explanation), the occupation number in the distant wings becomes important, introducing time-dependent aspects which are not included by the much simpler approximation of Rubiño-Martín et al. (2008).

From Fig. 4 one can also clearly see that at low redshifts line scattering leads to an important additional increase of the escape probability. We confirmed that the result of the new radiative transfer model is in agreement with the previous computations of Rubiño-Martín et al. (2008) when neglecting line-scattering. The effect of line-diffusion is currently not included by HyRec, but does lead to a correction of $\Delta \tau_{\text{int}} \approx -0.3\%$ at $z \approx 1800$, however, for the analysis of CMB data from Planck with $Y_e = 0.24$ this effect can apparently be neglected (Shaw & Chluba 2011).

As next step, we also include the effects of the other lines on the effective escape probability of the $2^1P_1 - 1^1S_0$ line. If we only add the $2^3P_1 - 1^3S_0$ line $P_1$ line in the radiative transfer code, we obtain the solid black line in Fig. 4. For clarity, we plot the relative difference with respect to the case for which only the $2^1P_1 - 1^1S_0$ resonance is present. Although the $2^1P_1 - 1^1S_0$ line energetically lies below the singlet line, the escape probability is affected at the level of $\approx -0.35\%$ at $z \approx 2300$, a correction that is not accounted for in earlier feedback models. This correction is consistent with absorption of $2^3P_1 - 1^3S_0$ photons in the red wing of the $2^1P_1 - 1^1S_0$ resonance. Since the Sobolev optical depth of the $2^3P_1 - 1^3S_0$ resonance during helium recombination remains $\tau_S \lesssim 3$, reabsorption of $2^3P_1 - 1^3S_0$ photons close to the $2^1P_1 - 1^1S_0$ line center in the blue wing of the $2^3P_1 - 1^3S_0$ line – an effect that could increase the $2^3P_1 - 1^3S_0$ escape probability – is negligible.

However, the modification to the escape probability caused by the presence of $2^3P_1 - 1^3S_0$ photons is smaller than the effect of feedback from the upper resonances with $n > 2$. Accounting for those lines, we find changes at the level of a few percent with respect to the isolated line case (see curve labeled ‘interaction with all lines $n < 11$’). Obviously, here the main effect arises from the feedback of photons from the triplet of lines with $n = 3$. Again one can observe that at $z \lesssim 2300$, feedback ceases since photons are efficiently absorbed in the H i Lyman continuum before being able to reach the $2^3P_1 - 1^3S_0$ line.

In Fig. 4 for comparison we also show the modification in the escape probability obtained with the feedback model of CosmoRec v1.3, which is based on the approximations of Chluba & Sunyaev (2010a). Although qualitatively similar, the feedback term is underestimated in this model. This is likely caused by an overestimation of the H i continuum opacity between the resonances of the third and second shell, as in CosmoRec v1.3 it is only computed roughly, using the Kramers’ approximation for the cross section. Also, the red wing absorption of $2^3P_1 - 1^3S_0$ photons by the $2^3P_1 - 1^3S_0$ line is not accounted for. With the new radiative transfer module these approximations can be avoided.

### 3.2 Partially overlapping triplets of lines with $n > 2$

In this section we illustrate the modifications to the escape probabilities of the upper, partially overlapping triplets of resonances with $n > 2$. Figure 4 shows the effective escape probabilities of the different lines for $3 \leq n \leq 5$. For the intercombination lines, one can clearly see that the simple isolated line approximation of Chluba & Sunyaev (2010a) is in rather good agreement with the result of the full radiative transfer calculation carried out here. Only at low redshifts, where the effect of H i continuum absorption is becoming significant, the difference is large. This is because in the simplified version of the treatment described in Chluba & Sunyaev (2010a) which is used in CosmoRec v1.3 this correction was neglected to accelerate the computation. Similarly, for the $n^1P_1 - 1^1S_0$ singlet lines at early times we find very good agreement with the approximations of Chluba & Sunyaev (2010a), with differences only showing up at the end of helium recombination.

For the $n^1D_2 - 1^1S_0$ quadrupole lines the overall differences relative to the radiative transfer calculation are much larger. We attribute these differences to details in the radiative transfer between the partially overlapping resonances, as well as the validity of approximations made in Chluba & Sunyaev (2010a). For example, for $n \geq 3$ the quadrupole and singlet lines are sufficiently close to each other to start communicating directly. Photons close to the $n^1P_1 - 1^1S_0$ resonance can be absorbed in the blue wing of the $n^1D_2 - 1^1S_0$ line. Similarly, absorption of photons in the cores of $n^1D_2 - 1^1S_0$ lines can occur via the red wing of the $n^1P_1 - 1^1S_0$. The latter process is more important, since the optical depth of the $n^1P_1 - 1^1S_0$ lines is orders of magnitude larger than for the corresponding quadrupole lines. The line therefore remains optically thick up to larger distances from the line center, comparable to the distance between the lines (see Table 1). The isolated line approximation breaks down for these lines, and at larger $n$ even the partial overlap with the intercombination line should start mattering.

We also confirmed numerically that line scattering itself is only responsible for a small part of the difference. Given that the differences are rather small, implying small changes in the recombination history, we did not attempt to further isolate the physical reasons for the deviations. Overall, for the upper levels the main improvement of the new radiative transfer module with respect to CosmoRec v1.3 is the consistent inclusion of line scattering and the absorption of photon in the H i continuum, which both become im-

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7 This implies that there is no direct line feedback to the $2^3P_1 - 1^3S_0$ resonance by redshifting.
transfer code of C (2010a). The blue are obtained using the isolated line approximation of Chluba & Sunyaev combination lines (upper panel), the

Figure 5. Comparison of escape probabilities for the $n^1P_1 - 1^3S_0$ intercombination lines (upper panel), the $n^1D_2 - 1^3S_0$ quadrupole lines (middle panel), and the $n^1P_1 - 1^3S_0$ singlet lines (lower panel) with $3 \leq n \leq 5$. The red/dashed lines give the Sobolev approximation; the solid/black lines are obtained using the isolated line approximation of Chluba & Sunyaev (2010a). The blue/dashed-dotted lines are computed with the radiative transfer code of CosmoRec v2.0.

Figure 6. Effect of electron scattering on the effective escape probability of the $2^3P_1 - 1^3S_0$ intercombination line. For the first three curves the $2^3P_1 - 1^3S_0$ resonance is treated as isolated line, while for the latter three curves all resonances with $n < 11$ are included.

important only at the end of helium recombination. However, because the upper levels contribute much less to the total recombination rate, the impact of these modifications remains small.

3.3 Electron scattering

In this section we investigate the effect of electron scattering on the escape probabilities. In particular, we demonstrate that the simple Fokker-Planck approximation for the redistribution of photons over frequency is a rather poor description of the problem. Unless stated otherwise, for the results presented here we use the helium radiative transfer code of CosmoRec v2.0.

3.3.1 Effect of electron scattering on the $2^3P_1 - 1^3S_0$ line

We first consider the isolated $2^3P_1 - 1^3S_0$ intercombination resonance. Figure 6 presents the effective escape probability for different runs with electron scattering enabled. The solid/blue line represents the result from Fig. 2 for comparison. The dotted/red curve is obtained using the Fokker-Planck approximation for the redistribution function of electron scattering. Comparing to the case without electron scattering, at $z \gtrsim 2100$ electron scattering causes a decrease in the escape probability, while at low redshifts it allows more photons to escape. If we compute the effective escape probability using the electron scattering kernel, as explained in Sect. 2.3 we obtain the black/dot-dash-dashed curve in Fig. 6. Here one can see that electron scattering only affects the escape probability at high redshifts and that the effect is significantly smaller than in the Fokker-Planck approximation. This curve is also in very good agreement with the result of Fig. 4 in Hirata & Switzel (2008), which was computed with a Monte Carlo approach (see Sect. 4.3 for more details), which confirms the validity of the approximations used in our implementation of this process.

To better understand the cause for these differences, in Fig. 7 we present the spectral distortion introduced by the $2^3P_1 - 1^3S_0$ intercombination resonance at $z \approx 2395$ as an example. Without any electron scattering, the spectrum has a steep step close to the resonance frequency of the $2^3P_1 - 1^3S_0$ line at $x \approx 0.988$. As the line
Below the resonance photons are efficiently scattering towards the blue wing, where a photon deficit is found.

Again taking a close look at the distortion across the Doppler core of the line shows that the support of the $2^3P_1$ level by photons is increased with respect to the no-scattering solution, however, not as much as in the Fokker-Planck approximation. This explains why the effective escape probability is decreased by a smaller amount than for the Fokker-Planck case (cf. Fig. 6).

At low redshifts ($z \lesssim 2000$), we find that the kernel approach for the spectral distortion in the cases with and without electron scattering practically coincide in the Doppler core, while in the Fokker-Planck approximation we find a rather large reduction of the line intensity. This does explain the different behaviour in the escape probability (cf. Fig. 6), however, what is the physical reason for this? When using the Fokker-Planck approximation, the distortion again is strongly smeared out in the vicinity of the Doppler core, and photons from within the Doppler core diffuse into the wings. This time also absorption of photons by neutral hydrogen becomes important, so that there are far fewer photons on the red side of the resonance, which in principle could return to the line center by electron scattering. Before they can do so they are likely absorbed in the H i Lyman continuum. Similarly, photons that scatter from the line center to the blue side of the resonance are likely to be absorbed by hydrogen before they return to the core to feedback. Therefore diffusion of photons out of the Doppler core wins and the total intensity is reduced with respect to the case without electron scattering. In the kernel approach, the smoothing of the spectral distortion is basically negligible, and the equilibrium of emission and absorption terms in the core remains largely unaltered by rare electron scattering events.

If we finally also include the effect of the other lines in the computation of the effective escape probability of the $2^3P_i - 1^3S_0$ resonance, we obtain the other three curves shown in Fig. 6. Again we give the result without electron scattering (see Fig. 5) for comparison. In the Fokker-Planck approximation at high redshifts, the total escape probability is slightly smaller than for the case without electron scattering, and at $z \lesssim 2000$, where the $2^3P_i - 1^3S_0$ resonance practically behaves as an isolated line because of H i continuum absorption, we again see a significant increase in the escape probability. The overall behaviour of the escape probability in the Fokker-Planck approximation is in good agreement with our understanding of the problem without the other lines included: at early times electron scattering delays recombination in comparison to the case without electron scattering.

When using the kernel approach, something more surprising happens. Instead of additionally decreasing the escape probability at high redshifts, we can observe a small increase at $z \gtrsim 2500$. Comparing with the previous behaviour for which no additional lines were present, this is not expected. One reason for this is that photons approaching the $2^3P_i - 1^3S_0$ resonance from the blue side occasionally scatter directly to the red side of the Doppler core, essentially jumping over the line without feeding back. In the Fokker-Planck approximation, this process is much less important, since large jumps in frequency are not accounted for. However, the more crucial effect is actually a reduction of the escape probability from the $2^3P_i - 1^3S_0$ singlet line once electron scattering is included (see Sect. 3.3.2); this reduces the spectral distortion on the blue side of the $2^3P_i - 1^3S_0$ resonance (cf. Fig. 5) and hence decreases the
feedback correction. As shown in Fig. 2 at $z \simeq 2900$ the escape probability of the $2^1P_1 - 1^1S_0$ line is reduced by $\simeq 14\%$ when accounting for electron scattering in the kernel approximation. This implies that the feedback correction should also be reduced by a similar amount. We find this being in agreement with the result presented in Fig. 3. To confirm this statement we in addition ran the radiative transfer code but including electron scattering only in the vicinity of the $2^1P_1 - 1^1S_0$ line. In this case the feedback correction was much closer to the case without electron scattering.

3.3.2 Effect of electron scattering on the $2^1P_1 - 1^1S_0$ singlet line

We carried out a similar exercise for the $2^1P_1 - 1^1S_0$ singlet line. The results are shown in Fig. 2. Let us first focus on the curve obtained using the Fokker-Planck approximation (red/dash-dot-dotted line). We can see that with respect to the case without electron scattering the escape probability is increased at all redshifts. At low redshifts, where H I continuum absorption is important, this is expected like for the $2^1P_1 - 1^1S_0$ intercombination line, however, escape from the $2^1P_1 - 1^1S_0$ singlet line is not affected by as much (in the case of the intercombination line the relative difference was $\sim 3 - 4$ times larger). Here it is important to mention that the $2^1P_1 - 1^1S_0$ singlet line itself is very optically thick, so that the spectral distortion is controlled by the line over a much broader range of frequencies (roughly a few hundred Doppler widths as opposed to only a few Doppler widths for the intercombination line). Therefore the effect of electron scattering on the redistribution of photons is limited much more to the line wings.

To understand the increase in the escape probability at high redshift, we ran the isolated line case for the $2^1P_1 - 1^1S_0$ resonance, but including electron scattering using the Fokker-Planck approach. This showed that the main increase in the escape probability is actually due to modifications in the transfer problem of the line itself, rather than being a consequence of changes in the blue wing feedback correction. Again diffusion of photons out of the line center into the wings is helping to reduce the total support of the $2^1P_1$ level, and hence increase the escape rate.

If we now turn to the results obtained within the kernel approach, we can see that at high redshifts photon escape is hindered, while at low redshifts again an increase can be seen, but not as large as for the Fokker-Planck approximation. Again we ran the case with all lines included (solid/black line) and when treating the $2^1P_1 - 1^1S_0$ resonance alone (red/dot-dash-dotted line). Comparing the two shows that changes in the solution for the line itself are causing the largest difference. At high redshifts, the main reason for the difference is that more photons from the red wing are scattered back to the Doppler core of the line and also into the blue wing when encountering a free electron. This delays photon escape. At later times, when hydrogen absorption starts becoming important, the extra broadening introduced by electron scattering mainly removes photons from the Doppler core, reducing the line intensity.

3.3.3 Effect of electron scattering for the upper lines

To close the discussion about the effect of electron scattering on the escape probabilities from the helium resonances, Fig. 10 shows the results of our computations for the $n^1P_1 - 1^1S_0$ intercombination lines with $n = 3, 4$ and 5. At high redshifts, like in the case of the $2^1P_1 - 1^1S_0$ line, we can see that the feedback correction is slightly smaller than in the case without electron scattering. As mentioned
above, this is mainly because the blue wing distortion of the $n^3\text{P}_1 - 1^1\text{S}_0$ lines is reduced by electron scattering, with photons close to the optically thick $n^3\text{P}_1 - 1^1\text{S}_0$ resonances returning to the Doppler core, where photon emission and absorption are effective.

For the quadrupole lines we find a similar effect, while for the higher $n^3\text{P}_1 - 1^1\text{S}_0$ singlet lines, the correction is in the opposite direction, again indicating that modifications to the radiative transfer problem of the line itself dominated. This is in agreement with the result found for the $2^3\text{P}_1 - 1^1\text{S}_0$ line (see Fig. 9).

3.4 Comparison with Monte Carlo computation

To confirm the results obtained with the He I radiative transfer module of CosmoRec, we compare the escape probabilities for several cases with those of a Monte Carlo implementation of the escape problem. The Monte Carlo follows an ensemble of photon trajectories subject to redshift, atomic and electron scattering, line absorption, and photon destruction in the H I continuum. The method used here is described in detail in Switzer & Hirata (2008a) and Switzer & Hirata (2008b), where it is shown that the probability distribution of photons in the Monte Carlo solves the continuous radiative transport equations. The simulation of line scattering draws from the probability distribution for the velocity of the scattering atoms (Lee 1982, 1977), and finds the outgoing photon frequency through kinematics that assume dipole angular redistribution and recoil. Electron scattering is treated identically, except that the electron velocity is normally distributed. True absorption and emission are simulated through the Voigt profile (with no memory of the incoming photon frequency).

Figures 11, 12, 13 compare the radiative transport and Monte Carlo solutions for the escape probability in the $2^3\text{P}_1 - 1^1\text{S}_0$, $2^3\text{P}_1 - 1^1\text{S}_0$, and $3^3\text{D}_2 - 1^1\text{S}_0$ transitions, respectively. The ordinary Fokker-Planck approximation breaks down for electron scattering because that scattering produces jumps in photon frequency which are both large and rare compared to scattering through the line. Because the Monte Carlo fully simulates these scattering processes, it demonstrates that electron scattering is treated accurately in the kernel approach developed here. Agreement in transport through $2^3\text{P}_1 - 1^1\text{S}_0$ also shows that the Fokker-Planck approximation is accurate for coherent atomic scattering. The Monte Carlo is computationally expensive, and cannot currently form the basis for a practical recombination code, but corroborates the fast kernel method in CosmoRec v2.0.

We also computed the effective escape probability for the isolated $3^3\text{P}_1 - 1^1\text{S}_0$ singlet line finding excellent agreement with the Monte Carlo result. This further confirms the validity of the approximations used in the new radiative transfer module of CosmoRec.

3.5 Changes in the ionization history and power spectra

In this section we discuss the effect of different processes on the dynamics of helium recombination. We start by switching off line and electron scattering, directly comparing CosmoRec v2.0 with CosmoRec v1.3. The results for the modifications in the ionization history for $Y_e = 0.24$ are presented in Fig. 14. The improvement over CosmoRec v1.3 is mainly due to consistent inclusion of time-dependent aspects, line-scattering, a precise modelling of H I con-
Radiative transfer effects

Figure 14. Direct comparison of CosmoRec v2.0 with CosmoRec v1.3. The recombination history of CosmoRec v1.3 was used as reference and both line and electron scattering are switched off. The differences are because of time-dependent aspects and additional approximations made in the simplified feedback treatment of CosmoRec v1.3.

Figure 15. Convergence of the line scattering correction with number of included shells \( n \). We internally compare CosmoRec v2.0 with and without resonance scattering enabled. Line scattering leads to a slight deceleration of helium recombination at high redshifts, while at low redshifts it accelerates recombination. We also show the direct comparison of CosmoRec v2.0 with CosmoRec v1.3 (used as reference).

Figure 16. Convergence of the electron scattering correction with number of included shells \( n \). We internally compare CosmoRec v2.0 with and without electron scattering enabled, modelling it with the kernel approximation. Electron scattering leads to a deceleration of helium recombination at high redshifts, while at low redshifts it accelerates recombination.

As we just saw for the standard cosmology, the overall improvement of the helium recombination calculation are not very significant, and smaller or comparable to other corrections (e.g., because of two-photon and Raman processes (Switzer & Hirata 2008) during helium recombination), that are omitted here. The currently used approximations of CosmoRec v1.3 therefore already provide a precise description for the problem. However, when going slightly away from the standard cosmology the differences start increasing...

\( \Delta \left( \frac{N_e}{N} \right) \approx 0.04\% \). Such a difference leads to negligible changes in the CMB temperature and polarization power spectra and even for future CMB experiments can be omitted. However, as we discuss below when varying the helium abundance, larger differences appear.

Finally, in Fig. 16 we show the effect of electron scattering on the helium recombination history. Again the improvement over CosmoRec v1.3 for this case is negligibly small, and convergence with \( n \) is very rapid. Given that one full run with all high frequency helium lines up to \( n = 10 \) included and electron scattering modelled in the kernel approximation takes about \( \approx 5 \) hours on eight CPUs, this is fortunate. Neglecting electron scattering, a helium atom with 10 shells takes about 90 seconds on one CPU. Also, when electron scattering is modelled with the full kernel, but the correction is only included in the vicinity of the \( 2^2P_1 - 1^1S_0 \) and \( 2^2P_1 - 1^3S_0 \) lines, the run finishes in roughly 15 – 30 minutes on eight CPUs, depending on the total number of lines included. For numerical computations this simplification is very useful and we checked that the difference in the ionization history is negligible. The results presented below are calculated using this approximation.

To conclude, for the standard cosmological model, with helium abundance \( Y_p = 0.24 \), the improvement of CosmoRec v2.0 over CosmoRec v1.3 remains smaller than \( |\Delta N_e/N_e| \approx 0.05\% \), and the difference in the CMB power spectra computed with these two versions of the recombination code are expected to be negligible.

3.5.1 Dependence of the correction on the helium abundance

As we just saw for the standard cosmology, the overall improvement of the helium recombination calculation are not very significant, and smaller or comparable to other corrections (e.g., because of two-photon and Raman processes (Switzer & Hirata 2008) during helium recombination), that are omitted here. The currently used approximations of CosmoRec v1.3 therefore already provide a precise description for the problem. However, when going slightly away from the standard cosmology the differences start increasing...
above the level of a few times 0.1%. As mentioned in the introduction, recent constraints on the helium mass fractions from CMB alone derived using WMAP and ACT data (Dunkley et al. 2011) indicate a slightly larger helium abundance of $Y_p = 0.313 \pm 0.044$. Varying the helium mass fraction, we obtain the modification shown in Fig. 17. For all curves except the last electron scattering is neglected. The main reason for the discrepancy is that the precomputed correction function which mimics the effect of line scattering diffusion (Rubiño-Martín et al. 2008) in the escape probability from the $2^1P_1 - 1^3S_0$ line is calibrated at fixed $Y_p = 0.24$. Apparently CosmoRec v1.3 overestimates this correction slightly for larger values of $Y_p$. Electron scattering only affects the recombination dynamics at very early times, but does not have a significant effect on the CMB power spectra.

We computed the modifications for the CMB power spectra and found that for $Y_p = 0.4$ the difference between CosmoRec v1.3 and CosmoRec v2.0 exceeded $|\Delta C_l / C_l| = 0.05\%$ only at $l \gtrsim 3000$. The difference related to a full treatment of electron scattering was negligible. This indicates that the improvements of this work remain small, and even for rather extreme values of $Y_p$, CosmoRec v1.3 can be reliably used to compute the recombination history.

The importance of line scattering alone is shown in Fig. 18, we internally compare the results obtained with CosmoRec v2.0 switching the corresponding terms on and off. HvRsc does not account for the diffusion correction, however, as Fig. 17 shows, when moving further away from the standard cosmology the differences increase to $\Delta N_e/N_e \approx -0.63\%$ at $z \approx 1780$ for $Y_p = 0.4$. Still the modifications in the CMB power spectra turn out to be only marginally important at high $l$ (cf. Fig. 19), since the impact of helium recombination on the power spectra is much smaller than the signals generated close to maximal visibility at $z \approx 1100$.

3.5.2 Direct comparison with Recfast v1.5

Finally we compare the results for the CMB power spectra computed using the helium radiative transfer module of CosmoRec v2.0 with those obtained using Recfast v1.5, which is the current standard in Cmb (Lewis et al. 2000). The differences are shown in Fig. 20, where we again vary the helium abundance. We use the default parameters for the fudge functions of Recfast v1.5. These were calibrated on the recombination calculations of Rubiño-Martín et al. (2010). However, even for $Y_p \approx 0.24$ small differences in the CMB power spectra are visible at high $l$. These increase for larger values of $Y_p$, reaching $\approx 0.4\%$ at $l \approx 1000$ for $Y_p \approx 0.4$. This shows that when a wider range of parameters is allowed, simple fudging does not represent the detailed recombination physics well. In this case a detailed recombination code should be used. Alternatively, an iterative procedure that relies on recalibration of the fudge factors is possible (Shaw & Chluba 2011).

For clarity, we emphasize that the differences seen in Fig. 20 are not mainly because of the improved helium recombination model discussed here, but were already present in comparison with CosmoRec v1.3. Furthermore, we would like to mention that the final code comparison is still not completed, so that the final calibration of the Recfast fudge factors for the current best fit cosmological model cannot be reliably performed at this point.

We also mention that the differences found here with Recfast are responsible for only a very small part of the actual difference seen when going from $Y_p = 0.24$ to $Y_p = 0.313$ (the $TT$ power spectrum amplitude changes by $\approx 10\%$). Therefore the recombination corrections discussed here cannot cause the apparent slight tension of the ACT and SPT results for $Y_p$ with standard Big Bang nucleosynthesis.

3.6 Effect of electron scattering during H1 recombination

For completeness, we also computed the effect of electron scattering during hydrogen recombination. Figure 21 presents the results. We include all two-photon and Raman-processes up to $n = 8$ in the computations. The solid black line shows the correction using the Fokker-Planck approximation, which was already computed earlier (Chluba & Sunyaev 2009a). The red/dash-dotted line shows the result obtained using the Doppler kernel for the redistribution function. This kernel does not include electron recoil and Doppler
boosting, however, as one can see these terms lead to a rather different overall behaviour of the correction (blue/dashed line, which is computed using the Kompaneets kernel that includes those processes in lowest order). In particular electron recoil is important to get the total modification right. Still the effect of electron scattering can be completely neglected for the analysis of future CMB data.

Our result is very similar to the one obtained earlier by Ali-Ha¨ımoud et al. (2010), however, here we include the term due to Doppler boosting in addition, which leads to a small correction. Furthermore, in the work of Ali-Ha¨ımoud et al. (2010) electron recoil is included by enforcing detailed balance rather than using the corresponding terms in the kernel (≈ hv/m_e c^2). We checked that higher order relativistic corrections provided by the formulae of Sazonov & Sunyaev (2000) did not affect the result at any significant level (both for hydrogen and helium recombination).

3.7 Effect of quadrupole lines during H\_1 recombination

The effect of nd-1s quadrupole lines during hydrogen recombination has been considered by Grin & Hirata (2010). Since the transition frequencies of the nd-1s quadrupole lines and np-1s Lyman-series practically coincide, it is clear that photons emitted in a nd-1s quadrupole transition are directly reabsorbed by the optically thick Lyman-\(n\) resonance. Similarly, photons released in a Lyman-\(n\) transition allow re-exciting electrons to the nd-state.

The overall distortion introduced by nd-1s quadrupole transition is much weaker than the one from the Lyman-\(n\) resonance. Therefore the photon distribution around the transition frequencies \(\nu_{nd}\) is approximately given by the distortion introduced by the Lyman-\(n\) resonance. Since the nd-1s quadrupole lines are much less optically thick than the np-1s lines, the mean occupation number of photons in the nd-1s quadrupole lines is given by \(\bar{n} \approx N_{np}/3N_{1s}\), where \(N_{np}\) and \(N_{1s}\) denote the populations of the np and 1s level.

\[\bar{n} \approx N_{np}/3N_{1s}\]

Because the variation of the photon occupation number across the nd-1s
Therefore the net transition rate in the \( nd\)-1s quadrupole line can be approximated as \( \Delta R_{\text{net}} = A_{\text{d1s}}[N_\text{nd} - 5N_{\text{1s}}]\), where \( A_{\text{d1s}} \) is the \( nd\)-1s transition rate, which is typically of order \( 10^2 \text{s}^{-1} \) for small \( n \) (see Table 51).

This is the same approximation that was given by Grin & Hirata (2010) and it effectively results in a direct \( nd\)-np transition. We confirm this approximation by including \( nd\)-1s quadrupole lines into the radiative transfer computation. We find that in the \( nd\)-1s series, the main effect is due to the 3d-1s quadrupole line, with the approximation given above being in excellent agreement with the full numerical result.

In addition, we consider the effect of quadrupole lines among excited levels. The rates can be computed as explained in Appendix B. We compute effective recombination rates (applying the method of Ali-Haımoud & Hirata 2010, to helium) including all quadrupole lines with \( n \leq 300 \) (\( n \leq 100 \) is sufficient for convergence). We find that close to the maximum of the Thomson visibility function the quadrupole lines among excited levels are actually more important than the \( nd\)-1s transitions, however, the overall effect is completely negligible (see Fig. 22).

4 Conclusions

We have developed an improved helium radiative transfer module for CosmoRec, accounting for several subtle processes that previously have been omitted or treated more approximately. This module is now included in CosmoRec v2.0, and can be activated as needed for most settings without affecting the runtime significantly.

For the standard cosmology, the processes described here lead to small corrections in the ionization history at \( z \approx 2000 \), and the associated modifications in the CMB power spectra are negligible (see Sect. 3.5). However, when considering a wider range of cosmological parameters, the new helium recombination module allows more precise computation of the ionization history. For example, for \( Y_p \approx 0.4 \) the modification of the ionization history reaches \( |\Delta N_e/N_0| \approx 0.22\% \) at \( z \approx 1800 \), however, the overall correction to the CMB power spectra is small, remaining below \( |\Delta C_l/C_l| \approx 0.05\% \) at \( l \leq 3000 \). This means that the improvements of CosmoRec v2.0 with respect to CosmoRec v1.3 are negligible. We furthermore showed that the effect of line scattering during helium recombination is important only for large \( Y_p \) (cf. Sect. 3.5).

We also compare the results obtained with CosmoRec v2.0 to those computed using RecFast v1.5 and find that the parameters for the fudge functions probably should be re-calibrated once the final recombination code comparison has been completed (see Sect. 3.5).

Finally, we considered the effect of quadrupole lines during hydrogen recombination (Sect. 3.7) and found that these can be safely neglected in computations of the CMB power spectra.

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Appendix A: Recursions for Reduced Dipole Matrix Elements

The dipole transition rates (selection rule \( \Delta l = \pm 1 \)) for hydrogenic atoms are given by (e.g., see Storey & Hummer 1991)

\[
A^{(\pm)}_{\ell\ell'} = |Z\mu_{\text{el}}c\alpha^2| \frac{1}{n^2} - \frac{1}{n'^2} \max(|\ell,\ell'|) \left(\frac{1}{2\ell + 1}\right)^2 |X^{(\pm)}_{\ell\ell'}|^2 .
\] (A1)

Here \( Z \) is the atomic charge; \( \mu_{\text{el}} = [1 + m_b/M_\Lambda]^{-1} \) is the reduced mass of the atom with mass \( M_\Lambda \) divided by \( m_b \); \( \alpha \) is the fine structure constant and \( c \) the speed of light; \( \ell_0 \) denotes the Bohr radius (with out reduced mass effect); and \( X^{(\pm)}_{\ell\ell'} \) is the reduced matrix element of the transition between the levels \((n,\ell) \rightarrow (n',\ell')\).
The electric quadrupole transition rates (selection rule $\Delta l = 0$) with the initial condition $X_{n_{1}l_{1}=1}^{2(2)} = \frac{C_{n_{1}}^{2}X_{n_{1}l_{1}=1}^{2(2)}}{2(l + 1)C_{l_{1}l_{1}=1}}$, where we defined $C_{l_{1}l_{1}=1} = \sqrt{\frac{\hbar}{\lambda}}$. These recursions can be started at $l$ with initial conditions (for $n > n' \geq 1$)

$X_{n'n'=1}^{2(2)} = 2^{l+1} \left[ \frac{(n' n')^{2l+1} + (n + n')!}{(n - n' - 1)! (2n' - 1)!} \right]^{1/2} \left[ \frac{n - n'}{n + n'} \right]^{n'n'=2-2} X_{n'n'=2}^{2(2)}$ \quad (A4)

and $(1)X_{n'n'=0}^{2(2)} = 0$ for $l' \geq n'$.

### APPENDIX B: RECURSIONS FOR REDUCED QUADRUPOLE MATRIX ELEMENTS

The electric quadrupole transition rates (selection rule $\Delta l = 0$, $\pm 2$) for hydrogenic atoms are given by [Hey 2006; Grin & Hirata 2010]

$$A_{n'l'=n''l'=l}^{2(2)} = \frac{Z^{2} \mu_{\text{elect}}^{2} \varepsilon^{2}}{2l + 1} \frac{1}{2} \left[ \begin{array}{c} 0 \\ 1 \\ 0 \end{array} \right] \left( \begin{array}{ccc} 0 & 2 & l' \\ 0 & 0 & 0 \end{array} \right) \frac{C_{n_{1}}^{2}X_{n_{1}l_{1}=1}^{2(2)}}{2(l + 1)C_{l_{1}l_{1}=1}} X_{n'n'=2}^{2(2)} \right|^{2}, \quad (B1)$$

where $(2)X_{n'l'}^{2(2)}$ is the reduced quadrupole matrix element.

#### B1 Case $\Delta l = 0$

According to [Hey 2006] the matrix elements $(2)X_{n'l'}^{2}$ are determined by the recursion formula: \[ (2)X_{n'l'}^{2} = \frac{C_{n_{1}}^{2}X_{n_{1}l_{1}=1}^{2(2)} + 2(l + 1)X_{n_{1}l_{1}=1}^{2(2)}}{C_{l_{1}l_{1}=1}^{2(2)}}, \quad (B2) \]

with the initial condition $(2)X_{n'n'=0}^{2} = 0$, assuming that the dipole matrix elements are known. Electric quadrupole transitions between s-states are forbidden.

#### B2 Cases $\Delta l = \pm 2$

Following [Hey 2006] the matrix elements $(2)X_{n'l'=n''l'=l}^{2(2)}$ and $(2)X_{n'l'=n''l'=l}^{2(2)}$ are determined by the recursion formulae:

$$X_{n''l'=n'l'=n''l'=l}^{2(2)} = \frac{(2l - 1)C_{l_{1}}^{2}X_{n_{1}l_{1}=1}^{2(2)} + 2C_{l_{1}}^{2}X_{n_{1}l_{1}=1}^{2(2)}}{(2l + 1)C_{l_{1}l_{1}=1}^{2(2)} + 2(l + 1)C_{l_{1}l_{1}=1}^{2(2)}}, \quad (B3)$$

$$X_{n''l'=n'l'=n''l'=l}^{2(2)} = \frac{(2l + 3)C_{l_{1}}^{2}X_{n_{1}l_{1}=1}^{2(2)} + 2C_{l_{1}}^{2}X_{n_{1}l_{1}=1}^{2(2)}}{(2l + 5)C_{l_{1}l_{1}=1}^{2(2)} + 2(l + 3)C_{l_{1}l_{1}=1}^{2(2)}}, \quad (B4)$$

with the initial condition $(2)X_{n'n'=1}^{2(2)} = -2n' \left( \frac{C_{n_{1}}^{2}X_{n_{1}l_{1}=1}^{2(2)}}{C_{l_{1}l_{1}=1}^{2(2)}} \right) X_{n'n'=1}^{2(2)}$, assuming that the dipole and diagonal quadrupole matrix elements are known. We note that in comparison to [Hey 2006] we multiplied all matrix elements by the phase factor $(-1)^{n''-1}$. The $\Delta l = 0$ quadrupole matrix elements are in phase with this, however, for the off-diagonal elements a factor of $(-1)$ remains.

In Table B1 we give the electric quadrupole transition rates of hydrogen for $n \leq 5$ and $n' < n$. Transitions between states with $l = 0$ and $l' = 0$ are not allowed. The typical fractional contribution of quadrupole transitions to the total decay rate of the initial levels is about $10^{-7} - 10^{-5}$ for excited states with $n \gg 1$ and $l \leq 30$ (see Fig. A4). The lifetimes of d-states are affected by $\approx 10^{-5}$, while s- and p-states are only modified at the $\approx \times 10^{-7}$. Beyond $l \approx 30$ the effect of quadrupole lines on the total decay rate drops below $10^{-7}$. For highly excited levels at large $l$ and $l' \ll n$ the relative contribution to the total lifetime falls off like $l^{-2}$.

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