1. Introduction

The epoch of reionization (EoR) started with the formation of high-redshift quasars (QSO) (e.g., Fan et al. 2006), the universe was fully reionized by $z \sim 6$. WMAP 5-year results show that the optical depth for the Thomson scattering of CMB photons has an amplitude greater than 100 mK during the early reionization, up to 10 times greater than the typical amplitude of a few 10 mK obtained during the later reionization phase. More importantly, we find that even a rather high contribution from QSO-like sources only damps the absorption regime without erasing it. Heating the IGM with X-ray takes time. Our results show that observations of the early reionization will probably benefit from a higher signal-to-noise value than during later stages. After analyzing the statistical properties of the signal (power spectrum and PDF) we find three diagnostics to constrain the level of X-ray, hence the nature of the first sources.

Key words. dark ages, reionization, first stars – radiative transfer – HII regions – quasars: general – intergalactic medium – early Universe

We present simulations of the 21-cm signal during the epoch of reionization. We focus on properly modeling the absorption regime in the presence of inhomogeneous Wouthuysen-Field effect and X-ray heating. We ran radiative transfer simulations for three bands in the source spectrum (Lyman, UV, and X-ray) to fully account for these processes. We find that the brightness temperature fluctuation of the 21 cm signal has an amplitude greater than 100 mK during the early reionization, up to 10 times greater than the typical amplitude of a few 10 mK obtained during the later emission phase. More importantly, we find that even a rather high contribution from QSO-like sources only damps the absorption regime without erasing it. Heating the IGM with X-ray takes time. Our results show that observations of the early reionization will probably benefit from a higher signal-to-noise value than during later stages. After analyzing the statistical properties of the signal (power spectrum and PDF) we find three diagnostics to constrain the level of X-ray, hence the nature of the first sources.

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or higher. Indeed, simulating the absorption phase correctly, as we do in this work, requires a more extensive and more costly implementation of radiative transfer. We are exploring the direct implication of this improved physical modeling, and will turn to better mass resolution in the near future.

There are three bands in the sources SED that influence the level of the 21 cm signal: the Lyman band, the ionizing UV band, and the soft X-ray band. Lyman band photons are necessary to decouple the spin temperature of hydrogen from the CMB temperature through the Wouthuysen-Field effect (Wouthuysen 1952; Field 1958), and make the EoR signal visible. UV band photons are of course responsible for the ionization of the IGM, and soft X-rays are able to preheat the neutral gas ahead of the ionizing front, deciding whether the decoupled spin temperature is less (weak preheating) or greater (strong preheating) than the CMB temperature. While a proper modeling should perform the full 3D radiative transfer in all 3 bands, a simpler modeling has often been used in previous works. Indeed, for the usual source SEDs and source formation histories, once the average ionization fraction of the universe is greater than ∼10%, the background flux of Lyman-α photons is so high that the hydrogen spin temperature is fully coupled to the kinetic temperature by the Wouthuysen-Field effect (Baek et al. 2009). Thereafter, computing the Lyman band radiative transfer is unnecessary. In the same spirit, it has usually been assumed that the preheating of the IGM by soft X-ray was strong enough to raise the kinetic temperature much higher than the CMB temperature everywhere early in the EoR. However, both assumptions fail during the early reionization: the absorption phase. Even in the later part of reionization the second assumption may fail, depending on the nature of the sources. We will quantify this possibility in this paper. Computing the full radiative transfer in all three bands is necessary to study the absorption regime. Indeed, fluctuations in the local Lyman-α flux induce fluctuations in the spin temperature (while the Wouthuysen-Field effect is not yet saturated), which, in turn, modify the power spectrum of the 21 cm signal (Barkana & Loeb 2005; Semelin et al. 2007; Chuzhoy & Zheng 2007; Naoz & Barkana 2008; Baek et al. 2009). The same is true for the fluctuations in the local flux of X-ray photons (Pritchard & Furlanetto 2007; Santos et al. 2008).

Let us emphasize however that, in modeling Lyman-α and X-ray fluctuations, Barkana & Loeb (2005), Naoz & Barkana (2008), Pritchard & Furlanetto (2007) and Santos et al. (2008) all use the semi-analytical approximation that the IGM has a uniform density of absorbers and ignore wing effects in the radiative transfer of Lyman-α photons. Semelin et al. (2007) and Chuzhoy & Zheng (2007) have shown that these wing effects do exist. Moreover, once reionization is under way, ionized bubbles create sharp fluctuations in the number density of absorbers (not to mention simple matter density fluctuations). In this work, for the first time, we present results based on simulations with full radiative transfer for both Lyman-α and X-ray photons.

What are the possible candidates as sources of reionization? Usually, two categories are considered: ionizing UV sources (Pop II and III stars), and X-ray sources (quasars). When we study 21 cm absorption, however, we must distinguish between Pop II and Pop III stars beyond the large difference in luminosity per unit mass of formed star. Indeed Pop II stars have a three times larger Lyman band to ionizing UV band luminosity ratio than Pop III stars. It means that the 21 cm signal will reach its full power (near saturated Wouthuysen-Field effect) at a lower average ionization fraction for Pop II stars than for Pop III stars. The relevant question is: how long do Pop III stars dominate the source population before Pop II stars take over? The answer to this question, related to the whole process of star formation, feedback and metal enrichment of the IGM, is a difficult one. At this stage, state of the art numerical simulations of the EoR use simple prescriptions in the best case (e.g. Iliev et al. 2007), or simply ignore this issue.

The other category of sources are X-ray sources. They may be mini-quasars, X-ray binaries, supernovae (Oh 2001; Glover & Brand 2003), or even more exotic candidates such as dark stars (Schleicher et al. 2009). The exact level of emission from these sources is a matter of speculation. The generally accepted view is that stars dominate over X-ray sources and are sufficient to drive reionization ( Shapiro & Giroux 1987; Giroux & Shapiro 1996; Madau et al. 1999; Ciardi et al. 2003). Recently, Volonteri & Gnedin (2009) supported the opposite view. While, in their models, X-ray sources are marginally able to complete reionization by z ∼ 6, they find a very low contribution from stars. Indeed they rely on Gnedin et al. (2008) who find, using numerical simulations, a negligible escape fraction for ionizing radiations from galaxies with total mass less than a few $10^{10} M_{\odot}$, who should actually contribute to 90% of the ionizing photon production during the EoR (Choudhury & Ferrara 2007). While the physical modeling in their innovative simulations is quite detailed, this surprising behavior of the escape fraction definitely needs to be checked at higher resolution and with different codes. For the time being the best simulations can only explore a plausible range of X-ray contributions, and quantify the impact on observables. When the observations become available we would like to be able, using simulation results, to derive tight constraints on the relative level of emission from ionizing UV and X-ray sources. This work, exploring the 21 cm signal for a few different levels of X-ray emission, is a first step toward this goal.

The paper is organized as follows. We present the numerical methods in Sect. 2 and describe our source models in Sect. 3. In Sect. 4, we show the results and analyze the differences between the models. We discuss our findings and conclude in Sect. 5.

2. Numerical simulation

The numerical methods used in this work are similar to those presented in Baek et al. (2009, hereafter Paper I). The references to previous and some new validation tests are presented in the Appendix. The dynamical simulations have been run with GADGET-2 (Springel 2005) and post-processed with UV continuum radiative transfer and further processed with Ly-α transfer using LICORICE. The same cosmological parameters and particle number are used and we refer the reader to Paper I for details related to the numerical methods and parameters. The main improvement on the previous work is using a more realistic source model including soft X-ray and implementing He chemistry.

We have run seven different simulations, all of which use the same $100 h^{-1}$ Mpc box, density fields, and star formation rate, but with different initial mass functions (IMF), chemistry (with helium or without), X-ray fraction of the total luminosity or X-ray spectral index. S1 is the reference model. S2 has a top-heavy IMF (Salpeter IMF restricted to a 100–120 $M_{\odot}$ range), while the others uses a Salpeter IMF in a 1.6–120 $M_{\odot}$ range. Only S3 contains helium. In all other models, helium is replaced by the same mass of hydrogen. X-ray radiative transfer is included in S4, S5, S6 and S7. They have either different X-ray fraction of the total luminosity or X-ray spectral index. The basic parameters of these simulations are summarized in Table 1.

The simulations are controlled by a few parameters. We adopted the same value as in Paper I for the maximum value
of the number of particles per radiative transfer cell in the adaptive grid: \( N_{\text{max}} = 30 \). The resulting minimum radiative transfer cell size is 200 h\(^{-1}\) kpc at \( z = 6.6 \). Between two snapshots, i.e. \(~10\) Myr, we cast 3 \( \times 10^6 \) photon packets for photoionization (all in the UV for models S1 to S3, half in the UV and half X-rays for models S4 to S7), and 3 \( \times 10^5 \) photons for Lyman-\( \alpha \) transfer. At the end of the simulations (\( z \approx 6 \)), the number of sources reaches \(~15\) 000, so the number of ionizing photon packets per source is only 200. However, at this final stage the sources are highly clustered and very large and ionized regions surround the source clusters. So the clustered sources cooperate to reduce the Monte Carlo noise at the ionization fronts. In addition, the adaptive grid responds better than a fixed grid to sampling issues: big cells where there are few photons, small cells where there are many. Maselli & Ferrara (2005) presents convergence tests for a Monte-Carlo radiative transfer code very similar to ours. Their convergence tests suggest that the typical convergence tests for a Monte-Carlo radiative transfer code are to reduce the Monte Carlo noise at the ionization fronts. In addition, the adaptive grid responds better than a fixed grid to sampling issues: big cells where there are few photons, small cells where there are many. Maselli & Ferrara (2005) presents convergence tests for a Monte-Carlo radiative transfer code very similar to ours. Their convergence tests suggest that the typical level of noise in our ionization and temperature cubes is \(~10\)%.

We accept as a reasonable value, especially since, having run the Iliev et al. (2009) comparison tests, we are confident that our ionization fronts propagate at the correct speed. We use 1000 frequency bins in each of the photoionizing-UV and X-ray spectra. For Lyman-\( \alpha \) transfer, we sample the frequency at random between Lyman-\( \alpha \) and Lyman-\( \beta \).

2.1. X-ray radiative transfer

The main difference between the cosmological radiative transfer of ionizing UV and X-ray is the mean free path of the photons, at most a few 10 comoving Mpc in the first case, possibly several 100 Mpc in the second case. A usual trick in implementing UV transfer is to use an infinite speed of light: do so with LICORICE (see Paper I). This is correct if the crossing time of the photons between emission and absorption points is much less than the recombination time, the photo-ionization time (Abel et al. 1999) and the typical time for the variation of luminosity of the sources. This is the case in most of the IGM during the EoR, except very close to the sources where the photo-ionizing rate is very high. Obviously, this is not the case for X-rays which have a much longer crossing time. Consequently, we implemented the correct propagation speed for X-ray photons. We propagate an X-ray photon packet during one radiative transfer time step \( \Delta t_{\text{reg}} \) (<1 Myr, same notation from Fig. 1 of Paper I) over a distance of \( c \Delta t_{\text{reg}} \), where \( c \) is the velocity of light. Then, the frequency of the photon packet and photoionizing cross sections are recomputed with the updated value of the cosmological expansion factor. The photon packet propagates during the next radiative transfer time step using these new parameters. If the photon packet loses 99% of its initial energy, we drop it. X-ray photon packets containing photons with an energy of several keV pass through the periodic simulation box several times before they lose most of their initial content. For each density snapshot, that is every \~10\) Myr, 1.5 millions of photon packets are sent from the X-ray sources. About half of them are absorbed during the computation on the same density snapshot when they were emitted, and the other half is stored in memory to be propagated through the next density snapshots. Some X-ray packets with very high energy photons still survive several snapshots later, so the number of stored photon packets grows as simulations progress. About 50 millions photon packets are stored in memory toward the end of the simulations.

It may seem that this memory overhead, which sets a limitation to the possible simulations with LICORICE, would not appear with radiative transfer algorithms which naturally include a finite velocity of light like moment methods. However, these methods would suffer from an overhead connected to the number of frequency bins necessary to correctly model X-rays, while it does not exist in Monte-Carlo methods. Including complete X-ray transfer in EoR simulations comes at a non-negligible cost, whatever the numerical implementation. Since the X-ray photons can propagate over several box sizes during several tens of Myr, the X-ray frequency can redshift considerably between emission and absorption. The cross-section of photoionization has a strong frequency dependence, so we have to redshift the frequency of the photons. At each radiative transfer time step \( \Delta t_{\text{reg}} \), we update the frequency of all the X-ray photon packets,

\[
\nu(t + \Delta t_{\text{reg}}) = \frac{a(t)}{a(t + \Delta t_{\text{reg}})} \nu(t),
\]

where \( a(t) \) is the expansion factor of the Universe.

The treatment of non-thermal electrons produced by X-ray will be described in Sect. 3.5.

2.2. Helium reionization

The intergalactic medium is mainly composed of hydrogen and helium, with contributions of 90% and 10% in number. Until now, we have run simulations with hydrogen only, but including helium is worth studying because the different value of the ionization thresholds and photoionization rates could affect the reionization history. We included He, He\(^+\), and He\(^{++}\) in LICORICE, and used Cen (1992) and Vermer et al. (1996) for various cooling rate and cross sections. When helium chemistry is turned on, the ionization fractions (H\(^+\), He\(^+\) and He\(^{++}\)) and the temperature are integrated explicitly using the adaptive scheme described in Paper I. More details on the numerical methods and a validation tests of the treatment of helium are presented in Appendix.

3. Source model

3.1. Computing the star formation rate

Our new source model needs the star formation rate for all baryon particles. We recompute the star formation in the radiative transfer simulations rather than to rerun the dynamical simulation. Here is why and how.

We adopted the procedure described in Mihos & Hernquist (1994), employing a local Schmidt law and an hybrid-particles algorithm to implement it in our code. Indeed, in our model, the star formation rate solely depends on the density, and we make the assumption that the star formation feedback (kinetic

Table 1. Simulation parameters.

| Model | IMF | Helium | \( L_{\text{star}} \) | \( L_{\text{QSO}} \) |
|-------|-----|--------|----------------|----------------|
| S1    | 1.6–120 M\(_\odot\) | No | 100% | 0% |
| S2    | 100–120 M\(_\odot\) | No | 100% | 0% |
| S3    | 1.6–120 M\(_\odot\) | Yes | 100% | 0% |
| S4    | 1.6–120 M\(_\odot\) | No | 99.9% | 0.1% |
| S5    | 1.6–120 M\(_\odot\) | No | 99.9% | 0.1% |
| S6    | 1.6–120 M\(_\odot\) | No | 99% | 1% |
| S7    | 1.6–120 M\(_\odot\) | No | 90% | 10% |

Notes. \( L_{\text{star}} \) is the stellar luminosity fraction and \( L_{\text{QSO}} \) is the X-ray luminosity fraction of the total luminosity.
and thermal) on the dynamics does not vary much from the fiducial simulation. LICORICE uses the classical Schmidt law:

$$\frac{df_*}{dt} = \frac{1}{t_*} \quad (\text{if} \quad \rho g > \rho_{\text{threshold}}), \quad (2)$$

where $t_*$ is defined by:

$$t_* = t_0 \left( \frac{\rho g}{\rho_{\text{threshold}}} \right)^{-1/2} \quad . \quad (3)$$

$\rho g$ is the gas density and $f_*$ is the star fraction. We set the parameters $t_0$ and $\rho_{\text{threshold}}$ so that the evolution of the global star fraction follows closely that of the S20 simulation ($20 \, h^{-1} \, \text{Mpc}$) in Paper I, and reionization completes at $z \sim 6$. In this way, we reuse the tuning made for the S20 simulation, and at high $z$, we get a similar star formation history as in the higher resolution (but smaller box size) S20 simulation. All simulations in the present work have a $100 \, h^{-1} \, \text{Mpc}$ box size. Following the above equations, a gas particle whose local density exceeds the threshold ($\rho_{\text{threshold}} = 5 \rho_{\text{critical}} \times \Omega_b$) increases its star fraction, $f_*$, where $\rho_{\text{critical}}$ is the critical density of the universe and $\Omega_b$ is the cosmological baryon density parameter.

### 3.2. Limiting the number of sources

We compute the increase in the star fraction for each particle, $\Delta f_*$, between two consecutive snapshots. Then the total mass of young stars formed in a particle is $m \times \Delta f_*$, where $m$ is the mass of the particle. To avoid a huge number of sources, we had to set a threshold on the new star fraction for the particle to act as a source. We used $\Delta f_* > 0.001$. We checked that this leaves out a negligible amount of star formation, about $0.4\%$. It happens that several source particles reside in the same radiative transfer cell, but we treated individually ray tracing for each source.

### 3.3. Choosing an IMF

With our mass resolution, this amount $m_{\text{gas}} \times \Delta f_*$ corresponds to a star cluster or a dwarf galaxy so an IMF should be taken into account. We choose a Salpeter IMF, with masses in the range $1.6-120 \, M_\odot$ or $100-120 \, M_\odot$ (model S2). The first range is used to model the SED of an intermediate Pop II and Pop III star population, and the other one is for pure Pop III stars.

### 3.4. Computing the luminosity and SED of the stellar sources

The next step is to make the link between the amount of created stars and the luminosity of the sources. Only the ionizing UV luminosity is considered, it is quite justified to use simple models. For example we can make it proportional to the mass of the host dark matter halo (Iliev et al. 2006b), or, as we did in Paper I, to the mass of the baryonic particles newly converted into stars. Things are more complicated when we consider both the Lyman band and ionizing UV. Indeed, since each particle is massive enough to contain a representative sample of the chosen IMF, and since each mass bin has a different life time, we should consider an SED evolving with the age of the star particle. This would be possible using pre-tabulated SEDs. However, unlike in Paper I, we decided to use hybrid particles which begin to produce photons as soon as a small fraction of the particle is turned into stars. This is useful to make the local luminosity less noisy in the early EoR when the source mass resolution is an issue. Including this star formation history for each particle and convolving with the time-varying SED would be extremely costly in terms of both memory and computation time.

We simplified the issue by considering the fact that in the Lyman and UV bands, most of the luminosity is produced by the massive stars, with a short life time comparable to the time between two snapshots of our simulations. So we decided to use a constant SED and luminosity during a characteristic life time. Both luminosity and SED are computed independently in each of the Lyman and ionizing band. To compute the luminosity and the SED for a star particle we use the data for massive, low metallicity ($Z = 0.004 \, Z_\odot$) stars in the main sequence (Meynet & Maeder 2005; Hansen & Kawaler 1994, see Table 2). The details of how this is done can be found in Appendix. The constant luminosity and characteristic life time, computed in the two spectral bands, are given in Table 3. We find characteristic life time of $< 8 \, \text{Myr}$ for the UV band. In the implementation of the UV transfer however, for technical reasons, the source fraction of the particle actually shines for a duration equal the interval between two snapshots. This varies between 6 and 20 Myr, so we recalibrate the luminosity to produce the correct amount of energy. The whole point of the procedure, is to take the different typical life time in the Lyman band into account, especially at $z$, when Lyman-\(\alpha\) coupling is not yet saturated. We should not concentrate the emission within a single snapshot interval, which is 3 times shorter than the source life-time, or we would artificially boost the coupling between the spin temperature of hydrogen and the kinetic temperature of the gas and alter the resulting brightness temperature. Consequently we let each newly formed star fraction of a particle shine for 3 consecutive snapshots, which is close to the typical life time in the Lyman band, and we still recalibrate the luminosity to produce the correct amount of energy. While we do not use a time-evolving SED, we believe that implementing different life times for the Lyman and UV sources with the correctly average luminosities is a substantial improvement in our source model.

We use an escape fraction $f_{\text{esc}} = 0.12$ for photoionizing UV photons and $f_{\text{esc}} = 1$ for Lyman-\(\alpha\).

### 3.5. X-ray source model

X-rays can have a significant effect on the 21 cm brightness temperature. The X-ray photons, having a smaller ionizing cross-section, can penetrate neutral hydrogen further than UV photons and heat the gas above the CMB temperature. This X-ray heating effect on the IGM is often assumed to be homogeneous because of X-rays’ long mean free path. In reality, the X-ray flux is

| Mass [$M_\odot$] | log($L/L_{\odot}$) | log($T_{\text{eff}}$) | $t_{\text{life}}$[Myr] |
|----------------|--------------------|----------------------|------------------------|
| 120            | 6.3                | 4.7                  | 3                      |
| 60             | 5.8                | 4.6                  | 4.5                    |
| 40             | 5.6                | 4.5                  | 6                      |
| 30             | 5.2                | 4.5                  | 7                      |
| 20             | 4.8                | 4.45                 | 10                     |
| 15             | 4.65               | 4.4                  | 14                     |
| 12             | 4.2                | 4.37                 | 20                     |
| 9              | 3.8                | 4.3                  | 34                     |
| 5.9            | 2.92               | 4.18                 | 120                    |
| 2.9            | 1.73               | 3.97                 | 700                    |
| 1.6            | 0.81               | 3.85                 | 3000                   |

Notes: $L$ is the bolometric luminosity, ($T_{\text{eff}}$) is the effective temperature and $t_{\text{life}}$[Myr] is the life time of the star.
stronger around the sources and the inhomogeneous X-ray flux can bring on extra fluctuations for the 21 cm brightness temperature (Pritchard & Furlanetto 2007; Santos et al. 2008). Moreover, patchy reionization induce further fluctuations in the local X-ray flux which can only be accounted for using a full radiative transfer modeling.

X-rays luminosity

First, we need to determine the luminosity and location of X-rays sources. We simply divided the total luminosity, \( L_{\text{tot}} \), of all source particles into a stellar contribution, \( L_{\text{star}} \), and a QSO contribution, \( L_{\text{QSO}} \). \( L_{\text{QSO}} \) depends on the star formation rate, since \( L_{\text{tot}} \) itself is proportional to the increment of the star fraction, \( \Delta f_{\star} \), between two snapshots of the dynamic simulation.

One reason for this approach is that X-ray binaries and supernova remnants contribute to X-ray sources as well as quasars and they are strongly related to the star formation rate. Following the work of Glover & Brand (2003), we took 0.1% of \( L_{\text{tot}} \) as the fiducial X-ray source luminosity, \( L_{\text{QSO}} \). However, considering that they assumed a simple and empirically motivated model we have run simulations with different values \( L_{\text{QSO}} \), 1% and 10% of \( L_{\text{tot}} \). Quasar luminosity fractions less than 0.1% are not of interest for us, since their heating effect will be negligible.

X-ray energy range and nature of the sources

First, we have to choose the photon energy range since hard X-ray photons have a huge mean free path which costs a lot for ray-tracing computations. The comoving mean free path of an X-ray photons have a huge mean free path which costs a lot for First, we have to choose the photon energy range since hard sources. We simply divided the total luminosity, \( L_{\text{tot}} \), of all source particles into a stellar contribution, \( L_{\text{star}} \), and a QSO contribution, \( L_{\text{QSO}} \). \( L_{\text{QSO}} \) depends on the star formation rate, since \( L_{\text{tot}} \) itself is proportional to the increment of the star fraction, \( \Delta f_{\star} \), between two snapshots of the dynamic simulation.

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QSO spectral index

We model the specific luminosity of our QSO-like sources as a power-law with index \( \alpha \);

\[
L_{\nu} = k \left( \frac{\nu}{\nu_0} \right)^{-\alpha}.
\]

\( k \) is a normalization constant so that

\[
L_{\text{QSO}} = \int_{v_1}^{v_2} L_{\nu} \, dv.
\]

where \( h\nu_1 = 0.1 \, \text{keV} \) and \( h\nu_2 = 2 \, \text{keV} \). The amount of X-ray heating can be altered by the shape of the spectrum but there exists a large observational uncertainty in the mean and distribution of \( \alpha \). We extrapolate from the measurement of extreme UV spectral index by Telfer et al. (2002) to higher energy. The index values are derived from fitting \( 1R_g < E < 4R_g \) and we extrapolated to 2 keV. The measured value by Telfer et al. (2002) is approximately \( \approx 1.6 \), but with a large gaussian standard deviation of 0.86. Scott et al. (2004) derived an average value of \( \alpha = 0.6 \) from a sample of FUSE and HST quasars. We choose \( \alpha = 1.6 \) as our fiducial case, and used \( \alpha = 0.6 \) for comparison.

Secondary ionization

X-rays deposit energy in the IGM by photoionization through three channels. The primary high velocity electron torn from hydrogen and helium atoms distributes its energy by 1) collisional

### Table 3. Averaged luminosities and life times of our source model for a baryon particle depending on the energy band.

| IMF mass range | Energy band | \( L_{\text{tot}} \) | Life time, [Myr] |
|---------------|-------------|--------------------|-----------------|
| 1–120 [M\(_\odot\)] | Luminosity [erg/s] \(^A\) | 6.32 \times 10\(^{41}\) | 20.36 |
| 100–120 [M\(_\odot\)] | Luminosity [erg/s] \(^B\) | 9.96 \times 10\(^{41}\) | 3.32 |

**Notes.** A values are from Salpeter IMF and \( B \) values are from top-heavy IMF.

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ionization, producing secondary electrons, 2) collisional excitation of H and He and 3) Coulomb collision with thermal electrons. The fitting formula in Shull & van Steenberg (1985) is used to compute the fraction of secondary ionization and heating. These are taken into account when computing the evolution of the state of the IGM.

Then, it is legitimate to ask whether the lyman-α electrons resulting from the collisional excitations are important for the Wouthuysen-Field effect. The simple answer is that, in our choice of models, the energy emitted as X-ray is at most 10% of the UV energy, itself 3 times less than the Lyman luminosity. Moreover at most 40% of the X-ray energy is converted into excitations (Shull & van Steenberg 1985), and only ∼30% of the excitations result in a Lyman-alpha photon (Pritchard & Furlanetto 2006). So, in the best case, Lyman-α photons produced by X-ray represent only 0.3% of the photons produced directly by the sources.

4. Results

4.1. Ionization fraction

The evolution of the averaged ionization fraction tells us about the global history of reionization. We plot the mass and volume weighted average ionization fraction in Fig. 2. Including quasar (S4, S5, S6 and S7, not plotted) does not change the global evolution of the ionization fraction much, because of its small fraction of the total luminosity. The total number of emitted photons is similar to S1. The S3 simulation has also the same number of emitted photons, but S3 reaches the end of reionization a little bit earlier (Δz ≈ 0.25) than S1. Unlike other simulations, S3 contains helium which occupies 25% of the IGM in mass. Including helium, the total number of atoms is reduced by 20%. At the same time, unless X-ray are the dominant source of reionization, most of the helium is only ionized once while z > 6, due to the higher energy threshold for the secondary ionization(He++). Therefore the number of emitted photons per baryon is higher in S3 than S1, and it results in an earlier reionization.

On the other hand, S2 has the same number of photon absorbers as S1, but the total number of emitted photons is much higher than in S1. Using a top-heavy IMF, it produces 10 times more photons (see Fig. 1 and Table 3), and results in a Δz ≈ 1 earlier reionization.

In all three cases, volume weighted values are less than mass weighted values, since gas particles in dense regions around the sources are ionized first. The volume occupied by each particle is estimated using the SPH smoothing length.

We computed the Thomson optical depth for all simulations, the values are τ = 0.062, 0.076, 0.064 for S1, S2 and S3. The other simulations (S4-S5) have the same τ as S1, since they follow the same evolution of ionization fraction. They are somewhat lower than the Thompson optical depth derived from WMAP5 (Hinshaw et al. 2009), τ = 0.084 ± 0.016, only the S2 value is within 1σ of the WMAP5 value. A variable escape fraction, decreasing with time, would allow the IGM to start ionization earlier and increase τ, without terminating ionization after z = 6.

4.2. Gas temperature

The main goal of this study is to investigate the effect of inhomogeneous X-ray heating on the 21-cm signal. If the Ly-α coupling is sufficient, and X-rays can heat the gas above the CMB temperature TCMB, the 21-cm signal will be observed in emission.

However if the X-ray heating is not very effective, particularly during the early phase of the EoR, we will observe the signal in absorption.

We plot in Fig. 3 the averaged gas temperature of the neutral IGM with redshift. The neutral gas is chosen so that its ionization fraction is less than 1%. We chose the criterion of xHII < 0.01 to evaluate properly the average temperature of neutral regions, and verified that xHII < 0.001 gives a very similar average temperature. We have checked that even for model S7 which has the highest level of X-rays, at z > 7.5, 99% of the neutral IGM has indeed an ionization fraction less than 1%, so we have not excluded a significant fraction of the 21 cm emitting IGM from our average. This neutral gas is mostly located in the voids of the IGM. In
fact, we have to consider Ly-α heating as well as X-ray heating since a few K difference can reduce the intensity of $\delta T_b$ by up to 100 mK. We recompute the gas temperature to include Ly-α heating as a post-treatment using the formula from Furlanetto & Pritchard (2006). This was detailed in Paper I. The temperature of all simulations in Fig. 3 decreases until $z = 12$ because of the adiabatic expansion of the universe. Then S7, which has the highest $L_{QSO}$, starts to increase first and reaches the CMB temperature at redshift $z \approx 8.8$. Our fiducial model, S4, which contains 0.1% of total energy as X-rays, shows very little increase with respect to S1, a simulation without X-rays. Even for S7, the gas temperature of neutral hydrogen in the void is still under the $T_{CMB}$ until $z \approx 8.8$. This means that the X-ray heating needs time to heat the IGM above $T_{CMB}$. Even with a rather high level of X-ray, the absorption phase survives and produces greater brightness temperature fluctuations than the subsequent emission phase (the delay in the absorption of X-ray connected to the long mean free path is partly responsible for this). It will be important to keep this result in mind when choosing the design and observation strategies for the future instruments.

There is a large observational uncertainty in the mean and distribution of the quasar spectral index $\alpha$. Our fiducial model assumes $\alpha = 1.6$ but we run a simulation (S5) with $\alpha = 0.6$ for comparison. The total emitted energy is fixed, but the S5 simulation has more energetic photons which penetrate the ionization front further than those of S4. However, the difference of the gas temperature between two simulations is negligible. The temperature of S5 is slightly higher than S4, but the difference is less than 0.5 K at all redshifts.

We estimate that our 1% model yields around 5 times more X-rays than the fiducial model in Santos et al. (2008), although the source formation modeling is quite different and the comparison is difficult (we do not use the dark matter halo mass at all in computing the star formation rate). However, comparing their plots of the average temperature and ionization fraction evolution with ours, we can deduce that their average gas kinetic temperature rises above the CMB temperature around ionization fraction $x_{HI} \approx 10\%$ while in our case the same event occurs at $x_{HI} \approx 15\%$. We find several reasons for this apparent discrepancy. First we defined neutral IGM as $x_{HI} < 0.01$ and use this to compute the average gas temperature. Although this is not absolutely explicit in the paper, we believe they use $x_{HI} < 0.5$, thereby including warmer gas in the average. Then, they have a more extended reionization history, which reduces the effect of the delay in the X-ray heating (see next section). Finally the initial X-ray heating is shifted to higher redshifts, when the difference between the average neutral gas temperature and the CMB temperature is less.

### 4.3. Brightness temperature maps

We have run Lyman-α simulations as a further post-treatment to compute the differential brightness temperature $\delta T_b$. The $\delta T_b$ is determined by various elements, and it is expressed as (Madau et al. 1997):

$$\delta T_b \approx 28.1 \text{ mK} \ x_{HI} \ (1 + \delta) \ \left(1 + z\right)^{\frac{3}{2}} \ \frac{T_S - T_{CMB}}{T_S} ,$$

where $\delta$ is the baryon over density, $T_s$ is the spin temperature, $T_{CMB}$ is the CMB temperature, and $x_{HI}$ is the neutral fraction. The contribution of the gradient of the proper velocity is not considered in this work.

The spin temperature $T_s$ can be computed with:

$$T_{s}^{-1} = T_{CMB}^{-1} + x_s T_{CMB}^{-1} + \frac{x_s T_{CMB}^{-1}}{1 + x_s}$$

and

$$x_s = \frac{4P_x T_{S}}{27 A_{10} T_{CMB}}$$

and

$$c_s = \frac{C_{10} T_{S}}{A_{10} T_{CMB}}$$

where $P_x$ is the number of Lyman-α scatterings per atom per second, $A_{10}$ is the spontaneous emission coefficient of the 21 cm hyperfine transition, $T_s$ is the excitation temperature of the 21 cm transition, and $C_{10}$ is the deexcitation rate via collisions. Details on deriving these relations and computing $C_{10}$ can be found, e.g., in Furlanetto et al. (2006). The peculiar velocity gradients (Barkana & Loeb 2005; Bharadwaj & Ali 2004) is not considered in this work.

As we can see in Eq. (9), $T_s$ is coupled to the CMB temperature $T_{CMB}$ by Thomson scattering of CMB photons, and to the kinetic temperature of the gas $T_k$ by collisions and Ly-α pumping. Coupling by collisions is efficient only at $z > 20$, or in dense clumps, so Ly-α is the key coupling process in the diffuse IGM. The $\delta T_b$ maps are a good way to see how these different elements affect the signal. In Fig. 4 we show several $\delta T_b$ maps of the same slice from different radiative transfer simulations, S1, S2, S6 and S7. S4 and S5, which sets the X-ray luminosity at 0.1% of the UV luminosity show a trend very similar to S1 and are not plotted. The bandwidth of the slice is 0.1 MHz for all maps, which corresponds to 1.9 Mpc for the maps on the left column (a)–(d), 1.8 Mpc for (e)–(h) and 1.6 Mpc for (i)–(l).

The left four maps of $\delta T_b$ in Fig. 4a–d, are plotted when the mass averaged Ly-α coupling coefficient $x_{HI}$ is $\langle x_{HI} \rangle = 1$. This value is interesting because in this moderate coupling regime, fluctuations in the Ly-α local flux induce fluctuations in the brightness temperature which is not the case anymore when the coupling saturates. The corresponding redshifts are $z = 10.50$ for S2 and 10.13 for the others. The corresponding averaged ionization fractions are 0.005, 0.018, 0.005 and 0.005 for (a)–(d). Indeed, the averaged ionization fraction of S2 is higher than the others since it uses a harder spectrum. The ratio of the integrated energy emitted in the Lyman band (Ly-α $< E < \text{Ly-}$limit) with respect to the ionizing band, $\beta = E_{\text{Lyman}} / E_{\text{Ion}}$, is three times less for S2 than for the others. For a given number of emitted Ly-α photons, a harder spectrum produces a larger number of UV ionizing photons, therefore S2 has a higher ionization fraction when $\langle x_{HI} \rangle = 1$. S1 shows a deeper absorption region around the ionized bubbles than S2, and it is also due to the different ratio of the number photons in the Lyman band and the ionizing band. In the case of S1, the ionized bubble is smaller than the highly Ly-α coupled region. Since the kinetic temperature outside of ionized bubble is a few kelvin, which is lower than the CMB temperature $T_{CMB} \approx 30$ K), the neutral hydrogen has a strong 21-cm absorption signal. On the other hand, the highly Ly-α coupled regions in S2 mostly resides in the ionized bubbles, which are bigger than in S1. S7 has almost the same averaged ionization fraction and ionized bubble size as S1, but the gas around the ionized bubbles as well as in the void is heated by strong X-rays. The signal is still in absorption because the X-ray heating has not been able to raise the IGM temperature above $T_{CMB}$, but the intensity is reduced. Contrary to S1 and S2, the neutral gas around the ionized bubbles produces a weaker signal than in the void, because the gas around the bubbles is more efficiently heated by X-rays. S6 shows an intermediate behavior between S1 and S7.

The four maps in the middle of Fig. 4e–h, are for $\langle x_{HI} \rangle = 10$. The corresponding redshifts are $z = 9.03$ for S2 and 8.57 for
Fig. 4. Differential brightness temperature maps for different simulations. The thickness of the slice is ≈2 Mpc. The maps in the left column are when $\langle x_\alpha \rangle = 1$, the middles when $\langle x_\alpha \rangle = 10$ and the right when $\langle x_{\text{HII}} \rangle = 0.5$. The black contour separates absorption and emission region. (l) shows no absorption region.

The averaged ionization fractions are 0.043, 0.141, 0.043 and 0.040 for (e)–(h). These redshifts when $\langle x_\alpha \rangle = 10$ are interesting because the amplitude of fluctuation $\delta T_b$ reaches a maximum. If we do not consider the effect of Ly-$\alpha$ coupling and...
assume $T_\alpha \gg T_{\text{CMB}}$, which does not allow the signal in absorption, the largest fluctuations would appear around $\langle \delta T_{\text{HI}} \rangle = 0.5$ as noticed by Mellemma et al. (2006a) and Lidz et al. (2007a), but including the inhomogeneous Ly-$\alpha$ coupling and computing $T_K$ self-consistently, this maximum is shifted to an earlier phase of reionization. The ionization fraction and bubble size in S2 are still greater than in the other models, but the absorption intensity is lower than in S1. Here is why. The evolution of kinetic temperature in the void regions is dominated by the adiabatic cooling; the temperature drops as the expansion progresses. The kinetic temperature of S1 in the voids is lower than in S2 by 0.5 K due to the difference in redshift, which explains the stronger absorption intensity in S1. The neutral gas around ionized bubbles in S7 is heated by the high X-rays level above $T_{\text{CMB}}$, and starts to produce the signal in emission. The neutral gas in the voids is also affected by X-rays. However, it is not sufficiently heated yet so that the signal turns everywhere from absorption to emission. Nevertheless the intensity of the signal is reduced by the X-ray heating, and it shows the weakest signal among the four maps at $\langle \alpha \rangle$.

The four maps on the right of Fig. 4, (i)–(l), are for $\langle \alpha \rangle$ = 0.5. The corresponding redshifts are $z = 7.68$ for S2, 7.00 for S1 and S6, 6.93 for S7. The averaged Ly-$\alpha$ coupling coefficients, $x_\alpha$, are 138.3, 34.9, 138.75 and 187.75 for (i)–(l). Contrary to the above cases, the absorption intensity in S1 in the void region is weaker than that of S2. This is due to the Ly-$\alpha$ heating. Ly-$\alpha$ heating is negligible during the early phase, but the amount of Ly-$\alpha$ heating accumulated between $z \sim 12$ and $z \sim 7$ can heat the gas in the voids by several kelvins. In order to reach 50% ionization, S1 produces a larger number of Ly-$\alpha$ photons, which propagate beyond the ionizing front. The $x_\alpha$ of S1 is almost 4 times greater than that of S2, and the accumulated Ly-$\alpha$ heating increases the kinetic temperature by 3–4 K more than in S2. The intensity in absorption is very sensitive to the value of kinetic heating, and it shows the weakest signal among the four maps at $\langle \alpha \rangle$. The X-ray heating in S7 is strong enough to heat all the gas above $T_{\text{CMB}}$, and it shows the smallest power. However, the power of S6 falls down under the power of S7 when the hydrogen is 50% ionized. At this redshift, the rising temperature of neutral gas reaches $T_{\text{CMB}}$ in S6, while it is already much greater than $T_{\text{CMB}}$ in S7. This is also visible in Fig. 4. Later, when $\langle \alpha \rangle = 0.9$, all power spectra drop. Our S6 model agrees quite well with Santos et al. (2008) both in shape and amplitude for these two last stages. Indeed both the effects of Lyman-$\alpha$ coupling and X-ray heating reach a saturation in the determination of the brightness temperature, erasing the differences in our treatments. S2 has the largest power over all scale when $\langle \alpha \rangle = 0.5$ and $\langle \alpha \rangle = 0.9$. This is due to the near lack of Ly-$\alpha$ heating. Let us mention however that some sort of transition to Pop II formation should have occurred by then, providing some level of Ly-$\alpha$ heating. So S2 is probably not realistic during the late EoR.

In brief, the 21-cm power spectra of our models vary in the 10 to 1000 mK$^2$ range, in broad agreement with Santos et al. (2008) who included the inhomogeneous X-ray and Ly-$\alpha$ effect on the signal in a semi-analytical way, with moderate discrepancies at high-redshift and small scale due to wing effect in the Lyman-$\alpha$ radiative transfer. Quite logically our results differ at high-redshift from Mellemma et al. 2006b; Zahn et al. 2007; Lidz et al. 2007b; McQuinn et al. 2006) who focused on the emission regime. These authors found a flattening of the spectrum around $\langle \delta T_{\text{HI}} \rangle = 0.5$. It is interesting to notice that in the case of a strong X-ray heating (model S7) the spectrum is quite flat at all redshift (temperature fluctuation boost the power on large scales at high-redshift). In the future observations, this would be a first clue of larger-than-expected contribution from X-ray sources.

We now plot the evolution of the power as a function of redshift for 4 different $k$ values. The evolution of the power spectrum with and without X-rays is very different. S1 and S2, which do not have X-rays, show a single maximum on small scales ($k = 1.00$ h/Mpc and $k = 3.15$ h/Mpc) around redshifts 8.5 and 9.

**Fig. 5.** Power spectrum evolution of the $\delta T_{\text{H}}$ from the S1 (thin black), S2 (thick black), S6 (thin gray) and S7 (thick gray) simulations. S2 use top-heavy IMF whereas the others use Salpeter IMF. S6 has 1% and S7 has 10% of total luminosity for X-rays.
which correspond to the redshifts of $\langle x_\alpha \rangle = 10$ for each simulation. On large scales ($k = 0.07 h$/Mpc and $k = 0.19 h$/Mpc) the power spectrum shows two local maxima on large scale. The first peak is related to the Ly-\(\alpha\) fluctuations. The $\delta T_b$ fluctuations are dominated by Ly-\(\alpha\) fluctuations at high-redshift, but it decreases when the Ly-\(\alpha\) coupling saturates. Then it rises again. This time, the fluctuations are dominated by the fluctuations of ionization fraction. The second peak appears at the redshift when $\langle x_{\text{HI}} \rangle = 0.5$ for each simulations. The overall amplitude of $S_1$ and $S_2$ are similar, but the position of the local maximum peaks of $S_2$ are at higher redshift due to the faster reionization. The key to the single-double peak difference is that the contribution of the ionizing field fluctuation to the brightness temperature power spectrum increases during reionization on large scale but not on small scales (Hill et al. 2006b).

With X-rays (models $S_6$ and $S_7$), the evolution follows a different scenario. We find a pattern similar to Santos et al. (2008). On small scales (thin and thick gray in Fig. 6), the intensity of the signal increases up to the maximum as the spin temperature couple to the kinetic temperature. Then it decreases during the absorption-emission transition. As the fluctuation due to the ionization fraction comes to dominate, the power reincreases slightly or remains in a plateau until it drops at the end of reionization. The evolution of the power on small scale does not show a marked minimum. The evolution of large scales (thin and thick gray in Fig. 6) is the most interesting: it shows three maxima. From high-redshift to low redshift, each peak corresponds to the period where the fluctuation of the Ly-\(\alpha\) coupling, gas temperature and ionization fraction dominate. There exists a deep suppression between the second and the third peaks which does not appear without X-ray. It occurs when the X-ray heating raises the gas temperature of the neutral IGM around $T_{\text{CMB}}$, which dampens the signal. The second minimum in $S_7$ occurs earlier than in $S_6$, since the stronger X-ray heating of $S_7$ increase the gas temperature around $T_{\text{CMB}}$ at a higher redshift. We find a much narrower third peak in $S_4$ (not plotted), which uses a 10 times weaker X-ray heating than $S_6$. The position and amplitude of the peaks as well as the width depend on the intensity of X-ray heating. The width of the third bump (6.5 $< z < 7.5$) is the largest in $S_7$ (with 10% X-ray) and the smallest or negligible in $S_4$ (with 0.1% X-ray). The existence/position of this third peak and of the second dip in the evolution of large scale power spectrum will be measurable by LOFAR and SKA observations, and it will help us constrain the nature of the sources during the EoR.

4.5. Non-Gaussianity of the 21-cm signal

The non-gaussianity of the 21-cm signal has been studied in previous works. Ciardi & Madau (2003); Mellema et al. (2006b) shows the non-gaussianity of the 21-cm signal in numerical simulations by computing the pixel distribution function (PDF). Ichikawa et al. (2010) draws the history of reionization from the measurement of the 21-cm PDF. Harker et al. (2009) compute the skewness of the PDF and show how it could help in separating the cosmological signals from the foregrounds. However, all these works model the signal in emission only.

We present the 21-cm PDF from our simulations in Fig. 7, for several representative redshifts. To obtain the 21-cm PDF, we sample $\delta T_b$ within a 1 $h^{-1}$ Mpc resolution, an acceptable value for SKA. The 21-cm PDF from our simulations is highly non-Gaussian as expected, but is also quite different from Ciardi & Madau (2003); Mellema et al. (2006b); Harker et al. (2009) and Ichikawa et al. (2010). Our distributions extend to negative differential brightness temperature with a variety of shapes depending on the redshift.

The panel (a) of Fig. 7 is the 21-cm PDF at the beginning of reionization, $z = 14.05$. It is at the beginning of reionization that Ichikawa et al. (2010) find a 21-cm PDF closest to a Gaussian, but this is also when their model is the less relevant. In our case, all signals are found in absorption and their distribution is peaked around 0mK, completely non-Gaussian.

We show the PDFs at $z = 10.64$, still during the early EoR, in panel (b) of Fig. 7. The position of peaks are shifted around $-100$ mK $\sim -50$ mK which means that the spin temperature of the particles is decoupled from $T_{\text{CMB}}$ by Ly-\(\alpha\) photons. The PDF is much close to a gaussian distribution, extending to positive values (the smooth curves are the best gaussian fit to the PDFs). However, all of them are left skewed (toward negative temperature). The reason for this negative skewness is the same as the reason for the positive skewness in Mellema et al. (2006b); Ichikawa et al. (2010); Harker et al. (2009): it is due to the signal from high density regions seen in absorption for us and in emission for them.

Panel (c) in Fig. 7 shows the PDFs when $z = 8.48$. Here we find a bimodal distribution with a plateau in the absorption region between 100 mK and 0 mK, for models $S_1$, $S_2$ and $S_4$. The left peaks of the PDFs moves also toward higher $\delta T_b$ with increasing heating efficiency. In the case of $S_7$, the left peak merges with the right one, and the form is very similar to a gaussian.

The panel (d) of Fig. 7 is plotted when the ionization fraction is 50%. The width of the PDF of $S_1$ and $S_4$ is reduced because Ly-\(\alpha\) heating is well advanced. We find signals in emission in $S_6$ and $S_7$, since X-rays heat the gas in neutral regions above $T_{\text{CMB}}$. Indeed, these PDF forms are similar to Ichikawa et al. (2010). The PDFs of $S_6$ and $S_7$ could be fitted by the Dirac-exponential-Gaussian distribution used by Ichikawa et al. (2010). $S_2$ has broad PDF still, since the Ly-\(\alpha\) heating is 4 times lower than in the others and it retains the signal in strong absorption.

It is interesting to note that the PDF always shows a spike around $\sim 0$ mK. During the beginning of reionization, it is due to the large amount of neutral hydrogen whose spin temperature is still well coupled to the CMB temperature. As the reionization proceed, the spin temperature is decoupled from the CMB and the number of pixels at $\sim 0$ mK decreases, but the peak grows again with the increasing contribution from completely ionized regions. This feature is interesting since interferometers such as LOFAR or SKA only measure fluctuations in the signal and do not directly provide a zero point.
Fig. 7. The evolution of the 21-cm PDF. The redshifts are 14.05, 10.64 and 8.48 for (a), (b), and (c). The PDFs of panel (d) is chosen so that the ionization fraction is 0.5. The red curves on (b) are gaussian fits with the mean and variance of the PDFs.

Ichikawa et al. (2010) extract informations about the averaged ionization fraction from the 21-cm PDF. As could be expected our PDFs converge with their result when the contribution from X-ray sources is sufficient and the EoR somewhat
advanced. What we find is that a clear tracking of the nature of the ionizing source remains on the PDF when the absorption phase is modeled. A strong X-ray contribution produce unimodal PDFs while a weak X-ray contribution yield a bimodal PDF.

The evolution of the skewness is presented in Fig. 8. The skewness $\gamma$ is defined as

$$\gamma = \frac{\frac{1}{N} \sum (\delta T_b^i - \bar{\delta T}_b)^3}{\left[ \frac{1}{N} \sum (\delta T_b^i - \bar{\delta T}_b)^2 \right]^{3/2}},$$

where $N$ is total number of pixels in $\delta T_b$ data cube, $\delta T_b^i$ is $\delta T_b$ in $ith$ pixel, and $\bar{\delta T}_b$ is the average on the data cube.

At the beginning the skewness is highly negative for all simulations as we can expect from the panel (a) and (b) of Fig. 7. Then, in all models, the skewness rises to a local positive maximum when the average ionization fraction is a few percents. It is interesting to notice that the skewness of all simulations close to zero again when the neutral fraction is about 0.3. While this behavior could be used to provide a milestone of reionization, its robustness should first be checked. We can notice that two of the three models presented in Harker et al. (2009) show the same behavior. Those are however the two less detailed models.

Another interesting feature in Fig. 8 is that the skewness of S7 has two local maxima while others do not. Again, this could be used as a clue to a large contribution from X-ray sources.

5. Conclusions

We modeled the 21-cm signal during the EoR using numerical simulations putting the emphasis on how various types of sources can affect the signal. The numerical methods used in this work are similar to Baek et al. (2009). The N-body and hydrodynamical simulations have been run with GADGET2 and post-processed with UV continuum radiative transfer further processed with Ly-\(\alpha\) transfer using LICORICE, allowing us to model the signal in absorption. The main difference from the previous work is a more elaborated source model, including X-ray radiative transfer and He chemistry.

We have run 7 simulations to investigate the effect of different IMFs, helium, different spectral indexes and the different luminosities of X-rays sources. The reference simulation in this work, S1, using only hydrogen and stellar type sources, reached the end of reionization at $z \approx 6.5$ and showed a strong absorption signal until the end of reionization. Our top heavy IMF (model S2) produces $\sim$2.6 times more ionizing photons than the Salpeter IMF. S2 reached the end of reionization earlier than the others by $\Delta z \approx 1$. In addition the different SED changes the ratio of Ly-\(\alpha\) and to ionizing UV photon numbers, and it slows down the saturation of the Ly-\(\alpha\) coupling and the heating by Lyman-\(\alpha\) in the top heavy IMF case. This modifies the statistical properties of 21-cm signal.

The simulation with helium, S3 also has a slightly earlier reionization than the others since the number of emitted photons per baryon is higher. Except for the slightly lower kinetic temperature in the bulk of ionized regions due to the higher ionization potential than for hydrogen, the properties of the 21-cm signal from S3 is similar to S1. We chose QSO type sources with a power-law spectrum as X-ray sources in model S4 to S7. The spectral index $\alpha$ has large observational uncertainty, so we used two different spectral indexes. S4 and S5 have 0.1% of the total luminosity in the X-ray band. S5 uses $\alpha = 0.6$, while other simulations with X-rays use $\alpha = 1.6$. S5 showed very little difference on the gas temperature with respect to S4. S4, S6 and S7 have different luminosities in the X-ray band, keeping the same values for the other simulation parameters. Using a stronger X-ray luminosity indeed increased the gas temperature in the neutral hydrogen. Accordingly the 21-cm signal and its power spectra are modified. We found an increase of a few kelvin for the neutral gas temperature in our fiducial model, S4, in which X-rays account for 0.1% of the total emitted energy. The 21-cm signal in S4 was similar to S1, showing the maximum intensity in absorption, $\sim$200 mK, at $z \approx 9$. Stronger X-ray levels increase the gas temperature and reduce the intensity. We found that in S6 and S7, which uses 1% and 10% of the total luminosity for X-rays, the absolute maximum intensity in absorption decreases to $\sim$130 mK and $\sim$80 mK. The 21-cm power spectrum of our work is greater by two or three orders of magnitude than in works focusing on the emission regime (Mellema et al. 2006b; Zahn et al. 2007; Lidz et al. 2007b; McQuinn et al. 2006). However, the results are in broad agreement with the work of Santos et al. (2008), who modeled absorption using semi-analytical methods for X-ray and Lyman-\(\alpha\) transfer. We noticed that the 21-cm fluctuation is dominated by Ly-\(\alpha\) fluctuations during the early phase, X-rays later (or the gas temperature), and the ionization fraction at the end. This is visible on the evolution of the 21-cm power spectrum with redshift. The 21-cm PDF of our work was different from other work, since we do not assume that the spin temperature $T_s \gg T_{\text{CMB}}$.

The first most important conclusion from our work is that even including a higher than generally expected level of X-ray, the absorption phase of the 21-cm survives. Its intensity and duration are reduced, but the signal is still stronger than in the emission regime. Heating the IGM with X-rays takes time!

The second important result is that we found three diagnostics which could be used in the analysis of future observations to constrain the nature of the sources of reionization. (i) The first and maybe the most robust is the evolution with redshift of large scale modes ($k \sim 0.1$ h/Mpc) of the powerspectrum. If reionization is overwhelmingly powered by stars, this evolution should have one local minimum (two local maxima). However, if the energy contribution of QSO is greater than $\sim 1\%$, a second local minimum (third maximum) appears. The higher the X-ray level, the broader the third peak. (ii) The second simple diagnostic is the bimodal aspect of the PDF which disappears when the X-ray level rises above 1% of the total ionizing luminosity. (iii) Last is the redshift evolution of the skewness of the 21-cm signal PDF.
While all other models show a single local maximum at a few percent reionization, a very high level of X-rays (>-10% of the total ionizing luminosity) produces a second local maximum appearing around 50% reionization.

Modeling the sources in the simulation is complex. It involves taking the formation history, IMF, SED, life time, and more into account. Although detailed models are desirable for the credibility of the results, we believe that the effect in the 21-cm signal can be bundled in 3 quantities. The first is the efficiency: how many photons are produced by atoms locked into a star. This parameter must be calibrated to fit observational constraint: end of reionization between redshift 6 and 7, and Thomson scattering optical depth in agreement with CMB experiment. The two other quantities which contain most of the information are two box-averaged ratios: the energy emitted in the Lyman band to the energy emitted in the ionizing band ratio and the same ionizing UV to X-ray ratio. In this work we explored values of 0.32 (model S2) and 0.75 (all other models) for the former and 0.001, 0.01 and 0.1 for the latter. Once additional physics is included in the simulation and using a higher resolution to account for all the sources, it will be interesting to explore the value of these quantities systematically.

We mentioned in the introduction that the minimum boxsize for reliable predictions of the signal is 100 h⁻¹ Mpc. It is important to realize that this value (confirmed by emission regime simulations, e.g., Iliev et al. 2006b) is estimated based on the clustering properties of the sources and applies to the topology of the ionization field. It may be underestimated when we study the early absorption regime, when only the highest density peaks contain sources. Their distribution is the most sensitive to possible non-gaussianities in the matter power spectrum. Moreover, they are distant from each other and, consequently, produce large scale fluctuations in the local flux of Lyman-α and X-ray photons. We intend to extend our investigation to larger box sizes in a future work.

A few final words on additional physics not included in our model. Shock heating from the cosmological structure formation is ignored, but it could have the potential to affect the 21-cm signal by increasing the gas temperature above the CMB temperature. However, it is not sure whether shocks are strong enough in the filaments of the neutral regions to affect the 21-cm signal. Mini-halos (~10⁶–10⁸ Mₗ) form very early during the EoR and are dense and warm enough from shock heating during virialization to emit the 21 cm signal, but Furlanetto & Oh (2006) find that the contribution of mini-halos will not dominate, because of the limited resolution of the instrumentation. However, shock heating is worth investigating with coupled radiative hydrodynamic simulations with higher mass resolution. Also worth investigating is the effect of including higher Lyman lines in the radiative transfer.

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Appendix A: Implemented method for UV and X-ray

In this section, we explain the numerical methods that we use for radiative transfer with helium and X-ray. A description of the main methods used in LICORICE appears in Baek (2009)⁶.

A.1. Hydrogen and helium ionization

The radiation field is discretized into monochromatic photon packets which are emitted with random direction and frequency (with appropriate distribution) from the point sources. Photon packets propagate through radiative transfer cells and deposit photons and energy depending on the absorption probability of each cell. We need to compute absorption probabilities for each absorbers H⁰, He⁰ and He⁺⁷.

The probabilities of the photon being absorbed by each elements are given by

\[ \mathcal{P}_{\text{H}^0} = \frac{\tau_{\text{H}^0}}{\tau_{\text{H}^0} + \tau_{\text{He}^0} + \tau_{\text{He}^+}} (1 - e^{-\tau_{\text{H}^0}}), \]

\[ \mathcal{P}_{\text{He}^0} = \frac{\tau_{\text{He}^0}}{\tau_{\text{H}^0} + \tau_{\text{He}^0} + \tau_{\text{He}^+}} (1 - e^{-\tau_{\text{He}^0}}), \]

\[ \mathcal{P}_{\text{He}^+} = \frac{\tau_{\text{He}^+}}{\tau_{\text{H}^0} + \tau_{\text{He}^0} + \tau_{\text{He}^+}} (1 - e^{-\tau_{\text{He}^+}}). \]

where \( \tau_{\text{H}^0}, \tau_{\text{He}^0} \) and \( \tau_{\text{He}^+} \) are the optical depths for H⁰, He⁰ and He⁺ in a given cell, and \( \tau = \tau_{\text{H}^0} + \tau_{\text{He}^0} + \tau_{\text{He}^+} \). Therefore, the total absorption probability, \( \mathcal{P}_{\text{total}} \), for a photon packet arriving in a cell of optical depth \( \tau \) is given by

\[ \mathcal{P}_{\text{total}}(\tau) = \mathcal{P}_{\text{H}^0} + \mathcal{P}_{\text{He}^0} + \mathcal{P}_{\text{He}^+} = 1 - e^{-\tau}. \]

For each cell, the optical depth \( \tau \) is

\[ \tau = \tau_{\text{H}^0} + \tau_{\text{He}^0} + \tau_{\text{He}^+} = \left[ \tau_{\text{H}^0}(\nu)n_{\text{H}^0} + \sigma_{\text{He}^0}(\nu)n_{\text{He}^0} + \sigma_{\text{He}^+}(\nu)n_{\text{He}^+} \right] l, \]

where \( \sigma_A \) is the photoionization cross-section for absorber \( A \in \{ \text{H}^0, \text{He}^0, \text{He}^+ \} \), \( n_A \) is number density in the cell and \( l \) is the path length in a cell. We use photoionization cross section fits in Verner et al. (1996).

If \( N_\gamma \) is the number of photons in a photon packet arriving in a cell, then the number of photons absorbed in a cell, \( N_A \), is:

\[ N_A = N_\gamma \mathcal{P}_{\text{total}} = N_\gamma (1 - e^{-\tau}). \]

The contribution of photoionization and photoheating to the evolution of the ionization fraction and the temperature within a time step \( \Delta t \) is:

\[ \Delta n_{\text{H}^0} = \frac{n_0}{n_{\text{H}}} \Gamma_{\text{H}^0} \Delta t = \frac{N_A \mathcal{P}_{\text{H}^0}}{N_{\text{H}}}, \]

\[ \Delta n_{\text{He}^0} = \frac{n_0}{n_{\text{H}}} \Gamma_{\text{He}^0} \Delta t = \frac{N_A \mathcal{P}_{\text{He}^0}}{N_{\text{He}}}, \]

\[ \Delta n_{\text{He}^+} = \frac{n_0}{n_{\text{H}}} \Gamma_{\text{He}^+} \Delta t = \frac{N_A \mathcal{P}_{\text{He}^+}}{N_{\text{He}}}. \]

\[ \Delta T = \frac{2}{3k_B n} \left\{ -\frac{3}{2} k_B T \Delta n + n_{\gamma} \mathcal{P}_{\text{H}^0}(h \nu - h \nu_{\text{th, H}^0}) + n_{\gamma} \mathcal{P}_{\text{He}^0}(h \nu - h \nu_{\text{th, He}^0}) + n_{\gamma} \mathcal{P}_{\text{He}^+}(h \nu - h \nu_{\text{th, He}^+}) \right\}. \]

where \( N_\gamma \) and \( N_{\text{He}} \) is the total number of hydrogen and helium in a cell, \( h \nu_{\text{th, A}} \) are the ionization potential of the recombinted atom, \( n_{\gamma} \) is the \( N_\gamma \) in a unit volume, \( \Gamma_A \) are the continuous photoionization rates used to actually integrate the evolution of the ionization fractions. Indeed recombination, collisional ionization and

⁶ Available at http://aramis.obspm.fr/~baek/these.pdf

⁷ H⁰ denotes the neutral hydrogen and He⁰ denotes the neutral helium.
radiative cooling are treated as continuous process, with integration time step $\Delta t^*$ much less than $\Delta t$ using the following coupled equations

\[
\frac{dn_\text{He}}{dt} = \gamma_{1\text{He}}(T)n_{\text{He}^0}n_e - \alpha_{1\text{He}}(T)n_{\text{He}^+}n_e + \Gamma_{1\text{He}}n_{\text{He}^0},
\]

\[
\frac{m_{\text{He}}}{dt} \frac{dx_{\text{He}^+}}{dt} = \gamma_{1\text{He}^+}(T)n_{\text{He}^+}n_e - \gamma_{1\text{He}^+}(T)n_{\text{He}^{++}}n_e - \alpha_{1\text{He}^+}(T)n_{\text{He}^{++}}n_e + \alpha_{1\text{He}^{++}}(T)n_{\text{He}^{++}}n_e + \Gamma_{1\text{He}^+}n_{\text{He}^{++}},
\]

\[
\frac{dx_{\text{He}^{++}}}{dt} = \gamma_{1\text{He}^{++}}(T)n_{\text{He}^{++}}n_e - \alpha_{1\text{He}^{++}}(T)n_{\text{He}^{++}}n_e + \Gamma_{1\text{He}^{++}}n_{\text{He}^{++}},
\]

where $\alpha_i$ and $\gamma_i$ indicate the recombination and collisional ionization coefficients and $I \in \{\text{H}^+, \text{He}^+, \text{He}^{++}\}$. We use the values in Hui & Gnedin (1997) for the recombination coefficient and recombination cooling. For collisional ionization coefficients and other radiative cooling we use the values in Cen (1992).

The temperature is computed from the energy conservation equation. $E$ is the internal energy of the gas, $E = \frac{1}{2}nk_B T$.

\[
\frac{dE}{dt} = \frac{d}{dt} \left(\frac{3}{2}nk_B T\right) = \mathcal{H} - \Lambda.
\]  

$\mathcal{H}$ and $\Lambda$ are the heating and cooling function which account for the energy gained and lost in a unit volume per unit time.

Each photon packet keeps propagating until it exits the simulation box (if we use free boundary condition) or until the remaining photon content is much less than the initial photon content, $N_e < 10^{-9}N_e^{\text{initial}}$. We typically adopt $p = 4$ for the UV continuum.

Appendix B: Validation tests

B.1. Radiative transfer comparison test

A cosmological radiative transfer code comparison project was performed in Iliev et al. (2006a) trying to understand which algorithms (including various flavors of ray-tracing and moment schemes) are suitable for a given non-trivial problem as well as to validate each code by comparing the results with other codes. Five tests are run for radiative transfer in a static density field. We reproduced several of tests in Iliev et al. (2006a) with LICORICE. LICORICE shows good agreements with other codes: results and comparisons with other codes are shown in Baek (2009).

Three additional tests are presented in Iliev et al. (2009) for the coupled gas dynamical and radiative transfer evolution. LICORICE directly participated in this second project and shows good agreements with the other codes.

B.2. Comparison with CLOUDY I: helium

An analytic solution to the radiative transfer evolution of an homogeneous medium around a single source exists only in the isothermal case with only hydrogen. In order to validate helium ionization and spectrum hardening, we reproduced the Strömgren sphere test in Maselli et al. (2003), and compared our results with the 1-D radiative transfer code CLOUDY\(^8\) (C08 version of the code). A point source, emitting as a black body at $T = 60,000$ K with ionizing luminosity $L = 10^{38}$ erg s$^{-1}$, is located at the center of the simulation box in a homogeneous density field with $n = 1$ cm$^{-3}$ composed of hydrogen (90% by number) and helium (10% by number). The gas is initially completely neutral at a temperature of $T = 10^4$ K in the entire simulation box cube of $L_{\text{box}} = 128$ pc. The comparison is performed at a time $t = 6 \times 10^5$ yr $\approx 5t_{\text{rec}}$ (where $t_{\text{rec}}$ is the characteristic time scale for hydrogen recombination).

The script used for CLOUDY is the following:

```
blackbody, T=60,000K
luminosity 38
radius 0.01 60 linear parsecs
hden -0.0457
abundances all -15
sphere static
element abundance helium -0.9542
punch element hydrogen "hydrogen_caseA.dat"
punch element helium "helium_caseA.dat"
punch temperature "temperature_caseA.dat"
punch continuum units eV "spectrum_60pc.dat"
```

In Figs. B.1 and B.2, the comparison between CLOUDY (solid lines) and LICORICE (circles) is shown. The value of the different physical quantities is plotted as a function of the distance from the source, expressed in cell units, $\Delta x = 1$ pc. The points represent spherically averaged LICORICE outputs. LICORICE shows a good agreement with CLOUDY. The positions of various ionizing fronts agree within 1%. To obtain this good agreement we consider the effect of secondary ionization and heating by fast electrons as CLOUDY does. The secondary ionization and heating fits in Shull & van Steenberg (1985) are only accurate for the primary photoelectrons higher than 100 eV, so we follow fits in Furlanetto & Johnson Stoever (2010) which is valid for ones less than 100 eV.

The temperature profile shows also a good agreement, except for the warm tail extending beyond the ionizing front for CLOUDY. This warm tail may originate from heat conduction which LICORICE does not implement.

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\(^8\) http://www.nublado.org/
Treating properly the spectrum hardening is an important issue in the radiative transfer with helium. Since each absorber (H\(^+\), He\(^0\) and He\(^+\)) has a different ionizing potential, the emitted spectrum will be strongly depleted just above the different ionizing frequencies once it hits a sufficient amount of absorbers. We plot in Fig. B.3 the luminosity per unit surface at different distances and compare between CLOUDY (solid lines) and LICORICE (circles). Our results agree well with the ones of CLOUDY. The noise at higher frequencies (> 5 Ryd) is due to the nature of the Monte Carlo method: the tails of the distributions are poorly sampled. At 4 pc, all species are already ionized so the medium is transparent and the spectrum is close to the initial one. At 14 pc, only photons with energy higher than 4 Ryd are absorbed to double ionize He. Above 40 pc, we can observe also the depletion at ~1.8 Ryd, which corresponds to the single ionization of helium.

B.3. Comparison with CLOUDY II: X-ray

We have also performed a comparison test with CLOUDY to validate X-ray radiative transfer. The geometrical initial set up is the same as in the test above. Now a point source in the center emits X-ray photons with energy from 100 eV to 2 keV, and a power-law spectrum with \(\alpha = -1.6\) in Eq. (5). The total luminosity is the same as in the helium test, \(L = 10^{38}\) erg s\(^{-1}\). The gas is composed of pure hydrogen, and completely neutral at the beginning. The recombinant time is proportional to the inverse of electron density. But instead of a sharp ionization front we have a gradual decrease in the ionization fraction with increasing radius. At the boundary of the box, the equilibrium ionization fraction is only ~5% so the recombinant time is ~20 times larger than in the inner region. Consequently the integration with LICORICE was extended to \(t = 2500\) Myr.

The script used for CLOUDY is the following:

```plaintext```
interpolate (1. -10.) (7.34 -10.) (7.35 0.)
(147.35 -2.08) (147.4 -10.) (200. -10.)
luminosity 38
radius 0.1 100 linear parsecs
hden -0.0457
abundances all -15
sphere static
punch element hydrogen "hydrogen_Xray.dat"
punch temperature "temperature_Xray.dat"
```

In Figs. B.4 and B.5, the comparison between CLOUDY (solid lines) and LICORICE (circles) is shown. The value of the different physical quantities is plotted as a function of the distance from the source, expressed in cell units, \(\Delta x = 1\) pc. The points represent spherically averaged LICORICE outputs. LICORICE shows a very good agreement with CLOUDY. The (very) small difference in the ionization fraction at large radii is due to the fact that we stopped the integration at \(t = 2500\) Myr, which is not much more than one local recombinant time: the equilibrium is not fully established. We could integrate for a longer time, but then the problem would only be shifted to larger radii and smaller ionization fractions.

Appendix C: Luminosity, SED, and typical life time of the sources

Let us consider a source particle containing a population of stars. The number of stars \(dN\) in a mass interval \(dM\) is described by the IMF function \(\xi(M)\):

\[dN = \xi(M)dM.\]
The properties of a star of mass M are described by the bolometric luminosity $L(M)$, the effective blackbody temperature $T_{\text{eff}}(M)$, and its life time $t_{\text{life}}(M)$. We can write the emissivity of a single star $\epsilon_{\nu}(\text{erg s}^{-1} \text{ Hz}^{-1})$:

$$\epsilon_{\nu}(M) = L(M) \frac{B_{\nu}(T_{\text{eff}}(M))}{\int_{0}^{\infty} B_{\nu}(T_{\text{eff}}(M)) d\nu},$$

where $B_\nu$ is the Planck law. Then we can write the time dependent emissivity of the complete population $\epsilon_{\text{tot}}$ as:

$$\epsilon_{\text{tot}}(\nu, t) = \frac{M_{\text{source}}}{\int \epsilon(\nu, M) dM} \int H(t_{\text{life}}(M) - t) \epsilon_{\nu}(M) \xi(M) dM$$

where $H$ is the Heavy side step function and the integrals over the masses are bounded by the mass interval in which we want to apply the IMF. With this quantity, we define a characteristic life-time during which most of the energy has been emitted. From this we can define the typical constant luminosity $L$ of a single star emitted in the band as:

$$L = \int_{0}^{\nu_{2}} \epsilon_{\text{tot}}(\nu, t) d\nu.$$ 

This gives the constant SED to use during a characteristic lifetime, and the normalization is given by the relation:

$$\int_{\nu_1}^{\nu_2} \epsilon_{\text{tot}}(\nu) d\nu = L_{\nu_1,\nu_2}.$$ 

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