IONIZED ABSORBERS IN ACTIVE GALACTIC NUCLEI: THE ROLE OF COLLISIONAL IONIZATION AND TIME-EVOLVING PHOTOIONIZATION

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ABSTRACT

In this paper we explore collisional ionization and time-evolving photoionization in the X-ray–discovered, ionized absorbers in Seyfert galaxies. These absorbers show temporal changes inconsistent with simple equilibrium models. We develop a simple code to follow the temporal evolution of nonequilibrium photoionized gas. As a result several effects appear that are easily observable and that, in fact, may explain otherwise paradoxical behavior. Specifically, we find the following:

1. In many important astrophysical conditions (O VII, O VIII dominant and high \( \geq 10^{22.5} \text{ cm}^{-2} \) column density) pure collisional and photoionization equilibria can be distinguished with moderate spectral resolution observations, because of a strong absorption structure between 1 and 3 keV. This feature is due mainly to iron L XVII–XIX and neon K IX–X absorption, which is much stronger in collisional models. This absorption structure may be misinterpreted as a flattening of the intrinsic emission spectrum above \( \sim 1 \text{ keV} \) in low-resolution data.

2. In time-evolving nonequilibrium photoionization models the response of the ionization state of the gas to sudden changes of the ionizing continuum is smoothed and delayed at low gas densities (usually up to \( 10^5 \text{ cm}^{-3} \)), even when the luminosity increases. The recombination time can be much longer (up to orders of magnitude) than the photoionization timescale. Hence, a photoionized absorber subject to frequent, quick, and consistent changes of ionizing luminosity is likely to be overionized with respect to the equilibrium ionization state.

3. If the changes of the ionizing luminosity are not instantaneous, and the electron density is low enough (the limit depends on the average ionization state of the gas but is usually \( \sim 10^4 \) to \( 10^5 \text{ cm}^{-3} \)), the ionization state of the gas can continue to increase while the source luminosity decreases, so a maximum in the ionization state of a given element may occur during a minimum of the ionizing intensity (the opposite of the prediction of equilibrium models).

4. Different ions of different elements reach their equilibrium configuration on different timescales, so models in which all ions of all elements are in photoionization equilibrium so often fail to describe active galactic nucleus (AGN) spectral evolution.

These properties are similar to those seen in several ionized absorbers in AGNs, properties that had hitherto been puzzling. We applied these models to a high signal-to-noise ratio ROSAT PSPC observation of the Seyfert 1 galaxy NGC 4051. The compressed dynamical range of variation of the ionization parameter \( U \) and the ionization delays seen in the ROSAT observations of NGC 4051 may be simply explained by a nonequilibrium photoionization model, giving well-constrained parameters:

\[
\frac{n_e}{\text{cm}^{-3}} = (1.0 \pm 0.2) \times 10^8 \quad \text{and} \quad R = (0.74_{-0.80}^{+0.84}) \times 10^{16} \text{ cm} \sim 3 \text{ light-days}.
\]

Subject headings: atomic processes — galaxies: active — galaxies: Seyfert — X-rays: galaxies

1. INTRODUCTION

The detection and the study of ionized absorbers is more difficult than that of cold neutral absorbers but can yield much more detailed information about the nature of the absorbers and the state and geometry of the nuclear regions of active galactic nuclei (AGNs). If photoionization applies, then the electron density of the gas and its distance from the ionizing source can be estimated.

Absorption features from ionized gas are common in the X-ray spectra of Seyfert galaxies and some quasars.5 Deep oxygen VII and VIII absorption edges at 0.74 and 0.87 keV (rest) have been detected by the ROSAT PSPC and the ASCA SIS in quite a large number of Seyfert 1 galaxies (Reynolds 1997).

We do not know yet what is the origin of the gas ionization. Models to date have assumed the simplest equilibrium photoionization case (e.g., Fiore et al. 1993; Guainazzi et al. 1996; Reynolds et al. 1995). In this case, if the gas is con-
fined in a single cloud of constant density, and if the recombination time is smaller than the typical variability timescale, then the ionization parameter, and hence the ionization state of the gas, should follow closely the intensity of the ionizing continuum. This is not always observed. In two ASCA observations of MCG – 6-30-15 (Fabian et al. 1994; Reynolds et al. 1995) the best-fit ionization parameter is higher when the ionizing flux is lower, in contrast with the expectations of the simplest equilibrium photoionization model; the ASCA observations of MR 2251 – 178 show a roughly constant ionization parameter despite large variations in the 2–10 keV flux (Reynolds & Fabian 1995); finally, in a ROSAT observation of NGC 4051 (McHardy et al. 1995), the ionization parameter does not linearly track the luminosity but shows changes that are smoothed and delayed with respect to the luminosity changes. It seems clear that, at least in the above three cases (which are also the best studied), the simplest photoionization equilibrium model is inadequate. We clearly need more complete and consistent models to interpret the available data. The gas could well be distributed in an irregular region with varying density. Different ionization states would then apply to different regions of this gas. Other authors have adopted such multizone models (Otani et al. 1996; Kriss et al. 1996). In those models the authors assume that absorption features from different ions of the same element are imprinted on the spectrum by the transmission of the ionizing radiation through multiple distinct clouds of gas with different geometrical configurations, ionization states, and densities. This, of course, a possibility, but to us it seems rather ad hoc.

In this paper we discuss instead the additional physics of collisional ionization and of time-evolving photoionization in a single-zone model. Time-evolving photoionization models have been applied in the past to the extended ionizing continua of Seyfert galaxies; however, those models assumed that absorption features from different ions of the same element are imprinted on the spectrum by the transmission of the ionizing radiation through multiple distinct clouds of gas with different geometrical configurations, ionization states, and densities. This, of course, a possibility, but to us it seems rather ad hoc.

In the case of a single zone, the best-fit ionization parameter is higher when the ionizing flux is lower, in contrast with the expectations of the simplest equilibrium photoionization model; the ASCA observations of MR 2251 – 178 show a roughly constant ionization parameter despite large variations in the 2–10 keV flux (Reynolds & Fabian 1995); finally, in a ROSAT observation of NGC 4051 (McHardy et al. 1995), the ionization parameter does not linearly track the luminosity but shows changes that are smoothed and delayed with respect to the luminosity changes. It seems clear that, at least in the above three cases (which are also the best studied), the simplest photoionization equilibrium model is inadequate. We clearly need more complete and consistent models to interpret the available data. The gas could well be distributed in an irregular region with varying density. Different ionization states would then apply to different regions of this gas. Other authors have adopted such multizone models (Otani et al. 1996; Kriss et al. 1996). In those models the authors assume that absorption features from different ions of the same element are imprinted on the spectrum by the transmission of the ionizing radiation through multiple distinct clouds of gas with different geometrical configurations, ionization states, and densities. This, of course, a possibility, but to us it seems rather ad hoc.

In this paper we discuss instead the additional physics of collisional ionization and of time-evolving photoionization in a single-zone model. Time-evolving photoionization models have been applied in the past to the extended narrow line region of radio galaxies, to predict the evolution of the line ratios in low-density ($n_e = 1 \text{ cm}^{-3}$), low-ionization ($\log U \sim -3$), multizone photoionized gas after the switch-off of the ionizing source (Binette & Robinson 1987). We present here an iterative solution of the time-dependent ionization balance equations (accounting for all ions of the most abundant elements in gas with solar composition) and apply our models to a drastically different physical and geometrical configuration: a relatively high-density ($n_e = 10^6$–$10^9 \text{ cm}^{-3}$), medium-high gas ionization ($\log U \sim -1$, 1), geometrically thin ($\Delta R/R \sim 10^{-3}$), single-zone absorber along the line of sight, undergoing rapid and persistent variations of the ionizing flux.

As an example, we apply our models to the ROSAT data of the Seyfert 1 galaxy NGC 4051. A detailed analysis of the time behavior of ionized gas in Seyfert 1 galaxies, using the ASCA-SIS relatively high resolution data, is deferred to a future publication.

2. IONIZATION MODELS

The innermost regions of an AGN are likely to be “active,” in the sense that the gas there confined is expected to be involved in significant bulk motion, as in strong outflows (Arav et al. 1995) or inflows. In particular, the evidence for ionized outflows from the central regions of AGNs is strong (Mathur et al. 1994; Mathur, Elvis, & Wilkes 1995; Mathur, Wilkes, & Aldcroft 1997; Mathur, Wilkes, & Elvis 1998). Hence, sources of mechanical heating of the gas (for example, adiabatic compression by shock waves) may well be at work in the high-density clouds, making collisional ionization the dominant ionization mechanism. In the low-density clouds the gas could be far from photoionization equilibrium because the recombination timescale is greater than the X-ray continuum variability timescale. Hereafter, by photoionization time ($t_{\text{ph}}$) and recombination time ($t_{\text{rec}}$) we mean the time necessary for the gas to reach photoionization equilibrium with the ionizing luminosity during increasing and decreasing phases, respectively. We discuss, in turn, the importance of collisional ionization and time-evolving photoionization in determining the final transmitted spectrum that we observe in AGNs.

2.1. Collisional Ionization Models

To study collisional ionization we constructed a series of pure collisional ionization models and compared them with the equilibrium photoionization models. The models were created using CLOUDY (version 90.01, Ferland 1996), fixing the ionization parameter (i.e., the dimensionless ratio between the number of hydrogen-ionizing photons and the electron density of the gas) $U$ to $10^{-5}$ and calculating the spectra transmitted through clouds with a total column density, $N_H = 10^{22.5} \text{ cm}^{-2}$, constant density and temperature (the calculation of the physical condition of clouds in coronal or collisional equilibrium is one of the options of CLOUDY, Ferland 1996). The distribution of fractional abundances for the most important ions is essentially independent of the electron density value: for $10^{-3} < n_e < 10^{12} \text{ cm}^{-3}$; the variation is smaller than 13% for ions with fractional abundances greater than 0.01. For temperatures $T_e$ in the range $10^{5}$–$10^{7.1}$ the main edges imprinted on the spectra are roughly the same as those in the spectra from photoionized gas with $-1 < \log (U) < 1.5$. However, the distribution of the different ionic species is very different in the two cases, and so the relative optical depth of the edges differ markedly. High-quality X-ray spectra, where more than one edge is visible, could then discriminate between the two cases.

The different ionic abundances in the collisional and photoionized cases can be seen in Figures 1a and 1b and 2a and 2b. Here we plot the fractional abundances of Ne VIII–Ne X and O VI–O IX (Fig. 1) and Fe XIV–Fe X XV (Fig. 2), computed in the case of photoionization and collisional ionization, respectively, as a function of $U$ and $T_e$. As Fiore et al. (1993) and Mathur et al. (1994) have shown, the photoionization state of the gas depends strongly on the spectral energy distribution (SED) of the ionizing continuum, from radio to hard X-rays. To simplify the comparison between photoionization models and the data, we therefore used an ionizing continuum similar to the SED of NGC 4051 (Done et al. 1990).

First we note that the fractional abundances in photoionized gas are more smoothly distributed than those of collisionally ionized gas. Let us consider the regime in which the O VII and O VIII fractions are higher than ~0.2, so that both O VII and O VIII edges are present in the emerging spectrum, as sometimes found in Seyfert galaxies spectra (e.g., Otani et al. 1996; Guainazzi et al. 1996). In the photoionization case the range of $U$ where this occurs is 1.2–3.6, a factor of 3. This is twice as wide as the corresponding range of $T_e$: 1.5 × $10^6$ to 2.4 × $10^6$ K (a factor of 1.6: dotted lines in panels a and b of Fig. 1). Moreover, Ne IX is abundant, and carbon is almost fully stripped in the collisional case.
but not in the photoionized case.

The difference in the resulting spectra is shown in Figure 3a. Here we show the ratio between two power-law spectra emerging, respectively, from clouds of collisionally ionized ($T_e = 1.8 \times 10^6$ K) gas and photoionized ($U = 2$) gas with solar abundances (the choices of $T_e$ and $U$ were made by looking for similar O VII–O VIII abundances; see Fig. 1a). Below 1 keV there is a large feature due to the deep C VI K absorption edge in the photoionization case. The most important feature above 1 keV is the Ne IX absorption edge in the collisional case, but this feature is smoothed by the presence, in both cases, of deep O VII and O VIII absorption edges. Reynolds (1997) fitted both a simple O VII and O VIII K-edges model and a physical photoionization model to ASCA SIS spectra of a sample of Seyfert 1 galaxies known to host a warm absorber. Comparing the results, he found
that in some cases the best-fit two-edge model continues to show residuals at the energy of the C \textsc{vi} K edge, while residuals to physical photoionization model do not. This may point to photoionization as an ionizing mechanism. Spectra with higher resolution from 0.4 to 3 keV should therefore allow one to diagnose absorption by collisionally ionized or photoionized gas that has significant O \textsc{vii} and/or O \textsc{viii} abundances (e.g., Otani et al.
1996; Guainazzi et al.
1996).

The other important feature of collisional models is the inertia of the heavy elements (from Ne to Fe) against becoming highly ionized, even when the oxygen is almost fully stripped. In Figures 1a and 1b we have marked the range of \( U \) for which O \textsc{ix} is by far the dominant ion (\( n_{\text{Oix}} \gtrsim 0.8 \)) and O \textsc{vii} is the only oxygen edge in the emergent spectrum (\( \tau \lesssim 0.8 \), for an \( N_H \) of 10\(^{22.5} \) cm\(^{-2} \) and solar abundances). These same intervals of \( U \) and \( T_e \) are also shown in Figures 2a and 2b. In the photoionization case neon is almost fully stripped (Ne \textsc{xii}) and the dominant ions of the iron are Fe \textsc{xviii}–Fe \textsc{xxiii}. Instead, in the collisional case there is a range of temperature for which neon and iron are much less ionized, with the dominant ions being Ne \textsc{x}–Ne \textsc{x}, and Fe \textsc{xvi}–Fe \textsc{xx}. In particular, we note the large abundance of Fe \textsc{xvii} (Fe \textsc{xvi} and Fe \textsc{xx} are, respectively, Ne-like and He-like and thus very stable). The different ionization level of neon and iron implies a very different emergent spectrum between 1 and 3 keV.

This is illustrated in Figure 3b, which shows the ratio between spectra transmitted from collisionally ionized gas with \( T_e = 3.9 \times 10^6 \) K and photoionized gas with \( U = 10 \) (both with \( N_H = 10^{22.5} \) cm\(^{-2} \)). With these values of \( T_e \) and \( U \), \( n_{\text{Oix}} \sim 0.8 \) in both collisional ionization and photoionization models (see Fig. 1a). The ratio does not show any significant feature at \( E < 1 \) keV (implying similar ionization states of carbon, nitrogen, and oxygen), but between 1 and 3 keV, the spectrum from collisionally ionized gas shows a large and complicated absorption structure due mainly to iron L \textsc{xvii}–\textsc{xix} and neon K \textsc{ix}–\textsc{x} absorption (Fig. 3b). This absorption structure may be misinterpreted in moderate-quality spectra as a flattening above \( \sim 1 \) keV. Again, higher spectral resolution data with good signal-to-noise ratios (S/N) can distinguish between photoionization or collisional equilibrium.

That the Ne \textsc{ix} and Fe \textsc{xvii} edges are of similar depth is because of a coincidence of cross sections and abundances. The photoelectric cross section of the iron L-shell atoms ranges from \( \sim 2 \times 10^{-19} \) cm\(^2 \) for the Lp levels of Fe \textsc{xvi}–Fe \textsc{xxi} \([ E \sim (1.17–1.47) \text{ keV}] \), to \( \sim 3 \times 10^{-20} \) cm\(^2 \) for the Lp level of Fe \textsc{xxii} \([ E = 1.78 \text{ keV}] \) (Kallman & Krolik
1995).

Furthermore, in the collisional case the relative abundance of Fe \textsc{xvii} is in the range 0.2–0.7, compared with \( \lesssim 0.25 \) in the analogous photoionization case. The relative abundance of the Ne \textsc{x}, in the collisional case, is greater than 0.25, but Ne \textsc{x} \( \lesssim 0.4 \) in the analogous photoionization case, while 0.05 \( \lesssim (\text{Ne } \textsc{x})_{\text{coll}} \lesssim 0.40 \) and (Ne \textsc{x})\text{phot} \( \lesssim 0.05 \) (Figs. 1a and 1b). The K-edge energy of Ne \textsc{x} is 1.36 keV, the same as that of the Fe \textsc{xviii} Lp-edge. While the photoelectric cross section of neon is about a factor 4 lower than iron, the solar abundance of the neon is about a factor 4 greater than that of iron (Grevesse & Anders
1989). Furthermore, the cross section of Ne \textsc{x} is about a factor 3 higher than Ne \textsc{x}. As a result, spectra from a cloud of collisional ionized gas at \( T_e \sim (4–6) \times 10^6 \) K, will show a very deep \( (\tau_{\text{FeLp xvi}} \sim 1.0 \times N_H) \) edge at the Fe \textsc{xviii} Lp-edge energy (1.26 keV), a deep \( (\tau_{\text{FeLp xvi+NeKX}} \sim 0.7 \times N_{\text{H}_2}) \) edge at 1.36 keV due to the comparable contributions of the Ne x K-edge and the Fe \textsc{xviii} Lp-edge, and a similarly deep absorption edge at the Ne \textsc{x} K-edge energy \([ E_{\text{K}}(\text{Ne } \textsc{x}) = 1.196 \text{ keV}, \tau_{\text{NeKX}} \sim 0.6 \times N_{\text{H}_2}] \). In the corresponding photoionization case, the relative abundances of both Ne \textsc{x}–\textsc{x} and Fe \textsc{xviii}–\textsc{xvii} are too low (Figs. 1a and 1b) to imprint similar features on the spectrum.

When collisions are the dominant ionization process, \( (U \lesssim 0.01) \) we can estimate the minimum distance, \( R \), between the ionized gas cloud and the central source. For an ionizing luminosity \( L_{\text{ion}} \), we find \( R > 1.6 \times 10^{16}(n_{\text{e10}})^{-0.5}(Q_{52})^{0.5} \) (with \( n_{\text{e10}} \) being the electron density in units of \( 10^{10} \) cm\(^{-3} \) and \( Q_{52} \) the rate of photons ionizing hydrogen in units of \( 10^{52} \) photons s\(^{-1} \); \( Q = \int_{0}^{\infty} dE[L_{\text{ion}}(E)/E] \)). For typical AGN ionizing continuum shapes and luminosities \( (L_{\text{ion}} = 10^{42}, 10^{45} \text{ ergs s}^{-1}) \) and assuming \( n_{\text{e10}} = 1 \), we find \( R > 6.2 \times 10^{16} \) cm, \( 2 \times 10^{18} \) cm, respectively, similar to the size of the broad emission line region for such AGNs (Peterson et al.
1993).

2.2. Mixed Collisional and Photoionization Models

The above discussion concerns gas in pure collisional equilibrium. If the gas density is low enough and/or if the gas is close enough to the central X-ray source, then photoionization can be important too. We examined a number of models with a varying mixture of collisional ionization and

\( ^6 \) The photoelectric cross section of the Ls levels of the iron ion is about 1 order of magnitude lower than the corresponding Lp cross sections, and the Lp levels of the ions Fe \textsc{xviii}–Fe \textsc{xxvi} are not populated in ordinary conditions. (Ls and Lp indicate atomic states with quantum numbers \( n = 2 \) and \( l = 0, 1 \), respectively.)
photoionization. For $T_e = 2.5 \times 10^6$ we find that for values of $U \leq 0.1$ the main features of the fractional abundance distribution are still those of pure collisional ionized gas. Increasing $U$ to 0.3 causes a sharp change in the ionization structure. For $0.3 < U \leq 100$ (the exact value of the upper limit depending on the equilibrium temperature determined by collisional mechanisms) the ionization structure is determined by both processes and the transmitted spectra resemble those from pure photoionized gas of much higher ionization parameter. This is shown in Figure 4 where we plot in the lower panel three spectra from purely photoionized gas (column density $\log N_H = 22.5$ and log ($U$) = 0.7, 1.1, and 1.4) and in the upper panel five spectra from gas in which both mechanisms are at work ($\log N_H = 22.5$, $T_e = 2.5 \times 10^6$ and log ($U$) = $-2, -1, 0.3, 0.4$). The pure photoionization spectra with log ($U$) = 0.7, 1 (lower panel) are very similar to the mixed collisional and photoionization spectra, but with a $U$ that is a factor of 4–10 times lower [$\log (U) = -0.3, 0.4$; upper panel]. Fitting the mixed spectra with pure photoionization models would give a much more compressed range of $U$ than the real one, as found by McHardy et al. in NGC 4051. The above spectra in the two panels of Figure 5 are practically indistinguishable even at high resolution, because the shape of the fractional abundance distributions when both processes are working closely resembles those of a purely photoionized gas (see upper panels of Figs. 1 and 2). Fortunately, the two models predict quite different delay properties (§ 2.3).

**2.3. Time-evolving Photoionization Models**

If photoionization is the dominant process (the gas being purely photoionized or the density low enough to give $U \gtrsim 0.1$) the main features of the fractional abundance distribution are those of photoionized gas. However, if the X-ray source is variable, photoionization equilibrium will apply only if the density is high enough to make the ion recombination timescales shorter than the variability timescales.

There are regimes where photoionization equilibrium does not apply at all. Here we investigate the low-density case in some detail, beginning with a discussion of the relevant physics.

### 2.3.1. Equilibrium Photoionization

Let us suppose that a geometrically thin single cloud of optically thin gas is illuminated by an intense flux of ionizing radiation emitted by a variable source located at a distance $R$ from the cloud.

We can calculate the equilibrium distribution of the ionic species in the cloud by setting the photoionization rate equal to the radiative recombination rate (see, e.g., Netzer 1990):

\[
\frac{n_{X_i+1}}{n_{X_i}} = \frac{F_{X_i}}{\alpha_{\text{eq}}(X^i, T_e)n_e},
\]

and adding the condition for charge conservation $\sum_i n_{X_i} = 1$. Here $\alpha_{\text{eq}}(X^i, T_e)$ is the radiative recombination coefficient (in $\text{cm}^3\text{s}^{-1}$), which includes recombination to all levels, and $F_{X_i}$ is the photoionization rate of the ion $X^i$, which, for optically thin gas clouds, can be written (see, e.g., Netzer 1990)

\[
F_{X_i} = \frac{1}{4\pi R^2} \int_0^\infty dv \frac{L_i}{hv} \sigma_i(X^i).
\]

The lower limit of the integral is the threshold ionization frequency of the ion $X^i$; $\sigma_i(X^i)$ is the photoelectric absorption differential cross section of the ion $X^i$.

In this simple scheme the ionization state of the gas is completely determined, at equilibrium, by the value of the ionization parameter $U = F_{X_i}/n_e c$. 8

With these approximations the ionization parameter $U$ can be written as a function of the ratio between any two consecutive ionic species of the generic element $X$:

\[
U = \left( \frac{n_{X_i+1}}{n_{X_i}} \right) \frac{F_{X_i}}{F_{X_{i+1}}} \frac{\alpha_{\text{eq}}(X^i, T_e)}{c}.
\]

### 2.3.2. Time-evolving Photoionization: Equations

If the gas is not in equilibrium equation (3) is, of course, meaningless. However, it is still possible to use it formally by introducing an ionization parameter $U^{X_i,X_{i+1}}(t)$ that depends both on time and on the specific ionic species under consideration.

The time evolution of the relative density of the ion $i$ of the element $X$, considering only radiative recombination

\[
\tau_i^1 = \frac{1}{F_{X_i}^*},
\]

\[
\tau_i^2 = \frac{1}{\alpha_{r}(X^i, T_e)n_e}.
\]

Instead, in this paper we refer to the different quantities as photoionization time and recombination time; see eq. (5).
and photoionization, is then given by (Krolik & Kriss 1995)
\[
\frac{dn_i}{dt} = - [F_{X_i} + \alpha_{rec}(X^{i-1}, T_e)n_{e}]n_{X_i} + F_{X^{i-1}}n_{X^{i-1}} + \alpha_{rec}(X^i, T_e)n_e n_{X^{i-1}}.
\] (4)

The first term on the right-hand side of equation (4), is the destruction rate of the ion \(X^i\), both by photoionization \(X^i \rightarrow X^{i-1}\) and radiative recombination \(X^i \rightarrow X^{i-1}\), while the other two terms indicate the formation rate of the ion \(X^i\) by photoionization of the ion \(X^{i-1}\) and radiative recombination of the ion \(X^{i-1}\), respectively.

The solution of equation (4) is a system of \(N\) coupled integral equations in the \(N\) unknowns \(n_{X_i}\), which is analytically solvable only for \(N = 2\), with the addition of the charge conservation condition. These solutions define the timescale \(t_{eq}\) that measures the time necessary for the gas to reach photoionization equilibrium with the ionizing continuum. This time is given at any point of the light curve of the ionizing continuum by the inverse of the destruction rate of the ion \(X^i\) (following Krolik & Kriss 1995). A useful analytical approximation for \(t_{eq}\) is
\[
t_{eq}^{X_i X^{i+1}}(t \rightarrow t + dt) \sim \left[ \frac{1}{\alpha_{rec}(X^i, T_e) n_e} \right] \times \left\{ \left[ \alpha_{rec}(X^{i-1}, T_e)/\alpha_{rec}(X^i, T_e) \right]_{eq} + (n_{X^{i+1}}/n_{X^i})_{eq} \right\}^{t + dt},
\] (5)

where “eq” indicates the equilibrium quantities. During increasing ionization flux phases we call \(t_{eq}^{X_i X^{i+1}}\) the photoionization time \(t_{ph}\); during decreasing ionization flux phases we call \(t_{eq}^{X_i X^{i+1}}\) the recombination time \(t_{rec}\). These times are generally different from \(t_1\) and \(t_2\) in the equations in footnote 8.

Equation (5) shows that the time \(t_{eq}^{X_i X^{i+1}}(t \rightarrow t + dt)\) necessary for the gas to reach equilibrium depends explicitly on the electron density \(n_e\) in the cloud and on the equilibrium ratio between two consecutive ionic species calculated at the time \(t + dt\). This is the key result of this work, which has major consequences:

1. The timescale on which the gas reaches equilibrium with the ionizing continuum depends on the electron density, even when the continuum increases (§ 2.3.5).
2. Different ions reach their equilibrium relative abundances at different points of the light curve (§ 2.3.6).
3. If changes of intensity are not instantaneous \((dI/dt < \infty)\), the time behavior of the relative abundance of a given ion can be opposite to that of the ionizing source (Fig. 6).

All of these effects have been seen in ionized absorbers in AGNs. Using these effects, nonequilibrium photoionization models can strongly constrain the physical state of the absorbing gas. 3. Time-evolving Photoionization: Calculations

We created a program to solve the first-order differential equation system of equation (4) for all the ions of the elements H, He, Li, C, N, and O. The program uses an iterative method (see Gallavotti 1986) that permits the solution of any system of \(N\) first-order differential equations in the \(N\) unknowns \(x_i\), of the form \(\dot{x}(t) = f[x(t)]\), \((\forall t > 0)\), with the only conditions being that \(f_i \in C^\infty\) and a limited ensemble \(\Omega\) exists, such that \(x_i \in \Omega\).

We consider the photoionization from the K shell of each element, and radiative recombination to all levels for each ion. We use the recombination rates tabulated by Shull & Van Steenberg (1982) for the metals. We take the values of the recombination rate of hydrogen from Ferland (1996); we get the total recombination rate by summing over levels \(n = 1, 20\). We calculate the photoionization rate from the K shell of each ion, carrying out the integrals in equation (2), using for the spectral shape of the ionizing continuum, from the Lyman limit to \(\gamma\)-rays, a simple power law with \(\alpha = 1.3\) (similar to the observed SED of NGC 4051). We use the photoelectric K-shell cross section tabulated by Kallman & Krolik (1995).

We carry out the calculation of the time-evolving heating-cooling balance, as follows: We calculate with CLOUDY (Ferland 1996) a grid of models for 300 values of \(U\) (from 0.01 to 100), and build the curve \(U = U(T_e)\), using the technique described by Kallman & Krolik (1995). We then interpolate on these to obtain the initial self-consistent equilibrium electron temperature of the gas. The time evolution of the temperature in the cloud is carried out performing an iterative calculation of the time-dependent photoionization equations, using the definition of \(U^{N X_i X^{i+1}}(t)\) given above.

The inputs the program needs are (1) the initial equilibrium value of the ionization parameter \(U\); (2) the spectral shape of the ionizing continuum; (3) the light curve of the ionizing continuum; (4) the electron density \(n_e\); (5) and the ratio, \(p\), between the source intensity and the intensity the source should have in order to produce the observed degree of ionization, assuming equilibrium at the beginning of the light curve.

The output of the program is a list of the relative ionic abundances of the chosen element and the source flux at the illuminated face of the cloud as a function of time.

2.3.4. Limits of the Code

We neglect photoionization from the L shell. This affects the computation of the abundance balance when the gas is allowed to recombine to medium-to-low ionization states, with the Li-like ions (or lower) being highly populated. However, this is not likely to be the case for warm absorbers in AGNs, for which the mean degree of ionization is usually very high and the most abundant elements (C to Ne) are almost equally distributed between He-like and fully stripped ions (see Fig. 1a). Our code can provide accurate results only at medium-high ionization regimes, those of interest for the astrophysical problem discussed in this paper.

We have also neglected iron ions. This will likely cause an overestimation of the O VII–IX and Ne IX–XI K-shell photoionization, since all these ions compete for the same photons. By comparing the O VII–IX, Ne IX–XI equilibrium abundances obtained with our method with those obtained with CLOUDY for \(U\) in the range 0.1–50 we estimate that the percentage differences are smaller than 7%.

The iterative technique presented in the previous section is only an approximation to the correct self-consistent time-evolving heating-cooling calculation accounting for all the physical and dynamical heating-cooling mechanisms. We verified that for high gas electron densities (and therefore in a situation close to equilibrium) this method is rather accu-
rate. By comparing the equilibrium relative abundances of the main ions obtained with our method with that obtained using CLOUDY we estimate that for $U$ in the range 0.1–50 the method works with a precision better than 10%. In the opposite case, when the density is sufficiently low, the gas remains basically in the same state and does not respond to variation of the ionizing continuum. Of course, in this case our approximation is also very good, albeit uninteresting. In the intermediate cases, when the heating timescale may be longer than the typical photoionization timescale, the recombination rate coefficients may be greater than those estimated by our assumption. We have tried to estimate the magnitude of the uncertainty on the ion abundance distribution induced by this effect by performing the time-evolving calculations fixing the temperature at a constant value equal to the initial value and letting the ionizing intensity be free to vary by a factor of 10. For an average gas ionization state typical of AGN warm absorbers ($U = 0.1–50$), the percentage difference between the relative abundances of the main ions calculated with the iterative solution of the time-dependent photoionization equations and those calculated for a gas with a constant temperature are $\leq 15\%$–$20\%$. The difference is so small because of the rather small dynamic range spanned by the temperature in a photoionized gas with $U = 0.1–50$ ($3 \times 10^4 \leq T_e \leq 3 \times 10^7$) and of the weak dependence of the recombination rates on $T_e$ (about the square root of $T_e$, see Shull & Van Steenberg 1982).

Given the configuration of the gas we considered (a thin, plane-parallel slab obscuring the line of sight) and the range of electron and column densities, the light-crossing time is smaller than both the typical source intensity variability timescales and the associated photoionization recombination timescales. We can then neglect the effects of the light-crossing time through the cloud of photoionized gas (see Binette 1988).

### 2.3.5. The Step Function Light Curve

The simplest case is that of a two-state light curve. Let us suppose that the ionizing continuum intensity goes instantaneously from a “low” to a “high” state and comes back to the “low” state after a time $t_{\text{var}}$ (Fig. 5a). The time behavior of the ionization state of the gas irradiated by this continuum depends on the value of the ratio $t_{\text{var}}/t_{\text{eq}}^{X^\text{VIII}}(t \to t + dt)$ and the amplitude of the flux variation (here we adopt a factor 10 change in flux).

We considered the case of an optically thin cloud of gas with an initial ionization such that the most relevant ionic species of the oxygen are $\text{O VIII}$ and $\text{O IX}$ (corresponding to an equilibrium value of $U \geq 5$ with the adopted SED).

The time behavior of the relative density of $n_{\text{O IX}}$ is shown in Figure 5b. In both panels different lines identifies different values of the electron density, $n_e = 10^7$, $5 \times 10^7$, $10^8$, $10^9$ cm$^{-3}$. In the upper panel different $n_e$ imply four values of the distance of the cloud from the ionizing source.

The photoionization time $t_{\text{ph}}^{\text{O VIII}, \text{O IX}}$ (eq. [5]) of the gas becomes progressively longer as the density decreases: from about $3 \times 10^3$ s for $n_e = 10^6$ cm$^{-3}$, to $\sim 10^4$ s for $n_e = 5 \times 10^6$ cm$^{-3}$, to even longer timescales for $n_e = 10^7$ cm$^{-3}$ (for $n_e \leq 10^6$ cm$^{-3}$ the changes of $n_{\text{O IX}}$ during the first $10^3$ s are less than 10%). Note that this is not true for $t_1$ in equation (3), which is the definition of photoionization time usually found in literature.

Formally, the dependence of $t_{\text{ph}}$ on $n_e$ is introduced by

![Figure 5](image_url)

**Fig. 5.** Step light curve of the ionizing radiation, for four different values of the electron density ($n_e = 10^7$, $5 \times 10^7$, $10^8$, and $10^9$ cm$^{-3}$, which correspond to four different values of the incident flux; upper panel). Time behavior of the relative abundance of $\text{O IX}$ for the four values of $n_e$ (lower panel).

the choice of a particular set of boundary conditions when solving equation (4) and hence defining $t_{\text{eq}}^{X^\text{VIII}}(t \to t + dt)$. Physically, fixing the boundary conditions of the system means choosing a particular initial ionic distribution in the gas and hence its initial ionization state. Different values of $n_e$, given the initial ionization state of the gas (and hence the ratio between the flux of ionizing photons at the illuminated face of the cloud and the electron density), imply different distances of the gas from the X-ray source. This is clear in the upper panels of Figure 5, where the ionizing flux at the illuminated face of the cloud is plotted for different values of $n_e$.

Recombination times are generally longer ($t_{\text{rec}}^{\text{O VIII}, \text{O IX}} > t_{\text{var}}$) and can be orders of magnitude longer. At the highest density tested ($n_e = 10^9$ cm$^{-3}$) the ionization state of the gas is able to relax to the initial equilibrium state in less than $10^4$ s after the source switch-off. Recombination timescales for $n_e \leq 10^8$ cm$^{-3}$ are long, on the order of many times $10^4$ s. Since the source switch-off is instantaneous the relative density of $\text{O IX}$ never increases after the switch-off.

This case illustrates clearly how photoionization recombination timescales can have a strong effect on the changes observed in ionized absorbers and why the photoionization timescale depends on electron density. In the following section we present a more realistic light curve and discuss in detail the main features of our models.

### 2.3.6. The Gradual Rise and Decay Light Curve: dL/dt $< \infty$

We now consider a more realistic light curve. In this case the source intensity goes from a low state to a high state in a finite time (4000 s), and after 2000 s comes back to the initial low state with the same absolute gradient (Fig. 6a). The entire up and down cycle lasts 10,000 s. As in the previous case the change in flux is a factor 10. The corresponding light curves of the relative abundances of the fully stripped ions of three different elements, C VII, N VIII, and O IX, are shown in Figure 6b. In both panels different lines corre-
respond to different values of the electron density, \( n_e = 10^9, 10^{10} \text{ cm}^{-3} \).

The gradual changes of the ionizing continuum produce time delays between the source light curve and the relative ion abundance light curves. In the lower density case \( n_e = 10^9 \text{ cm}^{-3}, \text{solid lines} \), \( n_{\text{ox}} \) reaches its maximum value (well below the equilibrium value \( \sim 1 \)) around the minimum of the luminosity intensity (at \( \sim 10,000 \text{ s} \)). During the whole decreasing luminosity phase \( n_{\text{ox}} \) is slightly increasing or constant. X-ray spectra taken during the high-luminosity phase and the decreasing luminosity phase would show an O viii absorption edge anticorrelated with the intensity of the ionizing continuum, as seen in MCG –6-30-15 (Fabian et al. 1994; Reynolds et al. 1995). At this density C vii is able to reach its maximum equilibrium value (Fig. 6b), but even this ion does not relax to its initial equilibrium value for many times up and down cycle time. Figure 6b also shows that the ions reach their maximum values at different times; \( n_{\text{C vii}} \) reach its maximum value about 4000 s before \( n_{\text{ox}} \).

At higher densities \( n_e = 10^9 \text{ cm}^{-3}, \text{dashed lines} \), the fractional abundance of each of the three ions reaches its maximum equilibrium value during the first 6000 s and relaxes to its minimum equilibrium value during the following \( 2.4 \times 10^4 \text{ s} \), but with different timescales: O ix reaches its minimum equilibrium value after a time corresponding to two cycles, about one cycle after C vii. This could help explain why models in which all ions of all elements are in photoionization equilibrium so often fail to describe AGN spectral evolution. Spectra accumulated immediately after a very steep decreasing intensity phase could contain no significant absorption features at the energies of O vii–O viii K edge (the oxygen being completely ionized) but still show a deep absorption edge at \( E \sim 0.5 \text{ keV} \), because of the presence of a large amount of recombinant C vii–C vi in the absorbing gas. High-quality spectra would allow powerful tests of nonequilibrium photoionization models.

A general result is that the observation of any delays in the response of the absorber to flux changes on timescales of \( \sim 5000–1000 \text{ s} \) immediately implies photoionization with a density in a reasonably restricted range, \( 10^8 \lesssim n_e \lesssim 10^9 \text{ cm}^{-3} \) (depending on the average ionization state).

3. Modeling the ROSAT Data of NGC 4051

As an example of the applicability of our models we present here the case of a ROSAT PSPC observation of the low-luminosity, rapidly variable Seyfert 1 galaxy NGC 4051.

Both the soft and hard X-ray flux of NGC 4051 vary by large factors (up to 20) on a timescale of hours (e.g., Lawrence et al. 1985; Guainazzi et al. 1996) in a roughly correlated way. The presence of an ionized absorber in this source was first proposed on the basis of variations of the Ginga softness ratio correlated with the flux (Fiore et al. 1992). ROSAT PSPC observations, and subsequently observations with the higher resolution CCDs on board the ASCA satellite (Pounds et al. 1994; McHardy et al. 1995; Mihara et al. 1994; Guainazzi et al. 1996), also suggested the presence of a high column density, highly ionized absorber through the detection of a deep absorption edge at 0.8–0.9 keV.

McHardy et al. (1995) find that a simple single-zone equilibrium photoionization model can provide a reasonably good representation of spectra accumulated in 1000–3000 s, but the best-fit ionization parameter does not track the source intensity, as required by the model. This behavior cannot be explained in a single-zone ionization equilibrium model, and therefore NGC 4051 is a good target to test our time-evolving photoionization and collisional ionization models. We decided to compare our models with the ROSAT PSPC data acquired on 1991 November 16 and reported by McHardy et al. (1995), when the source count rate showed large and rapid variations (up to a factor of 6 in a few thousand seconds; see Fig. 5 of McHardy et al. 1995). Analysis of the ASCA observation is deferred to a future publication.

The data reduction and the timing analysis were performed using the PROS package in IRAF. The observation spanned 77 ks and contained 28.7 ks of exposure time. NGC 4051 gave a mean count rate of 1.6 s⁻¹. We accumulated eight spectra (a–h), using a 3' radius extraction region, accumulating contiguous data with similar count rates. Background counts and spectra were accumulated from an annulus of internal and external radius of 3.5 and 6', respectively.

3.1. Hardness Ratio Analysis: The Absorber Is Not in Photoionization Equilibrium

Independent of any spectral fit the behavior of the main physical properties of the absorber can be seen in a color–color diagram. In Figure 7 we plot the hardness ratios \( HR = H/M \) against the softness ratio \( SR = S/M \) from the count rates in three bands \( S = 0.1–0.6 \text{ keV}, M = 0.9–1.5 \text{ keV}, \) and \( H = 1.7–2.5 \text{ keV} \) for theoretical curves [for log \( (N_{\text{H}}) = 22, 22.5 \) and 23] obtained by folding the equilibrium photoionization models [for log \( (U) \) in the range \( -0.3 \) to +1.5 and Galactic \( N_{\text{H}} = 3.13 \times 10^{20} \text{ cm}^{-2} \); Elvis, Wilkes, & Lockman 1989] with the response matrix of the PSPC. We also plot the colors of the source in the eight spectra (a–h). All the data points lie in a region of this diagram corresponding to the high-\( U \) ends of the photoionization theoretical curves where both SR and HR decrease.
as $U$ increases, until all the ions in the gas are fully stripped and the gas is completely transparent to radiation of any energy. The colors of NGC 4051 are all consistent with the O VIII–O IX ions being dominant. Filled circles on the log $N_H = 22.5$ curve mark values of $U$ in the range 4.0–7. All the data points are between the two extreme values of $U$, a factor of ~1.5 change, while the source intensity varies up to a factor of ~6 in the eight spectra. The gas is clearly not responding to the continuum variations, a conclusion equivalent to that obtained by McHardy et al. (1995) using spectral fits with equilibrium photoionization models (see their Fig. 5). In the framework of the models discussed in this paper this behavior suggests three different possibilities (we do not take into account pure collisional ionization in this case because the observed spectral variations should be attributed to ad hoc changes of $T_e$ or $N_H$ on timescales as short as 2000–4000 s):

1. The gas is far from ionization equilibrium;
2. The gas has a distribution of densities;
3. Both collisional and photoionization processes are comparably important in the same physical region.

We investigated these possibilities in turn using detailed spectral fits.

3.2. Time-evolving Photoionization

From Figure 1 we see that a given ionization state can be roughly determined by the measure of at least two consecutive ion abundances, e.g., O VII and O VIII. The measure of a single edge in fact would not distinguish between “low” and “high” ionization solutions. The same measured feature could be produced by a lower $N_H$, lower mean ionization gas, or by a higher $N_H$, higher mean ionization state gas.

The best derived quantities to compare observed spectra with our several physical models are atomic edge strengths. Here we are mainly interested in the O VII and O VIII because they are the strongest and therefore the easiest to detect and measure. However, with the PSPC, O VII and O VIII edges are not individually discernible and we must resort to model fits with multiple components. Our choice is to use the components that can ensure an estimate of the O VII and O VIII $\tau$ that is as unbiased as possible.

3.2.1. A Three-Edge “Model Independent” Spectral Fit

We fitted the eight spectra with a model consisting of a power law reduced at low energy by the Galactic column density, the O VII and O VIII edges and another edge at 1.36 keV to account for possible spectral complexity in the 1–2 keV region (in particular, the Ne–X K and Fe XV–XX Lp absorption discussed in §2.1). Five parameters were at first allowed to be free to vary: the spectral index $\alpha_E$, the O VII and O VIII edges $\tau$, the 1.36 keV edge $\tau$, and the model normalization. The results are presented in Table 1. The fits with the three-edge model are acceptable in all cases. We stress that $\tau$ (1.36 keV) in Table 1 should not be regarded as a true measure of the optical depth of Ne K and Fe L ions.

| Spectrum | $\tau$ (0.74 keV) | $\tau$ (0.87 keV) | $\tau$ (1.36 keV) | $\alpha_E$ | $\chi^2$ (dof) |
|----------|------------------|------------------|------------------|-------------|---------------|
| $a$       | 0.5±0.4          | 0.84±0.32        | 1.0±0.4          | 1.3±0.1     | 0.84 (21)     |
| $b$       | <0.5             | 0.66±0.21        | 0.7±0.3          | 1.3±0.1     | 1.19 (23)     |
| $c$       | 0.23±0.19        | 0.66±0.21        | 0.71±0.18        | 1.3±0.1     | 1.14 (24)     |
| $d$       | 0.38±0.33        | 0.41±0.31        | <0.6             | 1.3±0.1     | 0.85 (22)     |
| $e$       | 0.5±0.3          | 0.86±0.28        | 0.5±0.3          | 1.4±0.1     | 1.30 (22)     |
| $f$       | <0.6             | 1.01±0.24        | <0.5             | 1.4±0.1     | 1.41 (23)     |
| $g$       | 0.54±0.20        | 1.02±0.23        | 0.44±0.14        | 1.3±0.1     | 1.42 (24)     |
| $h$       | <0.5             | 1.47±0.29        | <0.5             | 1.3±0.1     | 0.65 (22)     |
|          | <0.18            | 1.42±0.16        | <0.21            | 1.3±0.1     | 0.67 (23)     |
| $i$       | 1.2±0.3          | 0.79±0.35        | 0.9±0.3          | 1.2±0.1     | 0.72 (22)     |
| $j$       | 0.63±0.23        | 0.86±0.33        | 0.45±0.19        | 1.3±0.1     | 0.82 (23)     |
This feature provides only one of the possible parameterizations of the spectrum in the 1–2 keV range, a band in which a change in the continuum spectral index could also be present (the 2–10 keV Ginga and ASCA spectra of this source are typically flatter by ≈0.5 than the PSPC 0.1–2 keV spectra). As discussed in § 2.1, it is difficult to discriminate between Ne absorption and a real spectral flattening above 1 keV with instruments of moderate spectral resolution like the PSPC. We have performed a series of fits using a broken power law with break energy in the 1–2 keV band and the two oxygen edges, obtaining τ similar to those reported in Table 1. We are therefore confident that the estimation of the oxygen edge τ is robust, within the rather large uncertainties given in Table 1. In principle, a way to reduce the uncertainties is to fix the continuum spectral index to a common value. The results of this series of fits are again given in Table 1. The uncertainties on τ (O VIII) and τ (1.36 keV) are indeed smaller than in the previous case, but this is not true for the τ (O VIII) uncertainties. The reason is that there is a strong anticorrelation between x_E and τ (O VIII). A similar anticorrelation is present between x_E and τ (1.36 keV). In contrast, no correlation is present between x_E and τ (O VIII). This is illustrated in Figure 8, where we show the χ² contours of these parameters for spectrum g. Since the τ (O VII) values could be biased in the fit with fixed x_E by the τ (O VIII)-x_E correlation, we prefer to compare our time-evolving models to the O VII and O VIII optical depths obtained leaving x_E free to vary.

In all but one case (spectrum d) the depth of the O VIII edge is higher than or comparable to that of O VII, suggesting a “high” ionization solution (U > 4; see Fig. 1), consistent with the hardness ratio analysis of Figure 7.

3.2.2. Comparison between Models and the Oxygen Edge Depths: Evidence for a Nonequilibrium Photoionization Absorber?

We converted the best-fit τ into τ (O VIII) and O VIII relative abundances assuming a solar oxygen abundance and a total hydrogen column density N_H. An indication of N_H comes from the color–color diagram of Figure 7. Although calculated using a photoionization equilibrium model, the theoretical curves in this diagram suggest a value for log N_H between 22 and 23, and so we adopt log N_H = 22.5. The three panels of Figure 9 show the light curves of the source count rate (upper panel) and of the O VIII and O VII abundances (middle and lower panels, respectively).

The time evolution of the ionization structure of a cloud of gas photoionized by a variable source is complex, and its behavior sometimes counterintuitive. We then discuss first the simplest case: high electron density, for which each ion is close to its equilibrium state. We examine next the case of lower densities and hence nonequilibrium solutions.

The dotted curves on the middle and lower panels of Figure 9 show the n_e = 10¹⁶ cm⁻³, P = 1 (the ratio of the incident flux to that needed to produce the initial ionization distribution assuming photoionization equilibrium) curves, when the gas is close to equilibrium. While the equilibrium O VII curve tracks the count rate variations (it is strictly anticorrelated with the count rate light curve), the O VIII curve does not. The different O VIII behavior is due to the different balance in the destruction rates of O IX and O VIII. When the ionizing flux is at its maximum most of the oxygen is O IX. When the flux decreases from the maximum (from point b to d) O IX recombines to O VIII, increasing the O VIII abundance. When the flux decreases from point e to point f, at first O VIII increases again because of the high destruction rate of O IX, but after a certain point the amount of O VIII recombining to O VII starts to be higher than the amount of O IX recombining to O VIII, and so the total O VIII abundance starts to decrease. Instead, the amount of O VIII recombining to O VII is always higher than the amount of O VII recombining to O VI. This gives rise to a different behavior of the O VIII and O VII curves in response to the same ionizing flux variations. It is interesting to note that in this case, while the dynamical range of variation of the O VII curve is larger than that of the ionizing flux, the O VIII equilibrium curve shows a more compressed range of
Fig. 9.—Light curve of NGC 4051 during the 1991 November ROSAT-PSPC observation (upper panel). Best-fit \( n_{\text{O VIII}} \) and \( n_{\text{O VII}} \) abundances (points with error bars), obtained by fitting a power-law continuum plus three-edge model to the eight spectra and by using (middle and lower panels). 

\[ \log(\frac{N}{H}) > 22.5 \]

Solid lines in the two panels represent the best-fit nonequilibrium photoionization model to the data, respectively, while dashed lines are the solutions obtained using the 1 \( \sigma \) confidence interval on \( n_e \) and \( P \). Dotted lines are the equilibrium \( n_{\text{O VIII}} \) and \( n_{\text{O VII}} \) curves and were obtained using the nonequilibrium photoionization code by fixing \( n_e = 10^{15} \text{ cm}^{-3} \) and \( P = 1 \).

Variations (when the ionizing flux varies by a factor of 6) the O VIII and O VII abundances vary by a factor of 3 and 30, respectively; see Fig. 9). Therefore, variations of O VIII would be much more difficult to detect than variations of O VII, at least at these regimes of ionization. We reemphasize that the behavior of a single edge does not provide a unique interpretation of the data.

As explained in § 2.3.3, a grid of theoretical O VII and O VIII light curves was generated using our time-evolving photoionization code for 28 values of \( n_e \) from \( 5 \times 10^6 \) to \( 10^9 \text{ cm}^{-3} \) and 15 values of \( P \) from 0.5 to 2. We compared these curves with the measured relative abundances of O VII and O VIII and found the best-fit model using a \( \chi^2 \) technique. The thick solid lines in the lower and middle panels of Figure 9 represent the best-fit nonequilibrium O VII and O VIII curves. The agreement between the best-fit model and the observed O VII and O VIII abundances is good: \( \chi^2 = 1.17 \) for 14 dof. The best-fitting values for \( n_e \) and \( P \) are tightly constrained: \( n_e = (1.0^{+1.2}_{-0.7}) \times 10^8 \text{ cm}^{-3}, P = 1.5^{+0.4}_{-0.3} \).

From the best-fit \( n_e \) we can estimate the distance of the absorbing cloud from the central source. We obtain \( R = (0.74^{+0.40}_{-0.40}) \times 10^{16} \text{ cm} \) (3 light-days).

The dashed lines represent the solutions obtained using the 1 \( \sigma \) confidence interval on \( n_e \) and \( P \). The best-fit curves (and the 1 \( \sigma \) confidence intervals curves) show a compressed dynamical range of O VIII and O VII abundance variations and a delay between the source maxima and the ion abundance minima of 3000–6000 s. The compressed dynamical range is due to a mean overionization of the gas. While the best-fit \( P \) shows that the gas in the initial point \( a \) is near to equilibrium, it departs strongly from equilibrium during the later low-intensity states (spectra \( d-h \)). So, despite the fact that the source spends more time in low states than in high states, the gas density is sufficiently low that the gas does not have time to fully recombine after the few events when it suffers high illumination and becomes highly ionized.

The above results were obtained assuming a total hydrogen column density of \( \log(\frac{N}{H}) = 22.5 \). Assuming a higher (lower) column would imply a mean lower (higher) O VII and O VIII relative ion abundance. Therefore, in principle, the accurate measure of both edges would also constrain the total warm column density. The uncertainties on the PSPC determinations, however, preclude this possibility, and better resolution measurements are therefore needed. The energy resolution of the ASCA SIS, for example, is just sufficient to separate the O VII and O VIII absorption edges.

A quantitative test of the nonequilibrium photoionization model using the ASCA data and a comparison between ROSAT PSPC and ASCA data of NGC 4051 is beyond the scope of this paper and will be presented in a future publication. A much better separation, and therefore characterization, of the absorption features will be possible with the
high-resolution (factor of 10–30 better than ASCA SIS) gratings on AXAF and XMM.

3.3. Other Models

Despite the success of time-evolving photoionization models, alternatives do exist. We discuss two of them in the following.

3.3.1. Large Density Variations in the Absorbing Gas

If the gas is not confined to a single cloud with constant density but rather is distributed in a region with, say, an increasing density, then different ionization equilibria could apply to different regions in the cloud. Two extreme regions may exist: in the region with higher density, lower \( U \), collisional ionization will be the dominant ionization mechanism, and the spectra transmitted by this region would always show the same features, irrespective of the source intensity; the other region, with lower density and higher \( U \), is completely ionized (for carbon, oxygen, neon, and iron ions up to Fe \( \text{XXII} \)) when the source is in the high state, but when the source is in a low state the abundances of O \( \text{VIII} \) are sizeable and imprint the edges in the spectrum seen in low-intensity spectra. Here we are assuming that the density of the photoionized region is high enough for the gas to be in instantaneous equilibrium with the ionizing intensity. If not, the average ionization degree of the gas would be very high during the whole observation and the gas would be always almost transparent at the energies of the relevant absorption edges.

We tested this hypothesis by fitting the highest intensity spectrum (b) with a simple power-law model plus a collisionally ionized absorber, using the method of Fiore et al. (1993). The best-fit temperature and \( N_{\text{HI}} \) are \( 2.8 \pm 0.1 \times 10^6 \) K and \( 1.5 \pm 0.2 \times 10^{22} \text{ cm}^{-2} \), respectively. We then used the same model (with fixed continuum parameters, fixed temperature, and fixed \( N_{\text{HI}} \), but with variable normalization) with the inclusion of an additional O \( \text{VIII} \) edge to mimic a variable ionization state, equilibrium photoionization absorber. The fits are all acceptable. The worst fit is that of spectrum (d) (\( \chi^2 = 1.4 \), 21 dof, probability of 10.4%). In Figure 10 we plot \( \tau (\text{O VIII}) \) as a function of time. The dynamic range of variations of \( \tau \) is here larger than that on \( n_{\text{O VIII}} \) in Figure 9 but is still more compressed than that predicted by the equilibrium photoionization model (solid line). We can therefore exclude the possibility that a major part of this absorber could be in pure photoionization equilibrium with the ionizing intensity.

3.4. A “Hot” Photoionized Absorber

The other possibility is that both collisional and photoionization processes are important in the same physical region. In this case the transmitted spectra are very similar to those transmitted from purely photoionized clouds of gas with a much higher ionization parameter (§ 2.1; Fig. 4). The electron temperature of the gas is mainly determined by collisions and is higher than that expected in pure photoionization equilibrium. The ionization parameter \( U \) is no longer linearly correlated with the ionizing intensity, and then its dynamical range of variations is compressed by a factor of less than 2 compared with the pure photoionization case.

This could, in principle, explain the compression observed in the measured dynamical range of variation of \( n_{\text{O VIII}} \) but could not account for the delays observed on the response of the ionization state of the gas to source intensity variation. Unfortunately, there is no way to distinguish between “hot” photoionized absorber models and simple pure photoionization models only on the basis of the spectral analysis. Nevertheless, the delay observed in the response of the ionization degree of the gas between spectra b and c suggests that a nonequilibrium photoionization component is preferred.

4. CONCLUSION

We have investigated ionization models for AGNs in different regimes of gas volume densities and photoionization states. In particular, we focused on “high gas density, low photoionization parameter” gas clouds, where collisional ionization is likely to play a significant role in the gas ionization, and on low gas densities, where the photoionization may be far from equilibrium.

We presented detailed model calculations in both regimes. While the time-evolving photoionization models in § 2.3 are far from being complete or exhaustive, they are nevertheless instructive and reveal the main features of these kinds of models.

Our main findings can be summarized as follows:

1. In many important astrophysical conditions (O \( \text{VII} \), O \( \text{VIII} \) regime) the fractional abundances of the most important ions of O and Ne in photoionized gas are more broadly distributed in \( U \) than those of collisionally ionized gas are in \( T \).

2. In the collisional ionization case the heavy elements show a strong inertia against becoming highly ionized, even when lighter elements, like oxygen, are almost fully stripped. In this case the transmitted spectrum shows a large and complicated absorption structure between 1 and 3 keV, mainly due to iron L \( \text{XVII} \text{–XIX} \) and neon K \( \text{IX} \text{–X} \) absorption, which is much less visible in spectra emerging from photoionized gas with similar O \( \text{VII} \) and O \( \text{VIII} \) abundances. This absorption structure may be misinterpreted as a flattening of the spectra above \( \sim 1 \text{ keV} \), when fitting low energy resolution data with a photoionization equilibrium.
model. Higher spectral resolution and good S/N observations are therefore needed to distinguish between collisional ionization and photoionization.

3. In nonequilibrium photoionization models the response of the ionization state of the gas to sudden changes of the ionizing continuum is delayed even during increasing luminosity phases. The delays increase for decreasing electron densities, as changes of \( n_e \) require changes of the intensity of the ionizing flux, i.e., changes of the distance of the gas from the X-ray source (taking as fixed the initial ionization state of the gas).

4. The recombination timescale is generally much longer (up to orders of magnitudes) than the photoionization timescale, because of the dependence of \( t_{\text{rec}} = \frac{1}{S_{\lambda} n_e^{q}}(t \rightarrow t + dt) \) on the equilibrium ratio \( (n_{e+1}/n_e) \), evaluated at the time \( t + dt \). This means that a photoionized absorber undergoing frequent, quick, and consistent changes of ionizing luminosity is likely to be overionized with respect to the equilibrium ionization, a state that would be reached only after a sufficiently long low-intensity phase.

5. If the changes of the ionizing luminosity are not instantaneous and the electron density of the cloud is low enough, the ionization state of the gas could continue to increase during decreasing source luminosity phases. This means that we may measure a maximum in ionization state when the ionizing flux is at a minimum (opposite to what is expected in equilibrium models).

6. Different ions of different elements reach their equilibrium abundance on different timescales. This is again (opposite to what is expected in equilibrium models). This means that we may measure a maximum in ionization state during decreasing source luminosity phases. This is enough, the ionization state of the gas could continue to increase during decreasing source luminosity phases. This means that a photoionized absorber undergoing frequent, quick, and consistent changes of ionizing luminosity is likely to be overionized with respect to the equilibrium ionization, a state that would be reached only after a sufficiently long low-intensity phase.

We have tested the above models in the case of the Seyfert 1 galaxy NGC 4051. The ROSAT observations of NGC 4051 are not consistent with a simple equilibrium model but can be explained straightforwardly by our time-evolving photoionization models. The two main features in the nonequilibrium best-fit models are (1) the compressed range of variability of the measured O VII and O VIII relative abundances with respect to the amplitude of the source variations and to the amplitude of the variations of the abundances of these ions expected in equilibrium photoionization models; (2) the 3000–6000 s delay between the maximum intensity state of the source (spectrum b) and the minimum of the best-fit O VIII abundance curve, i.e., the maximum ionization state of the gas (spectrum c). As a result we were able to estimate the gas electron density, \( n_e = (1.0 \times 10^{9} cm^{-3}) \) (assuming log \( N_{\text{H}} = 22.5 \)) and hence the distance of the ionized gas cloud from the X-ray source in \( R = (0.74 \times 10^{16} \text{ cm}) \) (3 light-days). We explored alternative models, and we also explored alternatives that we find to be less likely; we discuss ways to distinguish between them conclusively.

We conclude that nonequilibrium photoionization and collisional models apply to wide zones of gas density and ionization, zones that are expected in AGN. These effects must be considered in understanding ionized absorbers, and seem likely to explain otherwise puzzling behavior, without resorting to ad hoc distributions of gas. Several clear diagnostics of these models exist so that decisive tests will soon be possible.

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