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# Tree-based solvers for adaptive mesh refinement code FLASH – IV. An X-ray radiation scheme to couple discrete and diffuse X-ray emission sources to the thermochemistry of the interstellar medium

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## ABSTRACT

X-ray radiation, in particular radiation between 0.1 and 10 keV, is evident from both point-like sources, such as compact objects and T-Tauri young stellar objects, and extended emission from hot, cooling gas, such as in supernova remnants. The X-ray radiation is absorbed by nearby gas, providing a source of both heating and ionization. While protoplanetary chemistry models now often include X-ray emission from the central young stellar object, simulations of star-forming regions have yet to include X-ray emission coupled to the chemo-dynamical evolution of the gas. We present an extension of the TREERAY reverse ray trace algorithm implemented in the FLASH magnetohydrodynamic code which enables the inclusion of X-ray radiation from  $0.1 \text{ keV} < E_\gamma < 100 \text{ keV}$ , dubbed XRAYTHESPOT. XRAYTHESPOT allows for the use of an arbitrary number of bins, minimum and maximum energies, and both temperature-independent and temperature-dependent user-defined cross-sections, along with the ability to include both point and extended diffuse emission and is coupled to the thermochemical evolution. We demonstrate the method with several multibin benchmarks testing the radiation transfer solution and coupling to the thermochemistry. Finally, we show two example star formation science cases for this module: X-ray emission from protostellar accretion irradiating an accretion disc and simulations of molecular clouds with active chemistry, radiation pressure, and protostellar radiation feedback from infrared to X-ray radiation.

**Key words:** astrochemistry – radiative transfer – methods: numerical – ISM: clouds – X-rays: general – X-rays: ISM.

## 1 INTRODUCTION

Molecular gas is subjected to radiation across the electromagnetic spectrum. Hard radiation, such as X-ray and gamma-ray radiation, can penetrate deep into molecular gas and drive the thermochemistry of dense gas (Spitzer & Tomasko 1968; Maloney, Hollenbach & Tielens 1996; Yan 1997; Wolfire, Vallini & Chevance 2022). X-rays provide an important source of ionization in dense gas, driving the ion-neutral chemistry and providing heating through photoelectrons (Lepp & Shull 1983; Maloney et al. 1996; Dalgarno, Yan & Liu 1999). Using the typical molecular gas photoabsorption cross-sections (Maloney et al. 1996), the  $\tau = 1$  surface for 1 keV photons is approximately  $4 \times 10^{21} \text{ cm}^{-2}$  [compared to  $\approx 10^{-18} \text{ cm}^{-2}$  for ultraviolet (UV) radiation]. However, the photoabsorption cross-sections scale roughly as  $E^{-2.5}$  (Mackey et al. 2019), so harder radiation penetrates much further into the cloud. Therefore, in regions near bright X-ray sources, the cloud structure can become

dominated throughout by the X-ray radiation. Regions in which the thermochemistry is regulated primarily through X-ray radiation are often denoted as X-ray dominated regions (XDRs; coined by Maloney et al. 1996), in analogue to photodissociation regions.

X-ray radiation drives ionization primarily through secondary, induced processes. While the primary ionization cross-sections are low, the resulting ejected fast electrons can produce a cascade of secondary ionizations and pumped far ultraviolet (FUV) radiation through the excitation and subsequent de-excitation of H and H<sub>2</sub> (Prasad & Tarafdar 1983; Dalgarno et al. 1999; Meijerink & Spaans 2005). These fast electrons can also provide heating through photoelectric heating of dust grains. In this way, X-ray radiation acts in a very similar manner as cosmic rays, and untangling the two contributions can be difficult (see e.g. Meijerink, Spaans & Israel 2006). However, due to the significantly larger interaction cross-sections (and the strong energy and temperature dependence), the effect of X-ray radiation on the heating and ionization fraction are still noticeably different (Meijerink et al. 2006). There have been a plethora of investigations on the impact of cosmic rays on interstellar gas (e.g. Dalgarno 2006; Ceccarelli 2011; Indriolo & McCall 2013; Bisbas, Papadopoulos & Viti 2015; Gaches, Offner & Bisbas 2019;

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Bisbas et al. 2023) showing that they play a crucial role in the chemistry of the interstellar medium. Due to the ubiquity of the inclusion of cosmic rays, it is relatively common to treat the chemical impact of X-rays through enhancing the cosmic ray ionization rate, although some studies separate the heating treatments (e.g. Harada, Herbst & Wakelam 2010; Viti et al. 2014; Walsh, Nomura & van Dishoeck 2015; Viti 2017; Wang et al. 2021).

Molecular clouds are immersed in a bath of X-ray radiation, with contributions from both external and internal sources. Externally, molecular gas can be irradiated through supernovae and their remnants (e.g. Yamane et al. 2018; Brose, Sushch & Mackey 2022), X-ray binaries (e.g. White, Stella & Parmar 1988; Remillard & McClintock 2006; Reig 2011; Mineo, Gilfanov & Sunyaev 2012; Lutovinov et al. 2013; Giacobbo, Mapelli & Spera 2018), nearby active galactic nuclei (e.g. Sunyaev, Markevitch & Pavlinsky 1993; Sunyaev & Churazov 1998; Harada, Thompson & Herbst 2013; Churazov et al. 2017; Mingozi et al. 2018; Cruz-González et al. 2020). Internally, young stellar objects, including embedded accreting protostars and more evolved T-Tauri stars (Calvet & Gullbring 1998; Feigelson & Montmerle 1999; Feigelson et al. 2007), and high-mass stars just reaching the main sequence can become X-ray bright (e.g. Cassinelli et al. 1994), whether through accretion or magnetic powered radiation, or coronal emission. Finally, gas heated through feedback processes, such as winds and supernovae, can become warm enough to emit X-ray radiation while they cool (Raymond & Smith 1977). Observational X-ray surveys of molecular gas and star-forming regions show substantial amounts of diffuse emission and a sizable number of point sources (Sunyaev et al. 1993; Feigelson et al. 2013; Townsley et al. 2014, 2019).

Despite their potential importance, their inclusion into simulations of molecular clouds has been sparse. There has been substantial focus on thermochemical models of protoplanetary discs (e.g. Glassgold, Najita & Igea 1997; Igea & Glassgold 1999; Ercolano et al. 2008a; Ercolano, Clarke & Drake 2009; Owen, Ercolano & Clarke 2011; Meijerink, Cazaux & Spaans 2012; Cleeves et al. 2017; Picogna et al. 2019; Waggoner & Cleeves 2019) and models of molecular gas near external sources or compact objects (e.g. Krolik & Kallman 1983; Lepp & McCray 1983; Draine & Woods 1991; García-Burillo et al. 2010; Hocuk & Spaans 2010; Meijerink et al. 2011; Odaka et al. 2011; Orlando et al. 2011; Mackey et al. 2019). These methods typically utilize Monte Carlo methods (Ercolano et al. 2008a; Odaka et al. 2011; Molaro, Khatri & Sunyaev 2016; Walls et al. 2016; Cleeves et al. 2017), or ray trace schemes and focus primarily either on the inclusion of point sources or external radiation fields (e.g. Wise & Abel 2011; Mackey et al. 2019; Khabibullin et al. 2020).

In this paper, we will present an X-ray extension of the reverse ray tracing scheme TREERAY (Wünsch et al. 2021), which allows for the inclusion of an arbitrary number of point sources and diffuse radiation. The module is a TREERAY extension of the diffuse X-ray module presented in Mackey et al. (2019), which enabled diffuse X-ray irradiation at the domain boundary. Our implementation enables up to 100 energy bins at arbitrary locations between 0.1 and 100 keV and temperature-dependent photoabsorption cross-sections. In Section 2, we give an overview of the X-ray TREERAY algorithm, called XRAYTHESPOT, and the coupling of it to the X-ray-driven chemistry. In Section 3, we show the performance of the module with different radiation transfer tests and a benchmark against the CLOUDY code. In Sections 4 and 5, we demonstrate the use of this module for protostellar emission irradiating a surrounding disc and in a star-formation simulation, respectively. Finally, in

Section 6, we discuss the future extensions and scientific applications of XRAYTHESPOT.

## 2 METHODS

Our new XRAYTHESPOT module is able to treat radiation from 0.1 to 100 keV. We describe below in detail the adopted photoabsorption cross-sections and the module's implementation within TREERAY.

### 2.1 X-ray cross-sections

X-ray radiation is attenuated as it propagates through gas via a combination of photoionization, at lower energies, and the Compton process, at higher energies. The previous module, described in Mackey et al. (2019), used the low-energy approximation for the X-ray cross-section,  $\sigma_x$ , from Panoglou et al. (2012):

$$\sigma_x = 2.27 \times 10^{-22} E_\gamma^{-2.485} \text{ cm}^2 \quad (1)$$

per H-nucleus, where  $E_\gamma$  is the photon energy. However, this cross-section is valid only for cold, neutral gas, and for solar metallicity. We include now, as input during run time, temperature-dependent cross-sections, which can be re-computed for problems with different metallicities. Decreasing the metallicity will primarily impact X-rays between  $300 \text{ eV} \leq E \leq 10 \text{ keV}$  due to the decrease in contributions from various metals. A benchmark comparison for different metallicities are presented below. For photoionization, we use the analytical fits from Verner & Yakovlev (1995),  $\sigma_{\text{pi}}$ , where

$$\sigma_{\text{pi}} = \sigma_0 F(E_\gamma/E_0), \quad (2)$$

where

$$F(y) = [(y-1)^2 + y_w^2] y^{-Q} \left(1 + \sqrt{y/y_a}\right)^{-P}, \quad (3)$$

$y = E_\gamma/E_0$ ,  $Q = 5.5 + l - 0.5P$ , and  $l = 0, 1, 2$  is the subshell orbital quantum number, and  $\sigma_0$ ,  $E_0$ ,  $y_w$ ,  $y_a$ , and  $P$  are fit parameters from the associated public ViZieR catalogue.<sup>1</sup> However, to utilize these cross-sections, the ionization level populations must be known. We assume collisional ionization equilibrium and use the CHIANTI package (Dere 2013), using version 9 of the CHIANTI atomic data base (Dere et al. 1997, 2019) to compute the ionization fraction as a function of temperature.

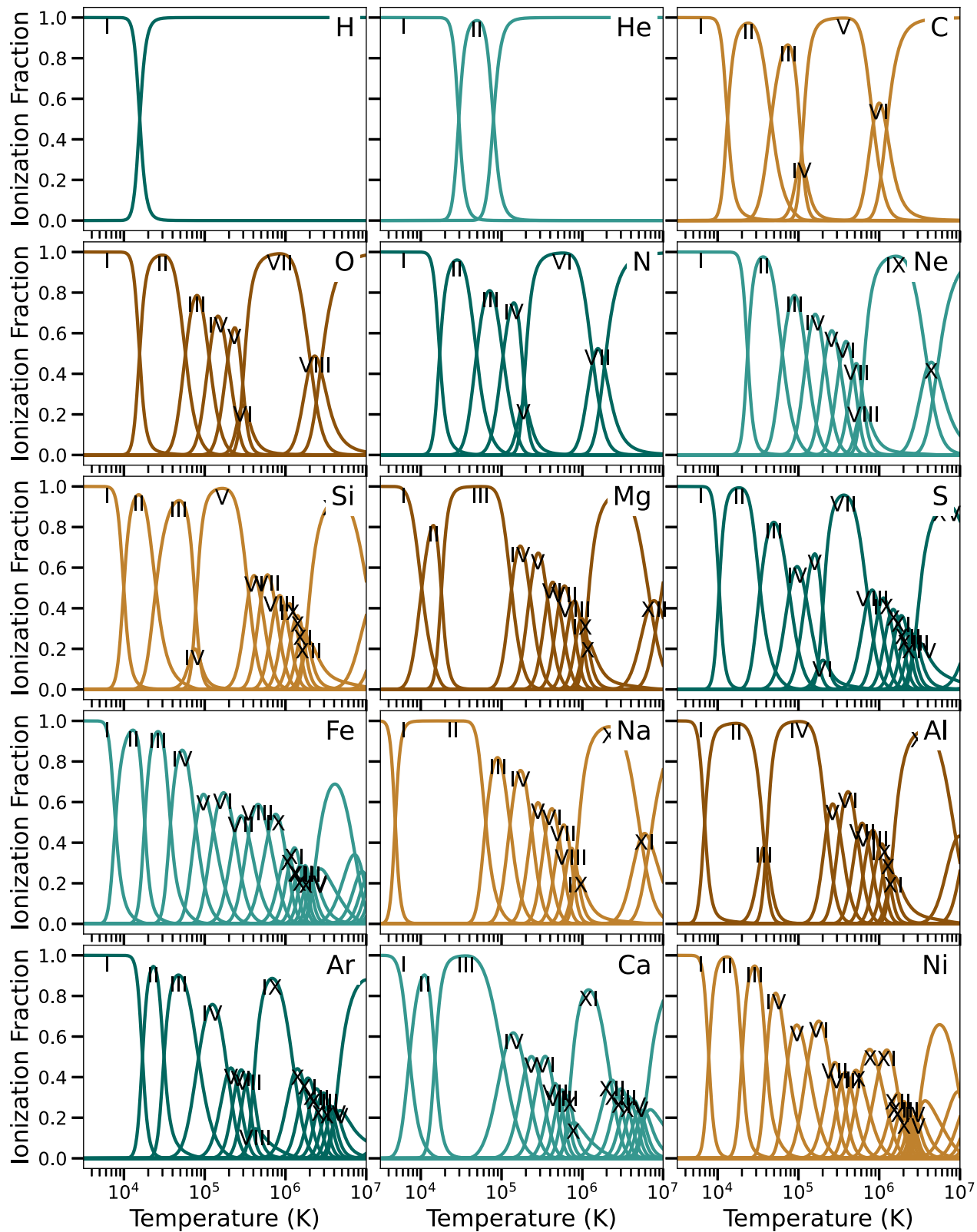
Fig. 1 shows the equilibrium ionization fractions as a function of temperature for the 15 different elements (see Table 1) we include in the cross-sections. These computations show that, particularly for  $T > 10^5 \text{ K}$ , there are multiple ionization states for metals which contribute to the photoionization cross-section.

We also include the cross-section for the Compton effect, which becomes particularly important at higher energies. We use the total Klein–Nishina (K–N) cross-section (Klein & Nishina 1929; Longair 2011),  $\sigma_{\text{KN}}$ ,

$$\sigma_{\text{KN}} = \pi r_e^2 x^{-1} \left\{ \left[ 1 - \frac{2(x+1)}{x^2} \right] \ln(2x+1) + \frac{1}{2} + \frac{4}{x} - \frac{1}{2(2x+1)^2} \right\}, \quad (4)$$

where  $r_e$  is the classical electron radius and  $x = E_\gamma/(m_e c^2)$ . While most applications will be in the limit of Thomson scattering, we include the full Compton cross-sections to enable more flexibility in the choice of energy bins. In the cross-section plots, we show the

<sup>1</sup><https://cdsarc.cds.unistra.fr/viz-bin/cat/J/A+AS/109/125>



**Figure 1.** Ionization fraction for different elements as a function of temperature. Annotated in the text are the peaks of different ionization levels for each element.

**Table 1.** Elements included in our photoabsorption cross-section calculation and their fiducial abundances,  $A_X$ , reported as  $A_X = \log(N_X/N_H) + 12$  (Asplund et al. 2009).

| Element | Abundance ( $A_X$ ) |
|---------|---------------------|
| H       | 12                  |
| He      | 10.986              |
| C       | 8.443               |
| O       | 8.783               |
| N       | 7.913               |
| Ne      | 8.103               |
| Na      | 6.353               |
| Mg      | 7.593               |
| Al      | 6.523               |
| Si      | 7.573               |
| S       | 7.193               |
| Ar      | 6.553               |
| Ca      | 6.383               |
| Fe      | 7.503               |
| Ni      | 6.283               |

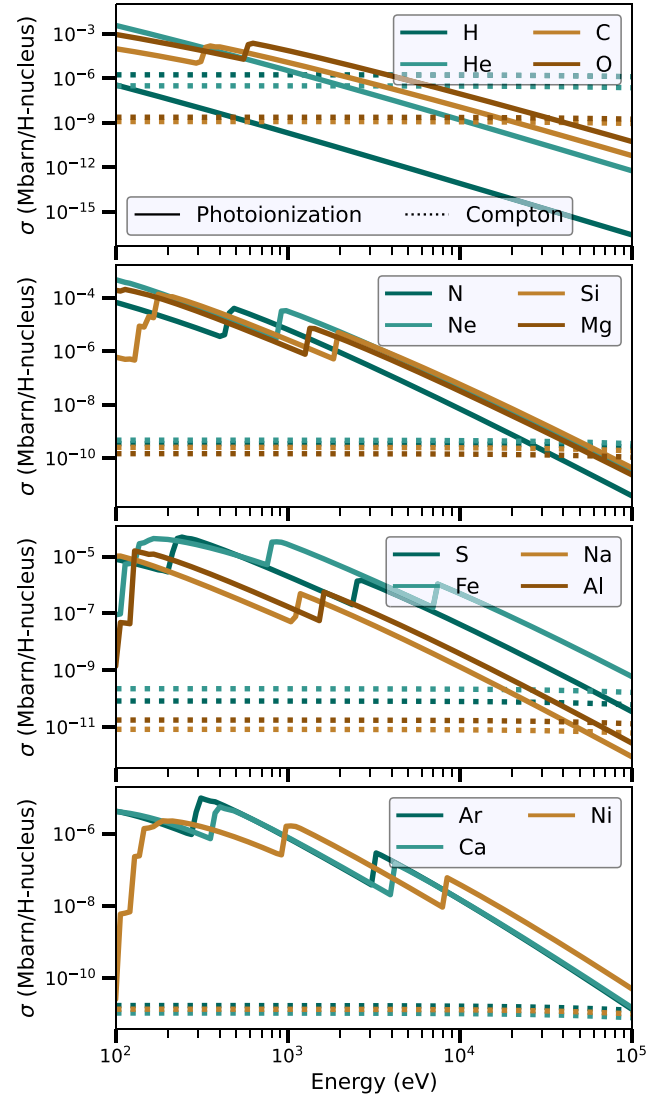
total Compton cross-section weighted by the number of free electrons contributed by each species. The total cross-section,  $\sigma_x$ , is thus

$$\sigma_x(E) = \sum_i^{N_{\text{elem}}} x_i \sigma_{\text{pi},i}(E) + \sigma_{\text{KN},i}(E), \quad (5)$$

where  $x_i$  is the abundance of element  $i$  with respect to hydrogen and the sum is carried out including the cross-sections for the  $N_{\text{elem}}$  elements. Table 1 shows the elements we include in the photoabsorption cross-section and their fiducial abundances relative to hydrogen.

Fig. 2 shows the photoionization cross-sections and free-electron contributions to the Compton cross-section as a function of energy for gas with  $T = 10^5$  K. As shown, for some elements, the Compton cross-section becomes more important than photoionization, in particular for hydrogen and helium above 1 keV, and for carbon and oxygen above 30 keV. For hydrogen, the Compton effect is dominant due to the negligible neutral fraction at  $T = 10^5$  K. Fig. 3 shows the total cross-section as a function of energy for  $T = 10^5$  K and each of the total elemental contributions. Here, the X-ray photoabsorption cross-section is dominated by helium (<0.3 keV), then carbon (0.3–0.6 keV), and oxygen (0.8–4 keV). At energies above 4 keV, the hydrogen and helium Compton cross-sections dominate with a contribution from the iron photoionization cross-section around 10 keV. However, many of the metals contribute equally to the total cross-section around 1 keV.

Finally, Fig. 4 shows  $\sigma_x$  as a function of energy and temperature from  $T = 10^4$  to  $10^7$  K. At low temperatures, we recover the analytical cross-section previously used, although around 10 keV there is an increase in the cross-section due to iron. However, the rather significant temperature dependence highlights the necessity of including a temperature-dependent cross-section: *at high temperatures, the higher thermal ionization state leads to a reduction of nearly two orders of magnitude in the cross-section*, thereby making the gas significantly more optically thin to the X-ray radiation, producing less heating and enabling more X-rays to escape. Below  $10^4$  K, the cross-section does not noticeably change since hydrogen is not significantly collisional ionized. Therefore, for the results of this paper, for colder gas, we use the  $T = 10^4$  K photoabsorption cross-section. The module though allows for the user to define their own temperature-dependent cross-sections across any temperature range.



**Figure 2.** Photoionization (solid) and Compton process (dashed) cross-sections for each element as a function of energy, assuming thermal ionization equilibrium at  $T = 10^5$  K. Each elemental contribution is weighted by the assumed abundance with respect to hydrogen.

For a given set of energy bins,  $\{(E_{l,i}, E_{r,i})\}$ , where  $i = 1$ , for  $N_{\text{bin}}$  bins, we define

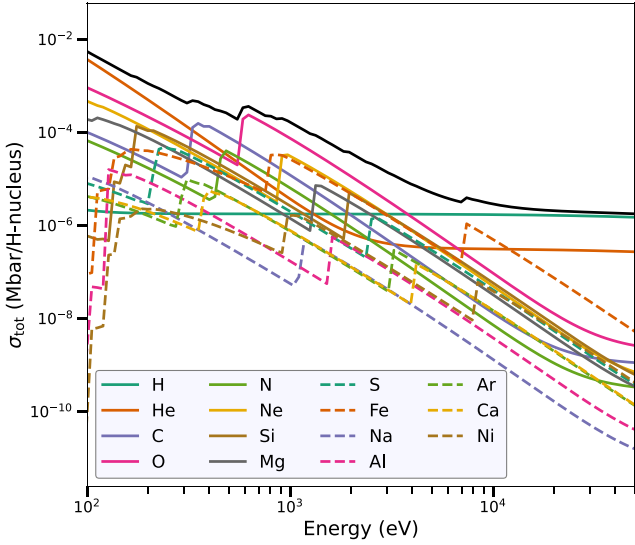
$$E_{c,i} = \frac{1}{2} (E_{l,i} + E_{r,i}), \quad (6)$$

where  $E_{l,i}$  is the left bound of the  $i$ th bin,  $E_{r,i}$  is the right bound, and  $E_{c,i}$  is the mid-point of the bin. We derive bin-averaged cross-sections, such that

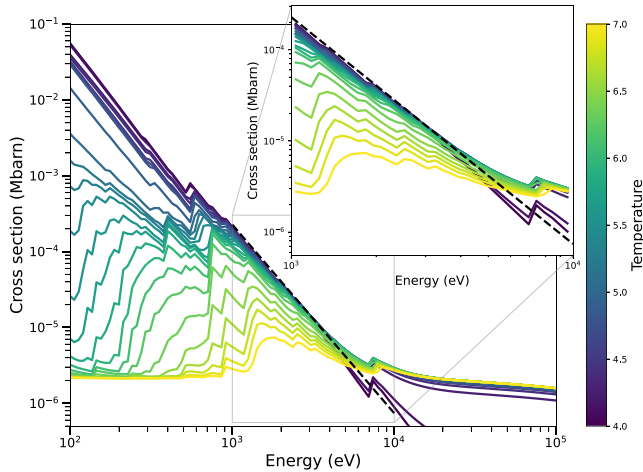
$$\exp\left(-\frac{\langle\sigma_{x,i}\rangle}{\sigma_c}\right) = \frac{1}{E_{l,i} - E_{r,i}} \int_{E_{l,i}}^{E_{r,i}} \exp\left(-\frac{\sigma_x(E_\gamma)}{\sigma_c}\right) dE, \quad (7)$$

where  $\langle\sigma_{x,i}\rangle$  is the bin-averaged cross-section for bin,  $i$ , and  $\sigma_c = \sigma_x(E_{c,i})$ . Our fiducial tests use  $N_{\text{bin}} = 8$  between 1 and 10 keV using logarithmically spaced bins.

All of these cross-section data, and the initialization and storage of the bins, and bin-averaged cross-sections are kept in a new FLASH module, XRAYCOMMON. FLASH is a highly module public magnetohydrodynamic code (MHD; Fryxell et al. 2000) written in FORTRAN and highly scalable with MPI. The scripts necessary to



**Figure 3.** Total photoabsorption cross-section (black) with each element contribution highlighted (colours) as a function of energy for gas at temperature,  $T = 10^5$  K. Each elemental contribution is weighted by the assumed abundance with respect to hydrogen.



**Figure 4.** Total photoabsorption cross-section as a function of energy and temperature (colour). The black dashed line shows the previously used analytical cross-section from Panoglou et al. (2012). Inset: Zoom-in to 1–10 keV.

compute the X-ray cross-sections are publicly available on GITHUB.<sup>2</sup> This module enables the coupling of X-ray physics to multiple other modules. Plasma models and the necessary X-ray data are also stored within this module, as a unified location.

## 2.2 TREERAY

TREERAY is a novel reverse ray tracing scheme, described fully in Wünsch et al. (2021), implemented in FLASH. Simply, TREERAY enables an efficient method to compute the contributions of radiation from every cell, for every cell. It does so through the combination of a reverse ray trace algorithm with a tree (Wünsch et al. 2018), which also currently is used in the gravity solver. Below, we describe briefly

<sup>2</sup><https://github.com/AstroBrandt/XRayCrossSections>

the different aspects of the TREERAY algorithm and XRAYTHESPOT extension and refer the reader to Wünsch et al. (2021) for more details.

### 2.2.1 Building the tree

The foundation of the TREERAY algorithm is an octree which stores all necessary variables for the various TREERAY modules. At minimum, the tree stores the mass and centre of mass coordinates for the respective cell, or leaf, or higher nodes. For XRAYTHESPOT, two further quantities are stored on to the tree: the bin-integrated X-ray luminosity in each energy bin and the gas temperature. While the bin-integrated X-ray luminosity is purely additive, the temperature is stored as a mass-weighted average of each set of eight subnodes (or leaves).

### 2.2.2 Ray structure

Before the tree walk is executed for a given cell, rays are generated by casting  $N_{\text{pix}} = 12N_{\text{side}}^2$  rays from each cell using directions defined by the HEALPIX (Górski et al. 2005) algorithm. HEALPIX tessellates the unit sphere into areas representing equal solid angles with a unit vector pointing to the centre of each of these surface areas from the sphere’s centre. TREERAY allows for  $N_{\text{side}} = 1, 2, 4, 8, \dots$ , with higher values representing higher angular resolution. The rays are split into  $N_r$  evaluation points, set by the grid resolution,  $\Delta x$ , the allowed length of the ray,  $L_{\text{ray}}$ , which is set to three-dimensional diagonal of the computational domain, and a free parameter,  $\eta_R$ .

Along each ray, the radial coordinate point of the  $i$ th evaluation point is

$$r_i = \frac{\Delta x i^2}{2\eta_R^2}, \quad (8)$$

leading to segments with increasing lengths. This behaviour coincides well with the geometric acceptance criterion described below for deciding whether or not to accept a tree node. The total number of evaluation points is

$$N_R = \eta_R \times \text{floor} \left( \sqrt{\frac{2L_{\text{ray}}}{\Delta X}} \right) + 1. \quad (9)$$

### 2.2.3 Tree walk

The mapping of the cells/nodes on to the rays requires two factors: a multipole acceptance criterion (MAC) and a weighting function to map from the tree on to the different radial evaluation points. When the MAC is met, a node is accepted and used. The simplest MAC is the Barnes–Hut geometric MAC (Barnes & Hut 1986), where a node of size  $h_n$ , at a distance  $d$ , from the cell is opened if

$$h_n/d < \theta_{\text{lim}}, \quad (10)$$

where  $\theta_{\text{lim}}$  is a user-defined opening angle with a sensible choice being  $\theta_{\text{lim}} = \sqrt{4\pi/N_{\text{pix}}}$ .<sup>3</sup> We also utilize the ‘Src MAC’ (Wünsch et al. 2021), where a node with sources is opened if

$$h_n/d < \theta_{\text{src}}, \quad (11)$$

<sup>3</sup>The resulting  $\theta_{\text{lim}}$  for  $N_{\text{side}} = 1, 2, 4$ , and  $8$  is  $1.0, 0.5, 0.25$ , and  $0.125$ , respectively. For the results of this paper, we adopted these recommended values of  $\theta_{\text{lim}}$  for the corresponding  $N_{\text{side}}$ . See Wünsch et al. (2018) for  $\theta_{\text{lim}}$  resolution tests in the context of TREERAY/OPTICALDEPTH.

where  $\theta_{\text{src}}$  is a user-defined parameter.

Quantities on the tree are mapped on to the radial evaluation points of a ray through the use of kernels. We utilize both a piece-wise third-order polynomial,  $W_p(\delta)$ , and a kernel derived to ensure it meets the requirements of the radiation transfer equation,  $W_f(\delta)$ , where  $\delta = (r_i - d)/h_n$  and  $d$  is the distance from the node centre of mass and the ray evaluation point (see Wünsch et al. 2021) and  $h_n$  is the node's linear size. The node quantities are weighted by the overlap of the volume of the ray segment and the node.

Following the tree walk, the rays from a cell outward store the mass, centre of mass, gas temperature, and bin-integrated luminosities. These provide all the necessary information to solve the equation of radiation transfer along each ray.

### 2.2.4 Solving the radiation transfer equation

Along the rays, the one-dimensional radiation transfer equation is solved:

$$\frac{dI_v}{ds} = -\epsilon_v + \alpha_v I_v, \quad (12)$$

where  $s$  is the distance along the ray, and  $\epsilon_v$  and  $\alpha_v$  are the emission and absorption coefficients. The band-integrated flux,  $J(E)$ , irradiating a cell  $i$  due to the band-integrated luminosity,  $L_X$ , emitting from node  $j$  can be simply written

$$J_{ji}(E) = L_{X,j}(E) \frac{e^{-\tau_x}}{4\pi r_{ij}^2}, \quad (13)$$

where  $r_{ij}$  is the distance between the centres of cell  $i$  to node  $j$  and

$$\tau_x = \int n_{\text{H}}(s) \sigma_x(E, T(s)) ds \approx \sum_k \frac{\rho_k}{\mu m_{\text{H}}} \langle \sigma_x(E, T_k) \rangle \delta s \quad (14)$$

is the X-ray opacity between cell  $i$  and node  $j$  and  $\rho_k$  is the density at evaluation point  $k$  along the ray,  $n_{\text{H}}$  is the hydrogen nuclei density, and  $\delta s = r_k - r_{k-1}$ . We store the solution as an energy density,  $\epsilon_x = J/c$ , where  $c$  is the speed of light, on to the grid to be used in chemistry, described below. The total energy density is the sum over the HEALPIX rays:

$$\epsilon_i = \sum_k^{N_{\text{pix}}} \epsilon_{ki}(E). \quad (15)$$

For the solution, we also store the cell mass and temperature on to the tree and map these to the rays using  $W_p$ . In our algorithm, no assumption is made with respect to what produces the X-ray energy density, enabling both point sources (with their radiation spread over their host cells) and diffuse emission produced via cooling of hot gas in the cell.

### 2.3 Pre-existing chemistry

We briefly describe here the previous treatment of X-ray radiation within the chemical network (see Mackey et al. 2019 for more details). The chemical network consists of 17 species, of which 9 are solved numerically and the rest are followed through conservation equations. We solve the non-equilibrium species  $\text{H}^+$ ,  $\text{H}_2$ ,  $\text{C}^+$ ,  $\text{CO}$ ,  $\text{HCO}^+$ ,  $\text{CH}_x$ ,  $\text{OH}_x$ ,  $\text{He}^+$ , and  $\text{M}^+$ .  $\text{CH}_x$  is a proxy species for simple hydrocarbons, e.g.  $\text{CH}$ ,  $\text{CH}_2$ , etc., and simple ions  $\text{CH}^+$ ,  $\text{CH}_2^+$ , etc. Similarly,  $\text{OH}_x$  is a proxy for  $\text{OH}$ ,  $\text{H}_2\text{O}$  and ions  $\text{OH}^+$ ,  $\text{H}_2\text{O}^+$ , etc.  $\text{M}$  is a proxy for metals that can become the primary source of electrons in shielded regions of molecular clouds, where reaction rates treat  $\text{M}$  as  $\text{Si}$ . The network is primarily based on the ‘NL99’ network

of Glover & Clark (2012), which uses the hydrogen chemistry from Glover & Mac Low (2007a,b) with the CO chemistry of Nelson & Langer (1999) including updated reaction rates from Gong, Ostriker & Wolfire (2017). For this work, all photodissociation rates have been updated using the KInetic Database for Astrochemistry (KIDA) astrochemistry data base (Wakelam et al. 2012).

X-ray radiation is coupled to the thermochemistry through the following primary processes (see also Mackey et al. 2019):

- (i) Dust heating, following the analytical prescription in Yan (1997).
- (ii) Primary ionization of a species by X-rays. Note though that is is relatively unimportant for our considered species, and plays a minor role in the heating and ionization for hydrogen species and helium.
- (iii) Secondary ionization through collisional ionization by fast electrons produced following primary ionizations (e.g. Dalgarno et al. 1999; Meijerink & Spaans 2005).
- (iv) Induced FUV radiation generated by  $\text{H}_2$ , which is collisionally excited by fast electrons and the subsequent ionizations and dissociations (Prasad & Tarafdar 1983; Gredel, Lepp & Dalgarno 1987; Maloney et al. 1996; Meijerink & Spaans 2005).
- (v) Coulomb heating of the gas via energy exchange between the produced fast electrons and other charged particles (Dalgarno et al. 1999).

These processes have all been generalized for the arbitrary number of energy bins and the temperature-dependent cross-sections. The input X-rays are computed by the XRAYTHESPOT module. Since we use band-integrated radiative variables, the heating parameter for a particular cell  $i$  due to the impinging X-ray radiation is

$$H_{x,i} = \sum_n^{N_{\text{bin}}} j_i(E_n) \langle \sigma_x(E_n, T_i) \rangle. \quad (16)$$

## 3 TESTS AND BENCHMARKING

Here, we show various numerical tests of the radiation transfer and a benchmark of the thermochemistry against CLOUDY. For our benchmarks, we fiducially use eight bins, logarithmically spaced between 1 and 10 keV, following Meijerink & Spaans (2005).

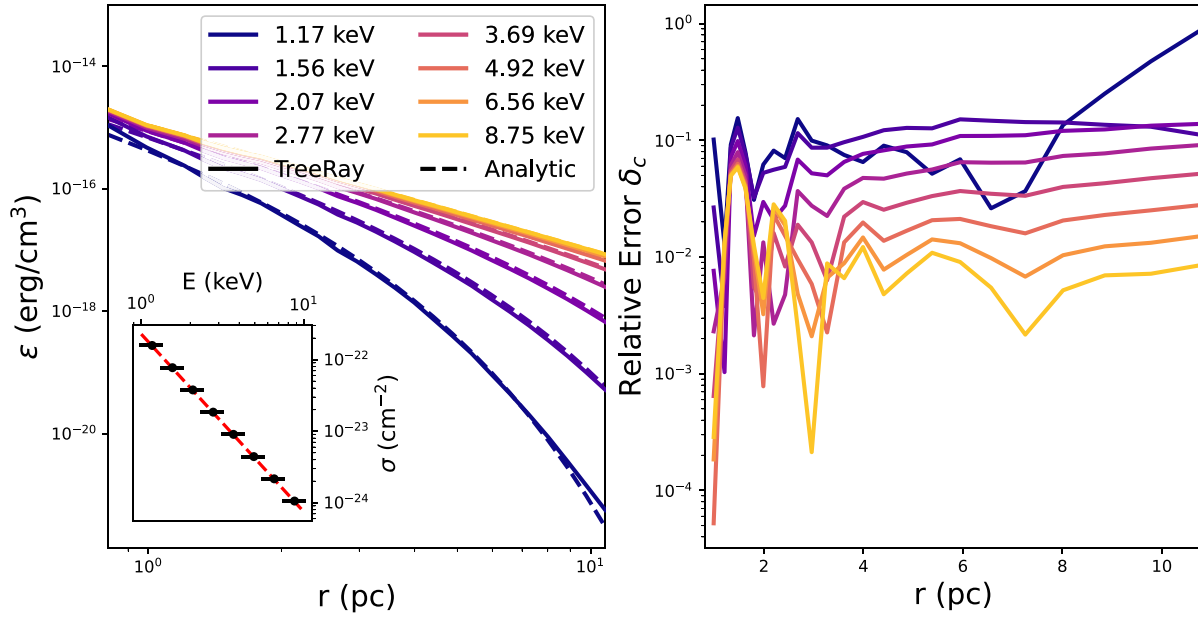
### 3.1 Point source test

Our first test is a single central point source with a constant luminosity distribution,  $L_{x,n} = 1 L_{\odot}$  for all  $N_{\text{bin}}$  bins, embedded in a volume with a uniform density of  $n(\text{H}) = 2 \times 10^3 \text{ cm}^{-3}$  and a spatially constant temperature  $T = 10 \text{ K}$  in a  $(30 \text{ pc})^3$  volume. We use a constant luminosity to better compare the solutions of different energy bins. We then compute the radial profiles of the energy density and compare against the analytical solution:

$$J(E, r) = L_x(E) \frac{e^{-\sigma_x(E)\rho r}}{4\pi r^2}, \quad (17)$$

where the energy density,  $\epsilon = J(E)/c$ .

Fig. 5 shows the performance of XRAYTHESPOT for a single bright point source as a function of radius. The results in Fig. 5 used  $\eta_R = 4$ ,  $N_{\text{side}} = 8$ , and  $N_{\text{block}} = 8$ , where  $N_{\text{block}}$  is the number of blocks of cells per spatial dimension, and one block consists of a cube of  $8^3$  cells. The radial range was chosen that for the lowest energy bin, the emission transitions from optically thin to strongly optically thick, with the maximum radius corresponding to  $\tau(E = 1.17 \text{ eV}) \approx 10$ . The left-hand panel shows the comparison between the ray trace



**Figure 5.** Radial profile test for a single source in a constant density and temperature medium. Left: Radiation density versus radius for each bin for the TREERAY (solid) and analytical solution (dotted). Left inset: Bin-averaged cross-sections (black points) and the analytical cross-section in equation (1). Right: Relative errors for each bin as a function of radius.

solution and the analytical solution. These solutions agree well with each other with the lines largely overlapping. The right-hand panel shows the relative error, defined as

$$\delta_c = \frac{|c\varepsilon - J(E, r)|}{J(E, r)}, \quad (18)$$

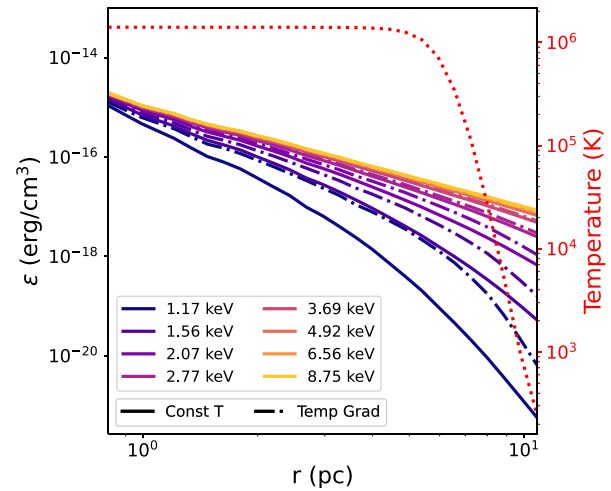
where  $\varepsilon$  is the solution from XRAYTHESPOT. The relative error is rather insensitive to the optical depth but more sensitive to how strongly the radiation field is coupled to the gas (e.g. the magnitude of the photoabsorption cross-section). The 10 percent error shown for the most optically thick bin at low energies is due to the mapping of the density structure on to the rays using the kernel. For X-ray optical depths greater than  $\tau_x \approx 10$ , the error starts to increase towards unity, but at these energy densities, the X-rays have a negligible impact on the thermochemistry. Therefore, these relative differences will have no discernible impact on the thermochemical evolution of the gas.

In order to highlight the differences of the new module with the previous cross-section implementation presented in Mackey et al. (2019), we perform a second calculation imposing a strong temperature gradient such that the radial temperature profile is

$$T(r) = 5 \times 10^5 [1 - \tanh(r - 8 \text{ pc})] + 100 \text{ K}. \quad (19)$$

This temperature is artificial and chosen such that the X-ray radiation transitions from optically thin to optically thick due to the change in cross-section. Fig. 6 shows the result of this comparison and as expected, the emission for the lower energy bins is up to an order of magnitude greater than the low-temperature solution and maintains an  $r^{-2}$  trend until a much greater radius.

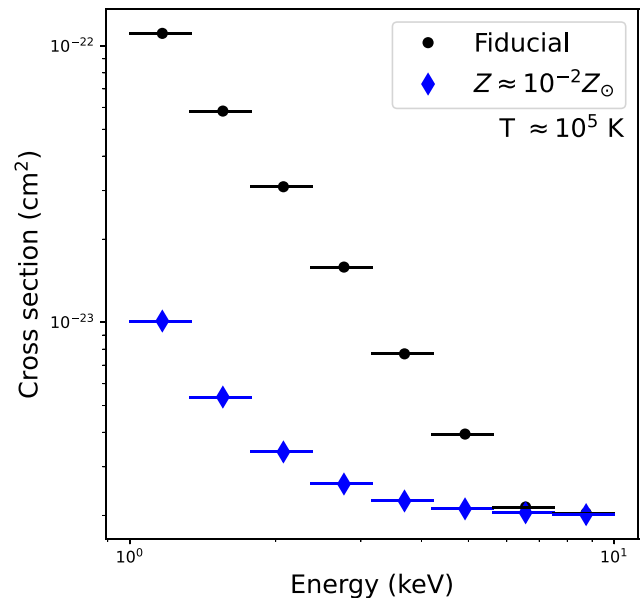
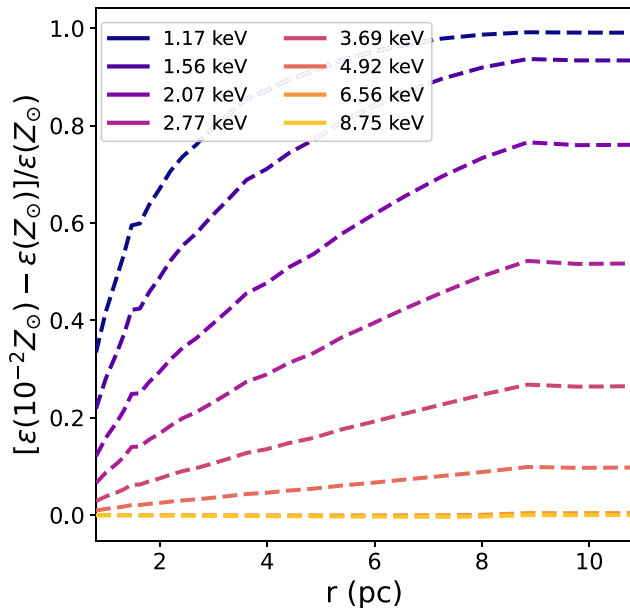
Fiducially, we assume the abundances shown in Table 1. We run an additional problem using the temperature profile above with a metallicity a factor of 100 lower. Fig. 7 shows the relative differences in the energy densities for the two different metallicities and the cross-sections at  $T = 10^5 \text{ K}$ . The emission at very low metallicity is significantly enhanced due to the lack of photoabsorption by metals



**Figure 6.** X-ray energy density versus radius for single point source. The solid line uses the constant temperature at  $T = 10 \text{ K}$  (same as Fig. 5) while the dash-dotted line uses the temperature profile shown by the red dotted line.

(see Fig. 3). For the highest energy bins, there is very little difference due to the cross-section being by Compton scattering.

Fig. 8 shows the performance of XRAYTHESPOT for a range of parameters, exploring both low- and high-spatial and ray resolutions. We find that grid resolution is the primary source of deviations at small radius, while the ray resolution increases the accuracy at larger radii. At large distances from the source, the solution tends to slightly underpredict for low ray and angular resolution due to overestimation of the column density. At small radii, the solution overpredicts the resulting flux. For optically thin radiation bins, the solution almost exactly matches the analytical. Therefore, the deviations come about due to mapping the mass from the cells and tree nodes on to the rays using the kernel.



**Figure 7.** Left: Relative deviations in the energy density between the fiducial abundances and a model with  $Z = 10^{-2} Z_{\odot}$ , defined as  $[\epsilon(10^{-2} Z_{\odot}) - \epsilon(Z_{\odot})]/\epsilon(Z_{\odot})$ . Right: The cross-sections at  $T \approx 10^5$  K for the fiducial abundances (black) and  $Z = 10^{-2} Z_{\odot}$  (blue).

For science uses, the number of blocks,  $N_{\text{block}}$ , is determined by the necessary resolution to resolve crucial gas dynamics (e.g. the Jeans length for gravity simulations). Increasing both  $N_{\text{side}}$  and  $\eta_R$ , while producing more accurate radiation transfer solutions, leads to substantially higher computational costs. Table 2 shows the computational time per processor for the initialization and per ray trace step for the models in Fig. 8. Between the lowest and highest accuracy tests,  $(N_{\text{block}}, N_{\text{side}}, \text{and } \eta_R) = (4, 2, \text{and } 2)$  and  $(8, 8, \text{and } 4)$ , respectively, the increase in cost of the initialization and ray trace was a factor of  $\approx 40$  and  $\approx 130$ , respectively. The time for the ray trace is dominated ( $\geq 95$  per cent) by the tree walk. We find using  $N_{\text{side}} = 4$  is the best balance of time and accuracy.

### 3.2 Shadow test

Our next test is a shadow test to verify the solution of the solver when sources are placed near dense regions. Here, we have a  $L_X = 10 L_{\odot}$  source with a spectrum,  $dL/dE \propto E^{-2}$ , placed near a dense core with a hydrogen-nuclei number density of  $n_{\text{H}} = 10^3 \text{ cm}^{-3}$ . We consider radiation between 1 and 10 keV, moving from optically thick bands to optically thin. Figs 9 and 10 show the results of this test, for both low- and high-ray resolution which use  $(N_{\text{block}} = 8, N_{\text{side}} = 4, \text{and } \eta_R = 2)$  and  $(N_{\text{block}} = 8, N_{\text{side}} = 8, \text{and } \eta_R = 4)$ , respectively. For both cases, the test reveals the expected results that the low-energy X-rays are absorbed by the dense core and this creates a wide-angle shadow, while higher energy X-rays are barely attenuated, producing smaller to no shadows. The high-ray resolution test also shows the expected drop in ray-tracing artifacts.

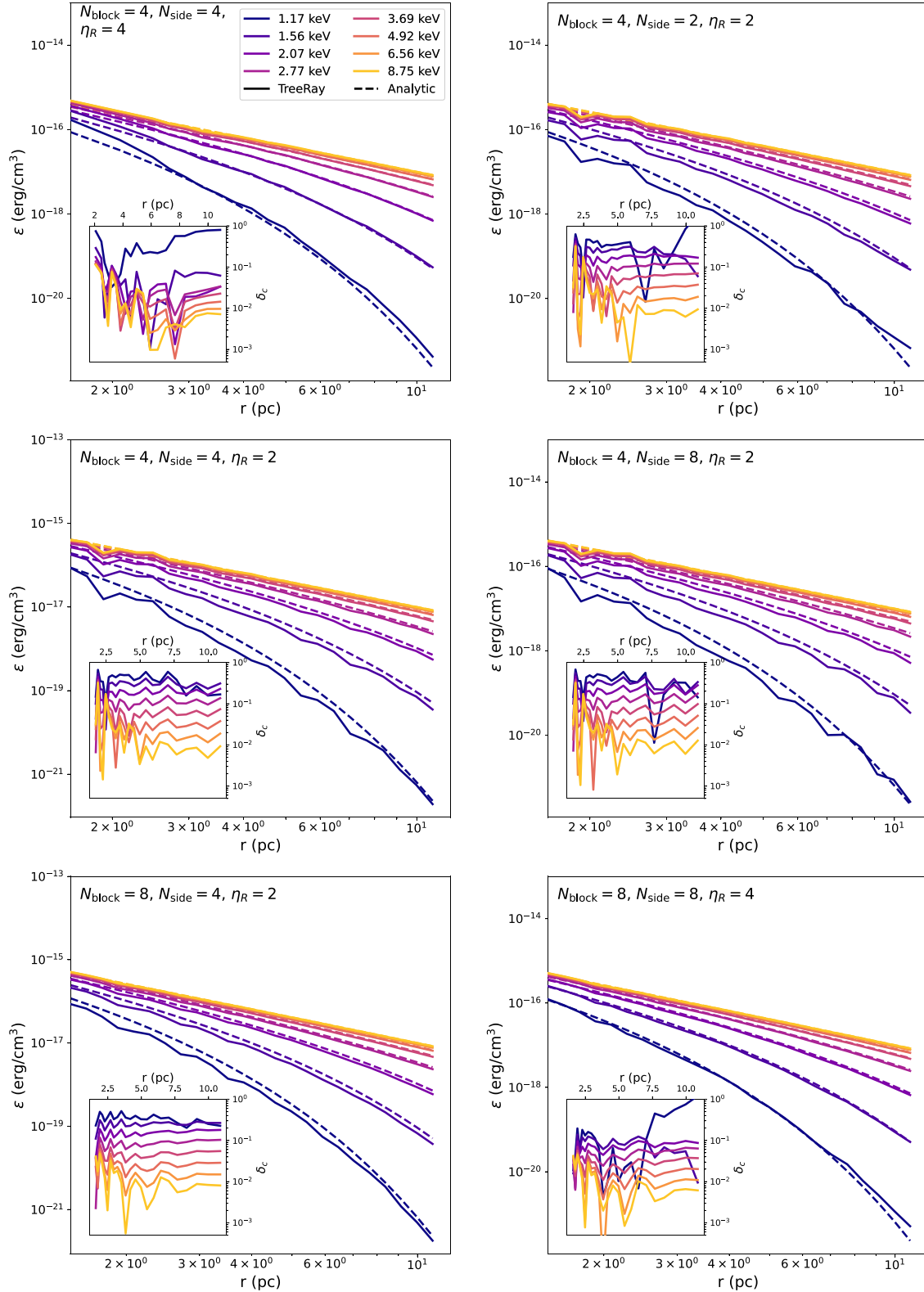
Fig. 11 shows a one-dimensional cut along the  $z$ -axis from the source through the dense blob of gas for the two ray resolutions compared. The figure shows that higher ray resolution leads to a smoother attenuation of the flux for the optically thick, lower energy bins while there is very little change for higher energy bins which are

substantially less attenuated. This is most pronounced for the  $E_c = 1.33$  keV bin.

### 3.3 Benchmark against CLOUDY

The final benchmark tests the X-ray radiation coupling to the thermochemistry. We place a point source with a physical size of  $10^{16}$  cm and total X-ray luminosity,  $L_{\text{XR}} = 10^{36} \text{ erg s}^{-1}$ , and a luminosity spectrum  $dL/dE \propto E^{-2}$  between 1 and 10 keV in a  $(1.3 \text{ pc})^3$  volume filled with a gas number density of  $n_{\text{H}} = 10^3 \text{ cm}^{-3}$ ,  $N_{\text{side}} = 4$ , and  $\eta_R = 2$  and compare with a one-dimensional model using the CLOUDY code. We also use a cosmic ray ionization rate of  $3 \times 10^{-17} \text{ s}^{-1}$  and a gas and dust temperature floor of 3 K to mimic the inclusion of a cosmic microwave background (CMB). The volume and resolution were chosen such the inner XDR is resolved by  $\approx 20$  cells while the optically thick regime is also traced. In particular, we use seven maximum adaptive-mesh resolution levels refining on the density and temperature, such that the maximal resolution is  $1.3 \times 10^{-3} \text{ pc}$ . The one-dimensional CLOUDY model used a ‘sphere’ geometry, an input power-law spectrum between 1 and 10 keV for the X-ray radiation source, a CMB and a cosmic ray ionization rate of  $3 \times 10^{-17} \text{ s}^{-1}$ . Further, we turn off grain physics, induced radiative processes, radiation pressure, radiation scattering, outward line radiation transfer, and molecule freeze-out, since the FLASH simulations do not have these processes. Finally, we set refractory metal abundances to zero, with the exception of silicon which FLASH uses as the proxy for metals for the chemistry (as described above). The CLOUDY script used is shown in Appendix A.

Since we use uniform-spaced grids, even with substantial adaptive mesh refinement (AMR) levels, it is in practice difficult to fully capture sharp thermochemical transition regions, such as that shown below as captured by CLOUDY. Further, the source encompasses several cells at the highest resolution, rather than an infinitely small point source. Capturing such ionization and dissociation fronts



**Figure 8.** Energy density versus radius for the different model parameters, annotated in the top left of each subfigure. Inset: Relative error,  $\delta_c$ , of the numerical solution against the analytical solution as a function of radius from the source.

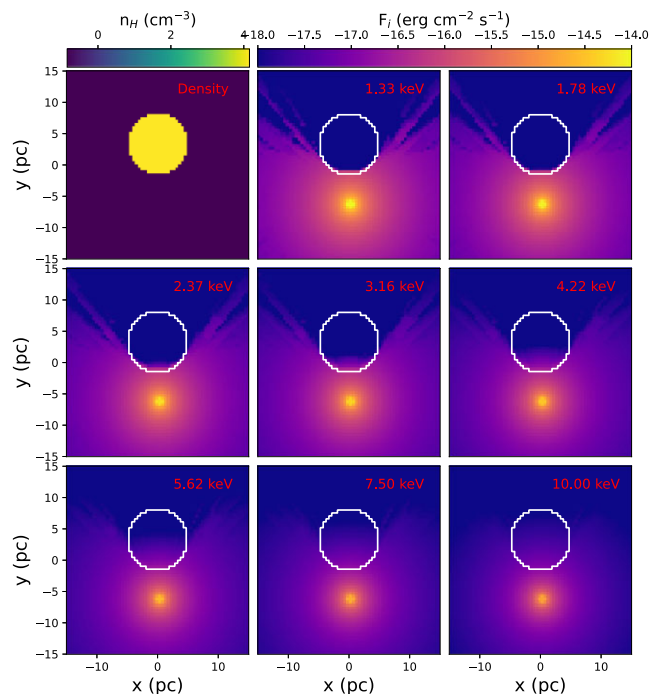
entirely is numerically intensive and generally requires the use of one-dimensional models tailored to do so (as with CLOUDY).

Fig. 12 shows the result of this benchmark. Near the source, the temperature and chemistry solutions well match the CLOUDY

solution. The FLASH and CLOUDY solutions qualitatively reproduce the chemical structure, although due to the larger cell size of the FLASH grids, the sharp HI transition seen at  $N(\text{H}) \approx 5 \times 10^{20}$  cm<sup>-2</sup> is not fully captured and is instead smoothed over a few cells.

**Table 2.** Timing for the point source test for the different runs in Fig. 8. Each row gives the model parameters of  $N_{\text{block}}$ ,  $N_{\text{side}}$ , and  $\eta_R$  and the time in seconds per processor for the initialization of the tree and a ray trace step.

| $N_{\text{block}}$ | $N_{\text{side}}$ | $\eta_R$ | Initialization (s/proc) | Evolution (s/proc) |
|--------------------|-------------------|----------|-------------------------|--------------------|
| 4                  | 2                 | 2        | 2.3                     | 1.3                |
| 4                  | 4                 | 2        | 5.5                     | 3.6                |
| 4                  | 4                 | 4        | 6.1                     | 4.6                |
| 4                  | 8                 | 2        | 22.1                    | 14.0               |
| 8                  | 4                 | 2        | 17.4                    | 29.3               |
| 8                  | 8                 | 4        | 94.0                    | 167.2              |



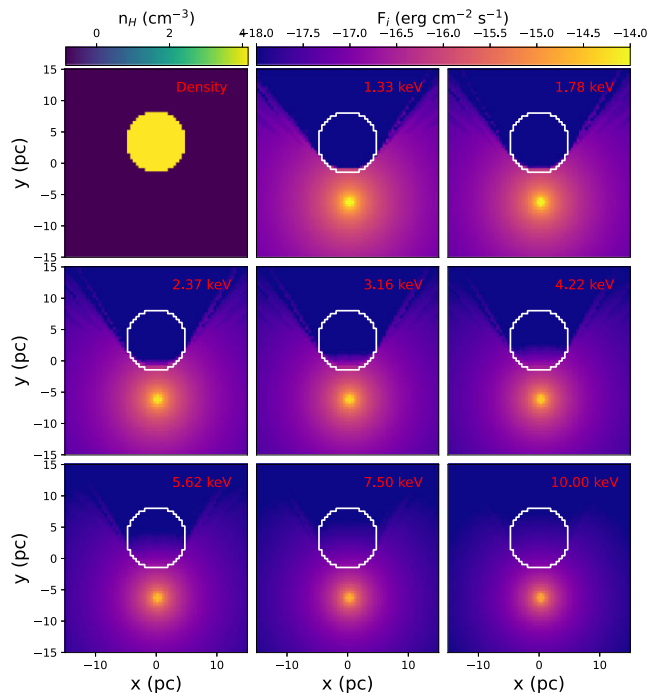
**Figure 9.** Shadow test, consisting of a point source illuminating a constant density core. Top left corner: Number density distribution for a  $z$ -axis slice. Others: X-ray flux in the given energy band for a  $z$ -axis slice using  $N_{\text{block}} = 8$ ,  $N_{\text{side}} = 4$ , and  $\eta_R = 2$ .

The temperature solutions agree within a factor of a few. However, CLOUDY solves the line cooling and level excitations in a much more robust manner than the included FLASH thermochemistry, including a full non-equilibrium solution with many more electronic and ionization states. Such inclusions though are not numerically feasible for *in situ* thermochemistry in three-dimensional MHD simulations. Further, CLOUDY solves the full radiation transfer solution from radio through X-ray radiation with substantially more bins.

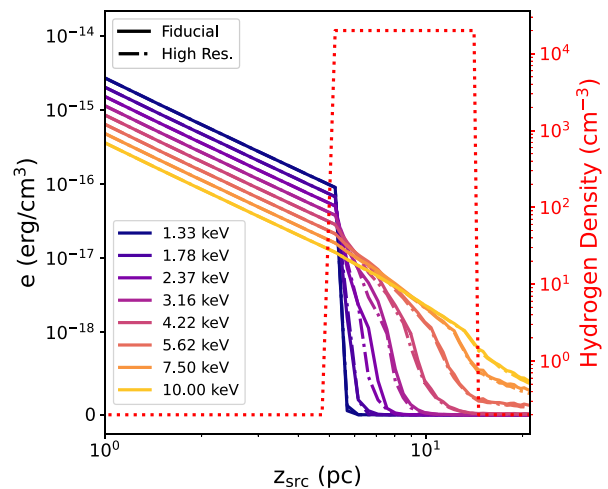
Fig. 13 shows a comparison of the heating rates. For FLASH, the X-ray heating is included via

$$\Gamma_X = \eta_X n_H H_X, \quad (20)$$

where  $\eta_X$  is the heating efficiency (Dalgarno et al. 1999). We calculate this heating term in post-processing for the FLASH runs, as the chemistry solver does not store this in real time in the simulation. For CLOUDY, the heating term output does not delineate heating caused by X-ray radiation. However, since we include only a low cosmic ray background with no other primary radiation heating source, the total heating rate will provide a close approximation. We find that our heating term is, on average,



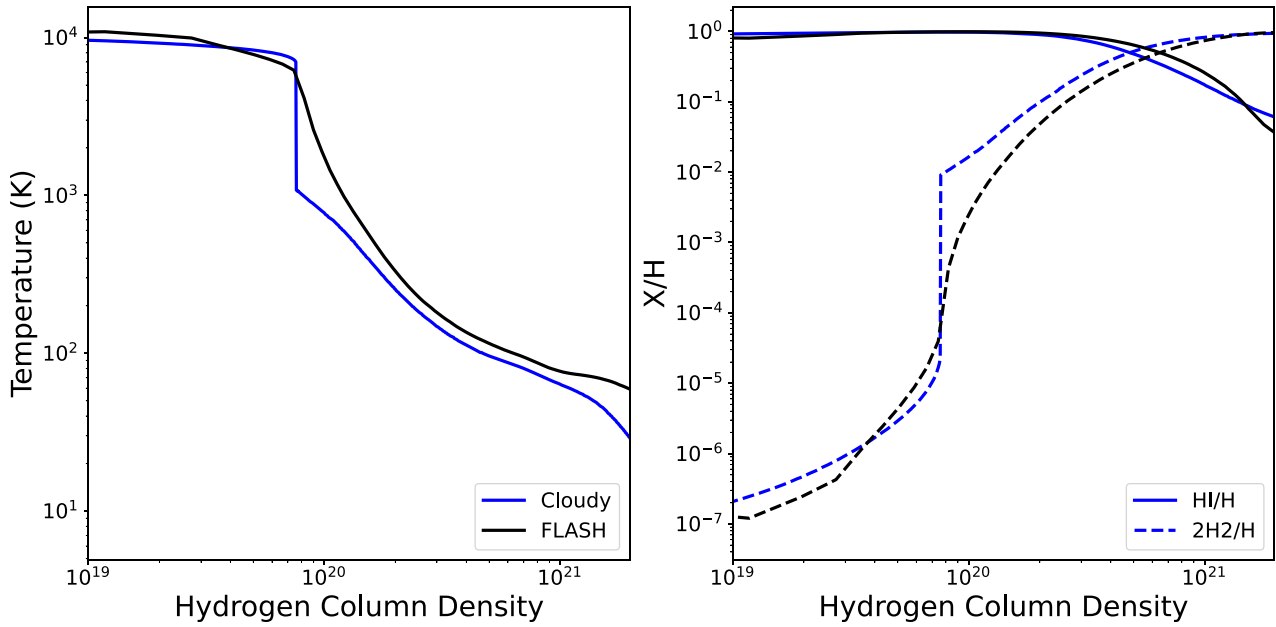
**Figure 10.** Same as Fig. 9, but with  $N_{\text{block}} = 8$ ,  $N_{\text{side}} = 8$ , and  $\eta_R = 4$ .



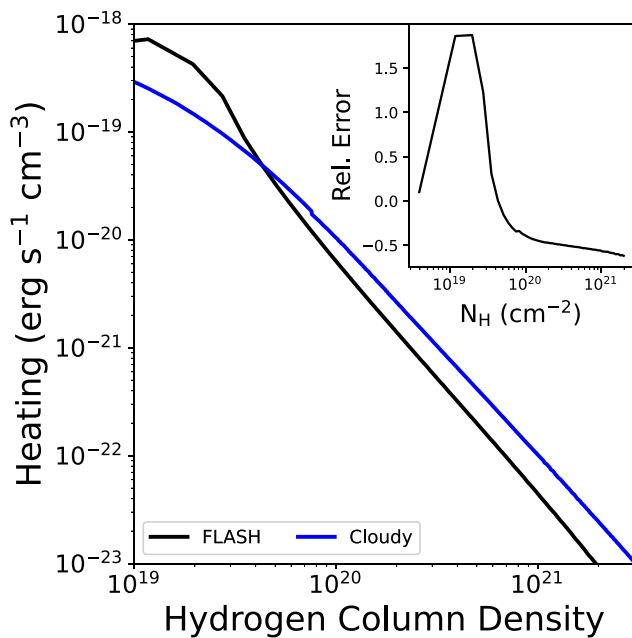
**Figure 11.** X-ray energy density versus distance along the  $z$ -axis from the source for the shadow test. The solid line uses the ray resolution in Fig. 9 and the dash-dotted uses the ray resolution in Fig. 10. The dotted red line shows the hydrogen nuclei density highlighting the location of the high-density blob.

within a factor of two of the total heating rate computed by CLOUDY.

Fig. 14 compares the relative abundances in the so-called carbon cycle,  $C^+ \rightarrow C \rightarrow CO$ , between the two codes. The results broadly agree, with FLASH showing an enhanced neutral carbon and rapid production of CO, which is a result of the greatly reduced network (Glover et al. 2010; Gong et al. 2017). CLOUDY uses a significantly larger network with detailed line radiation transfer for self-shielding (important for  $H_2$  and CO) while we use the average column densities of  $H_2$  and CO to compute the self-shielding factor (Walsh et al. 2015; Wunsch et al. 2018). Given the constraints of these physics, the found solution is deemed



**Figure 12.** FLASH versus CLOUDY benchmark. Left: Temperature versus hydrogen column density from the central point source for FLASH (black) and CLOUDY (blue). Right: Atomic (solid) and molecular (dashed) hydrogen abundances versus total hydrogen column density from the source, where  $H_{\text{tot}} = H^+ + H + 2H_2$ .

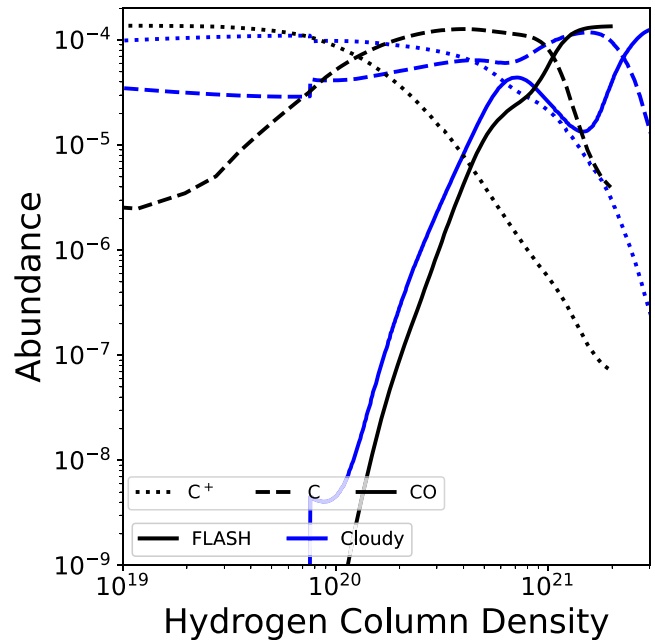


**Figure 13.** FLASH versus CLOUDY heating benchmark. Total heating from CLOUDY (blue) versus X-ray heating from FLASH. Inset: Relative difference between these.

to be adequate and matches the overall trends as determined by CLOUDY.

#### 4 PROTOSTELLAR DISC

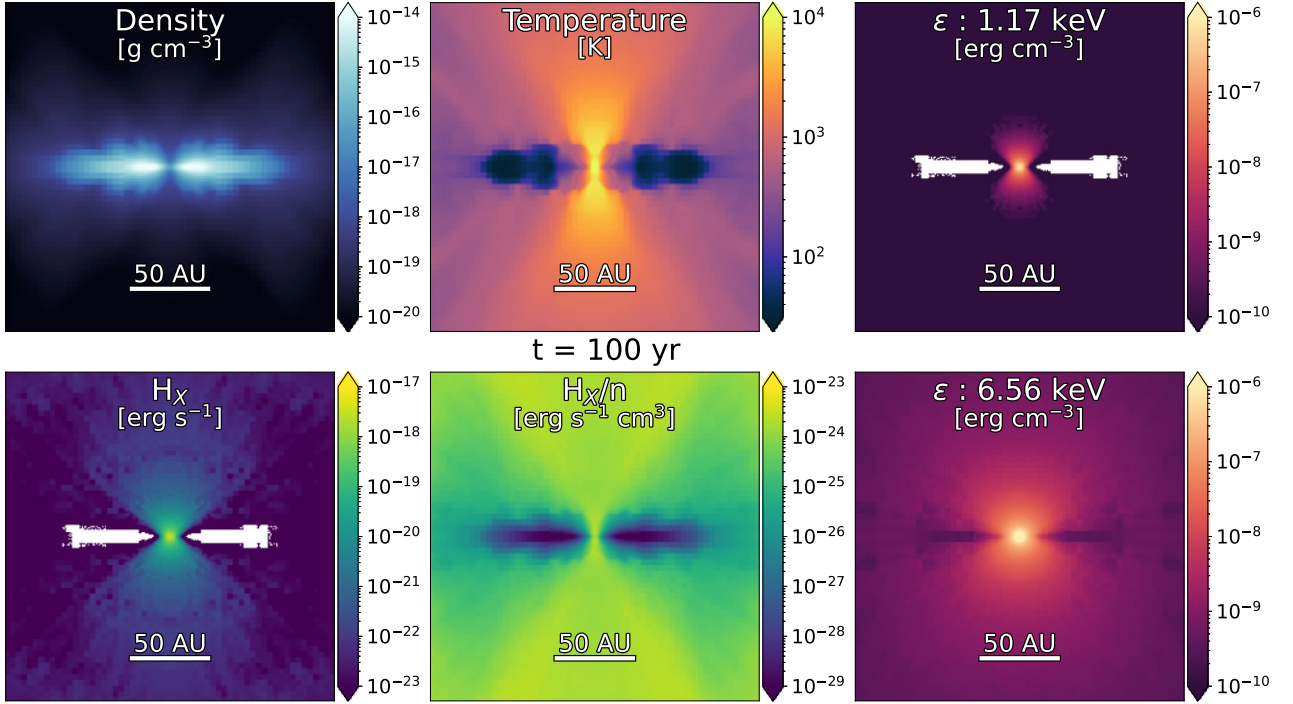
Evolved protostellar objects, in particular Class II objects in which the lack of a surrounding gaseous envelope leaves the central protostar and disc exposed, are known to be X-ray emitters. These X-rays can become important for disc dynamics and planet formation



**Figure 14.** FLASH versus CLOUDY benchmark. Abundances of  $C^+$  (dotted), C (dashed), and CO (solid) versus total hydrogen column density from the central point source for FLASH (black) and CLOUDY (blue).

(e.g. Ercolano et al. 2008b; Mohanty, Ercolano & Turner 2013). For these stars, the X-ray emission is thought to come from a combination of accretion and magnetospheric emission (Hartmann, Herczeg & Calvet 2016). As a first test science case, we model the X-ray radiation transport from a central protostar into a protostellar disc.

The surface density follows from the often used truncated power law (e.g. Lynden-Bell & Pringle 1974; Andrews et al. 2011; Cleeves



**Figure 15.** Protostellar disc example case usage. Top row: Slice plots at  $z = 0$  for the density (left), gas temperature (middle), and 1.17 keV radiation energy density. Bottom row: X-ray heating rate,  $H_X$ ,  $H_X/n$  diagnostic term (middle), and 6.56 keV radiation energy density.

et al. 2016):

$$\Sigma_g(R) = \Sigma_c \left( \frac{R}{R_c} \right)^{-\alpha} \exp \left[ - \left( \frac{R}{R_c} \right)^{2-\alpha} \right] \quad (21)$$

between an inner and outer radius,  $R_{\text{in}}$  and  $R_{\text{out}}$ , respectively,  $R_c$  is the critical radius where the surface density distribution becomes exponential,  $\alpha$  is the power-law index and  $\Sigma_c$  is the characteristic surface where the disc transitions to an exponential profile. For the initial conditions, we assume the gas is in hydrostatic equilibrium, such that the density follows

$$\rho_g(R, z) = \frac{\Sigma_g(R)}{\sqrt{2\pi}h} \exp \left[ - \left( \frac{z^2}{2h^2} \right) \right] \quad (22)$$

where  $h = c_s/\Omega$  is the disc scale height,  $c_s = \sqrt{\frac{\gamma k_B T_g}{\mu m_H}}$ ,  $k_B$  is Boltzmann's constant,  $\gamma = 5/3$  is the adiabatic index,  $T_g$  is the gas temperature,  $\mu = 2.33$  is the mean mass per particle for molecular gas,  $m_H$  is the mass of the hydrogen atom,  $\Omega = \frac{3}{4} \sqrt{\frac{GM_*}{R^3}}$  is the Keplerian rotational frequency, and  $M_*$  is the mass of the central protostellar object. For this fiducial test, we set  $M_* = 0.7 M_\odot$ ,  $\Sigma_c = 64 \text{ g cm}^{-2}$ ,  $R_c = 100 \text{ au}$ , and  $\alpha = 1$ . The temperature profile is given by

$$T(R) = \max \left[ T_0 \left( \frac{R}{1 \text{ au}} \right)^{-0.5}, 10 \text{ K} \right], \quad (23)$$

where we fiducially take  $T_0 = 50 \text{ K}$ . The disc is initialized to be rotating in Keplerian motion around the central protostellar object. We assume the disc is magnetized with an initial toroidal field such that the ratio of the magnetic to thermal pressure,  $\mu_M = 10^{-5}$ . We simulate the domain in a 240 au box with a maximal resolution of 1 au.

The central protostar is put in by hand, with active accretion. For the X-ray emission, we assume an accretion floor of  $10^{-9} M_\odot \text{ yr}^{-1}$ , similar to rates observed in young stellar objects (e.g. Ingleby et al. 2013). The simulation is run using the Bouchut-5 MHD solver, gravity, and XRAYTHESPOT. The X-ray emission is derived by assuming there is an accretion shock, with properties following ‘hotspot’ accretion (Hartmann et al. 2016) with accretion columns filling 10 per cent of the protostar surface, thermally emitting X-ray emission. The thermal X-ray emission is computed using a one-temperature Raymond–Smith plasma model (Raymond & Smith 1977). The implementation of a coronal model is left for a future work.

Fig. 15 shows a slice of the density, gas temperature, and X-ray emission at 1.17 keV (1st bin) and 6.56 keV (8th bin), the heating rate per H nucleus,  $H_X$ , and  $H_X/n$ , which is often used as a diagnostic for the importance of the X-ray heating (Wolfire et al. 2022). We find that the lowest energy X-rays are all absorbed near the protostar or escape through the outflow. However, the harder X-rays at 6.56 keV are able to permeate much of the domain. The  $H_X$  and  $H_X/n$  slices clearly show that the disc mid-plane is left relatively unheated by the X-rays, although the X-rays become important in the cavity and outer disc regions. In particular, most of the cavity exhibits very warm gas, even with only X-ray emission included, due to the rapid absorption of soft X-ray emission. The cavity heats to temperatures exceeding  $10^4 \text{ K}$ , potentially becoming bright in hydrogen recombination lines. The inclusion of extreme ultraviolet (EUV) radiation will heat the diffuse gas further, along with further ionizing the surrounding low-density cavity.

## 5 MOLECULAR CLOUD

We present an example application for XRAYTHESPOT, to demonstrate how all the different TREERAY energy bands work together: a

virialized, magnetized turbulent cloud. We consider a 2 pc region of a molecular cloud resolved with  $256^3$  cells. We produce an initial turbulent field by stirring the domain with a flat power spectrum between the largest wave modes  $k = 1, \dots, 3$  for 10 crossing times at a velocity dispersion of  $0.72 \text{ km s}^{-1}$ , consistent with the observed linewidth–size relationship (McKee & Ostriker 2007). During the stirring, we use periodic boundary conditions and chemistry to achieve more accurate initial conditions for the abundances before collapse. The choice of stirring for 10 crossing times is to ensure the chemistry has reached a more quiescent state, with the kinetic energy spectrum generally being reached after two crossing times (Federrath et al. 2010). We assume the cloud is nearly virialized, such that the virial parameter

$$\alpha \equiv \frac{5\sigma^2 R}{G\rho L^3} = 2, \quad (24)$$

where  $R = L$  is the box length, resulting in  $\rho = 5 \times 10^{-21} \text{ (g cm}^{-3}\text{)}$  and a total box mass of  $M = 590 M_\odot$ . Before stirring, we initialize a magnetic field in the  $z$ -axis with a magnitude such that the plasma beta,

$$\beta \equiv \frac{\rho c_s^2}{B^2/8\pi} = 10^3. \quad (25)$$

After the turbulence is initialized, gravity and source particles (stars) are turned on and the boundary conditions are changed to ‘diode’ such that gas can flow out of the domain. During the simulation, the cloud is irradiated by an FUV radiation field of  $\chi = 1.7$  in units of the Habing field (Habing 1968). The simulation is run using the chemistry described above, and all TREERAY modules:

(i) OPTICALDEPTH for the external radiation field (Wünsch et al. 2018). OPTICALDEPTH solves for the column density from a cell to the external boundary and attenuates a prescribes external radiation flux ( $\chi = 1.7$ ). In this study, it is only used for the FUV radiation, while (Mackey et al. 2019) implemented the ability to include an impinging X-ray flux.

(ii) ONTHESPOT for the EUV emission (Wünsch et al. 2021). This module solves for UV-ionizing radiation from arbitrary sources and iterates to convergence. The UV photon flux is coupled to the thermochemistry to model photochemistry.

(iii) RADPRESSURE to account for the thermal radiation and radiation pressure (Klepitko et al. 2022). This module enables the inclusion of thermal radiation from point and diffuse sources and the resulting radiation pressure. The thermal radiation is included in the chemistry through radiative dust heating.

(iv) XRAYTHESPOT, described above.

Sink particles representing protostars are injected when the density exceeds  $\rho_{\text{thresh}} \geq 4.59 \times 10^{-18} \text{ g cm}^{-3}$ . Further criteria are used: there are checks to ensure a local gravitational potential and a converging flow. The protostar evolution follows the Offner et al. (2009) model and implemented in FLASH (Klepitko et al. 2022). Protostellar emission consists of the intrinsic and accretion luminosities, where the total accretion luminosity is

$$L_{\text{acc}} = f_{\text{acc}} \frac{GM_* \dot{M}_*}{R_*}, \quad (26)$$

where  $M_*$  is the mass of the protostar,  $\dot{M}_*$  is the accretion rate, and  $R_*$  is the protostar’s radius and we take  $f_{\text{acc}} = 0.33$ . The X-ray spectrum was computed by assuming hotspot accretion, described above, which provides the temperature and the density of the accretion shocks near the protostellar surface (Calvet & Gullbring 1998; Hartmann et al. 2016) and a single temperature plasma model

(Raymond & Smith 1977). Due to the low resolution, we set a minimum of  $\dot{M}_* = 10^{-9} M_\odot \text{ yr}^{-1}$ . This is needed since when the protostar particles first form, the burst of accretion blows out H II regions, and the low resolution inhibits resolving the proper structure around the cores. The infrared to EUV spectrum, used for RADPRESSURE and ONTHESPOT is computed assuming the emission is composed of two blackbodies: one for the intrinsic spectrum of the protostar at the photosphere, such that

$$T_* = \left( \frac{L_*}{4\pi\sigma_{\text{sb}}R_*^2} \right)^{1/4}, \quad (27)$$

which is provided by the protostellar evolution model, and another assuming the accretion luminosity is reprocessed primarily as a blackbody with temperature  $T_{\text{acc}}$ , such that

$$T_{\text{acc}} = \left( \frac{L_{\text{acc}}}{4\pi\sigma_{\text{sb}}R_*^2} \right)^{1/4}, \quad (28)$$

where  $\sigma_{\text{sb}}$  is the Stefan–Boltzmann constant. Therefore, the total infrared luminosity from the protostar is described as

$$L_{*,\text{IR}} = f_{*,\text{IR}}(T_*)L_* + f_{\text{acc,IR}}(T_{\text{acc}})L_{\text{acc}}, \quad (29)$$

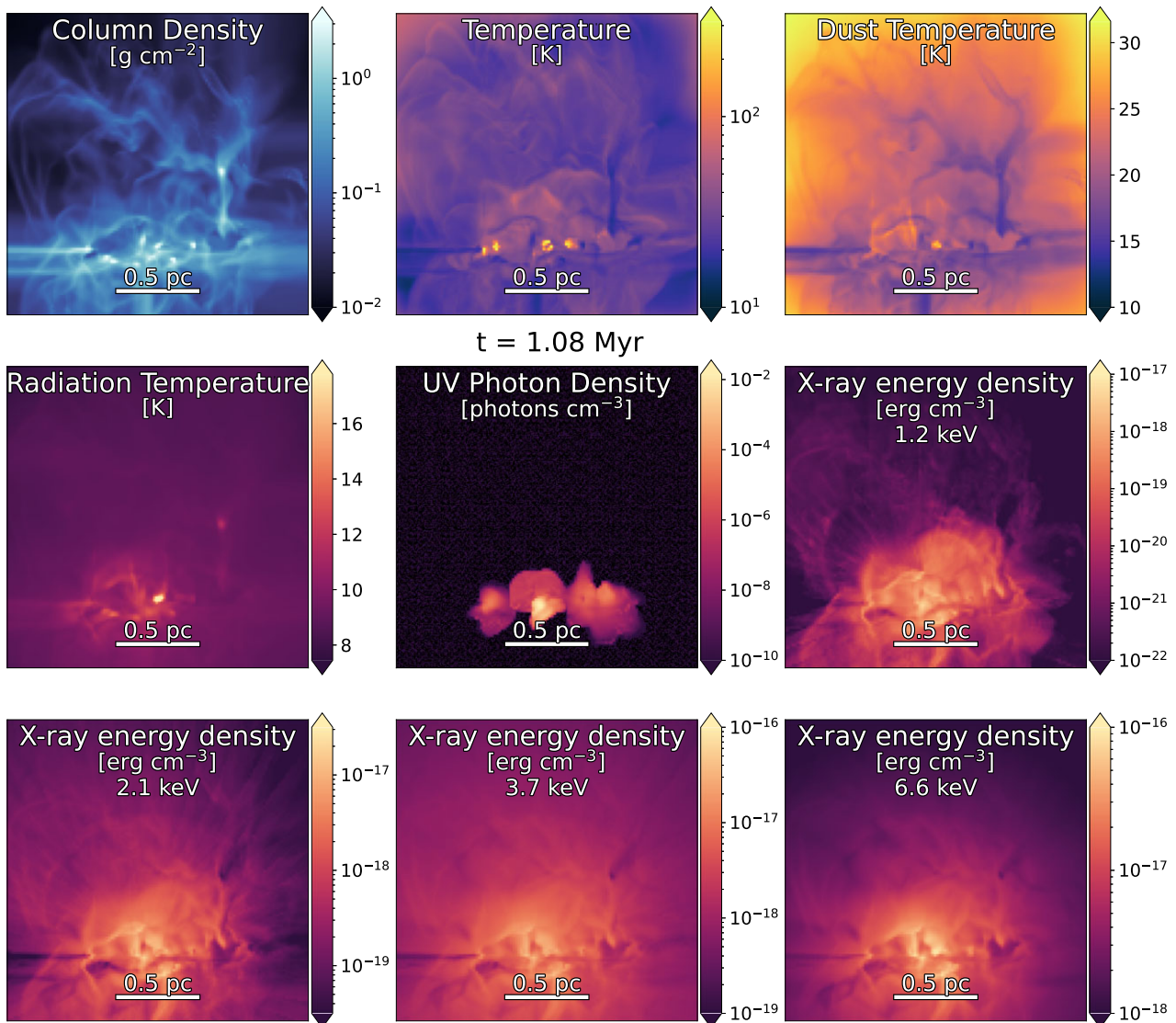
and the EUV luminosity as

$$L_{*,\text{EUV}} = f_{*,\text{EUV}}(T_*)L_* + f_{\text{acc,EUV}}(T_{\text{acc}})L_{\text{acc}}, \quad (30)$$

where  $f_{\text{IR}}(T)$  and  $f_{\text{UV}}(T)$  are the fraction of the blackbody emission in each of these bands ( $E < 13.6 \text{ eV}$  and  $13.6 \text{ eV} \leq E \leq 100 \text{ eV}$ , respectively). The X-ray emission was computed assuming the ‘hotspot’ model, described above. While there may be some double counting of emission by treating the total spectrum in the two different methods, we find this impact is marginal as the X-ray emission generally accounts for only a small fraction ( $\leq 10$  per cent) of the total protostellar luminosity.

Fig. 16 shows the column density, and density-weighted projections of the gas and temperature, radiation temperature, EUV photon density, and X-ray energy densities after  $\approx 1 \text{ Myr}$  of evolution with gravity. The star formation, as traced by heated knots of gas, is occurring along a main filament structure. The high temperatures here are primarily caused by the EUV photons, which are rapidly absorbed in the nearby gas. The X-ray emission is found to be highly absorbed along the main filament structure, and instead traces out the more diffuse turbulent structure of the molecular cloud. As expected, higher energy X-ray bands showcase more extended emission with the brightest emission in the 3.7 keV band. In all X-ray bands, the turbulent structure of the molecular cloud is seen in the density-weighted integrated emission.

Fig. 17 shows the surface densities of atomic and molecular hydrogen, atomic carbon, and carbon monoxide species in the cloud and Fig. 18 shows the  $\text{H}_2$  density-weighted density–temperature phase diagram. In this preliminary investigation, we see that there is an enhancement of neutral hydrogen roughly corresponding to where there are enhancements in the X-ray radiation field with some overlap of the UV-ionized  $\text{H}^+$ . Similarly, in the phase-diagram, there is a population of hot atomic gas, exceeded typical temperatures of H II regions (Haid et al. 2018) and an additional population of dense, molecular gas at temperatures  $10^2$ – $10^3 \text{ K}$ . These regions are likely X-ray heated gas close to X-ray emitting protostars, but a detailed investigation is beyond the scope of this work. This case study highlights the new capabilities of including protostellar radiative feedback from infrared to X-ray in star formation simulations.



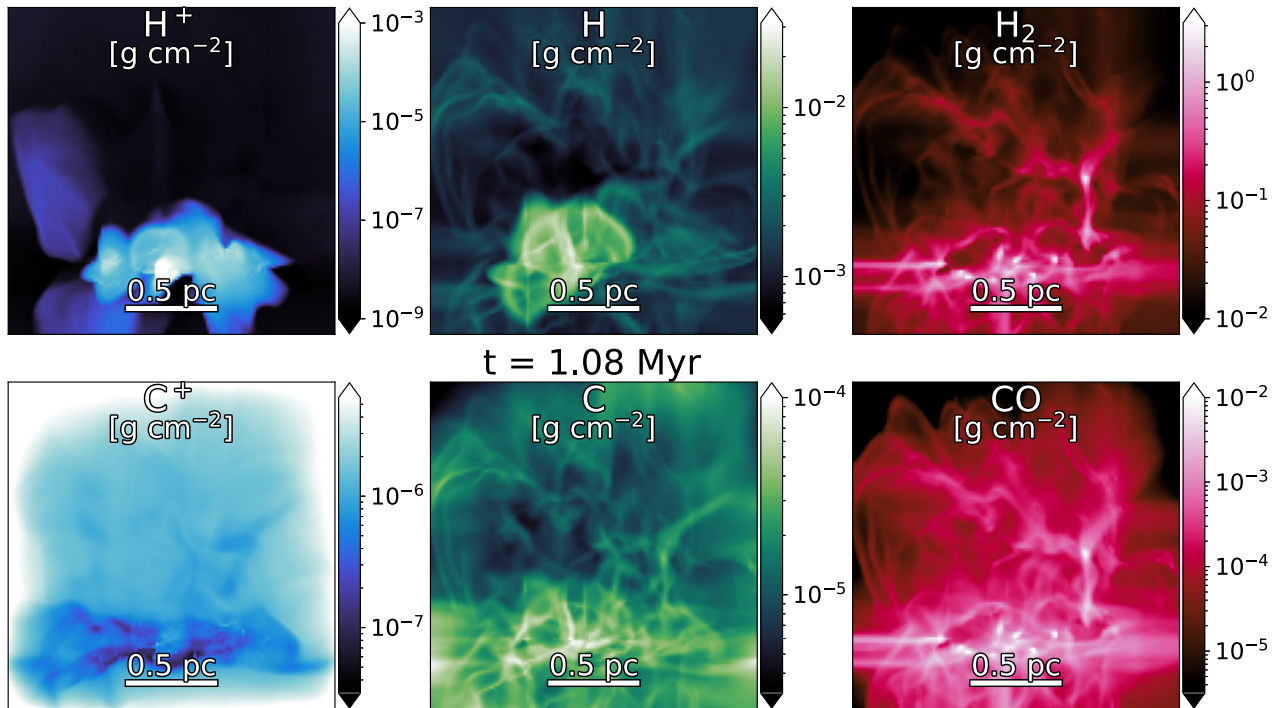
**Figure 16.** Panel plots highlighting the features of a 2 pc piece of a molecular cloud after  $t = 1.08$  Myr of evolution. For all fields except the column density, the panel is showing the density-weighted projection. All projections are along the  $z$ -axis. The figure shows a simulated molecular cloud after 1 Myr of gravitational evolution including protostellar sink particles and radiation feedback from infrared to X-rays. While the EUV radiation is rapidly absorbed (indicated by the black background colour), the infrared and X-ray emission is able to penetrate much further into the cloud.

## 6 DISCUSSION/FUTURE WORK

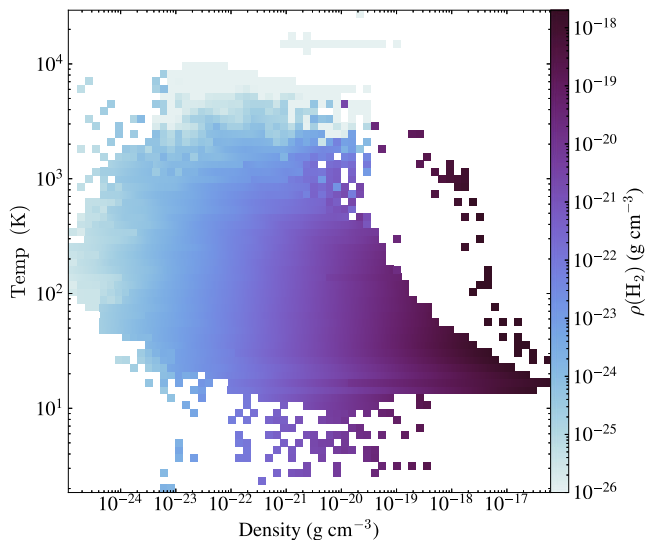
We have presented the new X-ray radiation transfer module, XRAYTHESPOT using the reverse ray-tracing scheme TREERAY implemented in FLASH (Wünsch et al. 2021). XRAYTHESPOT enables an arbitrary number of point or diffusive sources of X-ray emission, and an arbitrary number and position of energy bins. The module uses temperature-dependent cross-sections assuming gas in thermal ionization equilibrium. However, the module is flexible enough such that the user can provide their own cross-section data to be used. The module produces the expected behaviour for X-ray point sources and shadow tests and is able to reasonably reproduce the thermochemistry compared to CLOUDY, despite the significantly simpler treatment of X-ray chemistry and grain-processes in FLASH. Hence, the XRAYTHESPOT module allows the inclusion of, for instance, time-dependent feedback from protostars or X-ray binaries, or extended X-ray emission from hot, cooling gas with the X-ray

radiation transport coupled to the hydrodynamics, chemistry, and thermodynamics.

We demonstrated the utility of this module with two example science cases focusing on protostellar X-ray emission. First, we modelled the emission of an  $0.7 M_{\odot}$  protostar with an accretion rate of  $10^{-9} M_{\odot} \text{ yr}^{-1}$  through a protostellar disc. We find that soft X-rays are rapidly absorbed at the disc surface, with most of the emission escaping through the outflow cavity. However, harder X-rays are able to permeate the disc due to their significantly lower optical depth. The X-ray heating was also strong within the outflow cavity, with no X-ray heating towards the mid-plane of the disc, as expected. Secondly, we perform a low-resolution star formation simulation of a turbulent molecular cloud. In this simulation, protostars are self-consistently formed and the X-ray emission modelled on the fly. This simulation includes the entire range of different TREERAY radiation modules: diffuse FUV (OPTICALDEPTH; Wünsch et al. 2018), EUV (ONTHESPOT; Wünsch et al. 2021), thermal radiation



**Figure 17.** Panel plots highlighting the features of a 2 pc piece of a molecular cloud after  $t = 1.08$  Myr of evolution. The panels show the integrated gas surface density of  $\text{H}^+$ ,  $\text{H}$ , and  $\text{H}_2$  (top left, centre, and right) and  $\text{C}^+$ ,  $\text{C}$ , and  $\text{CO}$  (bottom left, centre, and right).



**Figure 18.** Density–temperature phase diagram, weighted by the  $\text{H}_2$  gas density. The phase diagram shows the typical qualitative behaviour of self-gravitating turbulent molecular clouds, with a population of atomic gas at  $T \approx 1.2 \times 10^4$  K and another population of dense, molecular gas with  $10^2 \leq T \leq 10^3$ , potentially resulting from X-ray heating.

and radiation pressure (RADPRESSURE; Kleitko et al. 2022) and X-ray emission from 1 to 10 keV. Since the X-ray emission in the simulation comes entirely from accretion on to the protostars, the X-ray emission is highly variable. Due to the lower resolution and the inclusion of ionizing radiation, the accretion occurs in bursts followed by the expansion of H II regions, which cut-off accretion. With higher resolution, accretion may still be able to occur through

discs, instabilities, and more porous density structures. In future work, we will perform higher resolution simulations to model star formation including chemistry and radiation feedback across the electromagnetic spectrum.

In this work, we focus primarily on point sources. However, XRAYTHESpot makes no differentiation between point sources versus extended more diffuse emission. Future studies will include diffuse X-ray emission from cooling hot gas and shocked gas. Our module currently includes the computation of X-ray emission from accretion on to protostars, and future work will include X-ray models for more types of point sources such as X-ray binaries. The module presented in this work will allow the first-generation of simulations of star formation and galaxies with the inclusion of a wide range of X-ray sources.

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by the DOE NNSA-ASC OASCR Flash Centre at the University of Chicago (Fryxell et al. 2000). The following PYTHON packages were utilized: NUMPY (Harris et al. 2020), SCIPY (Virtanen et al. 2020), MATPLOTLIB (Hunter 2007), YT (Turk et al. 2011), and CHIANTIPLY (Dere 2013).

## DATA AVAILABILITY

The data underlying this article will be shared on reasonable request to the corresponding author. The code used to compute the temperature-dependent cross-sections is publicly available at <https://github.com/AstroBrandt/XRayCrossSections>.

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## APPENDIX A: CLOUDY BENCHMARK SCRIPT

We present the CLOUDY script that was used for the X-ray benchmarking. The CLOUDY model consists of a uniform density medium with  $n_{\text{H}} = 10^3$  solved using the ‘sphere’ command. We turn off most induced and grain processes and set the abundances for most metals to zero to better match the methods used in our FLASH simulations.

```

1 title XDR source
2 ##radiation sources
3 CMB
4 ##Source
5 table SED "plaw.sed"
6 luminosity 35 range 73.5 to 735 Ryd
7 radius 16.1938
8 ##Density
9 hden 3.0
10 sphere
11 ##Stopping and iterate
12 stop H2 column density 24
13 stop temperature linear 3.0
14 iterate to convergence
15 ##Misc
16 abundances ISM
17 no grain qheat
18 no grain x-ray treatment
19 no induced processes
20 no radiation pressure
21 no scattering opacity
22 no grain molecules
23 no line transfer
24 element carbon abundance -3.853872
25 element helium abundance -1
26 element oxygen abundance -3.494850
27 element silicon abundance -7
28 element nitrogen off
29 element sulphur off
30 element neon off
31 element aluminium off
32 element phosphor off
33 element chlorine off
34 element argon off
35 element calcium off
36 element chromium off
37 element nickel off
38 element lithium off
39 element beryllium off
40 element fluorine off
41 element potassium off
42 element scandium off
43 element titanium off
44 element vanadium off
45 element manganese off
46 element cobalt off
47 element copper off
48 element zinc off
49 cosmic ray rate -16.523
50 ##output
51 set save luminosity old
52 save overview last "xdr.ovr"
53 save molecules last "xdr.mol"
54 save abundances last "xdr.abund"
55 save continuum last "xdr.cont"
56 save dr last "xdr.dr"
57 save PDR last "xdr.pdr"
58 save grain temperature last "xdr.dtemp"

```

Listing 1: Input file for CLOUDY benchmark.

This paper has been typeset from a  $\text{\TeX}/\text{\LaTeX}$  file prepared by the author.