HELIOS-K 2.0 OPACITY CALCULATOR AND OPEN-SOURCE OPACITY DATABASE FOR EXOPLANETARY ATMOSPHERES

SIMON L. GRIMM¹, MATEJ MALIK², DANIEL KITZMANN¹, ANDREA GUZMÁN-MESA¹, H. JENS HOEIJKMAKERS³, CHLOE FISHER¹,⁸, JOÃO M. MENDONÇA¹, SERGEY N. YURCHENKO⁴, JONATHAN TENNYSON⁴, FABIEN ALESINA⁵, NICOLAS BUCHSCHACHER⁵, JULIEN BURNIER⁶, DAMIEN SEGRANSAN⁷, ROBERT L. KURUCZ⁶, AND KEVIN HENG¹⁷

¹University of Bern, Center for Space and Habitability, Gesellschaftsstrasse 6, CH-3012, Bern, Switzerland. Emails: simon.grimm@csh.unibe.ch, kevin.heng@csh.unibe.ch
²Department of Astronomy, University of Maryland, College Park, MD 20742, USA
³National Space Institute, Technical University of Denmark, Elektrovej, 2800, Kgs. Lyngby, Denmark
⁴Department of Physics and Astronomy, University College London, London, WC1E 6BT, UK
⁵University of Geneva, Department of Astronomy, CH-1290 Versoix, Switzerland
⁶Center for Astrophysics | Harvard & Smithsonian, Cambridge, MA, 02138, USA
⁷University of Warwick, Department of Physics, Astronomy and Astrophysics Group, Coventry CV4 7AL, UK
⁸University of Bern International 2021 Ph.D Fellowship

ABSTRACT

Computing and using opacities is a key part of modeling and interpreting data of exoplanetary atmospheres. Since the underlying spectroscopic line lists are constantly expanding and currently include up to \( 10^{10} - 10^{11} \) transition lines, the opacity calculator codes need to become more powerful. Here we present major upgrades to the HELIOS-K GPU-accelerated opacity calculator and describe the necessary steps to process large line lists within a reasonable amount of time. Besides performance improvements, we include more capabilities and present a toolbox for handling different atomic and molecular data sets: from downloading and pre-processing the data to performing the opacity calculations in a user-friendly way. HELIOS-K supports line lists from ExoMol, HITRAN, HITEMP, NIST, Kurucz and VALD3. By matching the resolution of 0.1 cm\(^{-1}\) and cutting length of 25 cm\(^{-1}\) used by the ExoCross code for timing performance (251 seconds excluding data read-in time), HELIOS-K can process the ExoMol BT2 water line list in 12.5 seconds. Using a resolution of 0.01 cm\(^{-1}\), it takes 45 seconds—equivalent to about \( 10^7 \) lines per second. As a wavenumber resolution of 0.01 cm\(^{-1}\) suffices for most exoplanetary atmosphere spectroscopic calculations, we adopt this resolution in calculating opacity functions for several hundred atomic and molecular species, and make them freely available on the open-access DACE database. For the opacity calculations of the database, we use a cutting length of 100 cm\(^{-1}\) for molecules and no cutting length for atoms. Our opacities are available for downloading from https://dace.unige.ch/opacityDatabase and may be visualized using https://dace.unige.ch/opacity.

Keywords: planets and satellites: atmospheres

1. INTRODUCTION

Opacities² are the cross sections per unit mass of atoms, ions and molecules. They may be measured in the laboratory or calculated from first principles (by solving the Schrödinger equation), and are indispensable in the analysis of spectra across multiple subfields of astronomy and astrophysics. In the study of exoplanetary atmospheres, opacities are used in a number of fundamental ways:

1 Specifically, this term is used to describe specific forms of the cross section per unit mass involving photons/light/radiation and not electron or heavy-particle collisions.
Solving this computational problem requires the judicious design of software and its careful interfacing with hardware, including efficient memory management. In Grimm & Heng (2015), we took the first major steps towards producing an ultra-fast, open-source opacity calculator named HELIOS-K (https://github.com/exoclime/HELIOS-K), which is accelerated by the use of Graphics Processing Units (GPUs). In the current study, we report major upgrades to HELIOS-K that allow for further order-of-magnitude increases in computational speed.

As a service to the community and to promote scientific reproducibility and transparency, the computed opacities are publicly available on https://dace.unige.ch/opacityDatabase and https://chaldene.unibe.ch/. The former offers the possibility of obtaining customised opacities across user-specified arrays of temperature and pressure. The opacities may be visualised on the Swiss-run DACE database via https://dace.unige.ch/opacity, as we will describe. We note that the ExoMolOP database provides publicly available opacities as well (Chubb et al. 2020).

2. METHODOLOGY

The basic methodology of the HELIOS-K opacity calculator was published in Grimm & Heng (2015). The updates on the core methodology include mostly a new parallelization scheme and faster memory handling as well as new features of the code. In the following section, we summarize the supported line profiles in section 2.1 and the background theory in section 2.2, describe the difficult aspects of the usage of GPUs in section 2.3, and report the improvements and updates of HELIOS-K in section 2.4. In section 2.3, we describe in detail the handling of the different line list databases.

2.1. Line profiles

In most applications, molecular or atomic transition lines are approximated by a Voigt profile, which is a convolution of a Doppler and a Lorentzian profile. This convolution consists of an infinite integral, which is numerically expensive and requires optimized numerical methods to solve in a reasonable time. Besides the Voigt profile, sometimes also pure Doppler and Lorentzian profiles are used, and for comparison reasons we also implemented the “binned Gaussian integrated cross section” used by the ExoCross code (Yurchenko et al. 2018). HELIOS-K allows one to choose between these four profiles. Figure 1 shows an example of the available line profiles.

For some species, there are non-Voigt profiles available, e.g. Wcislo et al. (2016), but those are not implemented in the current version of HELIOS-K. However it is possible to replace individual transition lines with pre-calculated tables as necessary for certain alkali resonant lines.

![Figure 1. Available line profiles from HELIOS-K: Voigt, Lorentzian, Doppler and the binned Gaussian integrated cross section. The Voigt profile is numerically the most expensive to calculate.](image)

2.1.1. Removing the Plinth

HELIOS-K supports the option of removing the plinth (also called “base” sometimes) from transition lines. The height of the plinth is defined as the opacity value from a transition line at the cutting length position and depends on the line intensity and broadening parameters. (e.g. Clough et al. 1989; Paynter & Ramaswamy 2011; Ptashnik et al. 2012). Removing the plinth is necessary when opacities are used in combination with a background opacity which already includes the described plinth for each truncated transition line. This is the case, for example, when using the CKD (Clough et al. 1989), MT_CKD (Mlawer et al. 2012) or BPS (Paynter & Ramaswamy 2011) water vapor continuum as an opacity background. Not removing the plinth would lead to a double counting of it. An example of the plinth, and how it is removed from a transition line, is shown in Figure 2.

2.2. Background theory

We follow the definitions of the parameters described in Grimm & Heng (2015) and Heng (2017), and repeat only some of them here for a better overview. Molecular or atomic opacities can be approximated by Voigt line profiles, which can be described by two parameters, \( a \) and \( u \). These two

---

2 It should be noted that name of W. Voigt is usually mispronounced. The correct pronunciation is [fo:kt], i.e. the vowels are articulated “oo”, rather than the commonly used “oi”.

3 In the literature, these parameters are sometimes also called \( x \) and \( y \).
Figure 2. When truncated line wing opacities are used in combination with a far wing background, it may be necessary to remove the plinth from the opacities. The height of the plinth is defined as the opacity value at the cutting length position. Shown in blue is a full Voigt profile transition line, and in orange the transition line with the removed plinth. The plinth itself is shown in green.

Parameters for the Voigt line profiles are defined as

\[ a = \sqrt{\ln 2 \frac{\Gamma_L}{\Gamma_D}} \]  

(1)

and

\[ u = \sqrt{\ln 2 \frac{\nu - \nu_0}{\Gamma_D}}, \]  

(2)

where \( \nu \) is the wavenumber, \( \nu_0 \) is the line center in wavenumber in cm\(^{-1}\), and \( \Gamma_L \) and \( \Gamma_D \) are the Lorentz- and Doppler half-widths in cm\(^{-1}\), respectively. The Doppler half-width is given as:

\[ \Gamma_D = \frac{\nu_0}{c} \sqrt{\frac{2 \ln 2 k_B T}{m}}, \]  

(3)

where \( m \) is the mass in g (molar mass in g mol\(^{-1}\) divided by the Avogadro constant), \( k_B = 1.3806489 \times 10^{-16} \text{ cm}^2 \text{ g} \text{ s}^{-2} \text{ K} \) is the Boltzmann constant and \( T \) is the temperature in Kelvin.

The Lorentz half-width for HITRAN data is given as:

\[ \Gamma_L = \frac{A}{4\pi c} + \frac{T}{T_{ref}} \left[ \frac{\alpha_{\text{air}}(P - P_{\text{self}})}{P_{ref}} + \frac{\alpha_{\text{self}} P_{\text{self}}}{P_{ref}} \right], \]  

(4)

which is designed for studies of the terrestrial atmosphere so broadening is considered in two parts: air-broadening and self-broadening. For ExoMol data, the corresponding definition is (Tennyson et al. 2016):

\[ \Gamma_L = \frac{A}{4\pi c} + \alpha_{\text{ref}} \left( \frac{T_{ref}}{T} \right)^n \left( \frac{P}{P_{ref}} \right), \]  

(5)

where \( \alpha_{\text{ref}} \) gives the line broadening for the given broadening gas under consideration. In both cases, we have included the first term with the Einstein A-coefficient (which has physical units of s\(^{-1}\)), but it is generally negligible.

For atomic opacities, the Lorentz half-width consists mainly of a combination of the natural broadening coefficient and pressure broadening due to collisions with electrons (Stark broadening) and other atomic and molecular species (van der Waals broadening). Stark broadening depends directly on the abundance of free electrons and is, therefore, only important in hot stellar atmospheres. In this work, we only consider natural broadening for atomic opacities.

2.2.1. Natural broadening

The natural broadening of atomic transition lines is a consequence of the uncertainty principle \( \Delta E \Delta t \geq \hbar/2 \) (Kunze 2009, p. 156; Draine 2011, p. 57), and applies to both the upper and lower energy levels (Draine 2011, p. 57),

\[ \Gamma_{\text{nat},ij} = \sum_{E_j < E_u} A_{uj} + \sum_{E_j < E_l} A_{lj}. \]

(7)

For example,

\[ \Gamma_{\text{nat},31} = A_{31} + A_{32} \]

or

\[ \Gamma_{\text{nat},32} = A_{31} + A_{32} + A_{21}. \]

For Kurucz data we use

\[ \Gamma_L = \Gamma_{\text{nat}} = \frac{10^{\Gamma_u}}{4\pi c}. \]

(8)

The quantity \( \Gamma_R \) is tabulated in the Kurucz database.

2.2.2. Line intensity

Besides the Voigt line profile, the intensity \( S \) of the transition line is also needed. It is defined in units of cm g\(^{-1}\) as (e.g. Hill et al. 2013 or Chapter 5.3 of Heng 2017): 5

\[ S = \frac{A}{8\pi c \nu_0^3 Q(T)} \left( \frac{\nu - \nu_0}{\nu_0} \right) \left( 1 - e^{-\frac{\nu - \nu_0}{\nu_0}} \right), \]

(9)

where \( E'' \) is the lower state energy in cm\(^{-1}\), \( g' \) is the upper state statistical weighting and \( c = 2.99792458 \times 10^{10} \text{ cm s}^{-1} \) is the speed of light. Note that there is a typographical error in the first exponent of equation 17 of Grimm & Heng (2015), which we correct here.

5 Note that the equation in Hill et al. (2013) is defined in units of cm\(^{-2}\)/molecule, without the mass term in the denominator.
The partition function $Q(T)$ is in general available in a tabulated form, together with the line list data. If the partition function is not available, it may be computed using (e.g. Sharp & Burrows 2007)

$$Q(T) = \sum_{i=1}^{n} g_i e^{-\frac{hcE_i}{kBT}}, \tag{10}$$

where $g_i$ and $E_i$ are the statistical weight and the energy of the level $i$, respectively. Note that, as Sharp & Burrows (2007) write, this summation diverges formally for high pressure values. For pressure values that we are typically interested in, the sum is still accurate enough. For molecular species, the partition function would also include electronic, vibrational, and rotational terms (Sharp & Burrows 2007).

### 2.2.3. Decompose the $\alpha$-$\nu$ Voigt parameter plane

Following Letchworth & Benner (2007) and as already described in Grimm & Heng (2015), we split the $\alpha$ and $\nu$ parameter space into three different regions, called A, B and C, and use different methods and approximations for the Voigt profile calculations. For the A region, we use a first-order Gauss-Hermite quadrature approximation. For the B region, we use a third-order approximation. For the C region, we solve the Voigt profile up to machine precision, with an iterative method (Zaghloul & Ali 2011). In practice, we split the A-region into a left (AL) and a right part (AR) in order to avoid many zero calculations at the line center. We do not split the B region in the same way, because most of the time the C region is very narrow. Splitting B would introduce too many unnecessary memory accesses, especially when the $\alpha$-$\nu$ regions are applied to thousands of neighbouring transition lines, and the B and C regions can overlap. The used $\alpha$-$\nu$ regions are listed in Table 1, and illustrated in Figure 3. Figure 4 shows the fractional difference between the HELIOS-K Voigt calculation and the SciPy function wofz (scipy.special.wofz). The wofz function is similar to the algorithm in our region C. The largest error appears at the inner boundary of region B, where the Voigt function is approximated by a third order Gauss-Hermite quadrature. The precision of HELIOS-K could be improved if necessary by expanding region C to larger values on the $\nu$-axis.

Note that Letchworth & Benner (2007) split the C region into different subregions with higher order quadrature methods, or interpolation of pre-calculated tables. But we find that by using GPUs, the three regions from table 1 provide a good balance between efficiency of the method, parallel applicability and resource limitations on the GPU.

### 2.3. GPU implementation considerations

Using GPUs for numerical calculations is a very powerful approach. Modern GPUs can consist of more than 5000 CUDA cores, which allow us to parallelize the numerical problem at a high level. However, there are some limitations. First, not all CUDA threads are independent and threads in the same warp (a collection of 32 threads) must perform the same operation but can use individual data (single instruction, multiple threads, SIMT). This means that thread divergencies must be avoided. In our problem, thread divergencies could occur when neighbouring transition lines lie in different $\alpha$-$\nu$ regions and require a different algorithm to calculate the Voigt profile. To avoid those divergencies, we have to sort the data and organize them in a well-suited form.

A second limitation of the GPUs is the memory transfer bottleneck. Before data can be used on the GPU, it must be transferred to the GPU through the PCI (peripheral component interconnect) express, which can cause a large computing overhead time. However, the data transfer time can

| $\alpha$-$\nu$ Region | Limits | Method |
|----------------------|--------|--------|
| AL | $10^6 < \alpha^2 + \nu^2$ | $\nu < 0$ | first order Gauss-Hermite quadrature |
| AR | $10^6 < \alpha^2 + \nu^2$ | $\nu > 0$ | first order Gauss-Hermite quadrature |
| B | $100 < \alpha^2 + \nu^2 < 10^6$ | | third order Gauss-Hermite quadrature |
| C | $\alpha^2 + \nu^2 < 100$ | | Algorithm 916 (Zaghloul & Ali 2011) |

Table 1. The $\alpha$-$\nu$ plane is split into different regions, where different approximations on the Voigt profile calculations are applied. We find this splitting to be very efficient for GPU calculations, where branch divergencies should be avoided, and memory access should be minimized.

Figure 3. Example of the $\alpha$-$\nu$ regions, applied on a single transition line, with a cutting length of 1 cm−1. In practice, thousands of neighbouring transition lines are grouped together and the regions, especially B and C, can overlap.
be hidden by using asynchronous and interleaved memory transfer. This is possible because the GPUs have separated copy- and compute engines, and both can work simultaneously. Besides the memory transfer, CUDA also allows us to overlap kernel execution with calculations performed on the CPU. In our problem, we can therefore hide data reading and data transfer behind the opacity calculations on the GPU. But that only works if the GPU part takes long enough. That is typically the case when using high wavenumber resolution calculations ($\Delta \tilde{\nu} \approx 0.01 \text{ cm}^{-1}$). If this is not the case, then the full computing time is dominated by pure data reading.

A further limitation is the slow global memory access. Nvidia GPUs have different levels of memory, mainly: global memory, shared memory and registers. Most of the time, memory access goes through global memory first and is then stored temporarily in shared memory or registers. Global memory access is slow and should be avoided. This can be achieved by designing the parallelization scheme in such a way that data can be reused multiple times from registers or shared memory.

2.4. Improvements and Upgrades to the HELIOS-K opacity calculator

This section describes details of the parallelization and new implementations of HELIOS-K. Readers who are not interested in these coding details can skip this section and continue with the different line list databases in section 2.5.

2.4.1. Reading the line list data

Before the opacity calculation can be started, all of the line list data need to be read from the CPU and transferred to the GPU. We use a binary data format to store pre-processed line lists to speed up this reading process. How the line lists are pre-processed is described later in section 2.5. To read the data from the hard disk sounds simple, but actually one has to be very careful not to lose too much performance. In fact, when using low to moderate resolution in wavenumber, reading the line list can take more time than the opacity calculation itself. When the code is called many times, e.g. for different values of temperature and pressure, reading and writing of data can become the bottleneck. We use the following strategy to improve the reading and data transfer process:

- Avoid reading data line by line. Reading the line list, e.g. by an `fscanf` command line by line, would not generate enough work to hide the overhead time and the entire memory bandwidth would not be used. Instead, we read a bigger block of data, consisting of a few thousand transition lines at once by using the `fread` command and a memory buffer.
- Use asynchronous and interleaved memory copy to transfer the data to the GPU. In this way, one block of data is transferred to the GPU at the same time as the CPU reads in the next block.
- We overlap the reading and memory transfer with the opacity calculation of the previous block of data. Here we use the fact that the compute engine can work simultaneously with the copy engine and that the CPU can work in parallel to the GPU (see section 2.3). In order to make this work, a clever scheduler of the opacity-kernel calls and memory transfer commands is needed. This is described in detail in section 2.4.6 and illustrated in Figure 6.

2.4.2. Code parameters for the Voigt profile calculation

In practice, we do not use the parameters in equations (1) to (4) in their stated form, but instead cancel out the $\sqrt{\ln 2}$ factor in the numerator and denominator and directly store the inverse of the Doppler half-width, because the calculation of a product is faster than a division. The parameters used are then:

$$x = (\tilde{\nu} - \tilde{\nu}_0) \cdot \alpha_D^{-1},$$

and

$$y = \Gamma_L \cdot \alpha_D^{-1},$$

with

$$\alpha_D^{-1} = \frac{c}{\tilde{\nu}_0} \sqrt{\frac{m}{2k_B T}}.$$  

The parameters $x$, $y$ and $\alpha_D^{-1}$ correspond to the parameters $u$, $a$ and $\Gamma_D$ from equations (1) – (3). We introduce here

---

6 Nvidia GeForce cards use typically the same copy engine for data transfer for host-to-device and for device-to-host. Tesla cards have separate copy engines for host-to-device and for device-to-host, and can work simultaneously.
a new notation for the parameters which closely follows the implementation in the code. Since an ideal code implementation reduces the number of operations in each equation to the minimum, it is necessary sometimes to redefine parameters to be slightly different from their theoretical derivations.

When a constant resolution $\Delta \tilde{\nu}$ is used, then the parameter $x$ can be calculated very fast with a single fused-multiply-add operation as

$$x = d + i \cdot b,$$  \hspace{1cm} (14)

with a thread index $i$, and the variables $d = (\tilde{\nu}_{\text{min}} - \tilde{\nu}_0) \cdot \alpha_D^{-1}$ and $b = \Delta \tilde{\nu} \cdot \alpha_D^{-1}$. The quantity $\tilde{\nu}_{\text{min}}$ is the starting point in wavenumber.

Assuming a constant resolution in wavenumber is a natural choice, because the wavenumber points follow nearly the positions of the transition lines. In HELIOS-K, it is possible to use a different wavenumber resolution for different bands. In that case the variable $x$ in equation (14) is stored in a pre-calculated array.

In summary, the calculation of the Voigt profile needs a total of five parameters, which depend on the transition line and the wavenumber: the line intensity $S$, the quantities $b$, $d$ and $y$ as defined above, and an additional quantity $\tilde{\nu}_{\text{cut}}$ which handles the cutting length. The result of the opacity calculation is written into an array $K$. The major difficulty is how to handle these five parameters efficiently within GPU memory and use them in a parallel way.

2.4.3. Calculating the line parameters

As described in section 2.4.2, we need five line parameters, $S$, $b$, $d$, $y$ and $\tilde{\nu}_{\text{cut}}$. The calculation of these parameters is split into four different parts.

1. The line intensity (Equation 9) may be written as

$$S = \frac{S'}{Q(T)} e^{-\left(\frac{\tilde{\nu} - \tilde{\nu}_0}{\tau} \right)^2} \left(1 - e^{-\left(\frac{\tilde{\nu} - \tilde{\nu}_0}{\tau} \right)}\right),$$  \hspace{1cm} (15)

where the term $S'$ does not depend on the temperature or pressure,

$$S' = \frac{g' A}{8\pi c \tilde{\nu}^2 m}.$$  \hspace{1cm} (16)

This quantity $S'$ can be calculated beforehand and is part of the pre-processing routine, which is described below in section 2.5. In this step, the partition function $Q(T)$ has to be determined by reading and interpolating the tabulated values provided by the line list databases.

2. The next step takes place after a block of line list data is transferred into a memory buffer on the GPU. The parameters $S$, $y$ and temporarily the reverse Doppler half-width $\alpha_D^{-1}$ are then calculated on the GPU. We use one thread per transition line for these calculations and the quantities are stored in arrays in global memory on the GPU.

3. The line list data obtained are mostly sorted at this point, but can be slightly out of order due to pressure shift effects. For that reason, we sort the quantities along $\tilde{\nu}$ using the CUDA Thrust library$^7$.

4. In the last step, the quantities $b$, $d$ and $\tilde{\nu}_{\text{cut}}$ are calculated, where $\tilde{\nu}_{\text{cut}}$ depends on how the cutting length is defined. Options include a fixed cutting length, a fraction of the Doppler half-width or a fraction of the Lorentzian half-width.

With this way of splitting the calculation of the parameters, we can reduce the sorting step to a minimum.

2.4.4. Parallelizing the opacity calculations

The opacity calculation can be parallelized in two dimensions: the wavenumber $\tilde{\nu}$ and the transition line index $id$. Therefore, we divide the $\tilde{\nu}$-$id$ plane into data tiles. In this way, we can efficiently use all GPU threads to first read in each thread block of data from global memory into registers and then iterate on the points in wavenumber with one or multiple threads per transition line. How many threads to use for this process is not only dependent on the parameters, but also on the GPU type and is therefore hard to estimate beforehand. For that reason, we use a self-tuning routine, which periodically checks for the best number of threads to use, and adapts the CUDA-kernel calls automatically.

The top panel of Figure 5 gives a schematic of the parallelization scheme for an example with four transition lines and four points in wavenumber (-0.5, 0.0, 0.5 and 1.0). When we use four threads in this example, each thread is assigned to a different transition line (different color in the Figure) and has to iterate four times around the points in wavenumber. It is important that each thread calculates a different wavenumber point during the same iteration, because otherwise it would generate a race condition, which would lead to incorrect results. To prevent that, we can shift the starting point by the line index and take the modulo operator as:

$$\tilde{\nu}\text{-index} = (\text{line-index} + \text{iteration-index}) \pmod{(\text{number of threads})}. \hspace{1cm} (17)$$

In the example from Figure 5, e.g. the thread 2 would then calculate the opacities in the order: (0.5, 1.0, -0.5 and 0.0). It is also possible to use more than four threads for the example above, as shown for eight threads in the bottom panel of Figure 5. But then we have to use separate arrays to collect the results of the different threads, and merge the results later, because e.g. iteration b of thread 1 happens at the same time as iteration b of thread 3, and both threads write to the same point in wavenumber. Using eight threads instead of

$^7$ https://developer.nvidia.com/thrust
four increases the amount of parallel work performed by the GPU, but also increases the fraction of reading vs. computing, which means global memory access cannot be hidden behind the calculation time. Which version is the fastest depends on the individual case. Our algorithm self-tunes to find the best option.

Listing 1 shows a pseudocode for a GPU kernel which reads the transition line parameters and calls the Voigt profile function. The parameter NBx sets the number of iterations each thread has to perform. The parameter nl is the number of transition lines in a block of data. The parameters iil and nstart define the starting points of the transition line index and wavenumber index of the current block of data. The kernel is launched on a two dimensional grid. The x-dimension of that grid is defined as the maximum range of wavenumber points of the current data block divided by NBx. The y-dimension is used to launch multiple blocks of data together, in order to reduce the kernel launch overhead time. The y-dimension is on the order of 16 or 32.

In lines 12 - 16 of Listing 1, the five transition line parameters from section 2.4.2 are loaded from global memory into registers. The values in those registers can be reused into registers. The values in those registers can be reused

are merged into a single array, which can then be written back into global memory, and at the very end copied back to the CPU.

Listing 1. Pseudocode of the opacity calculation kernel.

### 2.4.5. Calculating the \( \bar{\nu} \) limits

As described before, the transition lines are grouped together into blocks of data, where each block consists of a few thousand lines. In order to call the Voigt calculation kernel, we have to determine the range of wavenumbers of each block. This range is defined as either the maximum or the minimum of the cutting range, or the limits of the \( a-u \) regions described in Table 1. The limits \( \Delta \bar{\nu} \) can then be calculated with the equation

\[
a^2 + u^2 = l^2,
\]

where \( l \) is either 10 or 1000, for the regions B and C. Inserting equations (1), (2) and (13) leads to

\[
\frac{\Gamma_L^2}{\alpha_D^2} + \frac{(\bar{\nu} - \nu_0)^2}{\alpha_D^2} = l^2
\]

and

\[
\Delta \bar{\nu} = (\bar{\nu} - \nu_0) = \sqrt{l^2 \cdot \alpha_D^2 - \Gamma_L^2}.
\]

Therefore the limits of the B and C regions for an individual transition line are \( \bar{\nu}_0 \pm \Delta \bar{\nu} \). The limits of the A region is

![Figure 5. Schematic for parallelizing the opacity calculation for an example with four transition lines and four points in wavenumber. In the top panel, four threads are used and each thread has to iterate through the points a, b, c and d. In the bottom panel, eight threads are used and each thread has to iterate through the points a, b, c and d. In the second case, the results have to be collected in two different arrays to avoid race conditions. The transition lines are cut at a distance from the line center of 0.9 cm \(^{-1}\).](image)
simply $\tilde{\nu}_0 \pm \tilde{\nu}_{\text{cut}}$, where $\tilde{\nu}_{\text{cut}}$ can depend on the Doppler or Lorentz half-width. In order to know the $\tilde{\nu}$-limits of an entire block of transition lines, we use a parallel reduction method on the GPU by using warp shuffle instructions.

2.4.6. The core workflow

Typically, the calculation of the Voigt function is the longest part of the GPU calculation, and we want to use that execution time to simultaneously read the next block of data from the hard disk and transfer it to the GPU. This is possible because different GPU engines can work at the same time, and because GPU kernel calls are asynchronous, meaning that the workflow goes back to the CPU immediately after the kernel is called and does not wait for completion. (See section 2.3). The resulting workflow is shown in figure 6 and can be described as the following:

1. First, the line properties have to be calculated. This operation scales with the number of transition lines and is split into several different kernels, but can be described mainly with two operations: 1) Calculate the line intensity (“L”, green box in Figure 6) and 2) organization of the tiles described in section 2.4.4 (“I”, yellow boxes). The “I” operation has to wait until “L” is finished, but by using multiple streams, parts from “L” and “I” can overlap on the GPU. This is indicated in Figure 6 by stacking boxes on top of each other.

2. The calculation of the Voigt profile (blue boxes) is split into four different parts (A left, A right, B and C; Table 1). Before the calculation starts, the CPU determines the total number of threads for the next block of data by using the information from the “I” operation. This part is indicated in the figure as “S”. Immediately after “S”, the GPU kernel starts, and the CPU can be used for the next part. At the end of the four “K” parts, temporary results are collected (“A”, red boxes).

3. While the GPU is busy, we use the CPU to read future data blocks from the hard disk. We have to interrupt this operation as soon as the “K” kernel on the GPU is finished. In practice, we use a CUDA event query for this check.

4. The new data is transferred asynchronously to the GPU (“C”, orange boxes) and its execution time is hidden almost completely behind the reading operation or GPU execution time.

5. During the kernel execution, the self tuning routine is called periodically to optimize the performance of the GPU, not shown in Figure 6.

The overall performance highly depends on if the data reading is faster (scenario a) or slower (scenario b) than the Voigt profile calculation in the GPU, (Figure 6 panels a and b). Scenario b can happen when the resolution in wavenumber $\Delta\tilde{\nu}$ is small, the GPU is very fast or the data access from the hard disk is limited. Data access speed can in general be improved by moving the data closer to the calculation unit, and ideally keeping data files in memory. If this is not possible, then opacities for multiple points in pressure should be calculated at once, in order to reduce memory access. Some performance results using different data storage locations are listed in section 3.5.

2.5. Preparation of spectroscopic line list data for computation

For the opacity calculations, we use line lists from several databases, which all use a different data format and support a different way of downloading the data. In order to use HELIOS-K in the most efficient way, we define a uniform binary data format for the line lists, and provide scripts for automatically downloading the data and converting them to this format. Besides the data format, we also define a parameter file for each species, which contains the meta data of the species.

Our procedure is to process all transition lines from the given line lists. We do not use a truncation of weak lines. The line intensities are calculated in double precision floating point numbers. However, the final opacity function is reduced to single precision to reduce storage space, which truncates the opacity function to about $10^{-50} - 10^{-40}$ cm$^2$ g$^{-1}$. If a truncation of weak lines is required to reduce the total number of transition lines, it could easily be included in the HELIOS-K preparation scripts.

In this section, we describe the file structures that we use for HELIOS-K and outline how the data from different databases can be pre-processed and used.

2.5.1. Parameter files for species

For each species (atom or molecule) we define a parameter file, which contains all necessary meta data and file structure information. These parameter files can be produced automatically and allow HELIOS-K to process the given data in a simple way, especially when the data from a database is split into many different subfiles. An example of a species parameter file for HITRAN H$_2$O is given in Table 2. The file lists all isotopologues or isotopes of the species with the corresponding abundances, molecular masses and partition function file names.

2.5.2. Binary file structure

The different databases provide their data files in different structures and also with a different number of parameters. We define a binary data format for the atomic or molecular transition lines. Depending on the database, these binary files contain different values, as indicated in Table 3. The possible values are described in Table 4. In the binary files, we pre-process the data in such a way that the final opacity calculation can be performed as fast as possible. One important
Figure 6. Simplified workflow of the code for two cases: a) fast memory access and b) slow memory access. The boxes indicate the compute time for all involved operations on the CPU (light and dark grey) and the GPU (colored). When boxes are stacked on top of each other, then the two operations are executed in parallel at the same time. In case a) the compute time is dominated by the opacity calculation and in case b) it is dominated by reading the data from the hard disk. The case b) should be avoided.

part is the pre-calculation of the temperature- and pressure-independent part of the line intensity $S'$, defined in Equation (16).

2.5.3. ExoMol

The ExoMol line lists (Tennyson et al. 2020) can be accessed from www.exomol.com and are organized by molecules, isotopologues and line list names. HELIOS-K provides a Python script to scan the webpage and produce a list of all available species. The species from ExoMol can be identified by their full species name, which includes the isotopologue information and the line list name, e.g. “1H2-16O_BT2”. Each species has a “.def” file, which contains properties like the isotopologue mass, default values of Lorentzian half-widths, default values of temperature exponents, or the number of data files. It is recommended to check the number of files and maximal wavenumber of the line list. The data itself are separated into a “.states” file and one or multiple “.trans” files. This data format is very compact to store on a server, but for using it with HELIOS-K we need to produce a new line list, which contains $[\nu, S', E_L, A]$. This list is stored compactly as one or multiple binary files, which can be read by HELIOS-K. We provide a script that does the download and file conversion automatically. Some molecules contain the wavenumber of the molecular lines as a fourth column in the “.trans” files. However, it is not recommended to use that, but instead to calculate it through the difference in energy levels. Partition functions are given in “.pf” files.

For some species, ExoMol also provides pre-calculated cross-sections (xsec). When available, we compare our calculations with that data to verify our calculation (See section 3.2).

2.5.4. ExoMol super-lines

For some molecules, ExoMol provides super lines (Yurchenko et al. 2017a; Tennyson et al. 2020). Super lines are temperature dependent collections of line intensities, integrated over small wavenumber bins. By using super lines, the total number of Voigt profile calculation can be reduced dramatically. The drawback is that individual transition lines are not represented anymore, and all lines are calculated with the same default pressure broadening values. Practically, it means that the quantities $S$ and $S'$ from equations (15) and (16) do not need to be calculated anymore but can be read in directly. Only the mass term needs to be included in the super line intensities. For example, the use of superlines reduces the ExoMol POKAZATEL water line list from $5.7 \times 10^9$ transitions to $8.3 \times 10^6$ super lines.

2.5.5. HITRAN and HITEMP
Table 2. Example species-parameter file for HITRAN H\textsubscript{2}O. The HELIOS-K species-parameter files contain all relevant meta parameters necessary for the opacity calculation. The files can be generated automatically with the provided scripts.

| Parameter | Description | Unit | Type |
|-----------|-------------|------|------|
| ID        | species identity | char4 |      |
| \(\tilde{\nu}\) | center of the line | cm\(^{-1}\) | F64  |
| \(S'\)   | Equation 16 | cm g\(^{-1}\) | F64  |
| \(E_L\)  | lower energy level | cm\(^{-1}\) | F64  |
| \(A\)    | Einstein A-coefficient | s\(^{-1}\) | F64  |
| \(\delta_{Air}\) | pressure dependent line shift | cm\(^{-1}\) atm\(^{-1}\) | F64  |
| \(\gamma_{Air}\) | air broadening coefficient | cm\(^{-1}\) atm\(^{-1}\) | F64  |
| \(\gamma_{Self}\) | self broadening coefficient | cm\(^{-1}\) atm\(^{-1}\) | F64  |
| \(n\)    | exponent of the temperature dependent Lorentz half-width | | |
| \(\Gamma_{nat}\) | natural broadening coefficient | s\(^{-1}\) | F64  |

More details can be found at https://hitran.org/docs/definitions-and-units/

Table 3. Values contained in the HELIOS-K binary data files, depending on the database used. The parameters are described in Table 4.

The HITRAN database\(^8\) (Gordon et al. 2017; Rothman et al. 2010) provides line lists in standard formatted text files (*.par files). These files can be downloaded for a single isotopologue or a mix of different molecules and isotopologues. In addition to the line list, HITRAN provides partition functions and molecular meta data in a webpage table. HITRAN supports a programming interface, HAPI (Kochanov et al. 2016), which allows data access through a Python script. However, several tests showed that the data in HAPI were not always updated and therefore we opt to use the content of the webpage table directly. For usage with HELIOS-K, we extract only the data needed from the line lists and store them in more compact binary files which contain \([ID, \tilde{\nu}, S', E_L, A, \delta_{Air}, \gamma_{Self}, n]\).

2.5.6. Kurucz

The Kurucz database\(^9\) (Kurucz 2018, 2017) provides line lists and partition functions for a large number of neutral and ionized atoms. In this work, we use the GFNEW lines in wavenumber (gfllw08oct17.dat), which includes for some species isotope fractions and hyperfine splittings. HELIOS-K provides a Python script to download the data file and the available partition functions and to extract the relevant data for the opacity calculations. The Kurucz database includes natural broadening (radiation dampening), as well as Stark and van der Waals broadening coefficients. The

\(^8\) www.hitran.org
\(^9\) http://kurucz.harvard.edu
Stark broadening coefficient in the Kurucz database is given at a temperature of 10,000 K, in units of the electron number density. The van der Waals coefficients, on the other hand, are stated also for a temperature of 10,000 K, but in units of the atomic hydrogen number density. Van der Waals broadening in the Kurucz database is based on calculations, employing a modified version of Unsöld’s approximation (Unsöld 1938; Aller 1963). The tabulated coefficients can be adapted to collisions with He and H₂ by scaling them with a factor of 0.42 or 0.85, respectively.

In the current version, we only include natural broadening in the opacity calculation. The resulting atomic opacities are therefore suitable for environments where collisional broadening is negligible—such as atomic absorption lines originating from upper atmospheres.

**Partition functions** — For most species, partition functions are available from the Kurucz database. If this is not the case, then we calculate the partition functions according to equation (10) and the energy levels from the NIST database (see section 2.5.7).

**Isotope fraction correction** — During the data processing, we noted an issue in the isotope fractions with the file gfallwn08oct17.dat. When a transition line consists of multiple isotopes, but only one isotope is split up into its hyperfine structure, then the file contains an incorrect isotope fraction of the remaining isotopes. We show an example of a potassium transition line in Table 5. Potassium consists of three natural isotopes: ³⁹K (93.3%), ⁴⁰K (0.0117%) and ⁴¹K (6.7%). In the example, isotope ³⁹K is split in its hyperfine structure, while isotopes ⁴⁰K and ⁴¹K are combined into an entry with an isotope fraction of 1.0, which is not correct. The combined isotope fraction should instead be 6.7117%. If this factor is not corrected, then the resulting opacity can be significantly overestimated. Our HELIOS-K data preparation script scans all transition lines for these types of missing factors and corrects them.

**Natural broadening coefficients** — Natural broadening coefficients are provided for many species, but not for all. We compute the missing natural broadening coefficients according to equation (7). The natural broadening coefficient Γ_{nat,ij} between an upper state i and a lower state j includes the Einstein A-coefficients from all transitions between two other levels k and l with either

\[
\begin{align*}
\langle E \rangle_i &= \langle E \rangle_k \\
g_i &= g_k \\
\langle E \rangle_j &= \langle E \rangle_k \\
g_j &= g_k,
\end{align*}
\]

where, \( \langle E \rangle \) are the energy states averaged over the hyperfine sublevels and \( g_{j,k} \) are the statistical weights from the Kurucz database. We compare our calculations to the tabulated values from Kurucz and with published values from resonance lines (Morton 2003, 2008), as shown in Figure 7 for Fe. For most lines, our values correspond well with those from Morton (2003, 2008), but for a few transition lines, there are differences. Our calculated values are close to the ones from Kurucz, but for many lines we cannot exactly reproduce them. The reason could be that we do not have full access to all relevant Einstein A-coefficients or line configurations. The described comparison allows us to estimate the quality of calculated natural broadening coefficients for species where there are no tabulated values available.

| \( \tilde{\nu} \) [cm⁻¹] | isotope | hyperfine fraction | isotope fraction |
|--------------------------|---------|-------------------|-----------------|
| 5554.11                  | 39      | 0.031             | 0.933           |
| 5554.11                  | 39      | 0.437             | 0.933           |
| 5554.11                  | 39      | 0.062             | 0.933           |
| 5554.11                  | 39      | 0.156             | 0.933           |
| 5554.11                  | 39      | 0.156             | 0.933           |
| 5554.11                  | 0       | 1.0               | 1.0 (wrong)     |
| 5554.11                  | 39      | 0.156             | 0.933           |

Table 5. Example of a Potassium transition line with a wrong isotope fraction in the Kurucz database. The missing fraction must be calculated to get correct opacities. The isotope fraction should be 0.067117. Similar issues occur for several species and various transition lines. The numbers are rounded and taken from the file gfallwn08oct17.dat from http://kurucz.harvard.edu.

**Figure 7.** Comparison of the natural broadening coefficients Γ_{nat} of Fe between tabulated values from Kurucz (light blue and dark blue dots), calculated values using the Einstein A-coefficients (grey error bars and crosses) and published data (Morton 2003, 2008) for resonance lines (blue circles).
2.5.7. NIST

The NIST database\(^{10}\) (Kramida et al. 2019) provides line lists and energy levels for a large number of neutral and ionized atoms. The database can be accessed through a web interface by selecting the desired species and additional configuration options. The web access is intuitive and simple, but it is not very practical for accessing in an automated way, and it can be very time consuming to download multiple line lists by hand. Therefore, we provide a Python script that navigates the web page automatically and fills in the necessary web forms to download the line lists and energy level files. The script also extracts the needed quantities from the files and converts them into the standard HELIOS-K format.

A potential problem of the NIST database is that energy levels and transition lines can be listed multiple times. We noted this especially for the hydrogen atom, where energy levels are listed for individual angular momentum quantum numbers, besides the total averaged energy levels. Simply processing the entire line list would lead to duplicated transition lines, and thus it is necessary to filter the line lists for such duplicated entries.

Partition functions — We calculate the partition functions by the available energy levels according to equation (10). As Sharp & Burrows (2007) note, this summation diverges formally for high pressure values. For pressure values that we are typically interested in, the sum is still accurate enough.

Natural broadening coefficients — The natural broadening coefficients are not provided by the NIST database, therefore we use equation (7) to calculate them by summing over all relevant Einstein A-coefficients. The natural broadening coefficient \(\Gamma^\text{nat,ij}\) between an upper state \(i\) and a lower state \(j\) includes the Einstein A-coefficients from all transitions between two other levels \(k\) and \(l\) with either

\[
\begin{align*}
\text{configuration}_i &= \text{configuration}_k \\
\text{term}_i &= \text{term}_k \\
J_i &= J_k
\end{align*}
\tag{21}
\]

or

\[
\begin{align*}
\text{configuration}_j &= \text{configuration}_k \\
\text{term}_j &= \text{term}_k \\
J_j &= J_k
\end{align*}
\tag{22}
\]

where “configuration”, “term” and \(J\) are given in the NIST database.

We compare our values with published values for resonance lines due to Morton (2003, 2008) as shown in Figure 8 for Fe. Differences between the NIST data and Morton (2003, 2008) could occur because the underlying transition line data could be different.

The pre-processing of the NIST data requires the following steps:

1. Download the energy levels for all species. This can either be done manually through the web interface, or by a script, which navigates the web page automatically.

2. Calculate the partition function. Alternatively, the partition function would be available through the web form, but must be queried for each temperature individually.

3. Download the line lists for all species. Similar to the energy levels, this can either be done manually through the web page or by automated script.

4. Calculate the natural broadening coefficients, and generate the HELIOS-K binary files.

2.5.8. VALD3

The VALD3 database\(^{11}\) (Ryabchikova et al. 2015; Pakhomov et al. 2019) provides line lists for a large number of neutral and ionized atoms. After undergoing a registration process on the website, the database can be accessed through a web interface. Similar to our process for the NIST database, we provide a script to navigate the website automatically. For every data request, VALD sends an email with a link to download the data. By requesting multiple species, this process can result in a large number of emails and data-links. To

\(^{10}\)https://www.nist.gov/pml/atomic-spectra-database

\(^{11}\)http://vald.astro.uu.se/
simplify the download, our script can be used without opening all the sent emails, and it converts the line lists to the standard HELIOS-K format.

**Partition functions** — Partition functions are not available in the VALD database. Therefore, we use the partition functions calculated from the NIST database.

**Natural broadening coefficients** — Natural broadening coefficients are provided for many species, but not for all. We compute the missing natural broadening coefficients as for the Kurucz database according to equation (7). The natural broadening coefficient $\Gamma_{\text{nat},ij}$ between an upper state $i$ and a lower state $j$ includes the Einstein A-coefficients from all transitions between two other levels $k$ and $l$ with either

$$
\begin{align*}
E_i &= E_k \\
g_i &= g_k
\end{align*}
$$

or

$$
\begin{align*}
E_j &= E_k \\
g_j &= g_k
\end{align*}
$$

where the energy $E$ and the statistical weight $g$ are given by the VALD database. We compare our calculations to the tabulated values from VALD3 and with published values from resonance lines (Morton 2003, 2008) as shown in Figure 7 for Fe. For most lines, our values correspond well with those from Morton, but for a few lines there are differences. Our calculated values are close to the ones from VALD3, but for many lines we cannot exactly reproduce them. The reason could be that we do not have full access to all relevant Einstein A-coefficients or line configurations. The comparison described allows us to estimate the quality of calculated natural broadening coefficients for species where there are no tabulated values available.

The pre-processing of the VALD3 data requires the following steps:

1. Request the data download for all species. This can either be done manually through the web interface, or by a script, which navigates the web page automatically.

2. Use the partition functions from the NIST database.

3. Calculate the natural broadening coefficients, and generate the HELIOS-K binary files.

**2.5.9. Alkali resonance lines**

The wings of the resonance lines of the alkali metals sodium and potassium are known to deviate from the usual Voigt profile. Especially their far-wing line profiles exhibit strong non-Lorentzian behavior due to collisions with other molecules, such as in particular H$_2$ and He. Various descriptions of these line profiles have been provided in the past to characterize these non-Lorentzian line wings (see e.g. Tsuji et al. 1999; Burrows et al. 2000; Burrows & Volobuyev 2003; Allard et al. 2012). The most recent theoretical calculations of the resonance line wings broadened by collisions with H$_2$ have been published by Allard et al. (2019) for Na and by Allard et al. (2016) for K. These calculations are valid for perturber densities of up to $10^{21}$ cm$^{-3}$.

Figure 9 shows an example of the non-Lorentzian behavior of the sodium resonance line wings at a temperature of 800 K and an H$_2$ partial pressure of 10 bar. The cross sections are calculated from the Kurucz line list data, including van der Waals broadening, and shown for a temperature of 800 K and an H$_2$ partial pressure of 10 bar. We use the Kurucz line list to generate the cross sections, including the pressure-dependent van der Waals broadening by H$_2$. To convert the corresponding broadening parameters listed
in the Kurucz line list from H to H2, we use a scaling factor of 0.85. For the calculation involving the Allard et al. (2019) line wing description, we remove the two resonance lines from the line list, calculate the cross sections without them and then add the two broadened resonance lines that have been constructed using the data from Allard et al. (2019) to the results.

The results shown in Figure 10 clearly suggest that the resonance lines have a very strong non-Lorentzian behaviour and are distinctively asymmetric. They are super-Lorentzian close to the line center and become sub-Lorentzian in the far wings. Using a Voigt line profile for these two lines, thus, results in overestimating the line absorption in the far wings by orders of magnitudes, while strongly underestimating it near the line centers.

3. RESULTS

3.1. Comparison of atomic opacities

We compare the atomic opacities between the three databases NIST, Kurucz and VALD3. As mentioned in section 2.5.6, the calculations neglect pressure broadening. Therefore, the resulting opacities are suitable only for low-pressure environments with negligible collisional broadening. An example of four species (H, Li, S and Fe) is shown in Figure 11. Figures with more species are shown in appendix A and also in the HELIOS-K documentation. The 1H example of Figure 11 shows a good agreement between all three databases. The only difference is in the natural broadening coefficients from NIST. In the example of 3Li the agreement is not as good as for 1H and some of the smaller transition lines are different in all three databases. The more intensive transition lines agree better with the exception of the natural broadening coefficient of the largest line. The example of S shows that NIST has far fewer lines than Kurucz and VALD3, and between Kurucz and VALD3, there is a large difference in the natural broadening coefficients around $\nu = 84000 \text{ cm}^{-1}$. The overall agreement for Fe is better, but again NIST has far fewer lines, and Kurucz has more lines than VALD3. In general, NIST has significantly fewer lines than Kurucz and VALD3. Kurucz and VALD3 are similar for many species, but still they are not identical. For some species VALD3 has fewer lines than Kurucz. The greatest impact on the opacities comes from the natural broadening coefficients from large resonance lines.

It should be noted that use of a Voigt profile for atomic lines is not always suitable. Autoionization lines (Auger transitions) are a particular example that are not well-described by a Voigt profile. The wings of such a line are more appropriately described by a so-called Fano profile (Fano 1961) which decreases much faster than the Voigt profile. The Fano profile, however, depends on a parameter $g$ that cannot be easily calculated (see Merts & Magee 1972 for details). Therefore, using a Voigt profile for these lines will result in an overestimate of the absorption, creating an artificial continuum. This can be seen, for example, in Figure 11 for sulphur (atomic number 16). The opacities based on the Kurucz line list show an elevated continuum due to autoionization lines located in the FUV wavelength region. Since these lines are not included in the VALD3 and NIST databases, their opacities are not affected by the overestimated line wings.

Other problematic lines are resonance lines, where using a normal Voigt profile can also lead to over or underestimated absorption. In particular, the Lyman $\alpha$ wings of atomic hydrogen shown in the top left corner of Figure 11 are again overestimated by using a Voigt profile. Instead of following a Voigt profile, the actual line profile converges towards the Rayleigh scattering opacity far away from the line center (Dijkstra 2019).

3.2. Validation against ExoMol opacities

Figure 11. Opacity comparison of four atoms (H, Li, S and Fe) between the databases NIST, Kurucz and VALD3. The subplots in the left column show the entire wavenumber range, the subplots in the right column show a more narrow band. We use a temperature of 3000 K and no cutting length. The three databases contain a different number of transition lines, leading to differences in wavenumber coverage of the opacities. Figures of all other species may be found in the HELIOS-K documentation.

https://helios-k.readthedocs.io/en/latest/index.html
For many species, ExoMol provides pre-calculated binned Gaussian cross-sections, which we use to validate our calculations. Even if the validation is not done on a full Voigt profile, it confirms that our line list pre-processing and line intensity calculations are done correctly. In Figure 12, we show an example comparison of the H$_2$O POKAZATEL line list (Polyansky et al. 2018); and in Figure 13 CO (Li et al. 2015a). Individual points in wavenumber are not reproduced exactly, as shown in the bottom panel. One reason is that every point in wavenumber can consist of millions of individual transition lines, for which the intensity can vary by several orders of magnitudes. If the order of summation over all the transitions involved is changed, it can affect the differences.

In appendix B, we show a comparison for 45 molecules between binned Gaussian cross-sections from ExoMol and computed with HELIOS-K. The fractional difference ($\frac{\kappa_{\text{HELIOS-K}} - \kappa_{\text{ExoMol}}}{\kappa_{\text{HELIOS-K}}}$) reaches typically values between $10^{-7}$ and $10^{-1}$, which shows a good agreement between the two calculations. The difference can also reach higher values, which is most likely caused by division of very small numbers and limited numerical data precision.

Examples of molecular opacities (Full Voigt profile) for a temperature of 1500 K (500 K for SO$_3$), a pressure of 0.001 bar and a cutting length of 100 cm$^{-1}$ are shown in the Figures C8 to C11. The figures include TiO (McKemmish et al. 2019), VO (McKemmish et al. 2016), AIO (Patrascu et al. 2015), SiO (Barton et al. 2013), CN (Brooke et al. 2014b), CH (Masseron et al. 2014), CP (Ram et al. 2014), CS (Paulose et al. 2015), H$_2$O (Polyansky et al. 2018), CO$_2$ (Rothman et al. 2010), CO (Rothman et al. 2010; Li et al. 2015b), PN (Yorke et al. 2014), PO (Prajapat et al. 2017), PS (Prajapat et al. 2017), CH$_4$ (Yurchenko & Tennnyson 2014), NH$_3$ (Yurchenko et al. 2011), C$_2$H$_2$ (Gordon et al. 2017), HCN (Barber et al. 2013; Harris et al. 2006), SO$_2$ (Underwood et al. 2016a), SO$_3$ (Underwood et al. 2016b), SH (Yurchenko et al. 2018a), H$_2$S (Azzam et al. 2016), AlH (Yurchenko et al. 2018c), SiH$_4$ (Owens et al. 2017), SiH (Yurchenko et al. 2017b), SiS (Upadhyay et al. 2018), NS (Yurchenko et al. 2018b), NH (Brooke et al. 2014a), NO (Wong et al. 2017) and NO$_2$ (Rothman et al. 2010; Hargreaves et al. 2019).

During our analysis, we also spotted several issues in the ExoMol cross sections or data files, which were since corrected.

3.3. Comparison of ExoMol super-line opacities to full opacities

We compare the opacities computed with super lines to opacities from the full ExoMol line lists. Figure 14 shows an example of the POKAZATEL water line list. While individual transition lines are not represented in the super lines, the overall opacities agree well, with a fractional difference of less than 20%. By using super lines, the computational speed of the example in Figure 14 is increased by a factor of

![Figure 12](image-url)  
**Figure 12.** First panel: Comparison of our calculated cross-sections of the POKAZATEL line list with cross sections provided from the ExoMol database, showing a good agreement. Second panel: relative error between our work and ExoMol cross sections. Third panel: Zoom comparison between our work and cross sections from ExoMol. Individual points in wavenumber are not reproduced exactly due to different orders in summation processes. Fourth panel: Comparison between the binned Gaussian integrated cross section and the full Voigt profile. Calculations are done with a resolution of 0.01 cm$^{-1}$ and a cutting length 100 cm$^{-1}$.

3.4. The effects of line lists on atmospheric calculations

In the following section, we estimate the impact on atmospheric modeling when using different line lists. Herein, we compare two different line lists for water, the older BT2 water line list (Barber et al. 2006) versus the newer POKAZATEL water line list (Polyansky et al. 2018). Using those, we run models predicting atmospheric temperatures in radiative-convective equilibrium and corresponding planetary emission spectra. We choose two reference cases: (i) a typical hot Jupiter with a primordial H$_2$/He envelope with solar elemental abundances, and (ii) a super-Earth with a water-steam atmosphere.

The atmospheric radiative transfer is calculated with
the radiative-convective code HELIOS (Malik et al. 2017, 2019b). Details of the opacity and chemistry calculations are given in Malik et al. (2019a). The planetary parameters for HD 189733b and GJ 1132b are chosen as in Malik et al. (2017, 2019a). For simplicity, the host star is modeled as a blackbody in both cases.

The top row of Figure 15 shows the results for the hot Jupiter test case, in the form of vertical temperature-pressure (TP) profile, planetary emission spectrum, and relative difference in emitted flux. The latter is calculated as $|F_{\mathrm{POK}} - F_{\mathrm{BT2}}| / F_{\mathrm{POK}}$, with $F_{\mathrm{POK}}$ and $F_{\mathrm{BT2}}$ being the spectral fluxes from the POKAZATEL and BT2 models at a given wavelength. The spectra are calculated at the native resolution $R = 3000$ and also shown down-sampled to $R = 30$. Choosing different water line lists causes a negligible effect on the vertical TP profile, with the temperature difference $< 2$ K for any layer. However, due to the unequal presence and strength of spectral lines between the two line lists, the difference in the planetary emission spectrum is significant. The relative difference is higher than 100% (up to 10%) for $R = 3000$ (for $R = 30$) at certain wavelengths in the near-infrared. The large dependency of this result on the spectral resolution is not surprising considering that the impact of individual lines decreases with decreasing spectral resolution.

The water world case is shown in the bottom row of Figure 15. Here, the temperatures in the bottom atmosphere deviate between the two models by around 15 K. Consequently, and also because water is the only atmospheric absorber, the relative difference in the spectrum is larger than in the hot Jupiter case, namely over 30% for some wavelengths at $R = 30$. Since POKAZATEL includes more spectral lines than BT2, the emission of the POKAZATEL model is often smaller by the relative amount shown.

In terms of observational consequences, the absolute difference associated with the secondary eclipse depth is up to $\sim 40$ ppm for the hot Jupiter case and up to $\sim 2$ ppm for the water world case in the near-infrared range of wavelengths.

### 3.5. Performance

An important part of the opacity calculation is the reading of the line list files. The performance therefore depends highly on the memory access speed. To quantify this dependency, we first test how long it takes to read the pre-processed BT2 water line list files (Barber et al. 2006) (15 Gbytes, $\tilde{\nu}$ range 0–30000 cm$^{-1}$) from memory. We test three scenarios: reading the file over a network from a different server, reading from hard disk and reading from RAM. We also test three different systems:

- A desktop machine (3.6GHz) with a GeForce GTX980 GPU and 10 GByte of free RAM. This machine does not have enough RAM to store the entire BT2 line list.
- A GPU server (1.7 GHz) with four GeForce GTX1080 GPUs and 32 GByte of RAM.

![Figure 13. Same as Figure 12, but for CO.](image)

![Figure 14. Comparison of the ExoMol POKAZATEL water line list with ExoMol super lines, for $T = 1000$K, $p = 1$ bar, $\Delta\nu = 0.01$ cm$^{-1}$. The fractional difference between the opacity from all lines and the super lines is between 10 - 20% for most of the wavenumber range.](image)
**Figure 15.** TP profiles and emission spectra for a hot Jupiter with a solar abundance atmosphere (top) and a super-Earth with a water-steam atmosphere (bottom), calculated using the BT2 and the POKAZATEL line lists for the water opacities. The right panels show the relative difference in emitted flux between the BT2 and POKAZATEL models. We find relative differences of up to $10^{-40}\%$ ($R = 30$) and $> 100\%$ ($R = 3000$) for certain wavelengths in the NIR. These translate to an absolute difference $\sim 30$ ppm ($\sim 2$ ppm) in the secondary eclipse signal of HD 189733b (GJ 1132b) at certain wavelengths in the near-infrared.

- A cluster node (2.2GHz) on the Ubelix supercomputer with eight GeForce RTX2080ti GPUs and 256 GBytes of RAM.

For all tests, we use only a single GPU.

Shown in Table 6 is the time needed to read the entire BT2 water line list from different memory types. It is clear that only reading the data files can take a substantial amount of time, and it is important to carefully plan memory usage before doing large opacity calculations. Table 6 indicates that there can be very large differences associated with the memory speed, depending on the system configuration.

Shown in Table 7 is the time needed for the entire calculation to complete (including memory access) for different computer systems. Again the measured times show clearly that the performance depends highly on the memory speed. If the line list fits into the system RAM, then one can achieve a significantly higher performance than reading from a hard
disk or even over a network cable.

A way to further increase the performance is to combine opacity calculations from different pressure values into a single run. In this way, the amount of data reads can be reduced because the data remains on the GPU for the different pressure point calculations. This effect is shown in Table 8. By using ten pressure points in a single run, the overall performance of low resolution runs can be improved by about 60–70%. In high resolution runs, the effect is not as strong.

Shown in Figure 16 are timing measurements for different parts of HELIOS-K for the example of the BT2 water line list. One can see that for low resolution opacity calculations, the reading part of the line list dominates the entire calculation. The Voigt calculation depends on the resolution and cutting length, while the other parts depend only on the number of transition lines in the line list database. Also shown is the time needed to download the lines lists from the ExoMol website and to pre-process the line list into the HELIOS-K binary file format. These two timings depend on the used internet connection speed and the CPU clock rate of the used machine.

HELIOS-K can perform a typical opacity calculation with a resolution of $\Delta \tilde{\nu} = 0.01$ cm$^{-1}$ with $10^7$ lines in about 1 s.

For comparison, Yurchenko et al. (2018) lists the timing of the ExoCross code as 251 seconds, excluding the data reading, and by using approximations for the line wings. HELIOS-K performs the entire calculation in 12.5 seconds, including the data reading time. The traditional Humlicek approach takes 2775.6 seconds (excluding the data reading time).

Table 6. Time to read the entire BT2 water line list for different machines and memory types. The GPUs are not involved in this test, but are listed for a better overview of the system types. The measured times indicate that the memory type used can highly affect the performance.

| System | GPU | $\tilde{\nu}$ range in cm$^{-1}$ | data location | time |
|--------|-----|----------------------------------|---------------|------|
| Desktop | GTX 980 | 0 - 30000 | network | 252 s |
| Desktop | GTX 980 | 0 - 30000 | hard disk | 30.0 s |
| Desktop | GTX 980 | 0 - 30000 | partially RAM | 12.5 s |
| Server | GTX 1080 | 0 - 30000 | hard disk | 40.5 s |
| Server | GTX 1080 | 0 - 30000 | RAM | 1.9 s |
| Cluster | RTX 2080ti | 0 - 30000 | hard disk | 55.5 s |
| Cluster | RTX 2080ti | 0 - 30000 | RAM | 4.8 s |

Table 7. Performance of BT2 water line list for $\Delta \tilde{\nu} = 0.1$ cm$^{-1}$ and a cutting length of 25 cm$^{-1}$. An essential factor of the overall performance is the memory access of the line lists. For the best performance, the data must remain fully in RAM.

| GPU | $\tilde{\nu}$ range in cm$^{-1}$ | data location | time |
|-----|----------------------------------|---------------|------|
| GTX 980 | 0 - 5000 | over network | 172.6 s |
| GTX 1080 | 0 - 5000 | hard disk | 20.9 s |
| GTX 1080 | 0 - 5000 | hard disk | 26.3 s |
| GTX 980 | 0 - 5000 | RAM | 13.9 s |
| GTX 1080 | 0 - 5000 | RAM | 8.8 s |
| GTX 980 | 0 - 30000 | partially in RAM | 30.7 s |
| GTX 1080 | 0 - 30000 | RAM | 16.5 s |

Table 8. Performance of BT2 water line list from $\tilde{\nu} = 0$–5000 cm$^{-1}$, cutting length 25 cm$^{-1}$. nP indicates the number of pressure points in the calculation. At low resolution, using multiple pressure points can improve the performance.

We present an improved and more powerful version of the GPU opacity calculator HELIOS-K. For certain species, the code achieves a speedup of nearly two orders of magnitude compared to the original code as 251 seconds, excluding the data reading, and by using approximations for the line wings. HELIOS-K performs the entire calculation in 12.5 seconds, including the data reading time. The traditional Humlicek approach takes 2775.6 seconds (excluding the data reading time).

The code also supports more features and allows a simple use of different line list databases. During our work, we encountered several issues with multiple databases, which we reported to the maintainers. Our new code is fast enough to process the current largest molecular line lists from ExoMol.

While also the progress in the speed of modern GPUs helped to reduce the computing time, the largest increase in speed comes from the new parallelization algorithm, especially the usage of asynchronous and simultaneous data read and computation technique, which turn out to be very powerful. It is not only the GPU itself that makes the speedup, but the efficient interaction between CPU, GPU and memory as a full system.

But it is clear that when even bigger molecular line lists get published, new challenges could occur in both computing time and storage capacity. Atomic opacities are not as challenging in terms of computing power, but there we are faced with the incompleteness of theory. When atomic opac-

4. DISCUSSION

The code also supports more features and allows a simple use of different line list databases. During our work, we encountered several issues with multiple databases, which we reported to the maintainers. Our new code is fast enough to process the current largest molecular line lists from ExoMol.

While also the progress in the speed of modern GPUs helped to reduce the computing time, the largest increase in speed comes from the new parallelization algorithm, especially the usage of asynchronous and simultaneous data read and computation technique, which turn out to be very powerful. It is not only the GPU itself that makes the speedup, but the efficient interaction between CPU, GPU and memory as a full system.

But it is clear that when even bigger molecular line lists get published, new challenges could occur in both computing time and storage capacity. Atomic opacities are not as challenging in terms of computing power, but there we are faced with the incompleteness of theory. When atomic opac-


diagram.
Figure 16. Timing of different parts of HELIOS-K for the example of the BT2 water line list (0 - 30000 cm$^{-1}$): The Voigt function calculations are in dark blue, blue, and light blue for the cutting lengths of 500, 100 and 25 cm$^{-1}$. Reading the line list (from RAM) in orange and preparing the transition line properties in magenta. Also shown is the time needed to download and unpack the line list from the ExoMol servers and to pre-process the line list into the HELIOS-K binary format. For low resolution calculations, the reading part of the line list dominates the opacity calculation. Timings are measured with a Nvidia GTX 1080 GPU.

As line intensities are approximated by Voigt profiles with Lorentzian or sub-Lorentzian line wings, it can introduce significant errors in the opacity function. This is true especially for resonance and auto-ionization lines, where broadening mechanisms are not known well enough.

We computed opacity functions for several hundred species, which we store on our database and share with the community via the website www.opacity.world and our data server https://chaldene.unibe.ch or via https://dace.unige.ch/opacity and https://dace.unige.ch/opacityDatabase.

It is evident that opacity calculations are a complex, tedious undertaking. In order to ensure reproducibility, the computed opacities have been made freely available on the Swiss PlanetS platform DACE (https://dace.unige.ch). To aid the user in developing intuition, the opacities may be visualised via a graphical user interface (https://dace.unige.ch/opacity). Opacities may be downloaded at the full spectral resolution (0.01 cm$^{-1}$) in binary format (https://dace.unige.ch/opacityDatabase). Alternatively, the user may specify customised arrays of temperature and pressure, as well as the range of wavenumbers or wavelengths required, and interpolated opacities will be produced in the HDF5 format. In the future, we envision a “versioning” capability, where different implementations using the same generation of spectroscopic line list (e.g., by different researchers) or implementations using different generations of line lists, may be archived, so that differences in the subsequent models or simulations may be diagnosed.

It is our belief that these capabilities are critical for the exoplanet community as we move into the era of precision spectroscopy of exoplanetary atmospheres with the James Webb Space Telescope and the next generation of large ground-based telescopes.

Calculations were performed on UBELIX (http://www.id.unibe.ch/hpc), the HPC cluster at the University of Bern. This work has made use of the VALD database, operated at Uppsala University, the Institute of Astronomy RAS in Moscow and the University of Vienna. We acknowledge partial financial support from the Swiss National Science Foundation, the European Research Council (via a Consolidator Grant to KH; grant number 771620), the PlanetS National Center of Competence in Research (NCCR) and the Center for Space and Habitability (CSH) and the Swiss-based MERAC Foundation.
Yurchenko, S. N., Sinden, F., Lodi, L., et al. 2017b, Monthly Notices of the Royal Astronomical Society, 473, 5324
Yurchenko, S. N., & Tennyson, J. 2014, MNRAS, 440, 1649
Yurchenko, S. N., Williams, H., Leyland, P. C., Lodi, L., & Tennyson, J. 2018c, Monthly Notices of the Royal Astronomical Society, 479, 1401
Zaghloul, M. R., & Ali, A. N. 2011, arXiv e-prints, arXiv:1106.0151

APPENDIX
A. ATOMIC OPACITIES
Figure A1. Comparison of atomic opacities between the NIST, Kurucz and VALD3 databases. Elements 1 to 12. We use a temperature of 3000 K and no cutting length.
Figure A2. Comparison of atomic opacities between the NIST, Kurucz and VALD3 databases. Elements 13 to 24. We use a temperature of 3000 K and no cutting length.
Figure A3. Comparison of atomic opacities between the NIST, Kurucz and VALD3 databases. Elements 25 to 36. We use a temperature of 3000 K and no cutting length.
Figure A4. Comparison of atomic opacities between the NIST, Kurucz and VALD3 databases. Elements 37 to 48. We use a temperature of 3000 K and no cutting length.
B. MOLECULAR OPACITIES COMPARISON
Figure B5. Comparison between cross sections, calculated with HELIOS-K and from the ExoMol website. We use a wavenumber resolution of $\Delta \nu = 0.1 \text{ cm}^{-1}$ and $T = 1500 \text{ K}$ (1000 K for 12C2-1H4_.MaYTY).
Figure B6. Comparison between cross sections, calculated with HELIOS-K and from the ExoMol website. We use a wavenumber resolution of $\Delta \tilde{v} = 0.1 \text{ cm}^{-1}$ and $T = 1500 \text{ K}$ (500 K for 1H-14N-16O3_AIJS).
| Compound          | HELIOS-K | ExoMol | Fractional Difference |
|-------------------|----------|--------|----------------------|
| 28S-1H2__CATS     | HELIOS-K | ExoMol | HELIOS-K             |
| 28S-32S__UCTY     | HELIOS-K | ExoMol | HELIOS-K             |
| 31P-1H3__SAlTY    | HELIOS-K | ExoMol | HELIOS-K             |
| 31P-14N__YYLT     | HELIOS-K | ExoMol | HELIOS-K             |
| 31P-16O__POPS     | HELIOS-K | ExoMol | HELIOS-K             |
| 28Si-1H4__OY2T    | HELIOS-K | ExoMol | HELIOS-K             |
| 40Ca-16O__VBATHY  | HELIOS-K | ExoMol | HELIOS-K             |
| 32S-1H__GYT       | HELIOS-K | ExoMol | HELIOS-K             |
| 39K-35Cl__Barton  | HELIOS-K | ExoMol | HELIOS-K             |
| 45Sc-1H__LYT      | HELIOS-K | ExoMol | HELIOS-K             |
| 28Si-32S__UCTY    | HELIOS-K | ExoMol | HELIOS-K             |

Figure B7. Comparison between cross sections, calculated with HELIOS-K and from the ExoMol website. We use a wavenumber resolution of $\Delta \nu = 0.1 \text{ cm}^{-1}$ and $T = 1500 \text{ K}$ (1000 K for 32S-16O3__UYT2).
C. MOLECULAR OPACITIES OVERVIEW
Figure C8. Examples of molecular opacities: TiO (McKemmish et al. 2019), VO (McKemmish et al. 2016), AlO (Patrascu et al. 2015), SiO (Barton et al. 2013), CN (Brooke et al. 2014b), CH (Masseron et al. 2014), CP (Ram et al. 2014) and CS (Paulose et al. 2015). We use a temperature of 1500 K, a pressure of 0.001 bar and a cutting length of 100 cm$^{-1}$. 
Figure C9. Examples of molecular opacities: \( \text{H}_2\text{O} \) (Polyansky et al. 2018), \( \text{CO}_2 \) (Rothman et al. 2010), \( \text{CO} \) (Li et al. 2015a), PN (Yorke et al. 2014), PO (Prajapat et al. 2017), PS (Prajapat et al. 2017). We use a temperature of 1500 K, a pressure of 0.001 bar and a cutting length of 100 \( \text{cm}^{-1} \).
Figure C10. Examples of molecular opacities: CH$_4$ (Yurchenko & Tennyson 2014), NH$_3$ (Yurchenko et al. 2011), C$_2$H$_2$ (Chubb et al. 2020), HCN (Barber et al. 2013; Harris et al. 2006), SO$_2$ (Underwood et al. 2016a), SO$_3$ (Underwood et al. 2016b), SH (Yurchenko et al. 2018a), H$_2$S (Azzam et al. 2016). We use a temperature of 1500 K (500 K for SO$_3$), a pressure of 0.001 bar and a cutting length of 100 cm$^{-1}$. 
Fig. C11. Examples of molecular opacities: AlH (Yurchenko et al. 2018c), SiH₄ (Owens et al. 2017), SiH (Yurchenko et al. 2017b), SiS (Upadhyay et al. 2018), NS (Yurchenko et al. 2018b), NH (Brooke et al. 2014a), NO (Wong et al. 2017), NO₂ (Rothman et al. 2010; Hargreaves et al. 2019). We use a temperature of 1500 K, a pressure of 0.001 bar and a cutting length of 100 cm⁻¹.