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A Hydrodynamic Model of Alfvénic Wave Heating in a Coronal Loop and Its Chromospheric Footpoints

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Abstract

Alfvénic waves have been proposed as an important energy transport mechanism in coronal loops, capable of delivering energy to both the corona and chromosphere and giving rise to many observed features of flaring and quiescent regions. In previous work, we established that resistive dissipation of waves (ambipolar diffusion) can drive strong chromospheric heating and evaporation, capable of producing flaring signatures. However, that model was based on a simplified assumption that the waves propagate instantly to the chromosphere, an assumption that the current work removes. Via a ray-tracing method, we have implemented traveling waves in a field-aligned hydrodynamic simulation that dissipate locally as they propagate along the field line. We compare this method to and validate against the magnetohydrodynamics code Lare3D. We then examine the importance of travel times to the dynamics of the loop evolution, finding that (1) the ionization level of the plasma plays a critical role in determining the location and rate at which waves dissipate; (2) long duration waves effectively bore a hole into the chromosphere, allowing subsequent waves to penetrate deeper than previously expected, unlike an electron beam whose energy deposition rises in height as evaporation reduces the mean-free paths of the electrons; and (3) the dissipation of these waves drives a pressure front that propagates to deeper depths, unlike energy deposition by an electron beam.

Key words: magnetohydrodynamics – Sun: atmosphere – Sun: corona – Sun: flares – waves

1. Introduction

Alfvén waves, first predicted by Alfvén (1942) and verified by Lundquist (1949a, 1949b), occur throughout the solar atmosphere, from the chromosphere (De Pontieu et al. 2007) through the corona (Tomczyk et al. 2007; McIntosh et al. 2011), extending out into the heliosphere with the solar wind (Belcher & Davis 1971). The dissipation of waves has been shown to heat and accelerate particles in solar and stellar winds (Suzuki & Inutsuka 2005; van Ballegooijen & Asgari-Targhi 2016) as well as cosmic rays (Fermi 1949; Lazarian 2016). Their potential to heat the corona was quickly recognized (Alfvén 1947), and heating due to their dissipation is still considered a plausible solution to the coronal heating problem (Klimchuk 2006; van Ballegooijen et al. 2011; Reale 2014). Alfvén waves are generated in the convection zone, as well as in the chromosphere by shock collisions (Osterbrock 1961), where they propagate upward, possibly dissipating their energy in the chromosphere (Arber et al. 2016). A ponderomotive force due to the propagation of Alfvén waves acting on ions in the chromosphere has been suggested as an explanation for the first ionization potential effect (Laming 2015), as well as affecting the streaming of stellar winds (Belcher 1971), and the force due to downward waves can drive acoustic waves that can then cause sunquakes (Russell et al. 2016). The recent article by Russell (2017) summarizes some of the major developments in the 75-year history of Alfvén waves in solar physics.

In this work, we focus on Alfvén waves generated via reconnection events in the corona. Although it is expected that waves are generated during the reconnection events that drive flares and nanoflares (Parker 1991; Takeuchi & Shibata 2001; Shibata & Moriyasu 2003; Kigure et al. 2010; Jelínek et al. 2017), it is not currently known what fraction of the released energy they carry, with what frequencies they oscillate, or whether they cause heating that helps to power the emitted radiation. Simple arcade models suggest that the waves occur across a spectrum without an upper bound on frequency (Oliver et al. 1993; Tarr 2017). Fletcher & Hudson (2008) showed that Alfvén waves generated in the corona and propagating downward might carry a significant fraction of the released magnetic energy during a flare, later supported by simulations of three-dimensional reconnection (Birn et al. 2009), and they perhaps could cause acceleration of particles in the chromosphere. Haerendel (2012) examined the possibility that the auroral acceleration process may take place in solar flares, whereby a release of magnetic shear stress due to Alfvén waves is converted into kinetic energy of particles. A recent study with the New Solar Telescope found evidence for downward moving waves in a large solar flare, where their impact in the lower atmosphere possibly triggered a sunspot rotation (Liu et al. 2016).

The propagation of waves into the chromosphere has received significant attention in recent years. Haerendel (2009) showed that such waves carry sufficient energy to heat the chromosphere via turbulent phase mixing, and thus drive chromospheric evaporation. With a linearized magnetohydrodynamics (MHD) model, Russell & Stackhouse (2013) confirmed that Alfvén waves can deliver concentrated energy flux to the chromosphere. Russell & Fletcher (2013) developed a model of energy transmission as waves propagate beyond the transition region and found strong resistive damping in the chromosphere due to ion-neutral friction, with a strong
dependence on frequency. Ion-neutral friction is well established as one of the primary damping mechanisms of Alfvén waves (Piddington 1956), and in recent years the governing theory has received significant attention (Soler et al. 2013, 2015a; Leake et al. 2014; Khomenko 2017). The effectiveness of ion-neutral friction depends on the local field strength, and is most efficient when the Lorentz force is strong compared to other dynamic forces (Soler et al. 2015b).

In partially ionized plasma, the components of an electric current parallel or perpendicular to the magnetic field dissipate differently, due to collisions between the ionized plasma and neutral gas. This anisotropic dissipation has been extensively covered in a broad range of fields, such as ionospheric/thermospheric physics, astrophysics, as well as solar physics. While the governing physics remain the same, the breadth of application has led to different names and conceptualizations being used to describe the dissipation of perpendicular currents by plasma-neutral collisions: Cowling resistivity (Cowling 1956; Leake et al. 2005), Pedersen resistivity (Haerendel 2006; Goodman 2011), and ambipolar diffusion (Zweibel 1989; Martínez-Sykora et al. 2012, 2017) are all terms related to the dissipation of perpendicular currents by plasma-neutral collisions. The discussions by Zweibel et al. (2011) and Leake et al. (2014) explain the differences of origin and emphasis between the terms. The recent applications in solar physics include wave propagation and dissipation (de Pontieu 1999; De Pontieu et al. 2001; Leake et al. 2005; Brady & Arber 2016; Soler et al. 2016), mass and magnetic flux transport (Martínez-Sykora et al. 2012, 2017; Leake & Linton 2013), chromospheric heating (Goodman & Kazeminezhad 2010), and magnetic reconnection (Ni et al. 2015). Applied to our topic, the various terms are simply different names for the same anisotropic dissipation of electric currents. We will primarily use the terminology of ion-neutral friction or Cowling resistivity.

A few models have been developed to study the hydrodynamic evolution of solar plasma due to wave heating. Emslie & Sturrock (1982) developed a WKB formulation of resistive dissipation of Alfvénic waves launched in the corona in order to explain temperature minimum heating observed in large flares (Machado et al. 1978; Emslie & Machado 1979). Adopting this model with a correction for ambipolar diffusion, Reep & Russell (2016) showed that waves can also strongly heat the upper chromosphere and produce explosive evaporation. Kerr et al. (2016) confirmed these results, and further showed that chromospheric lines such as Mg II could distinguish between Alfvénic wave heating and electron beam heating in flares, offering an important observational test. In these studies, however, it was assumed that the waves propagate instantly from their injection location to the depths where they dump. The assumption is justified for heating by a beam of electrons, where electrons with energy $>10$ keV travel at speeds $\geq 0.2c \approx 60$ Mm s$^{-1}$. Waves, however, travel at the local Alfvén speed, which is typically $<10$ Mm s$^{-1}$ in the corona (although it may approach $0.1c$ in the corona of some active regions Fletcher & Hudson 2008; Russell & Fletcher 2013), and significantly slower elsewhere. The damping of waves affects the atmosphere on a similar time-scale as the wave travel times, so travel times should not be neglected.

The model in Reep & Russell (2016) required high frequencies ($\geq 1$ Hz) for effective dissipation of wave energy, and so the frequency spectrum of waves in the corona is important to study. DeForest (2004) directly observed frequencies as high as 0.1 Hz with TRACE in the 1600 Å passband. Eclipse observations have also shown the presence of intensity oscillations of the order of 1–10 Hz (Pasachoff et al. 2002; Rudawy et al. 2004), although it is possible that those fluctuations are not due to Alfvén waves (Rudawy et al. 2010). Microwave and radio bursts during solar flares have revealed oscillations of the order of 10–100 Hz (Kiplinger et al. 1983; Kaufmann et al. 1984). As these frequencies are at the limit of current instrumental cadence, it is unclear whether higher cadence instruments would reveal the presence of higher frequencies, and if so, their importance to the physical processes occurring in the solar atmosphere. The next generation of solar instrumentation, in particular, the suite of instruments to come online in 2019–2020 at the Daniel K. Inouye Solar Telescope (Elmore et al. 2014), will have the cadence and sensitivity to probe the super-hertz frequency range of MHD waves in the low corona, offering a first look at whether such waves are present in the corona, if they are generated by reconnection, and what their energetic importance may be.

In this work, we directly examine the propagation of Alfvén waves by implementing a ray-tracing code that follows the waves as they travel and damp along a coronal loop in a one-dimensional hydrodynamics simulation. We describe the implementation and necessary physics in Section 2. We compare this implementation to the MHD code Lare3D in Section 3. We then directly compare the results of heating due to propagating waves with the previous work that assumed instantaneous travel times, as well as with an electron beam, in Section 4.

2. Relevant Physics and Implementation

We have implemented traveling Alfvén waves in the HYDrodynamics and RADIation code (HYDRAD, Bradshaw & Mason 2003), which solves the equations of conservation of mass, momentum, and energy for a two-fluid plasma confined to an isolated magnetic flux tube (equations and details are listed in Bradshaw & Cargill 2013). The code includes adaptive mesh refinement of arbitrary order, important for resolving rapid jumps in the temperature or density profiles, which can lead to improper estimates of coronal properties if not properly treated (Bradshaw & Cargill 2013). Radiative losses are treated with a full calculation of emissivities from abundant ions using CHIANTI version 8 (Dere et al. 1997; Del Zanna et al. 2015), with the ability to simultaneously solve for non-equilibrium ionization populations of any desired element (Bradshaw & Raymond 2013). The code is quick, robust, and computationally inexpensive.

Alfvén waves have long been considered important to the dynamics of energy transport in the solar atmosphere. They cannot be treated in pure hydrodynamic (HD) codes, which are designed to study energy transport but do not solve the full set of MHD equations. On the other hand, MHD codes generally do not have a full treatment of thermodynamics, or the proper resolution to resolve the transition region, which often means it is difficult to properly understand the evolution of a loop’s hydrodynamics (Bradshaw & Cargill 2013). We wish to bridge the gap between these two approaches, and implement a method that allows us to study Alfvén wave dissipation in an HD code.
We therefore introduce a new method to describe waves as pulses of period-averaged Poynting flux propagating along a loop, which we trace with individual rays at various points along the pulse. Figure 1 illustrates this. We inject a pulse with a certain initial Poynting flux, comprised of a number of rays \( N \) that approximate the spatial extent of a pulse. In Figure 1, this initial pulse is represented by the solid red curve, which has been divided into seven rays, each with their own position and Poynting flux (red plus signs, offset by an arbitrary factor). In order to trace the motion of the whole pulse, each ray is individually advected along the loop in the direction of propagation at the local Alfvén speed (which may differ for each ray), and the Poynting flux is decreased according to the local damping length (see Section 2.1). The blue curve in Figure 1 represents the pulse at some time later, where the Poynting flux has decreased, and the spatial extent has narrowed due to a decreasing Alfvén speed. As a result, the rays are closer together, especially near the leading edge of the pulse.

In order to inject a pulse onto the loop, we initialize the rays one at a time. If the pulse is injected at time \( t_0 \), the first ray is created at time \( t_0 \), with a position \( z_0 \) corresponding to the location of injection and Poynting flux equal to the pulse’s initial flux \( S_0 \). Supposing that there are \( N > 1 \) total rays comprising the pulse, and the duration of the pulse is \( \tau_{\text{dur}} \), the next ray is created at time \( t_1 = t_0 + \tau_{\text{dur}}/(N - 1) \), the next at \( t_2 = t_1 + \tau_{\text{dur}}/(N - 1) \), and so on until the last ray is created at time \( t_{N-1} = t_0 + \tau_{\text{dur}} \). Between times \( t_0 \) and \( t_1 \), the first ray is appropriately advected, between \( t_1 \) and \( t_2 \), the first two rays are advected, etc.

The number of rays \( N \) comprising a pulse must be high enough to both capture the spatial extent of that pulse and to capture small-scale changes in the Alfvén speed and damping lengths, particularly in regions of large gradients. If the duration is longer, the spatial extent is greater, and so requires more rays to capture the full extent and sharp gradients. Furthermore, with the WKB method, we need not resolve the wavelength, but we do need to resolve changes in the damping length. We therefore choose the number of rays \( N \) to be 50 rays per second of injection duration, or \( N = 50 \tau_{\text{dur}} \), which we have found through simple tests to accurately capture the heating while not being too demanding computationally.

The input for the waves is general: the user specifies the number of pulses, and, for each one, the start time, duration, injection location along the loop, direction of propagation, initial Poynting flux, frequency, and perpendicular wave number (at the photosphere). The code then divides each pulse up into a number of rays that are traced in time and position.

### 2.1. Ray Tracing

At each numerical time step, the code first checks to see if any new rays need to be initialized, i.e., if the current time is between the starting injection time \( t_0 \) and the final injection time \( t_0 + \tau_{\text{dur}} \) for any pulse. If so, new rays are created at the specified location with the specified properties of the pulse to which they belong (as described in the previous section).

All currently existing rays are then advected at each time step. The rays travel at the local Alfvén speed \( v_A(z) \), from an initial position \( z_0 \) to a final position \( z_f \). In general, the pulse will have arbitrary spatial extent and span numerous grid cells, each with a distinct Alfvén speed, so that the rays propagate at different speeds. The ray positions are continuous, while the physical variables (and the local Alfvén speed defined by them) are defined on the discrete grid. We must therefore carefully treat each ray’s propagation and damping within the discrete system.

Suppose the bounds of a grid cell are given by \( z_1 \) and \( z_2 \), with Alfvén speed \( v_A \) between those bounds. We assume that the Alfvén speed is constant within a grid cell for simplicity, though a more general treatment could interpolate the value for any point within the cell. For a ray position \( z_i < z_{\text{ray}} < z_2 \), the time to traverse the grid cell is given by

\[
\tau = \frac{z_2 - z_{\text{ray}}}{v_A}.
\]

We have two cases then. In the first, the numerical time step \( \Delta t < \tau \), i.e., that the current time step is smaller than the time for the ray to move to the next grid cell, in which case the ray is advected by

\[
\Delta z = v_A \Delta t.
\]

In the second case, \( \Delta t > \tau \), i.e., that the ray moves into a new grid cell in the current time step, we use a recursive function to then find the new ray position. Numbering each successive grid cell from the starting position of the ray, the new ray position is then:

\[
\Delta z = v_{A, 0} \tau_0 + v_{A, 1} \tau_1 + ... + v_{A, N} \tau_N
\]

\[
= \sum_{i=0}^{N} v_{A, i} \tau_i
\]

which is continued recursively, while \( \Delta t > \sum_{i=0}^{N} \tau_i \). When that condition fails, we advance the ray in the final grid cell by the distance \( v_A \times (\Delta t - \sum_{i=0}^{N} \tau_i) \). All of the rays are advected appropriately at each time step with this method. Simultaneously, we calculate the reduction in their Poynting fluxes due to resistive dissipation according to the local damping lengths (given in Section 2.2). Specifically, the decrease of Poynting flux from \( z_0 \) to \( z_f \) follows the general...
equation
\[ S(z_f) = S(z_0) \exp \left( -\int_{z_0}^{z_f} \frac{dz'}{L_D(z')} \right), \]
which is applied to each ray as their locations are updated.

2.2. Heating

The heating terms follow the same basic equations as those in Reep & Russell (2016), except that the heated locations are now limited to those grid cells that contain a pulse at a given time. The code loops across the numerical grid, locating the start and end positions of each pulse (i.e., the positions of the first and last ray in a pulse). If a grid cell is located between the leading and trailing edge of a pulse, then the grid cell is assumed to be heated by the resistive dissipation of that pulse as it passes through. After the rays have been advected forward and their Poynting fluxes appropriately decreased, that lost energy is taken to heat the local plasma. The heating function is given in general by
\[ Q(z) = \frac{dS(z)}{dz} = \frac{S(z)}{L_D(z)} \]
where \[ S(z) \left( \frac{1}{L_e} + \frac{1}{L_H} \right) = Q_e + Q_H, \]
which defines the split between the electron heating term and the hydrogen heating term in terms of the damping lengths:
\[ Q_e = \frac{S(z)}{L_e}, \]
\[ Q_H = \frac{S(z)}{L_H}. \]
The damping lengths are given by
\[ \frac{1}{L_D} = \frac{1}{L_e} + \frac{1}{L_H} = \left[ \eta_e \frac{k_e^2 c^2}{4 \pi \nu_A} + \frac{\omega^2 c^2}{4 \pi \nu_A} \right] + \left[ \eta_c \frac{\omega^2 c^2}{4 \pi \nu_A} \right], \]
where \( k_e(z) \) is the local perpendicular wave number (which we assume varies as \( k_e(z) = k_{e,0} B_{e,0} \) for \( k_{e,0} \) and \( B_{e,0} \) the values at the apex of the loop), \( \omega \) the angular frequency, \( c \) the speed of light in vacuum, \( \nu_A(z) \) the local Alfvén speed, and \( \eta_e(z) \) and \( \eta_c(z) \) the parallel and Cowling resistivities (due to ion-neutral friction, Cowling 1956), given by
\[ \eta_{\parallel} = \eta_{\parallel e} + \eta_{\parallel c} = \frac{m_e (\nu_{ei} + \nu_{en})}{n_e e^2} + \frac{B^2 \rho_n}{c^2 \nu_{ei} \rho_t (1 + \xi^2 \theta^2)}, \]
where \( m_e \) is the electron mass, \( B(z) \) is the magnetic field strength, \( \rho(z) \) is the mass density, and \( \nu(z) \) is the collision frequencies. The subscripts \( i, e, n, \) and \( t \) refer to electron, ion, neutral, and total, respectively. As in Reep & Russell (2016), we define \( \xi = \frac{\rho_i}{\rho_t} \) and \( \theta = \frac{\nu_i}{\nu_t} \), and modify the Alfvén speed due to the presence of neutrals:
\[ \nu_A(z) = \frac{B}{\sqrt{4 \pi \rho_t}} \left( \frac{1 + \xi^2 \theta^2}{1 + \xi^2 \theta^2} \right)^{1/2}. \]
This expression reduces to the standard form of the Alfvén speed in the limit of a fully ionized plasma.

The damping lengths are calculated locally at each time step, and each ray has its Poynting flux decreased using the local length in each cell it traverses. The heating rate in a grid cell is then found by interpolating the Poynting flux at the center of the grid cell using the two nearest rays. Consequently, there is a small error associated with the finite width of the grid cells: the heating rate is more accurate for small grid cells.

In order to partition the energy between the two species, we can also rewrite the equations more directly in terms of the total heating rate:
\[ Q_e(z) = \frac{Q(z) L_D(z)}{L_e(z)}, \]
\[ Q_H(z) = \frac{Q(z) L_D(z)}{L_H(z)}. \]
Then, each term is added to its respective energy equation.

2.3. Time Step

As mentioned before, if the current time is between the initial injection time \( t_0 \) and final injection time \( t_0 + \tau_{\text{dur}} \) for any pulse, new rays must be created at an appropriate time step \( \tau_{\text{dur}} / (N - 1) \) for the number of rays \( N \) in the pulse. If \( N \) is large, then this time step becomes tiny and may become smaller than the numerical time step \( \Delta t \) of the code. When this happens, some rays might fail to be properly injected. To avoid this problem, we introduce a new timescale to HYDRAD to ensure that all rays are properly created.

At a given time, if there is at least one pulse being initialized, we set a new timescale to one-tenth of the time between each successive ray (which is rather cautious, for computational speed this could be increased up to one-half of the time between rays). That is, for a duration of injection \( \tau_{\text{dur}} \) and number of rays \( N \), the new timescale is \( \tau_{\text{rays}} = \frac{\tau_{\text{dur}}}{10N} \). Then, this timescale is guaranteed to be smaller than the time between the creation of each ray, and each one will be properly initialized.

This timescale is then calculated for each pulse currently being injected, compared to the other relevant timescales, and then the time step in the simulation is set to the minimum timescale, as is generally done with HYDRAD (Bradshaw & Cargill 2013). In this way, we insure that all rays are properly injected, and that the CFL condition is met.

3. Comparison to Lare

To validate that the ray-tracing method reproduces the correct propagation and dissipation, we first compare our implementation of the WKB wave-packets with the magneto-hydrodynamics Lagrangian–Eulerian Remap 3D code (Lare3D, Arber et al. 2001). We present brief results necessary for validation here, but postpone a detailed comparison of the HD and MHD codes for a later work. In the next section, we present the detailed results of propagating wave simulations with HYDRAD.

Lare3D is a parallel code that solves the full set of nonlinear MHD equations by first taking a Lagrangian step with each
computational cell and, second, conservatively remapping the result back to the original Eulerian grid. This method allows for shocks to be properly resolved and can be readily adapted to include most physical processes. We use the modified MHD equations, which include the effects of partial ionization and Cowling resistivity (Leake et al. 2005). We use this modified Lare code in 1.5D (only one dependent variable, but all components of the 3D vector are evolved) to model a one-dimensional coronal loop (similarly done by other authors, e.g., Johnston et al. 2017a, 2017b), and introduce an Alfvénic wave pulse in order to compare to our new method in HYDRAD. We use a grid spacing of 460 m in Lare, and do not include the effects of radiative losses or thermal conduction. The ray-tracing method presented here includes only Alfvénic perturbations. Therefore, in the Lare3D simulation that we use to verify and validate the method, we remove the nonlinear coupling of Alfvénic perturbations into longitudinal perturbations. Future studies will examine the fully nonlinear MHD system, along with mode-coupling and multi-dimensions, as well as the thermodynamics appropriate for the coupled chromosphere-corona.

We have tuned the initial conditions of the Lare simulation to match the HYDRAD ones as closely as possible. We show the initial conditions of both codes in Figure 2, from left to right and top to bottom: the temperature, number densities, and ionization fraction of hydrogen as a function of position. On the bottom, for the assumed magnetic field strength profile, are the Alfvén and sound speed, resistivities, and damping lengths of the waves (assuming frequency $f = 10$ Hz, perpendicular wave number $k_z = 0$, and a constant $B = 107.9$ G).

Figure 2. Comparison of the initial values of the simulations for one-half of a loop of length $2L = 60$ Mm. In all of the plots, HYDRAD values are shown in black, Lare in red. On top, from left to right, are the temperature, number densities, and ionization fraction of hydrogen as a function of position. On the bottom, for the assumed magnetic field strength profile, are the Alfvén and sound speed, resistivities, and damping lengths of the waves (assuming frequency $f = 10$ Hz, perpendicular wave number $k_z = 0$, and a constant $B = 107.9$ G).

![Figure 2](image)

Figure 3. Energy transmission coefficient at the transition region as a function of frequency in the Lare simulations for the chosen Alfvén speed profile. High-frequency waves pass mostly unimpeded through the TR, while low-frequency waves reflect a large portion of their Poynting flux. The results are comparable with Russell & Fletcher (2013).

Figure 3. Energy transmission coefficient at the transition region as a function of frequency in the Lare simulations for the chosen Alfvén speed profile. High-frequency waves pass mostly unimpeded through the TR, while low-frequency waves reflect a large portion of their Poynting flux. The results are comparable with Russell & Fletcher (2013).

![Figure 3](image)

Although HYDRAD and Lare are given essentially the same initial conditions, they are fundamentally different codes that solve different equations; therefore, the resulting dynamics will not always agree. One major point of comparison is that we have not implemented reflection into HYDRAD, which arises in Lare because it solves the full MHD equations. At the region of largest Alfvén speed gradient, around the transition region (Figure 2), waves with low frequency will reflect back into the corona. Figure 3 shows the energy transmission coefficient, measured as a ratio of the transmitted Poynting flux to the original Poynting flux, as a function of wave-packet frequency. While reflection at frequencies at or below 1 Hz dominate, it is reasonable to assume that for frequencies $\geq 2$ Hz, the use of the current HYDRAD implementation without reflections is reasonable, though the inclusion of reflection would be an important generalization of the current method.
We have therefore run a simulation with both HYDRAD and Lare, using a frequency $f = 10$ Hz and perpendicular wave number $k_z = 0$. To partially account for the absence of reflection, we use a Poynting flux of $10^8$ erg s$^{-1}$ cm$^{-2}$ in Lare, which we reduce by the reflection coefficient ($\approx 10\%$) in HYDRAD. In order to facilitate comparison, we do not allow the plasma to be heated, which would cause discrepancies in the temperature and ionization fraction between Lare and HYDRAD. In other words, the initial profile is maintained throughout the simulation to ease the comparison.

Figure 4 shows a few snapshots of this comparison simulation near the top of the chromosphere, shortly after the waves impinge upon it. The plot on the left shows the period-averaged Poynting flux as a function of position, at times 3 s (blue) and 4 s (red) into the simulation. HYDRAD is shown with points (each ray), and Lare as a solid line. The plot on the right shows the derived heating rate as a function of position. The slight discrepancies at the trailing edge of the pulse are due to reflection, where some of the Poynting flux has begun to propagate in the opposite direction in the Lare simulation.

While there are differences between the two simulations, primarily due to reflection, which has not been included in HYDRAD, we find that the ray-tracing method reproduces the approximate behavior of the MHD code. There are differences that can be addressed in future iterations to further improve the comparison: the aforementioned reflection, the ponderomotive force, a method to treat mode conversion, etc. Although these limitations exist, this method faithfully captures the propagation and resistive dissipation of waves through a field-aligned flux tube. In the Appendix, in order to demonstrate the current limitations of this method, we show a comparison of a low-frequency case, where the effect of reflection is more pronounced.

### 4. Results

The comparison to Lare validated our implementation of a WKB approximation for wave propagation and damping in a hydrodynamic code. We now perform a detailed examination for different wave-packet parameters, with and without accounting for travel times. We have run simulations with HYDRAD to examine the hydrodynamics. We use a loop length $2L = 60$ Mm, assumed semi-circular and oriented vertically relative to the solar surface. We use the magnetic field strength profile $B(z)$ as specified in Russell & Fletcher (2013) in all the simulations here: $B(z) = B(0) \left( \frac{P(z)}{P(0)} \right)^{0.139}$ for the initial pressure profile calculated from the hydrostatic equations, and a constant field strength of $\approx 100$ G in the corona. We assume a photospheric value of 1000 G, as in Reep & Russell (2016). This is an empirical fit intended to allow the field to vary smoothly across the chromosphere, though as Russell & Fletcher (2013) note, it is difficult to observationally measure the field strength across the chromosphere so that this is an assumed profile.

We first examine the damping of Poynting flux at various frequencies. Figure 5 shows the Poynting flux in three HYDRAD
simulations for one-second pulses with $S_0 = 10^9 \text{erg s}^{-1} \text{cm}^{-2}$, $f = [1, 3, 10] \text{Hz}$, $k_x = 0$, propagating to the left. The colors ranging from violet to red show different time steps, at a 2 s cadence, for a total of 100 s after the initial injection of the pulse. Each plus sign denotes the location and flux of a single ray, while the lines connecting them denote the interpolated Poynting flux at a given position. The pulses take just over 3 s to reach the transition region, where they begin to damp. As each pulse propagates down the loop, the Poynting flux decreases appropriately, with high-frequency pulses dissipating more rapidly, though even the 10 Hz case takes well over 10 s to fall to 10% of its initial Poynting flux. By way of comparison, we over-plot the damping locations determined by the method used in Reep & Russell (2016), shown as black dashed lines. In all cases, the instant-travel method damps slightly higher in the atmosphere than the ray-tracing method, and the travel time is too significant to be ignored. The difference in damping locations is exacerbated at high frequencies.

We now compare the new method to Reep & Russell (2016) in order to better understand the effect of propagation delays. Figure 6 shows the hydrodynamic variables in a simulation with initial Poynting flux $S(z_0) = 10^{10} \text{erg s}^{-1} \text{cm}^{-2}$, perpendicular wave number at the apex $k_{x,a} = 10^{-5} \text{cm}^{-1}$, and $f = 10 \text{Hz}$, which can be compared to the top row of Figure 1 in Reep & Russell (2016). The colors show snapshots of the simulation every second, ranging from violet to red. The $x$-axis is shown on a logarithmic scale to emphasize the chromosphere of the loop, which has a total length $2L = 60 \text{Mm}$, so that the apex is at a position 30 Mm. The first plot shows the total (electron + hydrogen) heating rate (erg s$^{-1}$ cm$^{-2}$) versus curvilinear position.
It takes roughly 3 s for the wave to reach the chromosphere (with the assumed field strength), so that there is only weak coronal heating before that time. After it impinges on the chromosphere, the pulse begins to dissipate, and the pulse slows as the Alfvén speed decreases due to an increasing density. It takes roughly 7 s to travel 1 Mm below the transition region, while the heating slowly grows with depth. Since the perpendicular wave number is small, the Cowling resistivity initially dominates the energy dissipation terms, so that the hydrogen temperature rises more than the electron temperature, though both only increase modestly. As the pulse propagates, the plasma is ionized, reducing the Cowling resistivity, so that at later times damping due to electron collisions becomes the dominant term. Figure 7 shows the resistivities and damping lengths in the simulation, demonstrating the reduction in Cowling resistivity. This indicates the importance of the ionization level in determining the depth and rate at which waves dissipate. The parallel resistivity also decreases slightly as the electron density increases due to the change in ionization. As with the simulation in Reep & Russell (2016), the bulk flow velocity evaporates gently. As the chromosphere heats, the ionization fraction rises, locally raising the electron density, while the weak evaporation carries a small number of both electrons and ions into the corona.

For this same simulation, we show the partition of the heat as a function of position and time for the simulation in Figure 6. The left plot shows the fraction of dissipated energy that heats hydrogen, the right plot shows the fraction that heats electrons. The background heating is assumed to go entirely into electrons. In this case, since the perpendicular wave number is small, a significant fraction of the energy goes into heating the hydrogen directly, reaching above 80% at the leading edge of the pulse, and remaining close to 50% trailing that.

In Figure 9, we show a simulation with a higher perpendicular wave number \( k_x = 4 \times 10^{-4} \text{ cm}^{-1} \) (but otherwise equal parameters), which increases the relative importance of parallel dissipation. Due to the rise in parallel dissipation (both ion–electron and electron–neutral collisions), the heating is stronger in both the corona and upper chromosphere, and peaks more sharply. As a result, the electron temperature rises all across the corona, spreading outward from the apex of the loop at \( z = 30 \text{ Mm} \) and into the chromosphere. The hydrogen temperature rises sharply here as well, driving strong, explosive flows, and is comparable to the same case in Reep & Russell (2016; middle row of Figure 1 in that paper). Since the energy is deposited higher in the chromosphere, the waves do not travel as deep before dissipating, so the rise in ionization does not occur as deep as in the previous case.

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**Figure 7.** Resistivities (left) and damping lengths (right) at the same times in the same simulation as Figure 6. As the pulse propagates downward ionizing the plasma, the Cowling resistivity is reduced, thus causing damping due to electron collisions to become dominant at later times at the same depths. As before, the colors show different times, at a 1 s cadence, from violet to red.

**Figure 8.** Partition of the heat as a function of position and time for the simulation in Figure 6. The left plot shows the fraction of dissipated energy that heats hydrogen, the right plot shows the fraction that heats electrons. The background heating is assumed to go entirely into electrons.
As mentioned before and shown in Figure 7, as a pulse propagates into the chromosphere and dissipates its energy, the ionization fraction and electron density grow, thus decreasing both the Cowling and parallel resistivities. The trailing end of a long duration pulse, or any subsequent pulses, therefore propagate to greater depths into the chromosphere than the leading edge (or pulse) because of that reduction in the resistivity. Effectively, earlier pulses bore a hole into the chromosphere that allows later ones to penetrate deeper and deeper, even at frequencies expected to strongly dissipate high in the initial chromosphere. Heating due to the dissipation, therefore, effectively propagates downward.

This behavior is opposite to that of an electron beam. As a beam deposits its energy in the chromosphere, driving evaporation into the corona, the mean-free path of any later electrons is significantly reduced due to the rise in coronal density. The result is that the location of energy deposition for long duration beams rises into the corona and becomes more and more localized near the injection site, as shown, for example, in Figure 3 of Reep et al. (2015).

There is therefore an important distinction in the heating profiles between the cases of Alfvén waves and electrons beams. Electrons propagating down a loop are stopped as they collide with ambient plasma, in particular, as the density rises in the chromosphere, thus shortening their mean-free paths. Before there is significant evaporation into the corona, electrons deposit their energy at an approximately constant depth, as their mean-free paths do not change (assuming there are no significant time-varying changes in the beam, such as an increase in the low energy cut-off). This is in stark contrast to the case of Alfvén waves, which propagate significantly more slowly than beams, and deposit energy across more of the chromosphere.

Figure 9. Similar to Figure 6, with \( k_{\text{rad}} = 4 \times 10^{-4} \ \text{cm}^{-1} \) and \( f = 10 \ \text{Hz} \) (compare to the middle row of Figure 1 in Reep & Russell 2016).
Figure 10. Comparison of the development of the pressure profiles of loops heated by an Alfvén wave (top) and electron beam (bottom). In the case of the wave, because it propagates increasingly deep, the spike in pressure propagates downward. In the case of the beam, the electrons reach essentially the same depth continually, so that the pressure spike does not move. The colors on this plot refer to the local velocity, colored blue where the plasma is up-flowing, red where it is down-flowing, and white where it is stationary. Down-flowing velocities are defined as positive in this and the next figure. The white dotted line shows the initial profile.

We can directly compare the behavior for beam and wave heating. In Figure 10, we show the electron pressure profiles as a function of position at three select times for an Alfvén wave simulation with \( f = 10 \, \text{Hz, } k_x = 4 \times 10^{-4} \, \text{cm}^{-1} \), and \( S(z_0) = 10^{10} \, \text{erg} \, \text{s}^{-1} \, \text{cm}^{-2} \) (top row) and an electron beam with low energy cut-off \( E_c = 10 \, \text{keV} \), spectral index \( \delta = 5 \), and energy flux \( E_b = 10^{10} \, \text{erg} \, \text{s}^{-1} \, \text{cm}^{-2} \) (bottom row), using the heating form derived by Emslie (1978). We have colored the locations by the local flows, blue where the plasma is up-flowing, red where it is down-flowing, and white where it is stationary. The white dots indicate the initial profile. The spike in the chromospheric pressure propagates downward in the case of the waves, whereas in the case of the beam it does not reach progressively greater depths. In both cases, a pressure front begins to rise into the corona and evaporate plasma, although it travels at a higher speed for the beam.

In Figure 11, we show the electron density profiles in a similar manner to the previous figure. As noted before, as Alfvén waves propagate to deeper depths, they cause a downward propagating ionization spike, thus raising the local electron density while simultaneously driving up-flows into the corona. In the case of a beam, however, the plasma at the depth of energy deposition becomes ionized, but there is no propagation of that ionization spike.

These differences in the ionization, flows, and pressures suggest stark differences in chromospheric line profiles. Using the instantaneous method, Kerr et al. (2016) found that the Mg II k line behaved differently in an electron beam simulation from an Alfvén wave simulation. Specifically, the central reversal of the line gradually disappears, and the line develops a strong red–blue asymmetry in the wave simulation, while the line remains roughly symmetric with a strong central reversal during the electron beam simulation. On the other hand, the Ca II 8542 Å line was similar in both simulations, with a small difference in the Doppler shifts. The inclusion of travel time effects would likely exacerbate the differences found by those authors, which warrants a deeper investigation. Other authors have similarly reported discrepancies in the modeling of the Mg II lines when modeled with an electron beam (e.g., Rubio da Costa et al. 2016), which warrants further examination. Because these properties all vary with chromospheric depth and with time, in principle, they could be probed by high cadence spectrometry \( (\lesssim 1 \, \text{s}) \) to track the propagation of these waves and to measure the hydrodynamic profiles in time. We plan to address these possibilities in future work.

5. Conclusions

In this work, we have updated the model from Reep & Russell (2016) to include travel time effects due to the propagation of Alfvénic waves. The method is sufficiently general to be implemented in any hydrodynamics model that does not solve the MHD wave equations. To test that this method is valid, we have compared it against the MHD code Lare3D, adapted for a field-aligned flux tube, and found generally good agreement.

We have drawn a number of important conclusions from this work:

(1) Alfvénic waves can heat all levels of the chromosphere, and the damping of these waves depends strongly on not only the wave parameters, but also the ionization level. Ion-neutral friction is extremely efficient at damping waves, so a high proportion of neutral atoms greatly
increases the damping of Alfvén waves, while a low proportion means that waves travel mostly unimpeded.

(2) Because waves ionize the plasma as they propagate through the chromosphere, early waves effectively bore a hole in the chromosphere through which later waves can more readily propagate without significant dissipation. Successive waves penetrate to ever greater depths, and high-frequency waves penetrate significantly deeper than expected from our previous paper. This contrasts directly with the well-known result for electron beams: as evaporation brings material into the corona, later electrons have a shortened mean-free path. The height at which beams deposit their energy therefore tends to travel upwards along the loop for long duration heating (see, e.g., Reep et al. 2015).

(3) The propagation of these waves causes pressure and ionization fronts to form in the chromosphere that propagate with time. This is in contrast to an electron beam, where the depth of the maximum pressure remains localized. To observe this difference would likely require high cadence spectroscopy, so both observational and modeling studies of chromospheric lines should be undertaken. A study including a full wave spectrum (e.g., Tarr 2017) might also provide insight about the heating processes occurring in the chromosphere.

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### Appendix

In order to demonstrate the current limitation of this method while neglecting reflection, we briefly examine one more comparison between HYDRAD and Lare. We examine a simulation with one pulse with frequency $f = 1\text{ Hz}$, where the reflection coefficient at the transition region is expected to be
around 45%–50% (Figure 3). We set the initial Poynting flux to \( S_0 = 10^9 \text{erg s}^{-1} \text{cm}^{-2} \) and the perpendicular wave number \( k_z = 0 \) as before. The initial conditions are the same as in Section 3.

The damping and heating profiles are shown in Figure 12, which can be compared to the weak reflection case shown in Figure 4. As before, blue shows time 3 s and red 4 s into the simulation, near the top of the chromosphere shortly after the pulse reaches the transition region. Between the two times, some of the pulse reflects in the Lare simulation, which reduces the Poynting flux at the leading edge, causing some to begin to propagate in the opposite direction. This is why the width of the pulse is larger and why the Poynting flux appears to have damped more than in HYDRAD. The derived heating rates are therefore slightly higher and occur over a smaller height range in HYDRAD than those in Lare.

The primary differences between the two simulations therefore appear to be due to the neglect of reflection, which would be an important generalization. While an empirical fit can give an estimate of the transmission coefficient at the transition region, a more general treatment needs to allow for reflection at any strong gradient in the Alfvén speed (for example, at shocks). It is not currently clear how to parameterize the reflection coefficient for any given gradient, so further work is required.

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**Figure 12.** Comparison between a wave simulation with HYDRAD (points) and Lare (lines) in the case of strong reflection, \( f = 1 \text{ Hz} \) and initial Poynting flux of \( 10^9 \text{ erg s}^{-1} \text{ cm}^{-2} \). Similar to Figure 4. The neglect of reflection causes a major discrepancy in the damping and heating rate, as some Poynting flux is expected to propagate back into the corona.