Efficient radiative transfer simulations for a broadband infrared radiometer—Combining a weighted mean of representative frequencies approach with frequency selection by simulated annealing

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1. Introduction

Radiative transfer (RT) simulations for broadband infrared (IR) radiometers are computationally expensive, because of the fine spectral structure inside the instrument bands. A typical example of the class of instruments we are concerned with here is the High Resolution Infrared Radiation Sounder (HIRS), which flies on the NOAA and MetOp series of satellites.

Even for the clear-sky case, simple RT simulations for this instrument are costly, because they require thousands of monochromatic RT calculations for each instrument channel, which are then integrated according to the instrument channel response function. In the presence of clouds this problem is even worse, since the individual monochromatic RT calculations then are computationally much more expensive.

Simulated annealing (SA) is a well-established method for solving global optimization problems, i.e., for finding the global maximum or minimum of a function [1]. It is particularly well suited for functions that depend on many variables and that are strongly nonlinear.
a case where many other optimization methods are not applicable.

In this article, we show how SA can be combined with a linear regression method to find representative frequencies and associated weights for HIRS RT simulations. The presented method is an alternative to the established correlated k method for efficient RT simulations in the IR spectral range. To our knowledge, there is only one other article Moncet et al. [2] that describes a similar approach to modeling broadband IR radiances.

We focus on the HIRS instrument to have a well defined test case to illustrate and test the method. However, the method itself is completely general and could be easily applied to other similar instruments. Furthermore, we focus on the clear-sky case, but the method is in principle equally applicable to cloudy radiances, provided that the RT model is sufficiently fast to calculate the required high spectral resolution reference simulations in a reasonable time.

The article is structured as follows: Section 2 describes the RT model that was used for this study, explains the weighted mean of representative frequencies approximation, and describes how simulated annealing can be used to find a near-optimal frequency grid. Section 3 investigates the methods performance and discusses its accuracy and limitations. Finally, Section 4 contains summary and conclusions.

2. Methodology

2.1. The ARTS model

The Atmospheric Radiative Transfer Simulator (ARTS) is a flexible and powerful radiative transfer model for the thermal radiation spectral range (microwave to infrared). It is written in the C++ programming language and freely available under a GNU public license from the web site http://www.sat.ltu.se/arts/. A general description of the clear-sky part of the model can be found in Buehler et al. [3]. For calculations in the presence of clouds, ARTS has two alternative scattering algorithms, a discrete ordinate iterative solution method [4] and a Monte Carlo method [5]. The model has been validated against other RT models in various intercomparison studies, for example Melsheimer et al. [6], Buehler et al. [7], and Saunders et al. [8].

ARTS has already been used in a large and diverse number of applications. It was for example applied to analyze sub-millimeter wave limb sounder data, both for operational instruments [9,10] and for planned future missions [11–13]. It was also used extensively to analyze operational meteorological microwave satellite measurements [14–17], to simulate future sub-millimeter cloud ice measurements [18–20], and to simulate radio occultation measurements [21]. Furthermore, it was used to investigate the impact of clouds on ground based measurements of the cosmic microwave background [22] and to simulate outgoing longwave radiation fluxes [23,24]. Other diverse applications include the analysis of ground based [25,26] and airborne [27] microwave data, the analysis of ground based Fourier transform infrared spectrometer data [28], the airborne retrieval of snow microwave emissivity [29], and the simulation of the full Stokes vector in inhomogeneous precipitation [30].

The strength of ARTS is that it can be applied to a wide range of flux calculations and sensor simulations, in different viewing geometries. A drawback of this flexibility is that it may require some work to set up the model for a particular application. To ameliorate this, we include setup files for some standard instruments. The basis for this article is the ARTS standard setup for HIRS simulations.

2.2. The HIRS instrument

The High Resolution Infrared Radiation Sounder (HIRS) is described for example in Smith et al. [31]. At present, data are available from 15 different HIRS instruments since 1978. HIRS measures radiation coming from Earth and its atmosphere primarily in the IR region of the spectrum. This includes both thermal IR (Channels 1–12) and near IR (Channels 13–19). Furthermore, there is one channel in the visible spectral range (Channel 20). We focus on the thermal IR channels here. They are widely spaced over a large part of the Planck curve for typical atmospheric temperatures. Fig. 1 shows the HIRS channel positions, relative to the atmospheric IR spectrum.

Figs. 2 and 3 show the spectral response of the different thermal radiation HIRS channels. They also show zenith opacity spectra of various trace gases, so that one can easily see which trace gas is important for which channel. Spectral response functions vary between different HIRS instruments (on different satellites). All figures in this section are for HIRS on NOAA 14.

2.3. Broadband infrared radiative transfer simulations

Broadband infrared radiometers, such as HIRS, have a typical bandwidth of 50 cm\(^{-1}\), much wider than the width of individual spectral lines (compare Fig. 2). The instrument measures the integral of the radiance spectrum, weighted by some broad channel response function.

The straightforward approach to simulating such measurements is to calculate the monochromatic radiance spectrum on a high resolution frequency grid, fine enough to resolve all spectral features, and then do the integration as a discrete sum over all frequencies with weights according to the instrument channel response. Explicitly,

\[
I_{\text{int}} = \int R(v)I(v)\,dv \approx \sum_i R(v_i)I(v_i)\Delta v_i,
\]

where \(I_{\text{int}}\) is the integrated radiance for an instrument channel, \(v\) is frequency, \(R(v)\) is the instrument channel response function, and \(I(v)\) is radiance. The radiances \(I(v_i)\) at the individual grid frequencies can be easily calculated by solving the monochromatic radiative transfer (RT) equation. It should be noted here that, to be consistent with the HIRS channel response functions, \(I\) should be in radiance units (W/(m\(^2\) sr Hz)), and not in brightness temperature units (K).

This approach is robust and exact, provided that the frequency grid is fine enough to resolve all spectral features.
Characteristics of a particular instrument, such as spectral response function shape and antenna pattern, can be simulated efficiently by multiplying the monochromatic radiances with a pre-calculated sensor response matrix [32]. Unfortunately, however, the straightforward approach typically requires thousands of individual RT calculations, one for each point in the frequency grid. For broadband IR instruments it is therefore only suitable for reference calculations, not for operational use.

2.4. The correlated k method

The correlated k method [33–35] is a method that is often used to reduce the number of frequency grid points for which monochromatic RT calculations have to be done. The basic idea of the method is to sort the frequency grid according to the absorption coefficient at the different frequencies. On the re-ordered frequency grid the spectrum is smooth and monotonic, so that it can be approximated with very few frequency grid points and linear interpolation between the grid points.

The advantages of the method are that it is efficient and conceptually easy. The disadvantage is that the exact sorting depends on pressure, temperature, and trace gas concentration. It is thus not straightforward to determine the optimal compromise for the frequency grid sorting, optimal in the sense that it minimizes the error over a
large ensemble of atmospheric cases and for different instrument viewing angles.

2.5. *The weighted mean of representative frequencies (WMRF) approximation*

The WMRF approximation introduced here is numerically quite similar to the correlated $k$ method, but its derivation and theoretical justification are different. The simple assumption underlying this method is that the radiance at a single frequency will be representative for the radiance at some other frequencies (where atmospheric optical properties are similar). Thus, the integrated radiance for a broad instrument channel can be approximated as

\[
I_{\text{int}} \approx \tilde{I}_{\text{int}} = \sum_{j=1}^{n} w_j I(v_j).
\]

(2)
with a set of \( n \) appropriately selected frequencies \( v_j \) and
associated weights \( w_j \). In contrast to the \( v_j \) in Eq. (1), the \( v_j \)
here can be reduced, so that there can be a considerable
saving of computational cost. (According to our results
presented later, less than approximately 20 \( v_j \) are
sufficient to accurately reproduce a reference calculation
with several thousand \( v_j \).)

We will call Eq. (2) the Weighted Mean of Representative
Frequencies (WMRF) approximation. The authors of
[2] also used Eq. (2) as a starting point for an RT algorithm
that they call Optimal Spectral Sampling (OSS). We are not
using that name here, since OSS refers to the entire RT
calculation strategy, including the frequency selection.
However, our method shares several key properties with
OSS, as will be discussed further below.

The obvious questions about Eq. (2) are how to select
the \( v_j \) and how to determine the associated \( w_j \). One
possible strategy would be to pick regular intervals in an
absorption-sorted frequency grid and assume linear
interpolation, in which case the method would be very
similar to the correlated \( k \) method.

If a set of frequency grid points \( v_j \) is given, then one can
easily determine the optimal weights \( w_j \) by multiple
linear regression over a set of reference calculations for an
ensemble of atmospheric states \( \alpha_i \). The optimal weights
are those that minimize a cost function that is some
measure for the error of the approximation.

Natural choices for the cost function are the absolute
RMS error \( E_{\text{abs}} \) or the relative RMS error \( E_{\text{rel}} \), defined as

\[
E_{\text{abs}}(v_j, w_j) = \text{rms}(I_{\text{int}}(\alpha_i, v_j, w_j) - I_{\text{int}}(\alpha_i)),
\]

\[
E_{\text{rel}}(v_j, w_j) = \text{rms} \left( \frac{I_{\text{int}}(\alpha_i, v_j, w_j) - I_{\text{int}}(\alpha_i)}{I_{\text{int}}(\alpha_i)} \right),
\]

where \( \text{rms}(\cdot) \) indicates the root mean square. As the
radiance varies strongly with frequency, the relative error
was chosen as the appropriate quantity to minimize.

Concerning the weights \( w_j \), we demand them to be
positive, consistent with our physical understanding of the
problem. Negative weights would mean partial
cancelation of terms, which is undesirable if the approx-
imation is to generalize well. In practice, this is imple-
mented by simply removing frequencies with negative
weights from the list of active frequencies.

As pointed out by [2], it is desirable that the sum of all
weights equals exactly one, so that Eq. (2) corresponds to
a true weighted mean. The actual sum of the weights
determined by regression is usually close to one, but not
exactly one. (The maximum deviation that we encoun-
tered for the final weights without normalization was
0.0002.) To force the weight sum to be exactly one, we
normalize the weights by dividing them by the actual
weight sum value. This differs from the approach of [2],
who do the regression only for frequencies 1 to \( n-1 \), and
assign weight \( w_n = 1 - \sum_{j=1}^{n-1} w_j \) to frequency \( n \).

Since we know how to find optimal weights \( w_j \) for a
given set of frequencies \( v_j \), we are left with the problem of
how to find the \( v_j \). Unfortunately, the number of possible
sets is combinatorially large, since there are

\[
\binom{N}{n} = \frac{N!}{n!(N-n)!}
\]

ways to select a set of \( n \) frequencies from a total of \( N \),
where \( \binom{N}{n} \) indicates the binomial coefficient. For example,
there are approximately \( 2.6 \times 10^{23} \) possibilities to select
10 frequencies out of a set of 1000. Hence, trying all
possible sets is impractical. At the same time, the cost
function \( E(v_j, w_j) \) is very nonlinear, so that analytical
methods to find the minimum are not applicable. Hence, a
stochastic method is required.

One possibility is to guess appropriate frequencies,
based on opacity, list of absorbing species, and other
factors, and then fine tune the initial selection by trial and
error. This was termed ‘pseudo-k’ approach by Frank
Evans, and used for example to create training data for the
proposed SIRICE and CIWSIR [18,19] sub-millimeter ice
cloud satellite missions.

A more formal approach is described in Wiscombe and
Evans [36], who apply the method to transmission instead
of radiance. The problem to select representative fre-
quencies arises similarly in that case. Their algorithm
starts with \( n = 1 \) and determines the best frequency \( v_1 \) by
an exhaustive search. Then more frequencies are added
successively, each time doing an exhaustive search for the
frequency that gives the largest improvement when added
to the existing set. Some special measures are
necessary to prevent the algorithm from being trapped in
local minima.

In the OSS article, [2] describe an algorithm which they
call Monte Carlo search, and which shares key properties
with our algorithm. We shall come back to it below.

2.6. Simulated annealing

The simulated annealing method is described for example in Kirkpatrick et al. [1]. It is a method of
randomly moving a simulated system through different
states, in order to find a global minimum in some cost
function, such as Eq. (3).

Assume that the system is initially in a state \( S_0 \) (for
example a random state). In our application of the method
each state \( S \) corresponds to a selection of frequency grid
points. To be fully explicit here, we first calculate
simulated spectra on a high resolution frequency grid
for an ensemble of atmospheric conditions. The state \( S \) of
the system is given by a selection of \( n \) frequencies among
all the available frequencies. We also call the frequencies
in \( S \) the active frequencies, and all others the inactive
frequencies.

The number \( n \) of frequency grid points \( v_j \) is fixed and
much smaller than the number \( N \) of frequency grid points
\( v_j \) in the high resolution reference calculations. The
optimization problem to be solved is to find the optimum
set of frequencies for a given \( n \). (Later, one can then
increase \( n \) until a desired accuracy is reached.)

The system is fully described by the set of active
frequencies \( S \), because the associated weights can be
calculated directly by linear regression over the ensemble.
of atmospheric conditions. In practice the regression is also over the set of possible instrument viewing angles, since the goal is to find a set of frequencies that works for all viewing angles.

To implement the simulated annealing method, one needs to define a function
\[ S' = \text{neighbor}(S), \]
which returns a new state \( S' \) of the system that is in some sense a neighbor of the original state \( S \). In our implementation of the method this function exchanges one of the active frequencies with a random inactive frequency. We will call the operation of changing \( S \) to \( S' \) a move of the system.

For the new frequency set \( S' \) the weights \( w_j \) are determined by multiple linear regression, and the cost \( E' \) is calculated according to Eq. (3). If \( E' \) is lower than \( E \) then the new frequency set \( S' \) is better than the original one \( S \), and the move is therefore accepted. However, if one were to accept only moves where \( E \) is decreased, the algorithm would run into a local minimum of \( E \) very quickly, instead of finding a global minimum.

The key feature of the simulated annealing method is that sometimes the new state is accepted even if it results in an increase of \( E \). This step is controlled by an acceptance probability function \( p(E-E', T) \). The classical choice of this function is
\[ p(E-E', T) = \begin{cases} 1 & E < E' \\ \exp((E - E')/T) & E \geq E. \end{cases} \] (6)

This choice of \( p \) function, and the notation used, suggest an analogy with the physical process of annealing in metallurgy, where heating and slow cooling is used to create a more regular crystal structure in a material. In the analogy, \( E \) corresponds to the energy of the system, \( T \) to the temperature, and \( p \) resembles the Boltzmann probability factor. Note that the ‘temperature’ \( T \) as defined by Eq. (6) has the same unit as the error (or energy) \( E \). Technically, the \( p \) function is used in our annealing code as follows: We do a test move and calculate \( p \). Then we create a uniformly distributed random number \( x \) in the open interval \([0,1] \) with the Matlab ‘rand’ function. The move is accepted if \( x < p \).

The algorithm starts with a high \( T \), so that nearly all moves are accepted. Then \( T \) is gradually reduced, so that the system is pushed towards states with lower \( E \), but without getting stuck in local minima. (The exact procedure is described in more detail below.) At the end, when \( T \) is very small, the system is effectively ‘frozen in’ and no more moves are possible. The algorithm will always lead to a state with very low \( E \), compared to the entire state space. We call this unrigorously ‘a global minimum’ here. It should be noted, however, that it is unlikely to find the true global minimum. But for the application described here (and many others) the true global minimum is only of theoretical interest, while in practice any state with very low \( E \) is acceptable.

Our algorithm shares key properties with the OSS algorithm described by [2]. In OSS, new frequency sets are also accepted with a certain probability that is adjusted as the algorithm progresses. However, OSS retains from the Wiscombe and Evans [36] algorithm the idea to use the state \( S(n) \) for \( n \) frequencies as the starting point for the search for the best state \( S(n+1) \). Our algorithm does not use this concept. For each \( n \) we start with a random state and a high ‘temperature’, and derive the solution by ‘cooling down’ the system, as described above.

For the study described here, the algorithm was implemented as a set of simple Matlab functions. The code is publicly available from our web page as part of the Atmlab software package (see Section 4 for address).

2.7. Example case HIRS

The HIRS instrument ideally serves to illustrate the method. High-resolution reference spectra for this instrument were calculated with the RT software ARTS for a set of 42 different atmospheric states. The atmospheric states were those from Garand et al. [37]. The Garand data set contains pressure, temperature, and concentrations of the trace gases \( \text{H}_2\text{O}, \text{O}_3, \text{CO}_2, \text{N}_2\text{O}, \text{CO}, \) and \( \text{CH}_4 \). The spectra were simulated for five different viewing angles, covering the viewing angle range of HIRS.

The frequency grid for the reference calculations contained 72,717 frequencies in total, equally spaced in frequency (but only within the instrument bands). The HIRS instrument setup was the one that is distributed along with ARTS. It contains channel response functions for all different versions of HIRS (on different satellites) as distributed by NOAA/NESDIS (http://www.star.nesdis.noaa.gov/smcd/spb/calibration/hirs/hirssrf.html). Figs. 2 and 3 show these functions for NOAA 14.

Since some channels overlap, it is theoretically most efficient to simulate all channels simultaneously, so that some frequencies are used for more than one channel. We can take this into account in the optimization process, so the system state \( S \) is a selection of frequencies for all channels, not just for one. However, in that case we impose the constraint that the frequencies contribute to the sum of Eq. (2) only if they are inside the same channel. This constraint is believed to increase the robustness and generalizability of the method.

The example results shown in Sections 3.1 and 3.2 use the simultaneous optimization approach. However, for the final set of optimizations for operational use, discussed in Sections 3.3 and 3.4, we decided to use the more primitive approach of optimizing the grid separately for each channel. That option leads to faster convergence of the algorithm, at the cost of slightly more frequencies in the final grid.

3. Results and discussion

3.1. Annealing algorithm performance

Fig. 4 shows how the error \( E \) between the fast calculation and the reference calculation develops with time (iteration count). This example calculation tries to
find the best combination of 133 frequencies to cover all HIRS bands for NOAA 14.

The chosen number of frequencies here is arbitrary, since the aim is only to illustrate how the method works. The annealing algorithm will optimize the frequency selection for any (fixed) number of frequencies. This implies no loss of generality, since the whole procedure can be easily repeated with a different number of frequencies. Later on (in Section 3.3) we present calculations where frequencies have been successively added until a desired accuracy was reached.

The annealing run contained almost 4 million iterations, which were executed in blocks of 10,000. After each block, the RMS error for the block was calculated, together with the maximum and minimum errors. The blue shaded region in the figure is the area between the minimum and maximum errors; the RMS error lies in between.

The evolution of the block RMS error is used to adjust the ‘temperature’ \( T \). If the error has not decreased between the last block and the block before, then \( T \) is reduced (in this case multiplied by 0.9). The idea of this cooling scheme is to only cool if the system has reached an equilibrium, marked by a steady error level. This avoids a too rapid cooling which would lead to the algorithm being trapped prematurely in a local minimum.

One practical problem is how to select the initial \( T \) at the start of the annealing run. We solve this by simulating one block of 10,000 iterations where we accept all moves and save their errors. Then we choose the initial \( T \) such that almost all of the moves would have been accepted. Specifically, it was set here so that 99% of the moves would have been accepted.

Choosing a too low initial \( T \) will lead to the algorithm not really exploring the full range of possibilities, and should therefore be avoided. Choosing a too high initial \( T \), on the other hand, will not affect the annealing result, but will waste computation time during which \( T \) is reduced to a more appropriate value. Our choice is quite conservative, which can be seen from the fact that at the beginning (at low iteration number) the error \( E \) decreases rapidly. This is due to the elimination of obviously bad frequency combinations, where for example some instrument channel has no frequency at all.

After this initial drop \( E \) changes only very slowly up to approximately iteration block 120, although \( T \) decreases steadily. From approximately iteration block 120 onwards the system ‘condenses’, because \( T \) is now so low that only rather good frequency combinations (that lead to low \( E \)) have a chance to be accepted.

Towards the end of the run the accepted moves get fewer and fewer. The frequency combination is near optimal, so it is unlikely to find a better one by random changes. At the same time \( T \) is now very small, so it is unlikely that a move that increases \( E \) is accepted. The algorithm is stopped if there are no more successful moves inside one entire block.

We find the convergence behavior that is seen in Fig. 4 to be general for all annealing runs that were carried out. It leads to a characteristic \( S \) shape in a log–log plot of \( E \) versus \( T \) (Fig. 5). The entire run took approximately 15 h on a PC.

![Fig. 4. Time development of ‘temperature’ \( T \) (green) and error \( E \) (blue) during one annealing run. Time is measured by iteration number. Parameters are evaluated and plotted once for every block of 10,000 iterations. The blue shaded region marks the area between maximum and minimum errors during each block. As an exception, the minimization was here done for the brightness temperature error in Kelvin, in order to have a more intuitive error measure. Due to this choice, the ‘temperature’ is also in Kelvin, since it has the same unit as the error that is minimized. Note that operational optimizations in Section 3.3 did not use this unit, but used the relative error in radiance. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image-url)
3.2. Example results for HIRS channel 12

We use the example of Channel 12 for HIRS on the NOAA 14 satellite to discuss the results of the annealing optimization. The top plot of Fig. 6 shows the high resolution atmospheric spectrum in blue. This is the spectrum for the near-nadir looking direction and the first of the 42 Garand atmospheric scenarios. The spectrum is dominated by water vapor absorption features in this frequency region. The spectral response function of HIRS Channel 12 on NOAA 14 is superimposed in green. The nine black vertical lines indicate the selected representative frequencies.

The selected frequencies cover nearly the entire range of observed brightness temperatures \( T_b \). What is more, they cover the \( T_b \) range quite uniformly, which is more apparent if the spectrum is sorted by increasing \( T_b \), as shown in the bottom plot of the figure. Numbers mark identical frequencies.

Here the relationship to a correlated k method is clearly visible. Both methods lead to regularly spaced representative frequencies in the re-sorted frequency grid. The difference is that in our case the spacing is the outcome of the optimization procedure, whereas for a correlated k method it is prescribed.

An obvious question about Fig. 6 is why the frequency pairs 3–8 and 4–9 are so close together in the bottom plot. Would it not be better to spread out the frequencies more homogeneously? The answer is that the sorting depends on the atmospheric conditions. Fig. 7 shows that for Garand scenario 2 the sorting is quite different from scenario 1 (and the other 40 are again different). The position of the WMRF frequencies is derived by minimizing the RMS error over all 42 scenarios, so they are a reasonable compromise for each individual scenario.

However, there is something else that is odd about frequencies 8 and 9. Their weights (shown by the green bars in the bottom plots of Figs. 6 and 7) are very small compared to the other weights. If the frequency set is really optimal in some sense, there must be a reason for the presence of these frequencies despite their small weights. The reason must be that these frequencies represent important aspects of the spectrum for certain atmospheric conditions. Indeed, a closer analysis reveals that they play an important role in some of the atmospheric scenarios, as shown in Fig. 8. For the conditions of Garand scenario 7, frequency 9 alone represents the most transparent regions of the spectrum, where \( T_b \) is the highest. For the conditions of scenario 12, frequency 8 alone represents the most opaque regions of the spectrum, where \( T_b \) is the lowest.

3.3. Operational optimization runs

WMRF frequency grids and weights were derived for a variety of HIRS sensors and are included with the ARTS model distribution. Specifically, we have calculated them for the HIRS instruments on the satellites TIROS N, NOAA 6–19, and Metop A.

The training data were based on the 42 Garand profiles, but contained each profile for three different surface emissivities, 1.0, 0.8, and 0.6, so that there are 126 training cases in total. Five different HIRS viewing angles were simulated for each training case.

Each channel of each instrument was optimized separately. The optimization was done iteratively, starting with 5 frequency grid points for each channel. An annealing run was carried out to minimize the relative error in radiance. If the final fractional relative error was above \( 10^{-4} \), the procedure was repeated with one frequency more. In other words, frequencies were added, until the desired accuracy was reached. For a scene brightness temperature of 300 K, the choice of \( 10^{-4} \) for the desired fractional relative accuracy corresponds to an absolute accuracy in brightness temperature of \( 10^{-4} \times 300 \text{ K} = 0.03 \text{ K} \), sufficient for accurate instrument simulation.

This iterative optimization procedure results in 5–22 frequencies for each channel. In the case of NOAA 14, the average number is 17 frequencies per channel.

3.4. Error analysis and test of generality

The derived operational frequency and weight sets for HIRS on NOAA 14 were used for testing the accuracy of the WMRF fast calculations. Over the 42 Garand scenarios, the RMS error of the WMRF fast calculations compared to the reference calculations in brightness temperature units was found to be 0.014 K. This very good agreement for the Garand scenarios is expected, since these are the scenarios used for training, so this is really only a test of our method’s self consistency.

A more ambitious test is to compare reference and WMRF calculations for an independent set of atmospheric scenarios. The diverse data set by Chevallier et al. [38] was
used for this purpose. More precisely, we started with the diverse Chevallier data sets for temperature, humidity, and ozone, which are a total of 15,000 atmospheric scenarios. From these 15,000 scenarios, 1,000 were drawn randomly for the test.

The Chevallier scenarios do not contain all trace gases of the Garand scenarios: they are missing CO₂, CH₄, N₂O, and CO. For the test calculations, these were set to VMR(CO₂) = 3.60e-04, VMR(CH₄) = 1.7e-6, VMR(N₂O) = 3.2e-7, and VMR(CO) = 0. These volume mixing ratios are typical tropospheric values for these species in the Garand scenarios, except for CO. For CO the assumption of a constant VMR value is not appropriate, so this species was simply neglected. This has no impact on the results, since CO is only spectroscopically active in the short wave HIRS channels (mostly Channels 13 and 14), but has no transitions within the thermal HIRS channels.

ARTS was used to simulate HIRS measurements for five different viewing angles for the 1,000 selected Chevallier scenarios. Both WMRF simulations and high resolution reference simulations were done. The WMRF simulations use on average 17 frequency grid points per channel, the reference simulations 3,827. The total run time on Intel Core2 8-core CPUs was approximately 6 min for the
WMRF simulations, and 150 h for the reference simulations. (These run times include the line-by-line calculation of absorption coefficient tables, which is done once at the beginning of each batch run. The fast calculation was done in a single batch run, the reference calculation was broken down into 10 batch runs so that it could be run on a computer cluster.)

Fig. 9 summarizes the results of this exercise. It shows for each channel the mean and the standard deviation of the difference between fast calculation and reference calculation, taken over the 1000 atmospheric cases and all viewing angles. Differences for most channels are small, below 0.05 K. Only for Channel 1 the mean difference is as large as 0.3 K. The reason for this discrepancy is that this channel has a very high opacity (compare Fig. 2), so that it receives radiation from the upper stratosphere. The Garand and Chevallier profiles are less consistent at these high altitudes than at lower altitudes.

3.5. Jacobians

The ARTS program can calculate analytical or semi-analytical Jacobians for various quantities, such as trace gas concentrations or temperature [3]. Each monochromatic radiance can be accompanied by monochromatic Jacobians, which are subject to the same post processing as the radiances, in order to simulate instrument antenna pattern and frequency response. In this context, it is interesting whether the selected WMRF frequency grid is suitable for Jacobian calculation, in addition to radiance calculation.
To check this we examine the Jacobians for water vapor for the first of the Garand scenarios. The left plot in Fig. 10 shows the Jacobians as calculated on the reference grid (solid curves) and on the WMRF grid (circles). Only HIRS channels sensitive to water vapor are shown. On first sight the agreement between fast and reference calculations appears excellent.

For a closer look, the right plot in Fig. 10 shows the relative difference ((fast-reference)/reference in percent). This plot reveals that at altitudes where Jacobians are small, relative differences approach 100%.

This result is not completely unexpected. Implicitly, the WMRF grid is optimized to represent those altitudes where changes have the strongest impact on the radiances. (Since the grid is selected to minimize the radiance error.) These are the altitudes where Jacobians are large. Frequencies, that represent altitudes where the Jacobian is small, tend to be optimized away.

One can conclude that the WMRF method as presented here is suitable for Jacobian calculations only if large relative errors outside the Jacobian peak area are acceptable. This issue should not be a problem.
for most retrieval algorithms, but the user should be aware of it.

A possible solution to get precise Jacobians throughout would be to include the Jacobians in the cost function (Eq. (3)) for the simulated annealing algorithm. However, this has obvious drawbacks: Firstly, it is expected to lead to a larger number of frequencies in the WMRF grid (for the same radiance accuracy). Secondly, different Jacobian species would have to be taken into account for the different instrument channels, since not all species are active in each channel.

3.6. WMRF versus correlated k

A quantitative comparison of the new method to a traditional correlated k method is beyond the scope of this article. However, a few qualitative remarks are in order.

The WMRF method was shown to give good calculation accuracy at significantly reduced computational cost, compared to reference calculations. On theoretical grounds, we believe it to yield a speed increase that is at least comparable to a traditional correlated k method, but we have not shown this explicitly.

It is likely that for most users practical considerations drive the decision on what method to use. Compared to the correlated k method and the pseudo-k method, our new method has the advantage that the procedure is completely automatic. However, it has the disadvantage that the simulated annealing procedure requires considerable computation time. The calculation of high resolution reference spectra also requires considerable computation time, but is necessary for both methods.

4. Summary, conclusions, and outlook

We presented a method to efficiently simulate broadband IR radiometer measurements. It uses two basic ideas. Firstly, the channel radiance can be approximated by a weighted mean of the radiance at some representative frequencies, where the weights can be determined by linear regression. (We term this the Weighted Mean of Representative Frequencies (WMRF) approximation.) Secondly, a set of representative frequencies can be found by simulated annealing.

The method was shown to yield good accuracy (better than 0.05 K) with only few representative frequencies (5–22 per HIRS channel).

Representative frequencies and weights for the HIRS instrument are available under a GNU public license as part of the ARTS radiative transfer program at http://www.sat.ltu.se/arts/. Currently included are data for the instruments on TIROS N, NOAA 6–19, and Metop A. We plan to add further instruments, as their characteristics become available.

The optimization method itself was implemented as a set of simple Matlab functions. The only inputs required are instrument channel response characteristics, and high resolution reference spectra for a set of atmospheric scenarios. Hence, the functions can be used together with any radiative transfer program, and for any sensor with broadly similar characteristics as HIRS. The method should also be suitable for fast radiation flux or cooling rate calculations, but we have not yet tried this.

The functions are distributed under a GNU public license as part of the Atmlab software package, and can be downloaded from http://www.sat.ltu.se/arts/tools/.

An interesting open issue is, whether the presented method performs equally well for the calculation of cloudy spectra, as for the investigated clear sky spectra. Intuitively we believe so, but it has not yet been demonstrated. To study this is beyond the scope of the present article, since we are lacking at the moment a good set of high resolution reference calculations for a diverse set of atmospheric states. This issue will be the subject of further study.
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References


