Precipitable Water at the VLA — 1990–1998

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Introduction

The amount of water vapor in the atmosphere above the VLA is an interesting quantity for two reasons: water vapor contributes significantly to the opacity of the atmosphere, and the fluctuations in water vapor are the dominant source of interferometric phase fluctuations. We currently have the capability to constantly monitor the phase stability of the atmosphere, with the site-testing interferometer (Carilli & Roy 1998), and we also have the capability to directly measure the opacity in any of the observing frequency bands of the VLA, with the array antennas themselves (through TIP scans - Butler 1996). However, TIP scans are done at essentially random frequency bands and times. Also, the site-testing interferometer has only been operating for a short time, so information over long time periods is not available from that instrument. If information on atmospheric water were available, then it could be used with atmospheric models to produce estimates of opacity quite reliably. The estimation of phase stability from total water is considerably less certain. Although it is generally true that more water vapor means more unstable phase conditions, this is not always strictly true. In the ideal case, it is desirable to know the full vertical distribution of water vapor in the atmosphere, but this is a quantity which is not easy to measure (it can be done with a variety of techniques, but we have no such capability at the VLA). However, even some crude indicator of the total amount of water vapor in the atmosphere is better than nothing. The total column of water is just such a beast. This is the equivalent depth of water which would result if all of the water vapor in the atmosphere were concentrated into a layer of liquid. It is also commonly referred to as the amount of precipitable water. While knowing the precipitable water yields little information regarding the phase fluctuations, it can be used to make a first order prediction of the opacity of the atmosphere.

Given measurements of surface temperature and dew point, a rough estimate of the precipitable water can be made. Uncertainties in the actual profile of water vapor make this a truly rough estimate, but it is probably good to a few 10's of %. Such measurements are available from the electronic versions of the observing logs which have been made for some time now. The logs from mid-1990 up to the present are currently available, so this is the time period that will be presented in this memo.

Deriving the precipitable water

Consider a column of liquid water with cross-sectional area A, and height h. This height (h) is the precipitable water. The mass of this column of liquid water is:

$$m_l = \rho_l \ A \ h \quad ,$$

where ρ_l is the density of liquid water ($\rho_l = 1 \text{ g/cm}^3$).

Now, the mass of the water vapor in an atmospheric column with cross-sectional area A is:

$$m_v = A m_w \int_0^\infty n_w(z) dz \quad ,$$

where m_w is the mass of each water molecule ($m_w = 18 \text{ amu}$), n_w is the number density of water molecules, and the integration is done over altitude z. This integration over altitude is why it is strictly necessary to have the full vertical distribution of water vapor in order to calculate the precipitable water. However, if the water vapor is distributed exponentially (similar to the bulk of the lower atmosphere) like:

$$n_w(z) = n_0 e^{-(z-z_0)/H}$$

where n_0 is the number density of water vapor at z_0 (the surface, practically speaking), and H is the scale height of the water vapor distribution (the e-folding distance), then the integral can be done analytically, resulting in:

$$m_v = A m_w n_0 H \quad .$$

To find the precipitable water, equate the mass of the vapor to that of the liquid (conservation of mass):

$$m_l = m_v \qquad \Rightarrow \qquad \rho_l \ A \ h = A \ m_w \ n_0 \ H$$

,

therefore,

$$h = \frac{m_w \ n_0 \ H}{\rho_l} \quad .$$

So, given a measurement of n_0 , and an estimate for H, the precipitable water can be estimated. From the ideal gas law, the number density of water molecules is related to the water vapor partial pressure (P_0) and the temperature (T_0) via:

$$n_0 = \frac{P_0}{k T_0} \quad ,$$

Making this substitution, the precipitable water is then:

$$h = \frac{m_w \ P_0 \ H}{\rho_l \ k \ T_0}$$

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The surface temperature, T_0 is measured and recorded regularly in the observing logs. In fact, it is measured and recorded on the visibility archive tapes as well, but getting at that data is logistically harder, and the accuracy and time resolution gain is not really needed. The surface water vapor partial pressure can be derived from the surface dew point (D) via (Clark 1987):

$$P_0 = e^{\left(1.81 + \frac{17.27 \, D}{D + 237.3}\right)}$$

where the dew point is in degrees C, and P_0 is in millibar. The dew point is also measured and recorded regularly in the observing logs. Again, electronic versions of these logs exist, and these can be parsed for the temperature and dew point quantities.

What to use for the scale height H? Formally, for an isothermal atmosphere in hydrostatic equilibrium the scale height is given by:

$$H = \frac{k T}{m_v g}$$

where g is the gravitational acceleration. If this were the right value for H, then substituting this would yield the following very simple equation for the precipitable water:

$$h = \frac{P_0}{\rho_l \ g}$$

Unfortunately, it turns out that the above formal expression for the scale height is not correct for water vapor. Given a typical surface temperature ($T = 10^{\circ}$ C), that expression would give a scale height of about 13 km. Observationally, the scale height of water vapor in the Earth's atmosphere is between 1.5 and 2 km (e.g. Ulich 1980). Therefore, the slightly more complicated expression (with scale height and surface temperature explicitly included) must be used. In this memo, a scale height of 1.5 km will be assumed. Since the derived precipitable water is linearly proportional to the assumed scale height, the results can be scaled as desired with little effort.

Results

Electronic versions of the observers logs from the beginning of September 1990 through the end of August 1998 were parsed for recorded values of temperature and dew point. Oddball values were excluded (quite often, for example, the dew point and temperature would be switched, or the dew point would not have a negative sign when it clearly should have). A total of 26659 valid combinations of temperature and dew point were found and used. The values were used to calculate the precipitable water (h) according to the above formula, and this value along with the date and time were recorded for analysis. Figure 1 shows a plot of the entire set of data obtained. The seasonal variation is readily apparent in the data. Figure 2 shows the data for 1997 only. The effects of weather systems can be seen clearly at this higher time resolution (variations on the scales of a few to 10 days).



Figure 1: All precipitable water data from September 1, 1990 to August 31, 1998.



Figure 2: Precipitable water data for 1997.

Monthly values

Figure 3 shows the monthly mean and minimum value (the absolute lowest value in all of the data for that month) for all of the data. Again, the seasonal variation is clearly evident. The wet summer months have a mean precipitable water which is more than twice what it is in the winter months. The typical mean precipitable water in the "winter" months (November - April) is about 5 mm, while in the middle of "monsoon" season (July and August), the typical mean precipitable water is as bad as almost 15 mm. The absolute very best conditions are about 1 mm of precipitable water in the months from December to April, about 2 mm in October, November, May, and June, about 3 mm in July and September, and about 6 mm in August.



Figure 3: Monthly mean (open stars) and absolute minimum (filled stars) values of precipitable water.

Hourly values

Figure 4 shows the hourly mean values for all of the data. No clear trend is apparent, which is somewhat surprising on first inspection. However, when looking at the winter data by itself, a clear diurnal variation of about 20% is seen (Figure 5). This must be because during winter night, a sizable portion of the atmospheric water freezes out, lowering the amount of vapor. During the summer, on the other hand, the atmosphere does not cool down enough to freeze out an appreciable fraction of the vapor, and so little diurnal variation is seen. Since the summer precipitable water values are larger than the winter ones, they mask the wintertime signature in the overall average.



Figure 4: Hourly mean values of precipitable water.



Figure 5: Hourly mean values for wintertime data only (November through April).

Comparison with measured opacity

Since water vapor is one of the primary contributors to the opacity at radio wavelengths, the opacity is expected to correlate very well with the amount of precipitable water. However, there is some disagreement about whether surface measurements can yield any reasonable estimate of the precipitable water (e.g. Reber & Swope 1972). In order to test whether the precipitable water derived via the technique outlined above has a good correlation with true opacity, I took the results of reliable TIP data taken at K-band (frequencies between about 21 and 25 GHz) over the last 3 years (177 TIP scans in total), and plotted the measured opacity (via the technique outlined in Butler 1996) against the estimated precipitable water (via the technique outlined above - the temperature and dew point are provided with each set of TIP data). The result is shown in Figure 6. A good correlation is seen, and a fit with a second order polynomial is also shown in Figure 6. This fit is of the type:

$$au = a_0 + a_1 \, h + a_2 \, h^2$$

where the three coefficients are: $a_0 = 3.8\%$, $a_1 = 0.23\%$, $a_2 = 0.065\%$. While individual data points can be significantly different from the fit, for the purposes of statistical analysis it seems quite valid to use the surface measurements to predict precipitable water (and hence opacity).



Figure 6: Measured opacity at K-band at the VLA (via TIP scans) compared with estimated precipitable water over the past 3 years.

Is this correlation unique to K-band, the VLA band where the opacity effects of atmospheric water are greatest? In order to test this, I took TIP data at Q-band (I used data only from frequencies between 42 and 44 GHz) from the same 3 year period (164 TIP scans in total), and plotted the same quantities (measured opacity against estimated precipitable water). The result is shown in Figure 7. Again, a good correlation is seen. The fit coefficients are: $a_0 = 5.5\%$, $a_1 = 0.19\%$, and $a_2 = 0.0026\%$.



Figure 7: Measured opacity at Q-band at the VLA (via TIP scans) compared with estimated precipitable water over the past 3 years.

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References

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