

Scientific objectives and instrumental requirements of the IR spectrometer VenSpec-H onboard EnVision

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ABSTRACT

Hooray! The ESA EnVision mission is adopted.

Onboard the spacecraft, there will be a suite of three spectrometers, VenSpec. One of these is called VenSpec-H where the H stands for high spectral resolution. Its scientific objectives consist in measuring variations of minor species' abundances in the atmosphere of Venus. H₂O, SO₂, CO and OCS will be measured to characterize the potentially ongoing volcanic activity. These observations will allow us to understand both the importance of volatiles in volcanic activity on Venus and their effect on cloud maintenance and dynamics. VenSpec-H will measure these molecules in nadir viewing geometry, in infrared transparency windows of Venus' nightside to probe the troposphere and in infrared spectral ranges on the dayside to measure the mesosphere. In this paper, the scientific requirements enabling our scientific objectives will be demonstrated. An intercomparison exercise was first led to reproduce modelled and observational reference spectra. The molecular vertical profiles, the aerosols' model and the CO₂ continuum contribution were validated for the different spectral windows. This enabled us to determine the spectral bands, their bandwidth and the resolving power necessary for our purposes. Along the way, we identified possible improvements and science avenues. Some of them impact the instrument design, such as the need for polarimetric measurements. Others are related to remaining uncertainties in the model and laboratory measurements that will complement the investigation.

Keywords: EnVision, VenSpec-H, radiative transfer theory, Venus, molecular species, volcanic activity, instrumental requirements

1. INTRODUCTION

Despite their similar size and bulk composition, Venus and Earth have followed different geological evolution paths. In three years of radar mapping (1990 to 1993), Magellan showed that Venus has abundant volcanic and tectonic features but was unable to detect ongoing geological activity. Thousands of volcanoes from 1.5 to 280 km in diameter were imaged without detecting any new lava flows in the landscape [1]. Fifteen years later, Venus Express was launched with a payload focusing on atmospheric science. The mission (2006-2014) was a success and provided hints of active volcanism. More specifically, three instruments onboard Venus Express brought insights to this investigation. Firstly the Visible and Infrared Thermal Imaging Spectrometer (VIRTIS) detected local anomalies in the emissivity maps [2]. The infrared radiation coming from three volcanic regions was different to that from the surrounding terrain and interpreted as coming from relatively fresh lava flows

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that had not yet experienced significant surface weathering. Although these flows were found to be less than 2.5 million years old, the study could not establish whether there is still active volcanism on the planet. Secondly, the "Spectroscopy for Investigation of Characteristics of the Atmosphere of Venus" (SPICAV) UV channel was used to measure the abundances of SO₂ above the cloud level [3]. Complementing the series of measurements done by UVIS onboard Pioneer Venus during the 1970s and 1980s [4–6], the SPICAV-UV measurements showed a sharp rise in the sulphur dioxide content of the upper atmosphere in 2006–2007, followed by a gradual fall over the following five years [7]. As suggested by Marcq et al. (2020), these episodic sulphur dioxide injections to the cloud tops may be caused either by periods of increased buoyancy of volcanic plumes, or, in the absence of active volcanism, by long-period oscillations of the general atmospheric circulation. Finally, the most striking evidence for active volcanism on Venus came from the Venus Monitoring Camera (VMC) in 2014. VMC spotted localised changes in the surface brightness between images taken only a few days apart in 2008 and 2009. These infrared "flashes" over the edges of the rift zone Ganis Chasma are thought to be caused either by hot gases and/or lava released from volcanic eruptions [8]. Since Venus Express, more investigations have been carried out and recently, Sulcanese et al., (2024) identified evidences of ongoing volcanic activities using the Magellan dataset [9].

To determine whether Venus is geologically active, a comprehensive payload is necessary: the surface and subsurface need to be characterized but information about the atmosphere is crucial. This is why EnVision will also carry a complete spectroscopy suite called VenSpec [10] including an infrared mapper, VenSpec-M [11], an ultraviolet spectrometer, VenSpec-U [12] and an infrared spectrometer, VenSpec-H [13].

2. SCIENTIFIC OBJECTIVES AND REQUIREMENTS

The science objectives of the VenSpec suite are to search for temporal variations in surface temperatures and tropospheric concentrations of volcanically emitted gases, indicative of volcanic eruptions; to study surface-atmosphere interactions and weathering by mapping surface emissivity and tropospheric gas abundances; and to map the variability of trace species, cloud and aerosol properties and to distinguish intrinsic from extrinsic (e.g. volcanic emissions) variabilities in the mesosphere. Observations will provide insight on the spatial distribution of trace gas which is essential to understand the main chemical cycles on Venus. A typical spatial resolution of 100 km is sufficient to resolve most of the features.

Observing Venus' troposphere in the infrared is challenging. On the dayside of the planet, the reflected sunlight on the cloud deck prevents measuring below the clouds. The nightside of the planet must then be targeted. Straylight from the dayside and high density of CO₂ rendering the atmosphere opaque limit the number of useful spectral ranges. Molecular species are probed in specific spectral windows called transparency windows centered at 1.0, 1.1, 1.2, 1.3, 1.7 and 2.3 μm [14]. They have been used extensively since the 1990s to retrieve SO₂, OCS, CO, H₂O and HCl [15–21] in the lower layers of the atmosphere but so far only vertical profiles of the H₂O could be determined.

The dayside measurements provide information about the mesosphere. In synergy with VenSpec-U the possible exchange occurring between the layers below and above the clouds will be investigated, especially in case of the detection of a plume. In preparation for the mission, potential impacts of a volcanic plume have been discussed [22] and modeling efforts are ongoing [23, 24].

Based on our knowledge so far, the science requirements for the mission were decided upon and those related to VenSpec-H are summarized in Fig. 1.

R2-C-10 (Sect. 3.5.1)	Map tropospheric gases at 0-45 km altitude: (1) H ₂ O partial column random error: < 10% (and < 20% below 15 km altitude) (2) HDO partial column random error: < 20% (and < 30% below 15 km altitude) (3) CO partial column random error: < 14% (4) OCS partial column random error: < 18% (5) SO ₂ partial column random error: < 30% at a horizontal resolution of < 100 km (and < 200 km for H₂O and HDO below 15 km altitude), to detect gradients related to surface-atmosphere and cloud interaction processes.
R2-C-20 (Sect. 3.5.3)	Map mesospheric gases at 65-80 km altitude: (1) SO ₂ partial column random error: < 20% (2) SO:SO ₂ partial column random error: < 25% at a spatial resolution of 5 (small-scale convection and vertical mixing) to 50 km at UV wavelength (3) H ₂ O partial column random error: < 20% (4) HDO partial column random error: < 20% (5) H ₂ O:HDO partial column random error: < 25% (6) SO ₂ partial column random error: < 20% (7) CO partial column random error: < 20% (8) OCS partial column random error: < 20% at a spatial resolution of <50 km at IR wavelength in order to assess coupling between surface (volcanic activity, topographic features), cloud and mesospheric measurements.

Figure 1. EnVision Science observation requirements summary extracted from the Red Book [25]. "In the left-hand column, the EnVision requirements number is noted, together with a reference to the sections describing further the requirement flow-down and how the science objectives are met applying the intended observation strategy and mission payloads." The requirement R2-C-20 (2) is associated to VenSpec-U [12, 26, 27].

Simultaneous observations of different species at a high spectral resolution will be the asset of our VenSpec-H experiment. Few investigations reported measurements of different molecular species in the same spectra. Nevertheless anti-correlation was highlighted between SO₂ and H₂O [28] or OCS and CO [20]. The new instrumentation will enable us to derive a map of these four molecular species altogether. This will be done by measuring a wide spectral range in the 2.3 μm window at high spectral resolution. As will be discussed, the bandwidth is limited though by the spectral resolution that needs to be as high as possible to be able to differentiate the ro-vibrational lines for each molecule. This requirement will be translated into an increase of the resolving power by a factor of 4-5 compared to VIRTIS-H/Venus Express (R=2000). A higher spectral resolution has already proven being decisive in highlighting variation of SO₂ in the 2.46 micron band [29]. Using radiative transfer tools and our expertise with the NOMAD-LNO instrument onboard ExoMars Trace Gas Orbiter [30], we describe the instrumental requirements of this new experiment capable of measuring trace gases at high resolution both in the troposphere and in the mesosphere of the Venus atmosphere.

3. MODELING THE VENUS ATMOSPHERE

3.1 RADIATIVE TRANSFER CODE

In order to define the instrumental requirements of a new experiment, such as the Signal-to-Noise ratio (SNR), the resolving power (RP) and the spectral ranges, our radiative transfer code ASIMUT-ALVL was used extensively. ASIMUT is a modular program for radiative transfer calculations in planetary atmospheres [31]. One of the main particularities of the software is the possibility to retrieve columns and/or profiles of atmospheric constituents simultaneously from different spectra, which may have been recorded by different instruments or obtained under different geometries. This allows the possibility to perform combined retrieval, e.g. of a ground based measurement and a satellite-based one probing the same air mass, or from spectra recorded by different instruments on the same platform [32]. This Radiative Transfer code derives the Jacobians analytically and includes the Optimal Estimation Method (OEM) [33], using diagonal or full covariance matrices.

Initially developed for the Earth atmosphere, its applicability has been extended to extraterrestrial atmospheres, such as those of Mars [34–36] and Venus [37]. ASIMUT has been coupled to SPHER/TMATRIX [38, 39] and LIDORT [40] codes to include the complete treatment of the scattering effects into the radiative transfer

calculations [41].

ASIMUT-ALVL can be used as a forward modeling tool and also as a retrieval module. It allows to fit simultaneously or sequentially different parts of one or more spectra, to fit the surface temperature, to fit column/vertical profiles for molecular species and for aerosols and to fully characterize the outputs (averaging kernels, errors, degrees of freedom, etc.).

3.2 MOLECULAR DATA

When simulating the Venus atmosphere in the infrared spectral ranges, eight molecular species are needed: CO₂, H₂O, HDO, CO, SO₂, HCl, HF and OCS. Each of them have their own absorption spectrum. Fig. 2 shows the line intensities directly downloaded from HITRAN Online (<https://hitran.org>). It shows how interesting the 2.3-2.5 micron spectral range is. The CO band is visible at one end and the SO₂ one at the other. HF lines are more spaced to one another, due to the rotational constant of the molecule. In terms of intensity, the SO₂ band is weak compared to the rest.

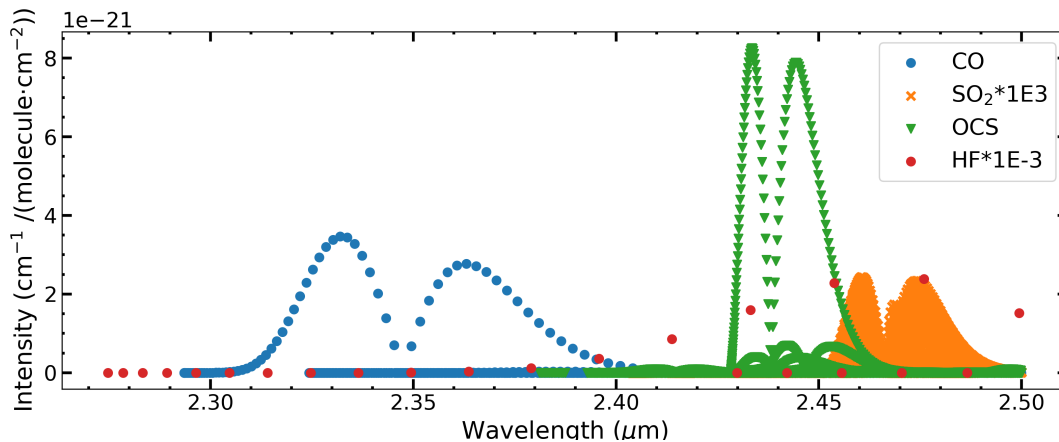


Figure 2. Line intensities of CO in blue, SO₂ in orange with a multiplicative factor of 10³, OCS in green and HF in red with a multiplicative factor of 10⁻³. The values have been downloaded from HITRAN Online on 24/07/2024 [42–74]. Water lines are not shown for better readability but they do absorb in that spectral range.

The line parameters were extracted from the HITRAN Online database [42] using HAPI [75]. The broadening line parameters have not been modified accounting for the CO₂ buffer atmosphere. This was a deliberate choice as in the frame of this investigation, our model still needed to be validated. Line parameters associated to a CO₂-atmosphere are available in the literature [76–79] and will be taken into account in further works [80]. HITEMP2010 [81] and HITRAN Online for H₂O were both considered for this investigation. The spectroscopic databases and their different versions are regularly the topic of validation papers [82–84] as none is perfect. We decided to use the most recent one at standard temperature for traceability reasons, firstly and also because the conclusion reached with this linelist should still hold with a more complete linelist, more consistent for the high temperature of the Venus atmosphere.

The vertical profiles of the different molecules are shown in Fig. 3.

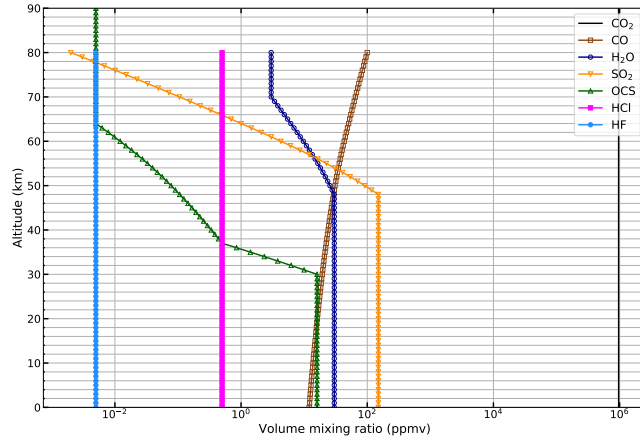


Figure 3. Molecular species profiles used in this work [85–92].

The high pressure of the Venus’ troposphere leads to collisional effects on molecular spectra [93]. No theoretical model is perfect yet and *ad hoc* solutions are used. Especially, two important effects of pressure were mitigated in our radiative transfer model. Firstly, to account for the far wings, each of the absorption lines is calculated on a 125 cm^{-1} spectral interval. A Voigt profile was used for all molecules except for CO_2 which requires a sublorentzian profile [94]. Secondly, the CO_2 continuum was prescribed using the empirical values based on the VIRTIS /Venus Express spectra [94–96]. This parameterization has proved efficient in previous investigations [21,94,97].

3.3 ATMOSPHERIC DATA

ASIMUT-ALVL was used to calculate synthetic spectra, using a line-by-line approach. Simulations of the day side and the night side of the Venus atmosphere were performed in the infrared spectral region spanning wavelengths from 1 to 2.6 microns. Eight molecular species were included, CO_2 , H_2O , HDO , CO , SO_2 , HCl , HF and OCS , as well as the aerosols and the CO_2 continuum contribution. The concentration profiles of the molecular species are based on the literature [85–88,92,98] and further refined into a common reference scenario defined at the mission level. Rayleigh scattering was also taken into account.

The temperature and pressure profiles were obtained from the literature [86,90], as shown in Fig. 4.

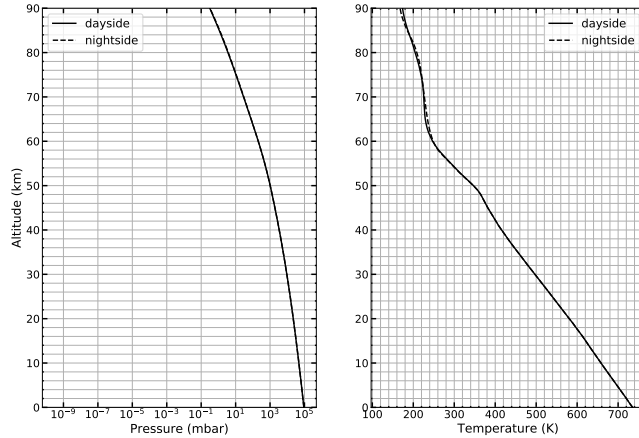


Figure 4. Pressure (on the left) in log scale and temperature (on the right) profiles in function of altitude, used in this work [86, 90].

The description of the cloud layer was based on former studies, namely [95, 99]. Because the particles are liquid and approximately spherical, and because the wavelength of the light is of the same order as the radius of the particles, a Mie code is appropriate for the calculation of their scattering properties. The particle number densities of four cloud modes were distributed from 0 to 100 km according to Fig. 12 in [95]. The modes themselves are described in Table 1. We assumed a lognormal distribution of sizes and the refractive indices were taken from [100].

Mode	mean radius [μm]	σ [-]
1	0.3	1.56
2	1.0	1.29
2p	1.4	1.23
3	3.65	1.28

Table 1. Mean radius and variance of the different modes of the Venus' aerosols.

4. INSTRUMENTAL REQUIREMENTS

Based on the quantities described in section 3, spectral simulations were performed. The radiance of the nightside comes from the thermal emission of the surface and atmosphere while on the dayside, where we considered a solar zenith angle of 30° , there is an additional solar reflectance contribution. We used our tool to determine:

- the limits of the spectral ranges and the resolving power ;
- the Signal to Noise ratio.

4.1 BANDWIDTH AND RESOLVING POWER

The baseline of our instrument is based on SOIR/Venus Express [37, 101] and NOMAD-LNO/ExoMars Trace Gas Orbiter [102, 103]. To retain much heritage from these successful instruments, the optical concept of our VenSpec-H spectrometer is based on an echelle grating coupled to a high-performance, actively-cooled detector [13]. The characteristics of the echelle grating that define the free spectral range and the instrument line profile, and of the detector, i.e. its size in pixels, need to be taken into account when defining the scientific requirements. Bandwidth and spectral resolution must be balanced out. The resolving power of a grating spectrometer with well corrected optics is independent of the wavelength. It defines the smallest difference in wavelength of two

spectral lines that can be distinguished from one another. The bandwidth is the wavelength range seen by the detector. They are defined by the following Eq. (1):

$$R_{max} = \frac{n_{pix} \times 10000}{\lambda \times BW} \quad (1)$$

where R_{max} is the maximum resolving power, n_{pix} is the number of pixels in the spectral direction and BW is the bandwidth.

As an example, let's imagine a detector of 500 pixels wide, in the spectral direction. To achieve a resolving power of 7000 in the 2.5 micron band, Eq.1 indicates that the bandwidth cannot be larger than 285.7 cm^{-1} . Besides, the minimum value of bandwidth assumes that the pixel pitch is the limiting factor and that the Full Width at Half Maximum of the instrumental line profile, i.e. the image of the slit for monochromatic illumination, is not wider than one pixel. If a slit is used with an equivalent width of more than one pixel, the maximum bandwidth is divided by the slit width in pixels. In other words, the higher the resolving power the narrower the bandwidth for a fixed number of pixels.

A trade-off must be found here. On one hand, measuring simultaneously CO and SO₂, so far apart in wavelength, as shown in Fig. 2 imposes a bandwidth of several hundreds of wavenumbers. On the other hand, a high resolving power is required to distinguish the spectral features of H₂O in the 1.17 micron band, as shown in Fig. 5. This led to strong constraints on the optical design [13, 104].

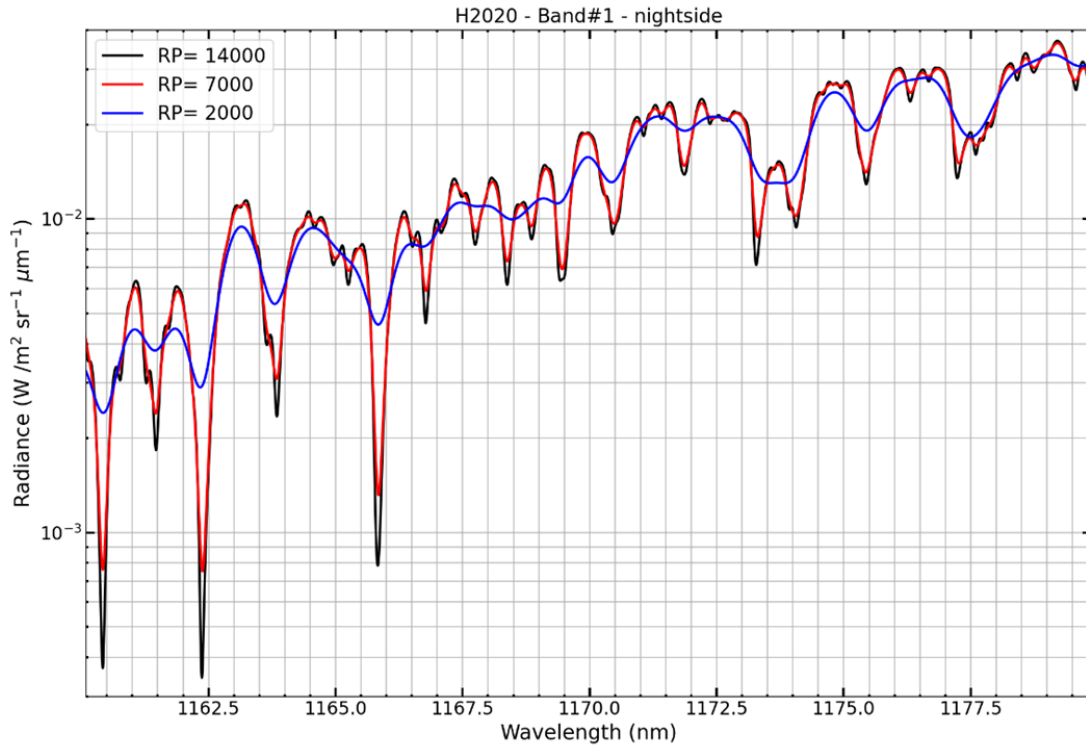


Figure 5. Simulated radiance of the Venus atmosphere at different resolving powers: 14000 in black, 7000 in red and 2000 in blue. To distinguish between the different spectral lines, a resolving power of at least 7000 is required.

The physical characteristics of the grating may deviate from the ideal band centers when deriving them only based on spectroscopy. One band center may be selected very precisely. The other wavelength ranges probed

will flow down from this first choice. Considering the scientific objectives of the mission and of the instrument, the cold overtone vibrational band $3\nu_3$ (003-000) of SO_2 , centered at $2.45 \mu\text{m}$, is considered as the primary target and the optical design was optimized accordingly.

4.2 SIGNAL-TO-NOISE RATIO

To determine the required Signal-to-Noise ratio (SNR), radiances were simulated at two resolving powers (RP= 7000 and RP= 8500) and noise was added before performing the retrievals with ASIMUT-ALVL. The science objective of 10% accuracy on each molecule was the target (except in the spectral band probing the near-surface contents), see Fig. 1.

The error budget on the retrieved abundances was estimated considering the following contributions:

- Random error: retrieval error directly from the fit
- Systematic errors: cross-sections (3%), raytracing (1%) and source term (night:1.2% - day:3.6%)

The considered systematic errors are originating from unavoidable uncertainties in the radiative transfer model. The uncertainties on the cross sections are linked to the choice of spectroscopy (HITEMP, HITRAN, with or without CO_2 -broadening, ...). All line parameters are not available yet for Venus' conditions. Besides the error on intensities for instance for H_2O varies from HITRAN error level 8 to 3, i.e., from less than 1% to above 20% relative error. The other molecules errors are mostly around 5 to 10% on their line intensities. In our error budget, we assumed a 3% error on the calculated cross-sections based on these HITRAN parameters. The raytracing error is defined based on what impact could a pointing error have in the optical path length. The source term errors on the nightside were calculated based on the impact on the Planck function when varying the surface temperature by 1K. For the dayside a factor 3 was added to this error to take into account the complexity of the incoming radiation.

The error budget in % was plotted against the SNR values, as shown in Fig. 6. The SNR value to meet the required accuracy was obtained by interpolation as shown by the dotted lines in Fig. 6. This was done for each molecule in each band.

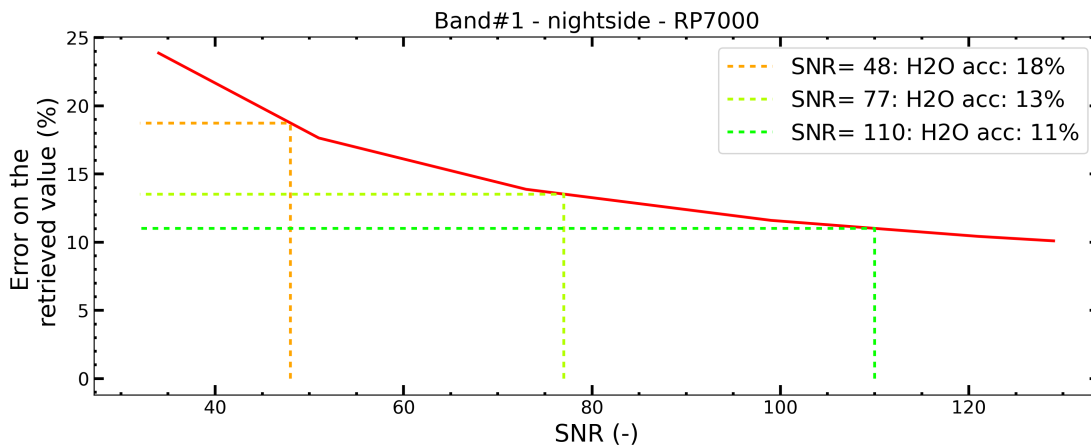


Figure 6. Error budget of Band#1 on the nightside for H_2O . As expected, the accuracy depends on the achievable SNR.

5. CONCLUSIONS

In this investigation, the molecular vertical profiles, the aerosols' model and the CO_2 continuum contribution were validated for the different spectral windows. This enabled us to determine the spectral bands, their bandwidth

and the resolving power necessary for our purposes. These values were confirmed as acceptable by performing retrievals for each spectral band. Diagnostics tools, such as the residuals of the fit and the degree of freedom for each species were analysed. The science requirements are driven by the lowest resolving power enabling the detection and the retrieval of the abundance of the different molecular species. The resolving power vs bandwidth trade-off has been investigated bearing in mind instrumental capabilities to reach a sensible solution, allowing us to resolve rovibrational lines with a high spectral resolution and cover an as wide as possible spectral range to measure simultaneously different molecular trace gases.

Table 2 summarizes the results obtained in this investigation.

Nightside						
	Band limits (nm)	Band limits (cm ⁻¹)	bandwidth (cm ⁻¹)	Required SNR	Molecular species	Altitude range (km)
Band#1	1160–1180	8474–8621	147	48	H ₂ O, HDO	0 – 15 km
Band#2a	2340–2420	4132–4274	142	126	CO, H ₂ O, HDO, OCS, HF	30 – 45km
Band#2b	2423–2507	3989–4127	138	126	SO ₂ , H ₂ O, HDO, OCS, HF	30 – 45km
Band#3	1704–1747	5724–5868	144	70	H ₂ O, HCl	20 – 30km
Dayside						
	Band limits (nm)	Band limits (cm ⁻¹)	bandwidth (cm ⁻¹)	Required SNR	Molecular species	Altitude range (km)
Band#2a	2340–2420	4132–4274	142	100	CO, H ₂ O, HDO, OCS, HF	55–80km
Band#2b	2423–2507	3989–4127	138	100	SO ₂ , H ₂ O, HDO, OCS, HF	55–80km
Band#4	1367–1394	7174–7315	141	100	H ₂ O, HDO	55 – 80 km

Table 2. Spectral bands, as defined after this investigation. The required Resolving Power is 7000.

In the near future, we will strengthen our model by comparing it to previous datasets from VIRTIS /Venus Express [96, 105]. Possible improvements are related to remaining uncertainties in the model such as for the CO₂ continuum treatment. A campaign of laboratory measurements that will complement the investigation has started [80] and will continue. New science avenues impacted the design, such as the need for polarimetric measurements [106].

ACKNOWLEDGMENTS

This work has been performed with the support of the Belgian Science Policy Office (BELSPO) contract 4000144206, with the financial and contractual coordination by the ESA Prodex Office. EM acknowledges support from CNES and ESA for all EnVision-related activities.

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