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Identification of short and long-lived atmospheric trace gases from IASI space observations

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Key Points:

- The whitening transformation allows analyzing anomalies in hyperspectral infrared spectra
- In a biomass burning plume we identify nine compounds including a first observation of glycolaldehyde
- On spectra observed a decade apart we identify atmospheric changes in eight long-lived halocarbons

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Abstract

In recent years, major progress has been made in measuring weakly absorbing atmospheric trace gases from high spectral resolution space observations. In this paper, we apply the so-called whitening transformation on spectra of the Infrared Atmospheric Sounding Interferometer (IASI), and show that it allows removing most of the climatological background from spectra, leaving a residual that contains those spectral signatures that depart from normality. These can subsequently be attributed to changes in the abundance of trace species. This is illustrated for two diverging cases: (1) a biomass burning plume from the 2019/2020 Australian bushfires, leading to the unambiguous identification of nine reactive trace gases, including a first observation of glycolaldehyde; (2) spectra observed a decade apart, from which changes in eight long-lived halogenated substances are identified; three of them never observed before by a nadir sounder.

Plain Language Summary

In recent years, several techniques have been developed for the detection of gases present in very small quantities in the atmosphere, which has significantly improved our knowledge on atmospheric composition and chemistry. In this paper, we describe a powerful complementary technique that transforms atmospheric spectra to highlight and attribute the spectral signatures of different species. We apply it on spectra measured by the spaceborne IASI (Infrared Atmospheric Sounding Interferometer) for two different cases: (1) a plume from the 2019/2020 Australian bushfires leading to the clear identification of nine rare gases including a first observation from space of glycolaldehyde; (2) spectra observed a decade apart, of which eight long-lived halogenated substances were identified; three of them never observed before by a nadir sounder. The detection of these reactive and long-lived trace gases is very promising for the monitoring of their temporal evolution and the attribution of their sources.

1 Introduction

The outgoing longwave radiation spectrum of the Earth-atmosphere system features strong absorption bands of CO_2 , H_2O , CH_4 , N_2O , O_3 and CO . Routine measurements of several of these strong absorbers are now available from nadir observing polar orbiting infrared sounders, which have led over the years to major advances in our understanding of their distribution (Crevoisier et al., 2014). Unexpectedly, hyperspectral nadir sounders have also proven most useful for the measurements of other, much weaker, absorbers. Harries et al. (2001) for instance demonstrated that infrared sounders can be used to monitor long-term changes of the ozone depleting halocarbons CFC-11 and CFC-12 (see also Coheur et al. (2003)). Beer et al. (2008) and Coheur et al. (2009) showed the first observations of NH_3 , C_2H_4 , HCOOH , CH_3OH and PAN in fire plumes and above highly polluted urban areas. Progress in retrieval techniques (see Franco et al. (2018) and references therein) has now allowed measuring these and other compounds also outside of highly concentrated plumes, in turn leading to major discoveries on their emission sources, e.g. Stavrakou et al. (2011) and Van Damme et al. (2018). Noticeable progress has been made using the Infrared Atmospheric Sounding Interferometer (IASI) on-board the Metop-A/B/C platforms (Clerbaux et al., 2009). The IASI instrument measures a very large continuous part of the thermal infrared ($645\text{--}2760\text{ cm}^{-1}$) and, thanks to its high radiometric performances, high spectral resolution (0.5 cm^{-1} , apodized) and bi-daily global coverage, has played a major role in the monitoring of weakly absorbing trace gases (e.g. Clarisse et al. (2011), Hilton et al. (2012) and Franco et al. (2019)). As a summary of past achievements, the list of 29 species that have been observed with IASI, either globally or in concentrated plumes is given in Table 1.

In recent years, remote sensing of weakly absorbing trace gases has benefited greatly from the discovery of improved algorithms for detecting very small features in spectra. In

particular, the method introduced by Walker et al. (2011) increased drastically the detection sensitivity over traditionally approaches and formed the basis of several quantitative retrieval techniques (Franco et al., 2018; Taylor et al., 2018; Clarisse et al., 2019). In this paper, we introduce a complementary technique based on ‘whitening’ spectra, which is suitable for analyzing spectra with unknown signatures. In the next section, we review briefly the detection method of Walker et al. (2011) and show how it naturally leads to the concept of whitening spectra. In section 3, we apply it on spectra from the 2019/2020 Australian fires and show that it enables, much easier than with alternative methods, to highlight and identify the different trace gas signatures, leading to the unambiguous identification of nine reactive trace gases. An entirely different application of whitening occurs when spectra are compared to those of a different period. As we show in section 4, whitening allows in this instance, highlighting small changes in a series of long-lived halogenated substances, most of which have until now never been identified in IASI spectra.

Table 1. Chemical species observed by IASI. The species reported in this paper which are observed for the first time with a nadir sounder are indicated in italics.

Greenhouse gases and ozone-related substances (14)	H ₂ O, CO ₂ , CH ₄ , N ₂ O, O ₃ , HNO ₃ , CFC-11, CFC-12, HCFC-22, CCl ₄ , CF ₄ , <i>HCFC-142b</i> , <i>HFC-134a</i> , <i>SF₆</i>
Air quality and VOCs (12)	CO, CH ₃ OH, HCOOH, CH ₃ COOH, CH ₃ C(O)CH ₃ , C ₂ H ₂ , C ₂ H ₄ , NH ₃ , HCN, PAN, SO ₂ , OCS
Concentrated plumes (7)	HCl, H ₂ S, C ₃ H ₆ , C ₄ H ₄ O, HONO, HCHO, <i>HOCH₂CHO</i>

2 Whitening transformation

The detection method presented in Walker et al. (2011) is based on a formula that quantifies in a single number the integrated spectral contributions of a target species, in a given spectral range. It relies on the mean $\bar{y}$ and the associated covariance matrix S of a set of spectra that are representative for IASI observations in the absence of enhancements of the target species. The pair $\{\bar{y}, S\}$ describes the distribution of spectra and how they are expected to vary (driven mostly by differences in temperature, surface emissivity, water vapor and interferences with other absorbers). The single number is sometimes referred to as a hyperspectral range index (HRI) (Van Damme et al., 2014) and is defined as

$$\text{HRI} = \frac{K^T S^{-1} (y - \bar{y})}{\sqrt{K^T S^{-1} K}}, \quad (1)$$

with K the Jacobian of the target species calculated from radiative transfer simulations (Van Damme et al., 2014). This formula was originally derived from the least squares estimate, and in that context the covariance matrix can be interpreted as a generalized noise covariance matrix, where all background variability is treated as noise. Note that, by construction, the HRI is normalized to provide a mean of zero and a standard deviation of one on the set of background spectra. Its value can therefore be interpreted in terms of standard deviations from the mean. Higher values (typically above 4σ in absolute value) usually indicate the presence of an anomalous abundance of the target species (or at least an anomaly in the spectrum that sufficiently resembles its Jacobian). The expression can be interpreted geometrically as the weighted projection of the Jacobian onto the zero-centered observed spectrum.

It turns out that Equation (1) has been known since the 1970s (Reed et al., 1974) under the name ‘matched filter’ and is used very frequently in the domain of hyperspectral imagery

(Manolakis et al., 2016). In this field, the covariance matrix is appropriately referred to as the background clutter matrix, the HRI as the signal-to-clutter or signal-to-noise ratio, and its expression is sometimes written as (Manolakis et al. (2016), p520)

$$\text{HRI} = \frac{\tilde{K}^T \tilde{y}}{\|\tilde{K}\|}, \quad (2)$$

with

$$\tilde{K} = S^{-\frac{1}{2}} K \quad (3a)$$

$$\tilde{y} = S^{-\frac{1}{2}} (y - \bar{y}) \quad (3b)$$

The operation $y \rightarrow \tilde{y}$, transforms the spectra into a set of spectra with a zero mean and an identity covariance matrix. As this transforms the generalized noise into white noise, the operation is referred to as whitening or pre-whitening (Scharf, 1991; Rodgers, 2000; Manolakis et al., 2016; Kessy et al., 2018). Whitening transforms the spectral channels to uncorrelated random variables with unit variance and zero mean. In this form, the HRI is simply the projection of the whitened spectrum $\tilde{y}$ onto the normalized whitened Jacobian $\tilde{K}$.

Instead of the HRI, one can also analyze the entire unprojected whitened spectra. Whitening removes most of the background signal. Therefore, on background spectra, each channel of a whitened spectrum has an expected mean value of zero and a standard deviation of 1 (this is a property inherited from the HRI). Conversely, a whitened spectral channel with an absolute value above 4 indicates a significant departure from Gaussian normality. Large negative values indicate an anomalous large absorption, while large positive values indicate either an emission signal or a decrease of a given trace gas with respect to background conditions (defined by the set of spectra used to generate $\tilde{y}$ and S). So rather than associating with each spectra a single number like the HRI, whitening a spectrum produces another spectrum in which all spectral aberrations are exposed. The strength of the whitening technique is that it allows highlighting anomalies without knowing the spectral signature (Jacobian) in advance. Once a candidate species is suspected, it can be unambiguously assigned by comparing the shape of the whitened Jacobian with the spectral signatures in the whitened spectrum. Unambiguous identification is another clear advantage over the HRI, which can be prone to false detections caused by a partial match of a spectral signature with the Jacobian of the target species. Note that by weighting with the inverse of the covariance matrix, the spectral components that normally exhibit the largest variability (e.g. the baseline, due to variation in the surface temperature and clouds, or the absorption features due to O_3 , H_2O or CO_2) will carry the least weight in the whitened spectrum.

When we talk about departure from normality, this is with respect to a set of spectra taken from a reference time period and/or geographic area of our choice. This freedom gives rise to at least two important applications: (1) when the covariance is representative for the entire set of IASI spectra, whitening can be used to study special events such as volcanic eruptions or large fires; (2) when the covariance matrix is representative for a specific time period and used on spectra from another time period, whitening, as we will show, can reveal small changes in the concentrations of long-lived compounds. A case study of each application is presented next.

As a side comment, the HRI formula can also be derived from standard linear discriminant analysis, and through the use of the covariance matrix, is also closely linked to principal component analysis (PCA). We refer to Clarisse et al. (2013) for a detailed discussion of this. Whitening can be seen as an alternative to PCA-based analysis of anomalies in spectra (Grainger et al., 2013).

Whitening can be seen as an alternative to analyzes based on principal components (PC) (Hurley et al., 2009; Grainger et al., 2013). In such approaches, the signatures of interest are contained in the orthogonal part to the PC. A key difference between the two

approaches, is that whitening disregards no components but instead weights them inversionally proportional to their weight in the covariance matrix (see also Clarisse et al. (2013)). As a consequence of this, it can more readily be used for the detection of species already present in the set of spectra that was used to create the covariance matrix, as illustrated in the following sections. Other statistical transformations of the spectra or residuals exist, such as the random projections technique used for quantifying OCS in the IASI spectra (see Camy-Peyret et al. (2017) and references therein).

3 Short-lived gas detection in Australian bushfires

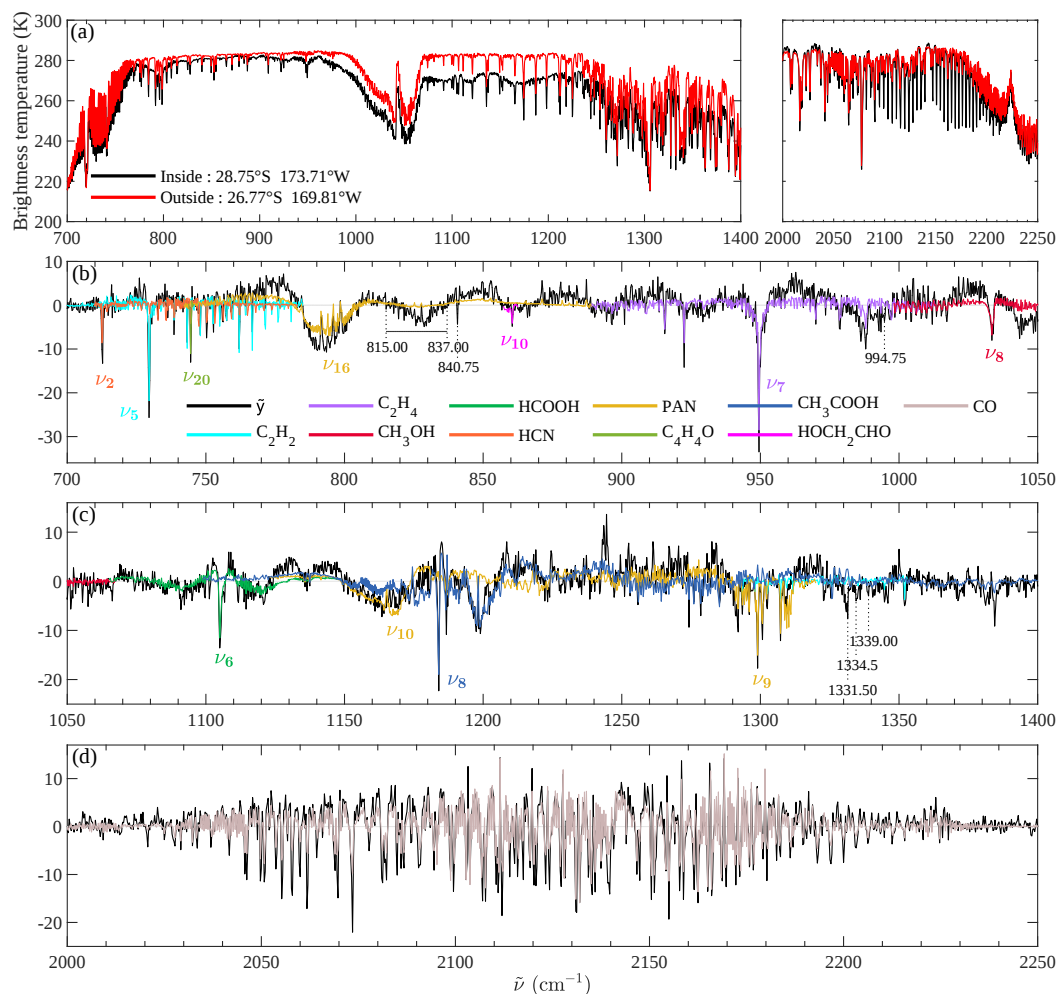


Figure 1. Spectrum from the Australian bushfires of 5 January 2020 revealing nine trace gas species and carbon monoxide. Panel a shows the spectrum as measured by IASI before the whitening transformation (in black), together with a spectrum measured just outside the plume (in red). The whitened spectrum is shown in panels b to d for selected spectral ranges.

This first case study concerns the massive bushfires which took place in Australia around the turn of year 2019–2020 (Nolan et al., 2020; Boer et al., 2020). IASI spectra of similarly large fires in Australia in 2009 were analyzed in Clarisse et al. (2011) with a physical retrieval approach, and revealed very strong spectral signatures of some well-known fire tracers. Here we will focus on a particular spectrum that was observed over the South

Pacific Ocean (28.75°S, 173.71°W) on January 5, 2020. This spectrum was selected on the basis of HRI-maps of C₂H₄ (see Figure A1 of Annex A). These indicate that the observation was made in the central part of a large fire plume originating from South-East Australia two or three days earlier. Calipso measurements (Winker et al., 2009) reveal that the plume was located between 4 and 6 km above sea level.

The chosen (untransformed) spectrum is shown in Figure 1a together with a spectrum measured outside the fire plume (their location is indicated in Figure A1 of Annex A). Immediately obvious is the strong absorption band of CO in the 2100–2200 cm⁻¹ region, and the large absorption in the atmospheric window between 1080 and 1230 cm⁻¹ due to smoke aerosols and other broadband absorbers. Figure 1b–d shows the whitened spectrum calculated with $\{\bar{y}, S\}$ built using a subset of around 10⁷ IASI spectra measured throughout 2013 and across the globe, from (Franco et al., 2018). From this spectrum, we identified nine species between 700 and 1400 cm⁻¹, namely acetylene (C₂H₂), ethylene (C₂H₄), methanol (CH₃OH), formic acid (HCOOH), hydrogen cyanide (HCN), peroxyacetyl nitrate (CH₃C(O)O₂NO₂, abbreviated as PAN), furan (C₄H₄O), acetic acid (CH₃COOH) and glycolaldehyde (HOCH₂CHO or hydroxyacetaldehyde). The identification was made with Jacobians obtained from a forward model (with temperature and main absorbers of the US standard atmosphere (NASA-USA, 1976), where each whitened Jacobian was scaled to match its maximum absolute value with the observed signature in the whitened spectrum. Note that for the figures, the scaling factor was slightly offset, so that the whitened spectra and the whitened Jacobian would not obscure each other.

The observation of HOCH₂CHO is the first from space to our knowledge. The detection of the Q-branch associated with the ν_{10} vibrational mode at 860.5 cm⁻¹ (Johnson et al., 2013) is promising for future exploitation, as HOCH₂CHO measurements are very limited at the moment (Treadaway et al., 2018). HOCH₂CHO, as well as all the other detected species, are well-known products of biomass burning (see e.g. Akagi et al. (2011); Andreae (2019) and references therein). In the troposphere it is formed from the oxidation of alkenes, in particular isoprene and several of its degradation products, and it is a key intermediate in the formation pathways of oxygenated compounds such as HCOOH and glyoxal (Paulot et al., 2009; Taraborrelli et al., 2012). Strong Q-branches of HCN, C₂H₂ and C₄H₄O are observed between 700 and 800 cm⁻¹, as well as several R lines for HCN and C₂H₂. Note that the strong ν_2 band of CO₂ present in this spectral region is completely removed by the whitening transformation. For C₂H₂, a weaker signal is also detected around 1300–1352 cm⁻¹. C₂H₄ is the species that exhibits the largest signal in the whitened spectrum with a value of -33 at 949.5 cm⁻¹ and several other features on both sides. The detection of the ν_8 band of CH₃OH is also straightforward with the removal of the large ozone band. At longer wavenumbers, HCOOH is observed in its ν_3 band around 1105 cm⁻¹. Finally, we observe the broadband absorbers PAN and CH₃COOH, which partially overlap between 1150 and 1170 cm⁻¹. PAN also has a remarkable broadband absorption between 780 and 800 cm⁻¹. The remaining part of the spectrum (1400–2760 cm⁻¹) was analyzed carefully, but apart from the large CO absorption features in the region at 2000–2250 cm⁻¹, no other spectral anomalies were identified. However, there are several notable enhancements in the 800–1350 cm⁻¹ range which we were unable to attribute to a specific species. The strongest features are identified by dotted lines in Figure 1, and in particular those that we could also observe in other spectra of the fire plume. Some are very sharp, such as the feature at 840.75 cm⁻¹; others are wider (815–837 cm⁻¹) and these could be due to heavier molecules or a combination of different absorbers. Note that in general, heavier molecules tend to have broader absorption features, for which unambiguous identification can become more difficult, especially when only one feature is identified or when its magnitude is weak.

We also analyzed the whitened spectrum of a fresh fire plume (measured at 39.28°S, 153.48°E on December 30, 2019), as displayed in Figure B1 of Annex B. The fresh plume exhibits very strong signatures of nitrous acid (HONO) and ammonia (NH₃) that are not seen in the transported plume. This makes sense as these species are known to be emitted

in very large quantities by fires (Whitburn et al., 2017; Theys et al., 2020), but are quickly degraded to form secondary products. A weak absorption feature compatible with the presence of propene (C_3H_6) is also detected. In contrast, HOCH_2CHO , HCOOH and PAN are only seen in the transported plume. Whereas these three species are well-known secondary products, HOCH_2CHO is also primarily emitted by fires; the fact that it is not observed in the fresh plume could be explained by the overlapping prominent HONO absorption feature.

4 Long-lived gas analysis over the Pacific Ocean

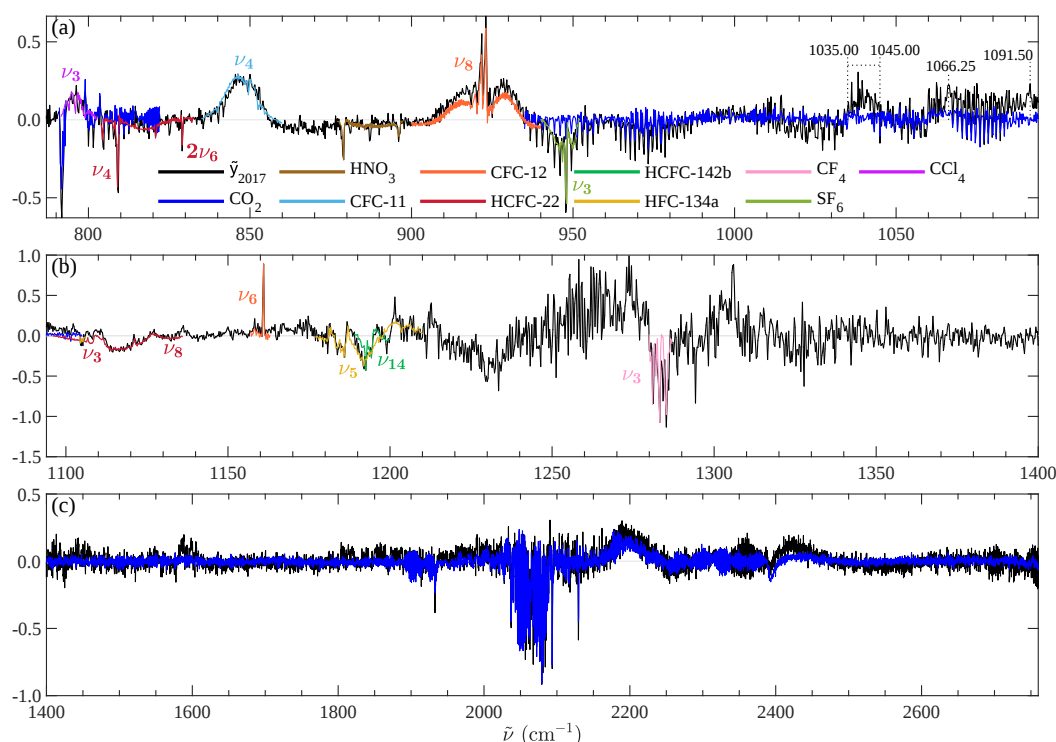


Figure 2. Whitenened spectrum of Pacific Ocean from October 2017 revealing eight halocarbons, carbon dioxide and nitric acid.

In this section, we show how small spectral changes that appear over longer time periods in mean spectra can be exposed with the help of the whitening transformation. Focusing over an area in the North Pacific Ocean (25° – 45°N ; 170° – 165°W), we chose 120,000 clear-sky spectra measured in 2008 to calculate a reference mean spectrum and associated covariance matrix. 2008 is the first complete year where IASI L1c reprocessed data are available (Bouillon et al., 2020). The spectrum on which we applied whitening was calculated from $N = 20,000$ clear-sky spectra measured in October 2017 over the same area. Using a mean rather than an individual spectrum is not strictly needed, but it helps focusing on long-lived compounds, as an individual spectrum might always exhibit anomalies compared to the reference due to natural variations in short-lived trace species. For instance plumes of volatile organic compounds are regularly observed in the North Pacific Ocean (Franco et al., 2018, 2019). The whitenened mean spectrum is shown in Figure 2. As we work with mean spectra, the maximum values observed in the whitenened spectrum are much lower than in the previous example. However, for the same reason, a significant value is in theory already obtained for values as low as $4/\sqrt{N} \approx 0.03$.

Changes in eight halocarbons were identified, these are: CFC-11 (CFCl_3), CFC-12 (CF_2Cl_2), HCFC-22 (CHF_2Cl), HCFC-142b ($\text{CH}_3\text{CF}_2\text{Cl}$), HFC-134a (CH_2FCF_3), carbon tetrafluoride (CF_4), sulfur hexafluoride (SF_6) and carbon tetrachloride (CCl_4). Halocarbons are synthetic halogenated compounds used in many industrial and domestic applications (Martinerie et al., 2009). Most of them are powerful greenhouse gases and contribute, for chlorinated and brominated compounds, to the depletion of stratospheric ozone and to the development of ozone hole (World Meteorological Organization, 2010). Of these eight species, five of them (CFC-11, CFC-12, HCFC-22, CCl_4 and CF_4) have been observed in IASI spectra before (Clerbaux et al., 2009; Liuzzi et al., 2016).

The absorption bands of the two CFCs, located around 849 cm^{-1} for CFC-11 and at ~ 923 and $\sim 1161\text{ cm}^{-1}$ for CFC-12 (here and also later, the wavenumbers refer to the position of strong Q-branches), have whitened signals of 0.3, 0.7 and 0.9 respectively. Those positive values indicate that their atmospheric abundance was smaller in 2017 than in 2008, which is consistent with independent satellite measurements (Chen et al., 2020), independent in-situ measurements (Prinn et al., 2018) and a direct consequence of the 1987 Montreal Protocol to limit the use and production of CFCs (World Meteorological Organization, 2018). The ν_3 band of CCl_4 shows up at $\sim 796\text{ cm}^{-1}$ with a whitened signal's value of 0.2. The production of CCl_4 has been controlled by Montreal Protocol as long as CFCs, explaining its decrease between 2008 and 2017.

The other halocarbons in Figure 2 are all detected with a negative value, indicating an increase in their atmospheric concentration between 2008 and 2017. Four absorption bands of HCFC-22 are observed: ν_4 ($\sim 809.25\text{ cm}^{-1}$), $2\nu_6$ ($\sim 829.00\text{ cm}^{-1}$), ν_3 ($\sim 1114.00\text{ cm}^{-1}$) and ν_8 ($\sim 1132.25\text{ cm}^{-1}$). In the spectral region between 1178 and 1200 cm^{-1} , HFC-134a and HCFC-142b are both absorbing. Although the detection of HFC-134a is unambiguous, the detection of HCFC-142b is less certain, but still very likely considering the presence of a sharp feature at 1192.50 cm^{-1} that would correspond to its ν_{14} Q-branch. Note that constraints on the production of both HCFCs and HFCs were only added in later amendments of the Montreal Protocol (World Meteorological Organization, 2018). The signatures of SF_6 and CF_4 are clearly observed at 947.75 cm^{-1} (Q-branch of the ν_3 band with value of -0.6) and around 1283.25 cm^{-1} (value of -1.1) respectively, and are consistent with increases observed from ground-based measurements (Prinn et al., 2018).

The region $1220\text{--}1370\text{ cm}^{-1}$ is quite complicated for a definite analysis of the individual trends of the atmospheric species contributing to the absorption: CH_4 , N_2O , H_2O , HDO , HNO_3 , O_3 . However, in this region the (negative) spectral signatures of CF_4 around 1283 cm^{-1} clearly indicate an increase of its abundance over the considered time frame i.e. 2008 (for the reference) compared to 2017 (for the Pacific data set).

Apart from the halocarbons, Figure 2 highlights several absorption lines of CO_2 ; the signal shows up at 791.5 cm^{-1} with a succession of rovibrational features around 800 , 950 , 975 , 1050 , 1070 and 2100 cm^{-1} . The negative values indicate an increase in the CO_2 signal, and thus a rise in concentration between 2008 and 2017. The CO_2 features in these spectral regions are less saturated than below 750 cm^{-1} , and are less affected by small changes in atmospheric temperatures. There are also absorption features of nitric acid (HNO_3) in its ν_5 and $2\nu_9$ bands: the corresponding signatures are stronger around the two Q branches centered at 879.00 and 896.00 cm^{-1} indicating an increasing atmospheric abundance of HNO_3 . This is consistent with a recent study that reported a small positive trend for HNO_3 (Bernath et al., 2020).

A number of smaller features remain unidentified in this spectrum, of which the strongest are spotted by dotted lines in Figure 2. Most of the observed features are found in the atmospheric window between 800 and 1300 cm^{-1} . The broad feature between 1215 and 1240 cm^{-1} could possibly due to the spectral contribution of COF_2 . As a caveat, some of these unidentified features could still be due to natural variations of short-lived trace gases. In addition, while the IASI L1c reanalyzed dataset was used for this study, small changes in

the IASI instrument that occurred in the last decade could also play a role. No major features were identified in the regions impacted by water vapor and CO₂ (645–790, 1300–2000, 2200–2400 cm⁻¹).

5 Conclusion

In this letter, we have shown that the whitening transformation allows analyzing anomalies in high resolution infrared spectra. It enables elimination of the large absorption features that usually dominate a spectrum and accentuates those channels which are incompatible with the background norm. We have presented two different applications: an analysis of an extreme event (the 2019/2020 Australian fires) and an analysis of long-term mean changes in IASI spectra. These exploit respectively the spatial variability of reactive species and the temporal variability of long-lived compounds. We have provided spectroscopic evidence of the occurrence of nine fire tracers and eight halocarbons. This includes four species (glycolaldehyde, HCFC-142b, HFC-134a, SF₆) which have never been observed before with a nadir sounder. Their detection is very promising, especially for the determination of their source emissions and the study of their temporal evolution.

The advantages of the whitening approach compared to more traditional techniques are numerous. For instance, Clarisse et al. (2011) analyzed spectra from the 2009 Australian fires using a fitting approach, which aims at reconstructing the observed spectrum by means of forward simulations. Especially in extreme cases, it is very difficult and time consuming to obtain reasonable fits of the corresponding spectra, and often fitting windows need to be restricted to narrow spectral ranges. In contrast, whitening is immediate, does not require an inverse model nor the ability of a forward model to account for all the observed features, and can be applied on spectral ranges of any size. Likewise, in a more traditional analysis of trends in high resolution infrared spectra (Brindley et al., 2015; Strow & DeSouza-Machado, 2020), it is very challenging to deal properly with variations of O₃, CH₄, CO₂ or H₂O due to e.g. natural temporal variations, variations in temperature or sampling biases related to clouds; whereas these are dealt with automatically using the whitening technique.

Appendix A

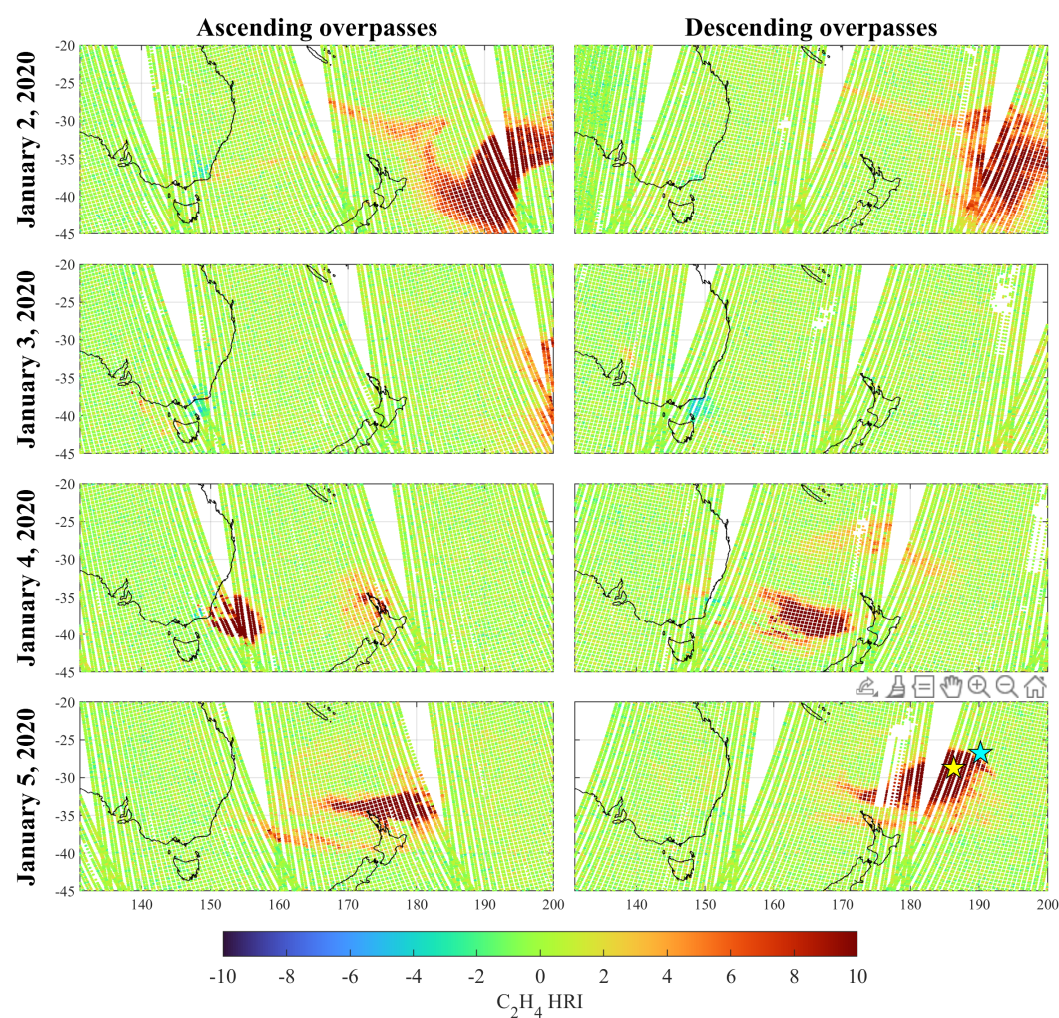


Figure A1. HRI maps of C_2H_4 of the 2020 Australian bushfires on selected days. The size of the IASI footprint has been slightly exaggerated for visualization purposes. ($\star$ and $\star$ respectively mark the spectra selected inside and outside the fire plume)

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Appendix B

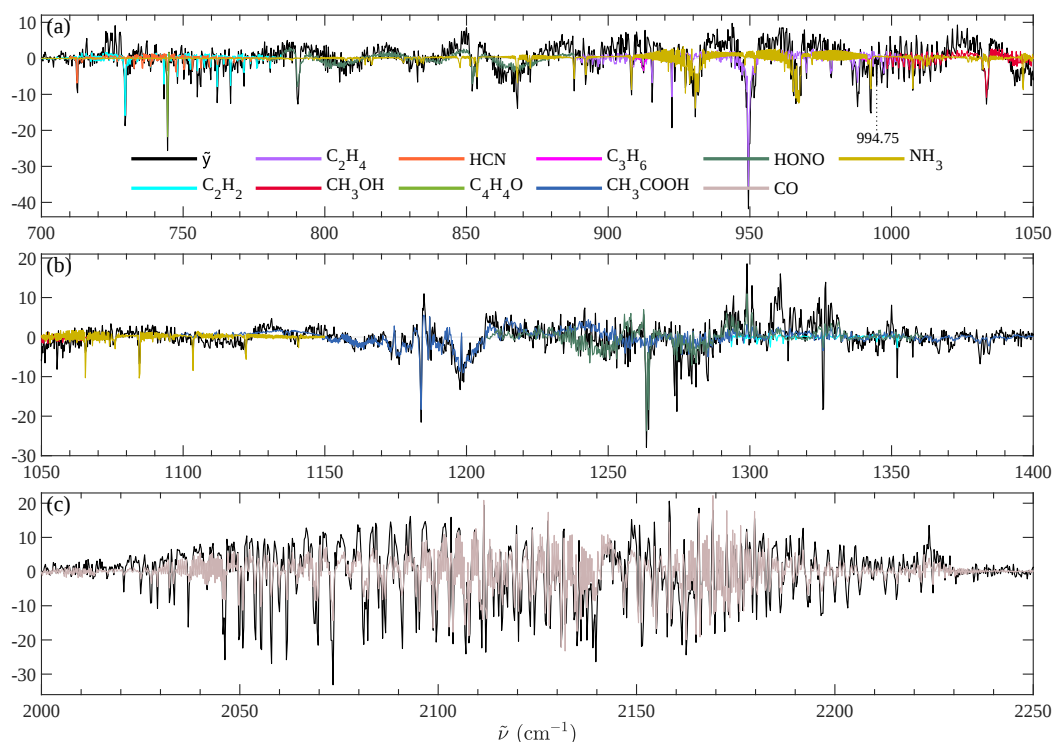


Figure B1. Whitened spectrum of Australian bushfires from December 30, 2019 revealing 8 trace gas species and carbon monoxide.

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Acknowledgments

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