Atmospheric carbon dioxide retrieved from the Greenhouse gases Observing SATellite; Method, comparisons and algorithm development

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by

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Declaration

I hereby declare that no part of this thesis has been previously submitted to this or any other University as part of the requirement for a higher degree. The work described herein was conducted by the undersigned except for contributions from colleagues as acknowledged in the text.

Austin J. Cogan
November 27, 2012
I would like to dedicate this thesis to my wife, Katya, and my parents, Christopher and Valerie, for all their support.
Carbon dioxide is the largest anthropogenic contributor to global warming and atmospheric concentrations have rapidly increased since the start of the industrial revolution. Networks of surface in-situ carbon dioxide sensors provide precise and accurate measurements of the global carbon dioxide concentration, including large scale temporal, seasonal and latitudinal variations. However, these observations are too sparse to allow the establishment of sub-continental carbon budgets, limiting the accuracy of climate change projections and the ability to mitigate future levels of atmospheric carbon dioxide.

Satellite observations can provide data with dense spatial and temporal coverage over regions poorly sampled by surface networks. Specifically, observations in the shortwave infrared region are well suited for constraining carbon fluxes as they can provide total column carbon dioxide with high sensitivity to the source and sink locations at the surface. The first dedicated greenhouse gases sensor, the Greenhouse gases Observing SATellite (GOSAT), was launched in January 2009 by the Japanese Aerospace eXploration Agency (JAXA) and has successfully started to acquire global observations of greenhouse gases, including carbon dioxide.

The University of Leicester Full Physics (UOL-FP) retrieval algorithm has been designed to estimate total column carbon dioxide from GOSAT shortwave infrared observations. The initial results were compared to coincident ground based measurements for a number of locations and compared on a global scale to a model. This showed an accuracy and precision that should provide improved surface flux estimates. Additionally, a bias correction scheme was developed that reduced observed geographical biases, allowing surface flux uncertainties to be potentially reduced further. To further develop the UOL-FP retrieval algorithm, a simulator capable of creating realistic GOSAT observations was built, allowing the investigation of different retrieval algorithm modifications, which may lead to reduced source and sink flux uncertainties and therefore aid future climate change forecasts.
PUBLICATIONS

Journal Articles


**Conference Proceedings**


Other Documents


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<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>4D-VAR</td>
<td>Four-dimensional variational assimilation</td>
</tr>
<tr>
<td>ACOS</td>
<td>Atmospheric CO$_2$ Observations from Space</td>
</tr>
<tr>
<td>AERONET</td>
<td>AErosol RObotic NETwork</td>
</tr>
<tr>
<td>AIRS</td>
<td>Aqua-advanced Infrared Radiation Sounder</td>
</tr>
<tr>
<td>AMSU</td>
<td>Advanced Microwave Sounding Unit</td>
</tr>
<tr>
<td>AOD</td>
<td>Aerosol Optical Depth</td>
</tr>
<tr>
<td>ARM</td>
<td>Atmospheric Radiation Measurement</td>
</tr>
<tr>
<td>ASCENDS</td>
<td>Active Sensing of CO$_2$ Emissions over Nights, Days and Seasons</td>
</tr>
<tr>
<td>ASTER</td>
<td>Advanced Spaceborne Thermal Emission and Reflection radiometer</td>
</tr>
<tr>
<td>AVIRIS</td>
<td>Airborne Visible InfraRed Imaging Spectrometer</td>
</tr>
<tr>
<td>BTD</td>
<td>Brightness Temperature Difference</td>
</tr>
<tr>
<td>CAI</td>
<td>Cloud and Aerosol Imager</td>
</tr>
<tr>
<td>CALIOP</td>
<td>Cloud-Aerosol Lidar with Orthogonal Polarization</td>
</tr>
<tr>
<td>CALIPSO</td>
<td>Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations</td>
</tr>
<tr>
<td>CAM</td>
<td>Monitor camera system</td>
</tr>
<tr>
<td>CARBONSAT</td>
<td>CARBON monitoring SATellite</td>
</tr>
<tr>
<td>CASA</td>
<td>Carnegie-Ames-Stanford Approach</td>
</tr>
<tr>
<td>CDIAC</td>
<td>Carbon Dioxide Information Analysis Center</td>
</tr>
<tr>
<td>CH$_4$</td>
<td>Methane</td>
</tr>
<tr>
<td>CLAUDIA</td>
<td>CLoud and Aerosol Unbiased Decision Intellectual Algorithm</td>
</tr>
<tr>
<td>CMS</td>
<td>Carbon Monoxide and Methane Spectrometer</td>
</tr>
<tr>
<td>CNES</td>
<td>Centre National d'Etudes Spatiales</td>
</tr>
<tr>
<td>CO</td>
<td>Carbon Monoxide</td>
</tr>
<tr>
<td>CO$_2$</td>
<td>Carbon Dioxide</td>
</tr>
<tr>
<td>DEM</td>
<td>Digital Elevation Model</td>
</tr>
<tr>
<td>DOAS</td>
<td>Differential Optical Absorption Spectroscopy</td>
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<tr>
<td>ECMWF</td>
<td>European Centre for Medium Range Weather Forecasts</td>
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<tr>
<td>EE</td>
<td>Earth Explorer opportunity missions</td>
</tr>
<tr>
<td>ENVISAT</td>
<td>ENVironment SATellite</td>
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<tr>
<td>ESA</td>
<td>European Space Agency</td>
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<tr>
<td>ESRL</td>
<td>Earth System Research Laboratory</td>
</tr>
<tr>
<td>ESSP</td>
<td>Earth System Science Partnership</td>
</tr>
<tr>
<td>FTS</td>
<td>Fourier Transform Spectrometer</td>
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</table>
FSI Full Spectral Initiation
FAR First Assessment Report
FFT Fast Fourier Transform
GEMS Global and regional Earth-system (Atmosphere) Monitoring using Satellite and in-situ data
GEOS Goddard Earth Observing System
GFED Global Fire Emissions Database
GOSAT Greenhouse gases Observing SATellite
GWP Global Warming Potential
H$_2$O Dihydrogen monoxide (Water)
HIRS High resolution Infrared Radiation Sounder
HITRAN HiGH-resolution TRANsmission molecular absorption database
IASI Infrared Atmospheric Sounding Interferometer
IFOV Instantaneous Field of View
IIR Imaging Infrared Radiometer
ILS Instrument Line Shape
IPCC Intergovernmental Panel on Climate Change
IWC Ice Water Content
JAXA Japan Aerospace eXploration Agency
L1B Level 1B
LIDORT Linearized Discrete Ordinate Radiative Transfer
LWC Liquid Water Content
MCT Mercury Cadmium Telluride (HgCdTe)
MEC Mass Extinction Coefficient
MET Surface Meteorological Instrumentation
MMR Mass Mixing Ratio
MOE Ministry Of Environment
MODIS MODerate RESolution Imaging Spectroradiometer
MSU Microwave Sounding Unit
N$_2$ Dinitrogen
NASA National Aeronautics and Space Administration
NDVI Normalized Difference Vegetation Index
NIES National Institute for Environmental Studies
NIR Near Infra-Red

XXXV
<table>
<thead>
<tr>
<th>Acronym</th>
<th>Full Form</th>
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<tr>
<td>NOAA</td>
<td>National Oceanic and Atmospheric Administration</td>
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<tr>
<td>O₂</td>
<td>Dioxgen</td>
</tr>
<tr>
<td>OCO</td>
<td>Orbiting Carbon Observatory</td>
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<tr>
<td>OPD</td>
<td>Optical Path Difference</td>
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<tr>
<td>P</td>
<td>Parallel Polarisation</td>
</tr>
<tr>
<td>PC</td>
<td>Photo-Conductive</td>
</tr>
<tr>
<td>PDF</td>
<td>Probability Density Function</td>
</tr>
<tr>
<td>PPDF</td>
<td>Path length Probability Density Function</td>
</tr>
<tr>
<td>RF</td>
<td>Radiative Forcing</td>
</tr>
<tr>
<td>RH</td>
<td>Relative Humidity</td>
</tr>
<tr>
<td>RI</td>
<td>Refractive Index</td>
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<tr>
<td>RT</td>
<td>Radiative Transfer</td>
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<tr>
<td>S</td>
<td>Perpendicular Polarisation</td>
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<tr>
<td>SAR</td>
<td>Second Assessment Report</td>
</tr>
<tr>
<td>SCIAMACHY</td>
<td>SCanning Imaging Absorption SpectroMeter for Atmospheric Cartography</td>
</tr>
<tr>
<td>SH</td>
<td>Specific Humidity</td>
</tr>
<tr>
<td>SGP</td>
<td>Southern Great Plains</td>
</tr>
<tr>
<td>SNR</td>
<td>Signal to Noise Ratio</td>
</tr>
<tr>
<td>SPRINTARS</td>
<td>Spectral Radiation-Transport Model for Aerosol Species</td>
</tr>
<tr>
<td>SSU</td>
<td>Stratospheric Sounding Unit</td>
</tr>
<tr>
<td>SWIR</td>
<td>Short Wave Infra-Red</td>
</tr>
<tr>
<td>SZA</td>
<td>Solar Zenith Angle</td>
</tr>
<tr>
<td>TANSO</td>
<td>Thermal And Near Infrared Sensor for Carbon Observations</td>
</tr>
<tr>
<td>TAR</td>
<td>Third Assessment Report</td>
</tr>
<tr>
<td>TCCON</td>
<td>Total Carbon Column Observing Network</td>
</tr>
<tr>
<td>TCM</td>
<td>Tropical Carbon Mission</td>
</tr>
<tr>
<td>THWAPS</td>
<td>Temperature, Humidity, Wind and Pressure Sensors</td>
</tr>
<tr>
<td>TIR</td>
<td>Thermal Infra-Red</td>
</tr>
<tr>
<td>TIROS</td>
<td>Television Infrared Observation Satellite</td>
</tr>
<tr>
<td>TOA</td>
<td>Top of Atmosphere</td>
</tr>
<tr>
<td>TOVS</td>
<td>TIROS Operational Vertical Sounder</td>
</tr>
<tr>
<td>UNFCCC</td>
<td>United Nations Framework Convention on Climate Change</td>
</tr>
<tr>
<td>UoL-FP</td>
<td>University of Leicester Full Physics</td>
</tr>
</tbody>
</table>
USA  United States of America
USGS  United States Geological Survey
UV  Ultra Violet
VFM  Vertical Feature Mask
VIS  Visible
VMR  Volume Mixing Ratio
WFC  Wide Field Camera
$X_{CO_2}$  Column averaged dry air mole fraction of $CO_2$
Chapter 1

Introduction

1.1 Climate Change

The Intergovernmental Panel on Climate Change (IPCC) defines climate change as any change in the mean state of a climate system over an extended period of time, such as decades or longer, whether it be caused by natural variability or due to human activity (IPCC 2001). The Earth’s mean climate is a consequence of incoming radiation from the Sun and outgoing radiation from the Earth, known as the radiation budget. Variations in solar radiation can also cause the climate to differ, which for instance can be caused by fluctuations in the radiation emitted from the Sun or due to variations in the Earth’s orbit. Also, various properties within the Earth’s atmosphere and surface can change the amount of incoming and outgoing radiation, such as reflection, absorption, emission and scattering of radiation.

Human activities have resulted in numerous changes of the atmosphere and surface that can alter the radiation budget of the Earth, see Section 1.2.1. For instance, anthropogenic emissions of greenhouse gases, such as carbon dioxide (CO$_2$), have increased rapidly over the last couple of centuries, leading to an enhancement of atmospheric absorption of outgoing radiation causing the Earth to become warmer (IPCC 2001). It is important to be able to project future climate changes so that humans can mitigate and adapt to the changing environments. The current knowledge of the Earth’s climate system is largely due to historical measurements, such as ice cores, and recent direct instrument observations (Houghton 2004). To this extent, recent climate variations have been observed and an overview of this is shown in Figure 1.1.

The recent anthropogenic emissions of greenhouse gases are considered geologically significant due to a rapid increase in temperature over a very short time frame (∼100
years), see Section 1.2. The present global warming trend is predicted to cause the temperature of the Earth to rise by 2-3 degrees to be similar to that of the mid-Pliocene where temperatures were higher (particularly in higher latitudes), polar ice cover was reduced and precipitation patterns were different (Robinson et al. 2008). Although the potential temperature increase over North America, Europe and Asia would increase agricultural productivity, this would unlikely compensate for substantial productivity losses in mid and low latitudes due to limitations of the terrain and soil (Haywood et al. 2008). As a consequence of higher temperatures, higher evapotranspiration would occur further reducing the productivity of agriculture (Haywood et al. 2008). In addition to changes in agriculture and precipitation, there may be many other consequences of global warming that would require adaptation, such as human migration (McLeman & Smit 2006). Thus, it is important to mitigate the temperature increase and one way to achieve this would be to reduce anthropogenic greenhouse gas emissions, which are the main drivers of global warming.

To attempt to achieve reduced emissions the Kyoto Protocol was adopted in 1997, which commits numerous countries to not exceed their assigned anthropogenic emission quantities and to reduce emissions by 5% below 1990 levels between 2008-2012 (NATIONS 1998). Since 1997 there have been many advances in the understanding of the climate, with the insight that it is more appropriate to limit CO₂ emissions based on a threshold of global temperature increase (Weaver 2011). The Copenhagen Accord (2009), proposed by the United Nations Framework Convention on Climate Change (UNFCCC), defined that an increase of 2 K above the preindustrial level would be the limit (NATIONS 2009). The second commitment of the Kyoto Protocol is proposed to include this and determine emission limits upon this (Weaver 2011). However, for countries to know how to reduce emissions it is important to understand what the sources and sinks are of each driver (e.g. gas, aerosol, land change) of climate change.
FIGURE 1.1. An overview of recorded climate variations between 1970 and 2004, showing temperature changes and locations of significant changes of physical systems (snow, ice and frozen ground; hydrology; coastal processes) and biological systems (terrestrial, marine, and freshwater). For each continental region the number of significant changes and those correlating with temperature are given in the 2 x 2 boxes. These regions include; North America (NAM), Latin America (LA), Europe (EUR), Africa (AFR), Asia (AS), Australia and New Zealand (ANZ), and Polar Regions (PR). This is also shown for a global-scale, Global (GLO), with it separated into Terrestrial (TER), Marine and Freshwater (MFW). Figure taken from IPCC (2001).
1.2 Global Warming

Global warming refers to the increases in mean global surface temperature, which provides one way to monitor climate change. Global surface temperatures have increased rapidly since a pre-industrial age, as shown in Figure 1.2, indicative of a changing climate.

![Figure 1.2](image)

**Figure 1.2.** Northern Hemisphere temperatures over the last two millenia. Shows temperature reconstructed from tree rings, corals, ice cores and historical records (blue) and recent instrumental data (red), with the data smoothed (black) and two standard error limits given (grey). Figure taken from IPCC (2001).

Whilst many different aspects could affect the surface temperature, measurements taken from ice cores have shown a strong correlation between temperature increases and atmospheric concentrations of CO$_2$ and CH$_4$ (IPCC 2001), see Figure 1.3 for example. This is likely due to the greenhouse effect, whereby atmospheric greenhouse gases, such as CO$_2$ and CH$_4$, cause the temperature of the atmosphere to increase (a more detailed explanation of this is given in Section 1.2.2).

Human activities such as fossil fuel combustion and land use change have led to dramatic increases in atmospheric greenhouse gas concentrations. Model simulations using
knowledge of the climate system predict that this will lead to increased global warming and future climate change. Model projections from the IPCC First Assessment Report (FAR) (IPCC 1990) and the Second Assessment Report (SAR) (IPCC 1996) estimated global mean temperature increases of $\sim 0.3$ K and $\sim 0.15$ K per decade respectively, with SAR including cooling effects from aerosols. The Third Assessment Report (TAR) (IPCC 2001) suggested a similar projection as SAR. These projections have been compared to recent observations that show an increase of $\sim 0.2$ K per decade, see Figure 1.4, showing the projections to be generally reasonable (IPCC 2007).
Figure 1.4. Observed global mean surface warming compared to model projected trends and ranges from the IPCC First (FAR), Second (SAR) and Third (TAR) Assessment Reports. B1 (blue), A1B (green), and A2 (red) represent the mean projected trends and ranges from multiple model scenarios and the commitment (orange) shows the mean predicted trend and range if greenhouse gas and aerosol concentrations were held constant from the year 2000. Figure taken from IPCC (2007).
1.2.1 Radiative Budget

The Earth absorbs and reflects radiation from the Sun and emits infra-red radiation towards space. For the Earth to be in dynamic equilibrium the incoming and outgoing radiation must be of equal energy, which would result in a constant planetary temperature. The incoming solar flux energy intercepted by the day-side of the Earth’s surface, $F_i$, per second can be written as

$$F_i = F_s (1 - A) \pi R_e^2$$  \hspace{1cm} (1.1)

where $F_s$ is the solar flux constant ($1368 \text{ Js}^{-1}\text{m}^{-2}$), $R_e$ is the radius of the Earth ($6.38 \times 10^6 \text{ m}$), and $A$ is the Earth’s albedo, representing the fraction of solar flux that is reflected from the Earth’s surface. The outgoing infra-red radiation $F_o$ per second from the entire Earth’s surface is

$$F_o = 4\pi R_e^2 \epsilon S T_e^4$$  \hspace{1cm} (1.2)

where $\epsilon$ is the surface emissivity, $S$ is Stefan’s constant ($5.67 \times 10^{-8} \text{ Js}^{-1}\text{m}^{-2}\text{K}^{-4}$) and $T_e$ is the temperature of the Earth. Balancing the incoming and outgoing radiation, using the Stefan-Boltzmann law, gives the temperature of the Earth as

$$T_e = \left[ \frac{F_s (1 - A)}{4\epsilon S} \right]^{1/4}$$  \hspace{1cm} (1.3)

If the Earth is assumed to be a black body, whose emissivity equals 1, with an albedo of 30%, then the temperature of the Earth would be about 256 K. However, observations have shown that the temperature of the Earth is closer to a value of 290 K. This elevation in temperature is caused by the greenhouse effect.

1.2.2 Greenhouse Effect

Excluding anthropogenic contributions, the greenhouse effect raises the temperature of the Earth to a level that is vital for many habitats. Essentially, the emitted infra-red radiation from the Earth’s surface becomes absorbed by atmospheric molecules and is re-emitted in all directions, such that some will be re-absorbed by the Earth’s surface causing the surface temperature to increase. These atmospheric molecules correspond to
gases with absorption bands in the infra-red wavelength range and are known as greenhouse gases, as described in Table 1.1. Water vapour is by far the most abundant greenhouse gas but has a very short lifetime of $\sim8$ days in the atmosphere (Trenberth 1998). However, all other greenhouse gases exist for many years in the atmosphere so can contribute to the greenhouse effect for prolonged periods (IPCC 2001).

Table 1.1. The main greenhouse gases and their quantities during a pre-industrial time (1700-1800) compared to 1998 (IPCC 2007).

<table>
<thead>
<tr>
<th>Gas</th>
<th>Chemical Formula</th>
<th>Pre-Industrial Level</th>
<th>2005 Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water Vapour</td>
<td>$\text{H}_2\text{O}$</td>
<td>$\sim0.03$</td>
<td>$\sim0.03$</td>
</tr>
<tr>
<td>Carbon Dioxide</td>
<td>$\text{CO}_2$</td>
<td>280 ppm</td>
<td>379 ppm</td>
</tr>
<tr>
<td>Methane</td>
<td>$\text{CH}_4$</td>
<td>715 ppb</td>
<td>1774 ppb</td>
</tr>
<tr>
<td>Nitrous Oxide</td>
<td>$\text{N}_2\text{O}$</td>
<td>270 ppb</td>
<td>314 ppb</td>
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<tr>
<td>Ozone</td>
<td>$\text{O}_3$</td>
<td>24 ppb</td>
<td>34 ppb</td>
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<td>Chlorofluorocarbons,</td>
<td></td>
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</tr>
<tr>
<td>CFC-11</td>
<td></td>
<td>0 ppt</td>
<td>251 ppt</td>
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<tr>
<td>CFC-12</td>
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<td>CFC-113</td>
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<td>0 ppt</td>
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<td>Hydrochlorofluorocarbons</td>
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<td>HCFC-22</td>
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<td>0 ppt</td>
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<td>HCFC-141b</td>
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<td>HCFC-142b</td>
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<td>Methyl Chloroform</td>
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<td>Sulphur Hexafluoride</td>
<td>$\text{SF}_6$</td>
<td>0 ppt</td>
<td>5.6 ppt</td>
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</tbody>
</table>

Since the start of the industrial revolution the atmospheric concentrations for some of these gases have rapidly increased leading to an enhanced greenhouse effect, hence higher planetary temperature. To demonstrate this, consider a single gas layer consisting of only greenhouse gases that is above a surface, as shown in Figure 1.5. Greenhouse gases are transparent to the incoming radiation that consists of ultra violet (UV) and visible (VIS) wavelengths, but absorb the outgoing infrared radiation. Consider the surface to be a black body that absorbs the incoming solar flux and emits infrared radiation.
The energy received by the surface is the sum of the solar energy and the energy directed back to the surface from the gas layer, it can be expressed as

\[ ST_g^4 = ST_s^4 + fST_a^4 \] (1.4)

where \( f \) is the fraction of outgoing infra-red radiation absorbed by the gas layer and \( T_g, T_s \) and \( T_a \) are the temperatures of the surface, solar flux and atmosphere respectively. The radiative balance of the atmosphere can be simplified to be

\[ fST_g^4 = 2fST_a^4 \] (1.5)

where the energy emitted from the surface is expressed by the left term of Equation 1.5 and the right term represents the sum of the energy that is absorbed by the gas layer and then re-emitted partly to space and partly to back towards the surface. By substitution the surface temperature is given as

\[ T_g = \left[ \frac{T_s^4}{(1 - f^2)} \right]^{1/4} \] (1.6)

If we assume \( f \) to be 0.75 then the surface temperature would be 30 K larger than without any gas layer. Additionally, if there was a higher concentration of greenhouse gases, then the fraction of outgoing radiation absorbed would be higher. For example, if
we assume $f$ to rise from 0.75 to 0.8 then the corresponding surface temperature would increase by 2 K.
1.2.3 Radiative Forcing

One way to assess and compare anthropogenic and natural drivers of climate change is through radiative forcing. The radiative forcing value is defined as the change in the average net irradiance at the tropopause (IPCC 2001) that occurs solely due to a driver of climate change. Radiative forcing, RF, can be related linearly to the global mean surface temperature change, \( \Delta T_g \), by

\[
\Delta T_g = \lambda RF
\]  
(1.7)

where \( \lambda \) is the climate sensitivity parameter (IPCC 2007). Thus, a positive radiative forcing value would indicate the surface warms on average and a negative value would indicate a cooling effect on the surface. The radiative forcing for human influenced drivers are shown in Figure 1.6, with greenhouse gases all being positive because they absorb part of the outgoing radiation from Earth to space. Notably, \( \text{CO}_2 \) gives the largest radiative forcing and its production is largely due to human activities.

However, radiative forcing does not take into account the temporal, spatial and vertical variations nor the lifetime of each driver of climate change. Therefore, to be able to evaluate the potential climate change caused by anthropogenic emissions an alternative method is required. Global Warming Potentials (GWP\(s \)) are one technique for assessing these.
Figure 1.6. Overview of the main components of radiative forcing with values representing the forcings in 2005 with respect to a pre-industrial age of about 1750. The black error bars give the range of uncertainty of the radiative forcing value. Figure taken from IPCC (2007).
1.2.4 Global Warming Potential

Strategies to limit anthropogenic climate change require a structure and comparative quantification of the emissions of climate drivers. Global Warming Potentials are the currently accepted method used to implement policies (NATIONS 1998) to allow multiple anthropogenic emitters (e.g. industrial companies or countries) to impose reductions of those emissions by the substitution of climate drivers, according to the specified emission constraints (IPCC 2007). Although alternative methods have been designed (Kandlikar 1996, Hammitt & Adams 1996, Tol 2002, den Elzen & Lucas 2005) these include some problematic issues so the simpler GWP method was developed and adopted for use in the Kyoto Protocol (IPCC 2007). This compares the time-integrated global mean radiative forcing of a 1 kg emission of the driver relative to that of CO$_2$, which can be defined by

\[
\text{GWP}_i = \frac{\int_0^{TH} RF_i(t) \, dt}{\int_0^{TH} RF_r(t) \, dt} - \frac{\int_0^{TH} a_i [C_i(t)] \, dt}{\int_0^{TH} a_r [C_r(t)] \, dt}
\]  

(1.8)

where $RF$ is the radiative forcing, $a$ is the radiative efficiency that is defined as the RF per unit mass increase in atmospheric abundance, $C(t)$ is the time-dependent abundance, $TH$ is the time horizon (i.e. 100 years in Kyoto Protocol), $i$ represents the climate driver considered, and $r$ is the reference (i.e. CO$_2$). This method clearly gives a GWP of 1 for CO$_2$, whilst for other gases such as CH$_4$ and N$_2$O have GWPs of 21 and 310 respectively. The GWP of Halocarbons varies from 90 to 11700 and the GWP of Sulphur hexafluoride is even higher with a value of 23900 (IPCC 2007). Although other substances have higher GWPs than CO$_2$ their abundances and emissions are far lower.

1.3 Role of CO$_2$

Other than water vapour, CO$_2$ is by far the most abundant greenhouse gas in the atmosphere contributing largely to global warming (IPCC 2007). In the absence of anthropogenic CO$_2$ emissions, the carbon cycle had periods of millennia where large carbon exchanges were in near balance, implying nearly constant reservoir contents (IPCC 2007). The addition of fossil fuels as anthropogenic CO$_2$ into the atmosphere has caused some carbon exchanges to become unbalanced. To this extent, atmospheric CO$_2$ concentrations have risen from a pre-industrial level of 280 ppm to a current level of about 385 ppm (IPCC 2007), as shown in Figure 1.7. Temperature has also increased over this period, as shown in Figure 1.2, correlating with increased anthropogenic emissions of greenhouse gases (IPCC 2007).
To improve global warming projection models, accurate knowledge of greenhouse gases is required, especially that of CO$_2$ which is the largest anthropogenic contributor. An overview of the current understanding of the carbon dioxide cycle is given in Figure 1.8.

The combustion of fossil fuels has increased rapidly since the start of the industrial revolution, emitting large amounts of anthropogenic CO$_2$. Fossil fuels are the remains of dead organisms that have undergone anaerobic decomposition. Coal is typically formed on land where waterlogged soil inhibits aerobic decomposition or where organisms are buried deep in soil quickly. Oil and natural gas are normally formed at the bottom of the ocean where the water transport is very slow. Over millions of years the pressure and heat created by the weight of the sediment above causes the remains to become compacted and takes the form of the fossil fuels. When fossil fuels are combusted they produce CO$_2$.
Figure 1.8. Diagram showing an overview of the main carbon dioxide cycle features.

and the simplest example of this is for pure coal where the chemical reaction is

\[ C + O_2 \rightarrow CO_2 + hv \] (1.9)

and \( hv \) represents the energy released. Another example is that of natural gas which has the reaction

\[ CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O + hv \] (1.10)

Cement production is also a large source of anthropogenic CO\(_2\), accounting for about 5% of global emissions (Pade & Guimaraes 2007). The production process requires fossil fuel combustion and the conversion of limestone to Calcium Oxide (CaO) and CO\(_2\) (Pade & Guimaraes 2007), called calcination. However, if cement products are exposed to the atmosphere then CO\(_2\) will diffuse back into the cement, forming Calcium Carbonate (CaCO\(_3\)) (Pade & Guimaraes 2007). This process is known as carbonation and over 100 years carbonation CO\(_2\) uptake can account for most of the CO\(_2\) released from calcination, acting as a sink of atmospheric CO\(_2\) (Pade & Guimaraes 2007).

Animals produce CO\(_2\) through respiration when the oxygen breathed in reacts with carbohydrates in the presence of enzymes in cells to produce CO\(_2\), H\(_2\)O and energy (hv), as shown by

\[ C_6H_{12}O_6 + 6O_2 \rightarrow 6CO_2 + 6H_2O + hv \] (1.11)

Photosynthesis of vegetation is essentially the reverse of this chemical reaction and cap-
tures CO$_2$. However, photosynthesis only occurs for a lot of vegetation in the spring/summer when they are growing, whilst in autumn/winter time they form litter on the ground which undergoes biological decomposition that returns most of CO$_2$ to the atmosphere. Also, some vegetation is eaten by animals, transferring the carbon to the animals which eventually will decay and contribute CO$_2$ to the soil and atmosphere.

The top layer of soil is composed of litter or detritus, in which carbon only has a lifetime of about 1 year due to microbial decay and erosion. The soil underneath comprises of ~5% living microbial biomass and ~95% humic material, which is a metabolic by-product of microbial activity. This has been found to be hundreds of thousands of years old, whilst the soil near the surface typically has a cycle of 30 years (Wigley & Schimel 2000). Soil carbon is released with increasing temperature and so acts as an additional global warming feedback (Fang & Moncrieff 2001). Thus, storage of carbon in soils generally increases from low to high absolute latitudes due to slower decomposition of dead plant material in colder environments. Additionally, the carbon stored in frozen soils and permafrost can be released as CO$_2$ by microbial aerobic respiration under warming temperature conditions (Lee et al. 2010). Flooded soils, where oxygen becomes depleted, have extremely low rates of decomposition and may accumulate large amounts of organic matter as peat (Wigley & Schimel 2000). Also, some soil is found as charcoals, especially where fires are frequent, although it is currently unknown how much carbon exists there (Ohlson et al. 2009).

The change in land use can cause CO$_2$ to be released into the atmosphere. Deforestation often includes the burning of vegetation and soil which produces CO$_2$, and removes the capture of CO$_2$ by photosynthesis. Carbon is released during logging from the decay of trees damaged in the harvest, decay of logging debris, and oxidation of wood products (Hoscilo et al. 2011). If the forest is allowed to regrow then the long term net carbon loss would be zero, providing no biomass burning takes place. However, sometimes when logging occurs artificial rivers are created to transport the logged trees, thus lowering the existing water table causing the top soil/vegetation to become dry. This in turn increases the probability of biomass burning events occurring, that produce even more CO$_2$ (Hoscilo et al. 2011). Agriculture both requires the land to be changed initially via deforestation, releasing CO$_2$, and then the soil repetitively undergoes tillage which causes additional CO$_2$ to be released from the soil (e.g. Paustian et al. 2000, Jr. et al. 2005).

Biomass burning can occur naturally and artificially, across large areas of the world. This has been observed by the National Aeronautics and Space Administration (NASA) Moderate Resolution Imaging Spectroradiometer (MODIS) which showed that biomass burning happened over $\gtrsim 2\%$ of land globally between July 2001 to June 2002 (Roy
The amount of global fires remained reasonably constant between 2002 and 2007, and declined during 2008 and 2009 by about 30\% (van der Werf et al. 2010). The main compound emitted from biomass fires is CO$_2$. For efficient fires CO$_2$ can account for $\sim$99\% of emissions, whilst smoldering fires could emit slightly less (\sim 90\%) CO$_2$ (Wooster et al. 2011). Carbon monoxide (CO) is the next dominant compound released from fire emissions which can further react to produce more CO$_2$ by

$$CO + OH \rightarrow CO_2 + H$$ (1.12)

The oceans contain $\sim$50 times that of atmospheric CO$_2$ and $\sim$95\% of it is in the form of inorganic dissolved carbon; bicarbonate (HCO$_3^-$) and carbonate ions (CO$_3^{2-}$) (Wigley & Schimel 2000). The rest is composed of organic carbon, consisting of living organic matter, particulate and dissolved organic carbon (Wigley & Schimel 2000). Terrestrial CO$_2$ can be transported to the oceans via the water flow in rivers. Additionally, CO$_2$ is transferred between the atmosphere and ocean via the air-sea interface, where it changes between atmospheric and dissolved CO$_2$ as

$$CO_2^{\text{Atmospheric}} \rightleftharpoons CO_2^{\text{Dissolved}}$$ (1.13)

The dissolved CO$_2$ converts to carbonic acid and vice versa by the reaction

$$CO_2^{\text{Dissolved}} + H_2O \rightleftharpoons H_2CO_3$$ (1.14)

The carbonic acid can convert to bicarbonate and carbonate ions by the ionizations

$$H_2CO_3 \rightleftharpoons H^+ + HCO_3^-$$ (1.15)

and

$$HCO_3^- \rightleftharpoons H^+ + CO_3^{2-}$$ (1.16)

CO$_2$ is transported into the intermediate and deep ocean by water mass transport where it can be stored for long durations. However, due to ocean circulations there is also a source created from upwelling water (Wigley & Schimel 2000). Phytoplankton exists near the ocean surface where it absorbs CO$_2$ via photosynthesis (Follows et al. 2007). Zooplankton feed on phytoplankton and in turn get eaten by larger marine organisms (Stige et al. 2011). These organisms will normally die and sink to the bottom of the ocean, but with a growing human population increasing numbers of these are fished and eaten by terrestrial animals, so the carbon can contribute to both the soil and atmosphere.
subsequently. Bicarbonate can be converted to marine shell material that eventually sinks to the ocean floor, fossilizing to become carbonaceous rock.

In some regions, such as arid and semi-arid climates, the underlying geology often consists of carbonate rock. All surfaces undergo erosion, whether it be from precipitation, wind blown particle impacts, contraction/expansion due to temperature, ice formation, geological factors, biological factors, or land use change. When carbonate rock is broken down it releases inorganic carbon (CaCO$_3$) into the atmosphere, this process is known as pedogenic carbonate which becomes precipitated back to the surface (Wigley & Schimel 2000). However, the pedogenic carbonate can be transported in the atmosphere with the prevailing winds to an area of non-carbonate terrain, providing a net sink of carbon from the atmosphere (Wigley & Schimel 2000).

CO$_2$ has been previously measured from volcanoes (Ryan 2001) and is known to be emitted in volcanic gas (Shinohara et al. 2011) as a consequence of decarbonation of underlying carbonate rocks (Frondini et al. 2004). There are a number of difficulties in measuring CO$_2$ from volcanoes. For ground based observations this mainly includes the inaccessibility and risk of taking measurements at the summits and the high level of background CO$_2$ concentrations (Spinetti et al. 2008). Whilst airborne observations overcome most of these problems, both techniques have trouble measuring CO$_2$ within a volcanic plume (Burton et al. 2000).

One method to reduce the current levels of CO$_2$ is the enhancement of natural sinks. Forests and soils could be increased to sequester CO$_2$, however this requires a large land mass (Chadwick et al. 2000) which would cause competition with agriculture. Also, the CO$_2$ captured by trees would only last for $\sim$100 years and that is provided that the area does not undergo deforestation, logging, biomass burning or that it becomes unsuitable for forestry due to increasing temperatures from climate change (Baines & Worden 2004). Agricultural techniques could be modified to enhance soil CO$_2$ sequestration but this would not be a permanent sink (IPCC 2007). Although these natural sinks could be increased, it is not a feasible method to use for large scale reductions in CO$_2$.

Geological sequestration is another way that is currently being used to reduce CO$_2$ concentrations. This method is capable of lowering fossil fuel emissions by over 80% (Freund 2003) but has two disadvantages; it requires energy (so fossil fuels are depleted quicker) and the CO$_2$ has the potential to leak back into the atmosphere, possibly leading to future climate change events (Keller et al. 2008). The rate of CO$_2$ leaking depends on where it is geologically stored (Freund 2003). Deep oceans will take centuries to leak. Whilst geological reservoirs have a much longer timescale of millennia and sequestration in thermodynamically stable minerals will take even longer to leak (Freund 2003,
Lackner 2003).
1.4 Observations of CO₂

Observations of CO₂ can be made on a local scale and the scaled up to estimate regional carbon fluxes, which are useful for land management and emission policies (Wofsy & Harriss 2002). This bottom-up method can be limited by the distribution of observations, and regional scaling can be difficult where multiple ecosystem types, heterogeneous terrain or urban areas exist (Desai et al. 2008, Dolman et al. 2009, Riley et al. 2009). Alternatively, regional estimates can be made using a top-down approach, by scaling down global surface flux estimates which are based on atmospheric observations of CO₂ (Ahmadav et al. 2009, Gurney et al. 2002a). Top-down observations have previously been limited to continental scales due to the complexity of regional carbon fluxes (Desai et al. 2010). However, with improvements in remote sensing technologies, top-down observations are approaching the point where they may be useful for the estimation carbon fluxes on a sub-continental scale, which this thesis focases on.

CO₂ has been measured in the atmosphere directly since 1957 (Wigley & Schimel 2000). The concentration and isotope records before then consist from evidence of ice cores, moss cores, packrat middens, tree rings, and isotopic measurements of plantonic and benthic foraminifera (Wigley & Schimel 2000). The longest record of atmospheric CO₂ concentrations is at Mauna Loa, which was started at the Institute of Oceanography in 1958. The National Oceanic and Atmospheric Administration (NOAA) has continued to measure in parallel here since 1974 until present (Tans & Keeling 2012). This is shown in Figure 1.9 which clearly shows the annual increase and seasonal cycle of CO₂.

In recent years there have been many in-situ measurements made from instruments such as ground based, tower, and aircraft observations. These have been collected into the GLOBALVIEW-CO2 product, as a Cooperative Atmospheric Data Integration Project (Maserie & Tans 1995). This project includes the work of many organisations and is coordinated by the Carbon Cycle Greenhouse Gases Group of the NOAA, Earth System Research Laboratory (ESRL). The measurement locations of this project are shown in Figure 1.10.

Airborne measurements have previously been obtained from commercial airliners typically, which normally provide data over short distances and mostly at a single altitude (Wofsy et al. 2010). However, the HIAPER Pole-to-Pole Observations (HIPPO) programme has taken a sequence of measurements, going from North to South with vertical information (Wofsy et al. 2010). These flights were made in 2009 and 2010, and have shown both the latitudinal and vertical gradients of a number of trace gases, specifically illustrating that CO₂ has a low variability in the Southern hemisphere (Wofsy et al.
However, due to the variability of CO$_2$ and lack of a comprehensive network of measurement locations, in-situ observations only allow inferred carbon fluxes on a sub-continental scale. Furthermore, their sparse and uneven distribution results in large uncertainties in the natural carbon cycle for key regions such as tropical and boreal regions (Hungershoefer et al. 2010, Gurney et al. 2002a). As shown by numerous synthetic studies, densely sampled satellite observations of CO$_2$ concentrations can help to reduce uncertainties in estimated regional carbon fluxes if the observations have a precision of 1-2 ppm on a regional scale with no regional to continental scale geographical biases (Rayner & O’Brien 2001, Houweling et al. 2004, Chevallier 2007b, Miller et al. 2007, Feng et al. 2009, Baker et al. 2010, Hungershoefer et al. 2010, Palmer et al. 2011). Additionally, Hungershoefer et al. (2010) investigated the concept of using a larger network of ground based measurement sites, costing $\sim$200 Million Euros (roughly the same as
a satellite mission) and found that the model error reduction was far higher if satellites are used, mainly due to the fact that remote sensing by satellite observations can provide a high density of measurements globally (Hungershoefer et al. 2010). However, Palmer et al. (2011) found that the benefit of the increased number of soundings of satellite observations is reduced with more correlated data. An example of the number of satellite observations globally can been seen in Figure 1.12.

Unlike most in-situ measurements, satellites observe throughout the atmosphere rather than at a single location and altitude, giving total column average concentrations. To validate these remotely sensed observations, the Total Carbon Column Observing Network (TCCON) of ground based Fourier Transform Spectrometers (FTS) was setup to retrieve accurate and precise column averaged abundances of greenhouse gases (Wunch et al. 2011a). The locations of TCCON stations are given in Figure 1.13.

Observations of CO$_2$ have been made from previous satellites. One notable example is the SCanning Imaging Absorption spectroMeter for Atmospheric CHartographY (SCIAMACHY) instrument. This was launched onboard the ENVISAT satellite in
March 2002 and has allowed continental and seasonal variations in CO₂ to be observed (Barkley et al. 2006a, Buchwitz et al. 2005, Scheising 2009, Schneising et al. 2008), as shown in Figure 1.14 which compares SCIAMACHY CO₂ with model CO₂ from the NOAA ESRL CarbonTracker model (Peters et al. 2007). Also, the Atmospheric Infrared Sounder (AIRS) instrument onboard the NASA Aqua satellite, launched in May 2002, has provided information about CO₂ in the upper troposphere globally, see Figure 1.15 for example. The main findings include a variability in the distribution of CO₂ with larger values observed over the northern hemispheric land masses and also a belt across the southern hemisphere, both of which were not previously incorporated in models (Chahine et al. 2008, Pagano 2011). Further details of previous satellite observations of CO₂ are given in Chapter 2.
**Figure 1.12.** Nominal ground track measurement locations of the Greenhouse gases Observing SATellite (GOSAT) for a 3-day orbital repeat cycle, with each measurement location is shown by a blue circle.

**Figure 1.13.** Current Total Carbon Column Observing Network (TCCON) sites located around the world. Figure taken from Wunch et al. (2011a).
Figure 1.14. A comparison of column-averaged dry air mole fractions of CO$_2$ retrieved from SCIAMACHY using the WFM-DOAS version 1.0 (WFMDv1.0) retrieval algorithm with CarbonTracker model calculations for three years. Figure taken from Scheising (2009).
FIGURE 1.15. Monthly average mid-tropospheric CO$_2$ retrieved from AIRS for July 2010. Figure taken from Pagano (2011).
1.5 Current understanding of the carbon cycle

It has been established in recent decades that over half the anthropogenic CO₂ emitted into the atmosphere has been sequestered into the oceans and terrestrial biosphere, leaving the remainder in the atmosphere to accumulate (see Figure 1.16). This has lead to a rise in atmospheric CO₂ which has been observed since the start of the industrial revolution and reflects the inability of the natural carbon sinks to uptake the increased volume of atmospheric CO₂.

![Diagram showing CO₂ budget between 1959 and 2006](image)

**Figure 1.16.** The budget of CO₂ between 1959 and 2006. The upper panel shows CO₂ emissions to the atmosphere as combination of fossil fuel combustion, land-use change and other emissions (primarily from cement production). The lower panel shows the resulting atmospheric, ocean and land accumulations of CO₂, where interannual variations are driven by the land sink. Figure taken from Canadell et al. (2007).
Although anthropogenic CO$_2$ emissions are consistently increasing, interannual CO$_2$ variations are observed (Figure 1.16), which are primarily controlled by changes of CO$_2$ flux between the atmosphere and terrestrial biophere (IPCC 2007). There is a lack of accurate quantitative knowledge of the carbon sinks, specifically their identity, strength and geographical distribution, as well as how they might change in the future and their potential feedback on the carbon-climate system (Houghton 2002, Fung et al. 2005, IPCC 2007). Also, the fraction of atmospheric CO$_2$ absorbed between the ocean and terrestrial sinks is highly uncertain (see Figure 1.17), with some studies suggesting the ocean sink to be dominant (e.g. Bopp et al. 2002) while others point to terrestrial reservoirs being dominant (e.g. Fan et al. 1998, Bousquet et al. 1999, Pacala et al. 2001). The uptake of carbon by the southern hemisphere oceans remains largely uncertain (Takahashi et al. 2002, McNeil et al. 2003), as does coastal ocean regions (Tsunogai et al. 1999, Muller-Karger et al. 2005). The geographical distribution of the northern hemisphere terrestrial sink has been suggested to be dominant in various regions, ranging from North America (Fan et al. 1998) to Asia (Bousquet et al. 1999).

Numerous chemistry transport models have been previously used to estimate surface-atmosphere CO$_2$ fluxes using inversion techniques, which assimilated in-situ CO$_2$ measurements from flasks, continuous sensors, tall towers and aircraft flights. The Atmospheric Tracer Transport Model Intercomparison Project (Transcom) provided an intercomparison of such models, giving estimates of the uncertainties in the net carbon fluxes from oceans and terrestrial reservoirs (Gurney et al. 2002b). The regional flux uncertainty estimates of the TRANSCOM experiment are summarised in Figure 1.18, showing that the uncertainties are large enough that most ocean and terrestrial regions could be net sources or net sinks (Gurney et al. 2002a). Furthermore, the experiment highlighted that the existing network of CO$_2$ measurements were too sparse and unevenly distributed, leading to an under sampled system and hence the requirement for more measurements.

Presently, large uncertainties exist for climate-carbon cycle feedback mechanisms (see Figure 1.19). It is expected that anthropogenic emissions will increase throughout the 21st century, leading to a reduced CO$_2$ uptake efficiency of oceans due to the carbonate buffering mechanism, and of land due to the saturation of terrestrial sinks (Friedlingstein et al. 2006). Additionally, the predicted change in future climate is estimated to suppress carbon sinks, therefore increasing the fraction of atmospheric CO$_2$ (e.g. Gurney et al. 2002a, Friedlingstein et al. 2006). This creates a positive climate change feedback that is highly uncertain, with current models estimating a rise in the rate of increase of CO$_2$ ranging between 4% and 44% (IPCC 2007).
Figure 1.17. Fractional partitioning of the predicted increase in total emissions that contribute to atmospheric CO$_2$. Variations of the emissions partitioning as simulated by the C$^4$MIP models up to 2000 are given by black letters and for the entire simulation period to 2100 are given in red letters, where each letter represents a different model. The dotted box shows a constraint on the historical carbon balance based which was based on records of atmospheric CO$_2$ increase, estimates of total emissions from fossil fuel combustion and land use change, and oceanic uptake of anthropogenic CO$_2$ (Sabine et al. 2004). The black and red diamond symbols give the mean carbon partitioning of all the models for the historical and entire simulation periods, respectively. The red line between these diamond symbols shows the mean model tendency towards an increasing atmospheric fraction through the 21st century, which is common to all models. Figure taken from IPCC (2007).
Figure 1.18. The mean estimated CO$_2$ sources and uncertainties for two different inversion models for various ocean and land regions. For each region the left-hand symbols are for a control inversion and the right-hand symbols are for an inversion without any background seasonal biosphere flux. The mean estimated flux is shown by a cross symbol, which includes all background fluxes expect fossil fuels. The circle symbols represent the mean estimated uncertainty across all models. The error bars give the standard deviation of the models’ estimated fluxes. The priori flux estimates and their uncertainties are indicated by the boxes (solid for land, dashed for ocean), where the central horizontal bar gives the prior flux estimate, and the top and bottom of the box shows the prior flux uncertainty range. Figure taken from Gurney et al. (2002a).
Figure 1.19. The uncertainties in carbon cycle feedbacks estimated from C4MIP models for the simulated period of typically 1860 to 2100. The top bar shows the range that the climate-carbon cycle feedback is estimated to have on the airbourne fraction of total emissions. Similarly, the middle 4 bars show the impact of climate change on the carbon cycle, and the lower 3 bars give the direct response to increasing atmospheric CO$_2$ concentrations. Figure taken from IPCC (2007).
1.6 Precision Requirements

CO₂ inferred from global satellite measurements has been shown through numerous sensitivity studies to improve carbon flux inversions provided the observations are precise (Baker et al. 2006b, Dufour & Breon 2003, Houweling et al. 2004, Moa & Kawa 2004, O’Brien & Rayner 2002, Rayner et al. 2002, Rayner & O’Brien 2001). These investigations conclude that precisions between 1 - 10 ppm will reduce uncertainties of CO₂ sources and sinks due to the dense global sampling of satellites. The spatial and temporal resolution of the satellite measurements and the surface flux inversion spatiotemporal scale effects the requirement for precision (Houweling et al. 2004). The greater the precision the more beneficial the data will be for determining sources and sinks, but this must also be balanced against what our current level of satellite technology, retrieval method and validation system can reproduce (Miller et al. 2007). The most realistic study (Rayner et al. 2002) has shown that satellites with a $8^\circ \times 10^\circ$ monthly mean column CO₂ precision of 2.5 ppm or less would surpass the capability of existing surface networks in determining sources and sinks on a continental scale (Rayner & O’Brien 2001) (see Figure 1.20). The ESA Climate Change Initiative (CCI) for the Essential Climate Variable (ECV) Greenhouse Gases (GHG) provides the latest accuracy and precision requirements for CO₂ satellite observations that are based on studies of synthetic data (e.g. Chevallier et al. 2005a, 2007a, 2009, Miller et al. 2007). It states a breakthrough requirement for satellite observations would be a $<0.3$ ppm accuracy (systematic error) and a precision (random error) of $<3$ ppm for a single observation and $<1$ ppm for monthly mean 1000 km$^2$ observations. Additionally, it gives the ideal satellite requirement of $<0.2$ ppm accuracy (systematic error) and a precision (random error) of $<1$ ppm for a single observation and $<0.3$ ppm for monthly mean 1000 km$^2$ observations (Buchwitz et al. 2011).
FIGURE 1.20. Comparison of global CO$_2$ uncertainty and precision when no observations are assimilated (dotted line), observations from 56 ground based measurement stations are assimilated (dashdotted line), simulated global coverage from satellite observations are assimilated (solid line), and simulated global coverage of oceans only from satellite observations are assimilated (dashed line). Figure taken from Rayner & O’Brien (2001).
1.7 Thesis Overview

The aim of this thesis is to develop an algorithm to estimate atmospheric CO$_2$ from space based observations and to assess the accuracy of these estimations by validating them with independent in-situ observations. Satellites measurements, if acquired with a high accuracy and precision, have the potential to reduce model flux uncertainties and improve our knowledge of the carbon cycle, allowing the identification of surface sources and sinks of CO$_2$.

This thesis describes the development of a new retrieval algorithm capable of retrieving CO$_2$ from space borne measurements, called the University of Leicester Full Physics (UoL-FP) retrieval algorithm. The second chapter provides the fundamental theories for remote sensing and how these have been used by previous satellite instruments to infer CO$_2$. The first satellite that is proposed to meet this requirement, called GOSAT, is then described. Chapter three explains the UoL-FP retrieval algorithm that was developed for use with GOSAT observations. The retrieved CO$_2$ estimates are validated against ground based remote sensing measurements in the fourth chapter to assess the accuracy of the algorithm. Chapter four also includes global comparisons to a global chemistry transport model providing the first steps towards source/sink flux inversions and compares to alternative retrieval algorithms to evaluate any obvious retrieval dependent biases. The framework of the algorithm is developed in the fifth chapter by the creation of a simulator that can be used to investigate multiple retrieval algorithm modifications, which may lead to the development of a new retrieval algorithm method with improved CO$_2$ precision. Chapter six summarises the thesis and provides an outlook for future expansion.
Chapter 2

Remote Sensing of Atmospheric CO$_2$

2.1 Introduction to Remote Sensing of Atmospheric CO$_2$

Satellite instruments can measure light at the top of the atmosphere that has come from the Earth. These measurements can be made using active or passive remote sensing techniques. Active sensing is where monochromatic light is sent from the satellite to the Earth’s surface and then the reflected light is measured. The difference between the original and measured light can then provide information about the content of the atmosphere. However, this method requires high levels of power and only measures light at specific wavelengths. Passive sensing measures Sun light reflected off the Earth’s surface, and the difference between the measured light and estimated light from the Sun can provide information about the Earth’s atmosphere. This method requires much less power, can measure multiple wavelength ranges and has a high accuracy.

The Sun emits light strongly in the visible and shortwave infrared (SWIR) range, but any radiation absorbed by the Earth will be re-radiated at wavelengths in the thermal infrared (TIR) range (the same range as the Earth’s surface naturally radiates at). Space based TIR measurements therefore provide information where absorption occurs throughout the atmosphere, especially in the upper troposphere where clouds are present. Satellite SWIR measurements also provide information throughout the atmosphere, but are more sensitive to reflection. Hence, SWIR observations are obstructed by thick clouds. However, for cloud free scenes, SWIR measurements are most sensitive to the lower troposphere, where there are more gas molecules, and can provide information about the atmosphere just above surface CO$_2$ sources and sinks.

As light travels through the atmosphere some of it gets absorbed by gas molecules and some light will be absorbed or scattered by aerosols and clouds. Each of the different
gases that make up the atmosphere will cause light to be absorbed at certain wavelengths specific to the structure of that gas. Hence, remote sensing of specific wavelengths can provide information about a particular type of gas, allowing its atmospheric concentration to be inferred.

Figure 2.1 illustrates absorption by different atmospheric gases for some example wavelength ranges, where lower radiance values typically mean absorption has occurred. The amount of absorption that has occurred (i.e. how much the radiance is decreased by compared to the continuum level) can be compared to tabulated information of a single molecule to estimate how much of that gas exists.

Figure 2.1. Example radiances for each of the TANSO-FTS SWIR bands showing absorption of different gases, where blue represents O$_2$, green represents CO$_2$, pink represents CH$_4$, and cyan represents H$_2$O.
2.2 Fundamentals of Radiation in the Atmosphere

2.2.1 Light Absorption

The absorption spectral lines observed in measurements of atmospheric light are created when light of a certain wavelength becomes absorbed by a molecule within the beam. If a photon with energy $\delta E = h\nu$ is absorbed then a molecule goes through a transition from its ground energy state to an excited energy state, where $h$ is Planck’s constant ($6.626076 \times 10^{-34}$ J$s$) and $\nu$ is the frequency of the electromagnetic radiation. The total energy of a molecule is given by the sum of the potential energy and kinetic energy, which can be represented by the Hamiltonian operator $\hat{H}$. The energy level, $E$, of a molecule must obey the time independent Schrödinger equation

$$E\Psi = \hat{H}\Psi \quad (2.1)$$

where $\Psi$ is the eigenvalue of the operator. During the absorption of light by a molecule, there are additionally four contributing aspects that must be considered; the electronic, vibrational, rotational and nuclear energies of the molecule. Assuming these to be independent (Born-Oppenheimer approximation), the Schrödinger equation can be solved for each energy separately and the total energy of a molecule can then be given as

$$E = E_{\text{Electronic}} + E_{\text{Vibrational}} + E_{\text{Rotational}} + E_{\text{Nuclear}} \quad (2.2)$$

where the eigenvalue is the product of each aspect:

$$\Psi = \Psi_{\text{Electronic}}\Psi_{\text{Vibrational}}\Psi_{\text{Rotational}}\Psi_{\text{Nuclear}} \quad (2.3)$$

In the SWIR wavelength range the nuclear energies are small, whilst the electronic, vibrational and rotational energies are larger, with the vibrational energy being comparable in energy to that of infrared radiation.

2.2.1.1 Rotational Transitions

Molecules can be considered as rigid structures that may rotate. This rotation can be described about three orthogonal axes that pass through the molecules centre of gravity. The axes A, B and C are defined such that the corresponding inertia, $I$, of the molecule rotating around each axis is given as $I_A \leq I_B \leq I_C$. The rotational energy can be given
by

\[ E_{\text{Rotational}} = \frac{1}{2} I_A \omega_A^2 + \frac{1}{2} I_B \omega_B^2 + \frac{1}{2} I_C \omega_C^2 \]  

(2.4)

where \( \omega \) represents the corresponding angular velocity around each axis. The energy is proportional to both inertia and angular velocity about each axis, which both depend on the structure of the molecule. Thus, molecules are categorized by their structure, as shown in Table 2.1.

Table 2.1. Molecule structure classification.

<table>
<thead>
<tr>
<th>Classification</th>
<th>Moment of Inertia I</th>
<th>Example molecules</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linear</td>
<td>( I_A = 0, I_B = I_C )</td>
<td>CO(_2), N(_2)O</td>
</tr>
<tr>
<td>Symmetric top</td>
<td>( I_A \leq I_B = I_C ) or ( I_A = I_B \leq I_C )</td>
<td>CFC(_3), NH(_3)</td>
</tr>
<tr>
<td>Spherical top</td>
<td>( I_A = I_B = I_C )</td>
<td>CH(_4), SF(_6), CCl(_4)</td>
</tr>
<tr>
<td>Asymmetric top</td>
<td>( I_A \neq I_B \neq I_C )</td>
<td>H(_2)O</td>
</tr>
</tbody>
</table>

Symmetric top molecules can be sub-catagorised as prolate\(^{\dagger}\) or oblate\(^{\ddagger}\) symmetric top molecules.

However, the assumption of rigid structures rotating is only an approximation and in reality there are clear deviations from this model.

### 2.2.1.2 Vibrational Transitions

Molecules may vibrate with a given number of modes. This is determined by the number of degrees of freedom, where a molecule of \( x \) atoms has \( 3x \) degrees of freedom. However, three of the degrees of freedom represent the translational motion of the molecule in each axis direction and another three represent the rotational motion about each axis. Therefore, \( 3x - 6 \) degrees of freedom is the number of vibrational modes for a non-linear molecule. For a linear molecule the rotational motion can be described by two degrees of freedom, so the number of modes becomes \( 3x - 5 \). Taking CO\(_2\) as an example, it is linear and comprises of 3 atoms so has 4 vibrational modes, which are shown in Figure 2.2.

The energy associated with each vibrational mode, \( i \), can be approximately described by the harmonic displacements of each atom in the molecule which vibrate at a certain characteristic frequency, \( v_i \). This is given by the equation

\[ E_{i,\nu} = \hbar v_i \left( v_i + \frac{1}{2} \right) \]  

(2.5)
where $\nu$ is the vibrational quantum number. However, in reality only harmonic displacements for low vibrational modes are valid. Higher vibrational modes are better characterised by anharmonic displacement, with the energy given by

$$
E_{i,\nu} = \hbar \nu_i \left( \nu_i + \frac{1}{2} \right) + \hbar \nu_i x_i \left( \nu_i + \frac{1}{2} \right)^2
$$

(2.6)

where $x$ represents the anharmonicity constant.
2.2.1.3 Vibrational-Rotational Bands

The energy needed for a molecule to rotate is much smaller than that for vibration. Therefore, when a vibrational transition occurs a simultaneous rotational transition will accompany it. There may be multiple rotational transitions occurring simultaneously from different energy levels. But because the rotational transitions are too close together to be resolved they are grouped together as a vibration-rotation band (Griffiths & de Haseth 2007). The vibration-rotation band is generally defined by three branches; P, Q and R branches. The P and R branches correspond to wavelengths higher and lower than the band centre, respectively. Whilst the Q branch occurs during a vibration bending mode. In the SWIR range, CO$_2$ contains mostly strong P and R branches (Toth et al. 2008).

2.2.1.4 Line Shape

The spectral line intensity depends on the absorption by the interfering molecule, which can be related as

\[ S = \int_{-\infty}^{\infty} C_v dv \]  

(2.7)

where \( S \) is the line intensity and \( C \) is the absorption cross section. The energy levels during transitions change slightly due to external influences on the molecule and the loss of energy in emission. Repeated energy transitions cause non-monochromatic radiances, therefore spectral lines have a finite width. This can be expressed in terms of the shape of the spectral line:

\[ k_v = S \Phi (v – v_0) \]  

(2.8)

where \( \Phi(v-v_0) \) represents the line shape factor of absorption at specific spectral distances from the line centre, \( v_0 \). The broadening of lines depends on the damping of molecular vibrations; a doppler effect resulting from differences in thermal velocities, and any collisions between molecules.

Natural broadening of lines is caused by the loss of energy in emission. This can be explained by the Heisenberg uncertainty principle, whereby the line width is inversely proportional to the lifetime of an excited state (\( \Delta t \)), which in turn is related to the energy of that quantum state (\( \Delta E=h\Delta v \)):

\[ \Delta t \approx \frac{1}{2\pi \Delta v} \]  

(2.9)

The line shape of natural broadening, \( \Phi_N \), can be given as a Lorentz profile (Lorentz...
1906):

\[ \Phi_N (v - v_0) = \frac{\alpha_L / \pi}{(v - v_0)^2 + \alpha_L^2} \]  

(2.10)

where \( \alpha_L \) represents the natural Lorentz shape at half-width-half-maximum (HWHM).

Pressure broadening occurs due to the collisions of molecules and therefore is more dominant in the lower atmosphere where more molecules exist. The line shape of pressure broadening is also described by a Lorentz profile:

\[ \Phi_P (v - v_0) = \frac{\alpha_L / \pi}{(v - v_0)^2 + \alpha_L^2} \]  

(2.11)

At higher altitudes there are less molecules that can collide so the doppler effect becomes important. If a molecule has a velocity in the line of sight and \( v \ll c \) then the doppler effect occurs:

\[ v = v_0 \left( 1 \pm \frac{v}{c} \right) \]  

(2.12)

In Kinetic theory, translational states are in thermodynamic equilibrium and the probability that the velocity is between \( v \) and \( v + dv \) is \( p(v)dv \). The maxwell distribution gives \( p(v) \) such that

\[ p(v) dv = \left( \frac{m}{2\pi kT} \right)^{\frac{1}{2}} e^{-\frac{mv^2}{2kT}} dv \]  

(2.13)

where \( k \) is Boltzmann’s constant, \( T \) is temperature and \( m \) is the mass. Combining Equations 2.12 and 2.13, the line shape of doppler broadening can be given as

\[ \Phi_D (v - v_0) = \frac{1}{\alpha_D \sqrt{\pi}} \exp \left[ - \left( \frac{v - v_0}{\alpha_D^2} \right)^2 \right] \]  

(2.14)

where the HWHM is \( \alpha_D \sqrt{\ln 2} \) and

\[ \alpha_D = v_0 \left( \frac{2kT}{mc^2} \right)^{\frac{1}{2}} \]  

(2.15)

In the upper atmosphere both pressure and doppler broadening are important to consider. Therefore, the line shapes of both are convolved to create a Voigt profile shape:

\[ \Phi_V (v - v_0) = \frac{1}{\pi^2 \alpha_D} \int_{-\infty}^{\infty} \frac{1}{\alpha_L} \exp \left[ - \left( \frac{v - v'}{\alpha_D^2} \right)^2 \right] dv' \]  

(2.16)

where the doppler shift is added to the pressure broadened line at wavenumbers \( v' = v_0 \).

It is also possible for line-mixing to occur, where multiple absorption lines overlap.
Recent studies have shown this must be taken into account for highly accurate retrievals of $X_{CO2}$ to be achieved (e.g. Hartmann et al. 2009, Lamouroux et al. 2010, Vangwichith et al. 2009).

2.2.2 Radiative Transfer

2.2.2.1 The Radiative Transfer Equation

Passive space borne SWIR remote sensing instruments measure the intensity of electromagnetic radiation that has travelled from the Sun and been reflected off the Earth’s surface towards the instrument optics, which is the focus of this study. This beam of radiation travels through the Earth’s atmosphere where a number of different physical processes can alter the intensity of the radiation. Figure 2.3 illustrates these processes which include; absorption of radiation, radiation scattered out of the beam, single scattering of direct solar radiation into the beam, emitted radiation into the beam, and multiple scattering of indirect diffuse radiation into the beam (both solar and emitted radiances). The radiative transfer equation describes the change in the radiance of the beam for a specified wavelength, $dI_\lambda$, over the distance travelled, $ds$, and can be given for simplicity as

$$dI_\lambda = -dI_{SO,\lambda} - dI_{AO,\lambda} + dI_{SI,\lambda} + dI_{MI,\lambda} + dI_{EI,\lambda} \quad (2.17)$$

where the first term on the right hand side is the intensity of radiation scattered out of the beam and is expressed as

$$dI_{SO,\lambda} = I_\lambda \sigma_{s,\lambda} ds \quad (2.18)$$

where $I$ is the intensity and $\sigma_s$ is the scattering extinction coefficient. The second term of Equation 2.17 gives the intensity of radiation absorbed throughout the beam and can be written as

$$dI_{AO,\lambda} = I_\lambda \sigma_{a,\lambda} ds \quad (2.19)$$

where $\sigma_a$ is the absorption extinction coefficient. The intensity of solar radiation directly scattered into the beam by a single particle is represented in Equation 2.17 as the third term and is given as

$$dI_{SI,\lambda} = \left[ \sum_k \left( \frac{\sigma_{s,k,\lambda} P_{s,k,\lambda,\mu,-\mu_s,\phi_s}}{4\pi} \right) \right] F_{s,\lambda} e^{-\tau_{\lambda,\mu_s}} ds \quad (2.20)$$

where $k$ represents different particles, $P$ is the scattering phase function that describes the angular distribution of scattering with directions, $F$ is the total irradiance (solar ra-
radiation) at the top of the atmosphere, and Beer’s law is included with the exponential term showing that the absorption of radiation increases exponentially with the optical depth, $\tau$. The fourth term of Equation 2.17 represents the multiple scattering of diffuse radiation into the beam and is given by

$$dI_{MI,\lambda} = \left[ \sum_k \left( \frac{\sigma_{s,k,\lambda}}{4\pi} \int_0^{2\pi} \int_{-1}^1 I_{\lambda,\mu',\phi'} P_{s,k,\lambda,\mu,\mu',\phi,\phi'} d\mu' d\phi' \right) \right] ds \quad (2.21)$$

The final term of Equation 2.17 represents the emitted radiation into the beam. The emission occurs in the infrared wavelength range whereby particles that have previously absorbed radiation re-emit infrared radiation. This can be expressed as

$$dI_{EI,\lambda} = \sigma_{a,\lambda} B_{\lambda,T} ds \quad (2.22)$$

where $B_{\lambda,T}$ represents Planck’s law, which gives the intensity emitted from a particle as a function of wavelength and temperature. Planck’s law is given by

$$B_{\lambda,T} = \frac{2hc^2}{\lambda^5 \left[ e^{(\frac{hc}{\lambda k_b T})} - 1 \right]} \quad (2.23)$$

where $h$ is Planck’s constant ($6.6256 \times 10^{-34}$ Js), $c$ is the speed of light ($2.9979 \times 10^8$ ms$^{-1}$ in a vacuum), $k_b$ is Boltzmann’s constant ($1.38 \times 10^{-23}$ JK$^{-1}$), and $T$ is temperature. Thus, substituting Equations 2.18 to 2.22 into Equation 2.17 the radiative transfer equation can be given as

$$\frac{dI_{s,\lambda,\mu,\phi}}{ds} = -I_{s,\lambda,\mu,\phi} (\sigma_{s,\lambda} + \sigma_{a,\lambda}) + F_{s,\lambda} e^{-\tau_{s,s}} \sum_k \left( \frac{\sigma_{s,k,\lambda}}{4\pi} P_{s,k,\lambda,\mu,-\mu_s,\phi_s} \right)$$

$$+ \sum_k \left( \frac{\sigma_{s,k,\lambda}}{4\pi} \int_0^{2\pi} \int_{-1}^1 I_{\lambda,\mu',\phi'} P_{s,k,\lambda,\mu,\mu',\phi,\phi'} d\mu' d\phi' \right) + \sigma_{a,\lambda} B_{\lambda,T} \quad (2.24)$$
Figure 2.3. Radiative transfer diagram, illustrating the unattenuated beam (left) and the beam with multiple processes interacting with it (right), where the beam orientation is $\mu$, $\phi$. The solar zenith angles of the beam and diffuse radiation is given by $\theta$ and $\theta'$, respectively. $\phi$ is the azimuth angle, $\phi'$ is the azimuth angle of the diffuse radiation, $\mu = \cos \theta$, and $\mu' = \cos \theta'$. For downward radiances $\mu$ is negatively assigned and for upward radiances $\mu$ is positively assigned. Adapted from Jacobson (2005) and Liou (2002).
2.3 Types of Scattering

Electromagnetic radiation becomes scattered when a particle is within the path of a light wave, causing light to become reflected, refracted or diffracted, changing the direction of the light path. The effect that the size of the particle has on scattering is defined by a term called the size parameter, which for a spherical particle is given as

\[ x = \frac{2\pi a}{\lambda} \]  

(2.25)

where \( x \) is the size parameter, \( a \) is the particle radius and \( \lambda \) is the wavelength of light.

Scattering can be defined with two types; Rayleigh scattering where \( x \ll 1 \) and Lorenz-Mie scattering where \( x \gg 1 \).

The polarisation state of an electromagnetic wave can be described using four linearly independent values referred to as the Stokes parameters:

\[ I = E_P E_P^* + E_S E_S^* \]  

(2.26)

\[ Q = E_P E_P^* - E_S E_S^* \]  

(2.27)

\[ U = E_P E_S^* + E_S E_P^* \]  

(2.28)

\[ V = -i (E_P E_S^* - E_S E_P^*) \]  

(2.29)

where \( I \) is the total intensity of the wave, \( Q \) and \( U \) give the linear polarisations, \( V \) gives the state of the circular polarisation of the wave, \( * \) denotes the complex conjugate, and \( i \) equals \( \sqrt{-1} \). These parameters are based on the wave being characterised by the electric field, \( E \), being parallel (P) and perpendicular (S) to the incident plane. These stokes parameters can be summarised by the Stokes vector

\[ \mathbf{I} = \begin{bmatrix} I \\ Q \\ U \\ V \end{bmatrix} \]  

(2.30)

The intensity of scattered light can be given in terms of the incident light (\( I_0 \)) by

\[ \mathbf{I} = \mathbf{P}(\Omega) \mathbf{I}_0 \]  

(2.31)

where \( \mathbf{P} \) represents the phase matrix as a function of scattering angle, \( \Omega \). The phase
matrix gives the phase function with polarisation, and can be defined as

\[
P(\Omega) = \begin{bmatrix}
P_{11}(\Omega) & P_{12}(\Omega) & 0 & 0 \\
P_{12}(\Omega) & P_{22}(\Omega) & 0 & 0 \\
0 & 0 & P_{33}(\Omega) & P_{34}(\Omega) \\
0 & 0 & -P_{34}(\Omega) & P_{44}(\Omega)
\end{bmatrix}
\] (2.32)

The phase matrix is defined by the phase function, P, which gives the intensity as a function of scattering angle. For spheres, \( P_{22} = P_{11} \) and \( P_{44} = P_{33} \).

When light is scattered by particles that are small compared to the wavelength of light or the photons are elastically scattered, Rayleigh scattering occurs. The scattering cross section for Rayleigh scattering, \( C_{\text{Rayleigh}} \), is given as:

\[
C_{\text{Rayleigh}} = \frac{8\pi r^6}{3} \left( \frac{2\pi n}{\lambda} \right)^4 \left( \frac{m^2 - 1}{m^2 + 2} \right)^2
\] (2.33)

where \( n \) is the refractive index of the surrounding medium (air), \( r \) is the radius of the particle and \( m \) is the ratio of the particle refractive index to \( n \). Note that the Rayleigh scattering cross section is proportional to \( r^6 \), so all air molecules will scatter and more scattering will occur for longer light paths through the atmosphere (hence at higher SZAs). Additionally, this means that larger particles have a larger scattering cross section, but are inversely proportional to wavelength, giving larger scattering cross sections at smaller wavelengths. Thus, Rayleigh scattering is small in the SWIR range. The phase function for Rayleigh scattering can be defined as

\[
P(\Omega) = \frac{3}{4} \left( 1 + \cos^2 \Omega \right)
\] (2.34)

Figure 2.4 shows the phase function for Rayleigh scattering, which scatters light equally in the forward and backward directions.

In the SWIR, Lorenz-Mie scattering dominates, where larger particles scatter light (\( x \gtrsim 1 \)). The direction of the scattered light becomes more towards the forward direction (see Figure 2.4). For Lorenz-Mie scattering, the off diagonal terms of the phase matrix are typically small, so polarisation will not affect intensity much. Thus, intensity can be approximated by \( P_{11} \), which can be defined as

\[
P_{11}(\Omega) = \frac{4\pi}{k^2 C_{\text{Lorenz-Mie}}} \frac{|S_1|^2 + |S_2|^2}{2}
\] (2.35)
where \( C_{\text{Lorenz-Mie}} \) is the scattering cross-section, \( k = 2\pi n/\lambda \), and \( S_1 \) and \( S_2 \) represent scattering amplitudes which are given as

\[
S_1 (\Omega) = \sum_{n=1}^{\infty} \frac{2n+1}{n(n+1)} [a_n \pi_n (\cos \Omega) + b_n \tau_n (\cos \Omega)]
\] (2.36)

\[
S_2 (\Omega) = \sum_{n=1}^{\infty} \frac{2n+1}{n(n+1)} [b_n \pi_n (\cos \Omega) + a_n \tau_n (\cos \Omega)]
\] (2.37)

where \( a \) and \( b \) are complex Mie coefficients that are expressed in terms of the spherical Bessel functions, and the Mie angular functions are defined as

\[
\pi_n (\cos \Omega) = \frac{1}{\sin \Omega} P_n^1 (\cos \Omega)
\] (2.38)

\[
\tau_n (\cos \Omega) = \frac{d}{d\Omega} P_n^1 (\cos \Omega)
\] (2.39)

where \( P_n^1 \) represent Legendre polynomials, where the number of terms is proportional to the size parameter. Additionally, the strength of forward scattering can be indicated by the asymmetry parameter which is given as

\[
Q = \frac{4}{x^2} \left[ \sum_n \frac{n(n+2)}{n+1} \text{Re} (a_n a_{n+1}^* + b_n b_{n+1}^*) + \frac{2n+1}{n(n+1)} \text{Re} (a_n b_n^*) \right]
\] (2.40)

where \( \text{Re} \) represents the effective radius.

Normally, the scattered light has the same wavelength as the incident light, this is known as elastic scattering and both Rayleigh and Lorenz-Mie scattering are examples of this. However, inelastic scattering can occur where a different wavelength is emitted. This is often referred to as Raman scattering, although fluorescence can also be considered an example of this.

The amount of radiation removed from a certain length of the light path by scattering and absorption is known as the optical depth. The total column optical depth is the total amount of light removed as it passes through the atmosphere towards the Earth’s surface and as it travels throughout the atmosphere after being reflected towards the satellite. The total optical depth is also the sum of the optical depth due to Rayleigh molecules, \( R \), aerosols, \( A \), and atmospheric gases, \( G \). As shown for Rayleigh and Lorenz-Mie scattering, the amount of scattering varies with wavelength, therefore the optical depth is wavelength dependent and can be expressed as

\[
\tau (\lambda) = \tau_R (\lambda) + \tau_A (\lambda) + \tau_G (\lambda)
\] (2.41)
Figure 2.4. Example scattering phase function in polar coordinates for a homogeneous spherical particle of different sizes and wavelengths. An example of Rayleigh scattering is shown in green. Whilst an example of Lorenz-Mie scattering is shown by the solid black line. Figure taken from (Steinacker et al. 2003).

The optical depths at different wavelengths (1 and 2) can be related by the angstrom coefficient:

$$\alpha = -\frac{\log \left( \frac{\tau_1}{\tau_2} \right)}{\log \left( \frac{\lambda_1}{\lambda_2} \right)}$$  \hspace{1cm} (2.42)

### 2.4 Atmospheric Scattering Particles

Although the type of scattering depends on the size of the particle, the direction and strength of scattering also depends on the shape of the particle and its refractive index as a function of wavelength in the Lorenz-Mie regime. In the atmosphere there is a large variety of particles, both natural and anthropogenic.

Cloud particles form when the temperature is low enough for water vapour to condense. However, condensation can only occur if there is a surface to condense onto.
In the atmosphere minute particles (aerosols) act as condensation nuclei allowing water vapour to condense to form clouds. Under normal conditions, water vapour will condense into water droplets, unless below 0°C when ice crystals form. However, under certain conditions water droplets can exist below 0°C and these are known as supercooled droplets. Since the type of cloud particle formed depends on temperature, water droplets are found at low altitudes (e.g. Stratus and Cumulus clouds), ice crystals at high altitudes (e.g. Cirrus, Cirrustratus and Cirrocumulus clouds) and a mixture of water and ice at middle altitudes (e.g. Altostratus and Altocumulus clouds). Water droplets are spherical in shape whilst ice crystals are uniaxial with a high variability (see Figure 2.6), and both may grow due to coalescence (collision of droplets as they fall through the atmosphere) (Lawson et al. 2001). Additionally, the Bergeron process may create larger cloud particles, whereby water molecules transfer from supercooled water droplets to nearby ice crystals due to the vapour pressure difference between them (Liou 1992). The refractive index of water and ice particles vary with wavelength as shown in Figure 2.5.

Water droplets typically have a radius between 1 and 40 µm and and due to their spherical shape, scatter mostly in the forward direction with a smaller backwards peak (see Figure 2.7). Ice crystals are larger in size, ranging between roughly 10 and 2000 µm, and scatter slightly less in the forward direction. There is also a large range in size for mixed phase (part water and ice) particles and these have similar scattering properties (Jourdan et al. 2003).

The other type of particles that scatter light are aerosols. Some aerosols are created naturally and these are generally well distributed around the world with sources covering large areas. Examples of this include; dust blown into the atmosphere from desert regions, sea salt spraying/blowing into the atmosphere from oceans, organic compounds from plant particles, and soot from biomass burning. These aerosols typically exist in the troposphere and have a lifetime of a couple of weeks due to water vapour condensing onto them and also due to coagulation of aerosols forming larger particles, both of which will cause the aerosol to fall to the surface (Williams et al. 2002). Another source of aerosols comes from volcanic eruptions which may throw up particles into the troposphere but also into the stratosphere where there is less water vapour for condensation to occur so aerosol lifetimes of 1-3 years are possible (Robuck et al. 2002). Urban aerosols tend to be anthropogenic and concentrated over small polluting regions in the lowest kilometer of the atmosphere. Urban aerosols are complex due to the mixtures of aerosols from various anthropogenic emission sources. Aerosols are highly variable in shape, size and refractive index (Williams et al. 2002), see Figures 2.8 and 2.9 for examples. Additionally, some aerosol types are hydrophobic whilst others are hydrophilic and may
Figure 2.5. Example of real and imaginary refractive index of liquid and ice cloud particles. The water refractive index values were obtained from Hale & Querry (1973) and the ice from Warren (1984).

change in size and shape with humidity (Shettle & Fenn 1979).
FIGURE 2.6. Example images of spherical water particles (left) and a few ice crystal shape variations (centre and right). Obtained from Lawson et al. (2001).

FIGURE 2.7. Measured and retrieved scattering phase functions and particle size distributions for water droplet clouds (a), mixed-phase clouds (b) and ice crystal clouds (c). Figure taken from Jourdan et al. (2003).
Figure 2.8. Example of real and imaginary refractive index of different three different aerosol types: dust, soot and sea salt. The refractive index values were obtained from Shettle & Fenn (1979).

Figure 2.9. Example images of aerosols from scanning electron microscopes courtesy of USGS, UMBC (Chere Petty, and Arizona State University (Peter Buseck). This shows a variety of aerosol shapes, including; volcanic ash (far left), pollen (centre left), sea salt (centre right), and soot (far right).
2.5 Previous Satellite Remote Sensing Observations of CO$_2$

The fundamental theories used for remote sensing described in the previous sections has allowed CO$_2$ to be inferred from satellite measurements for over three decades. A summary of these observations is given below.

The Television and Infrared Operational Satellite Next Generation (TIROS-N) Operational Vertical Sounder (TOVS) have been included onboard the National Oceanic and Atmospheric Administration (NOAA) polar meteorological satellites since 1978 (Chedin et al. 2003, Smith et al. 1979). This instrument comprises of three sounding units; the High-Resolution Infrared Radiation Sounder (HIRS-2), the Microwave Sounding Unit (MSU) and the Stratospheric Sounding Unit (SSU). Atmospheric and surface emissions can be inferred from HIRS-2 in seven channels; one at 15.0 $\mu$m, five channels around 4.3 $\mu$m, and one channel at 11.0 $\mu$m wavelength. The spectral bands around 15.0 $\mu$m and 4.3 $\mu$m observe radiances which highly depend on both atmospheric temperature and greenhouse gas concentrations, such as CO$_2$. Although CO$_2$ has been retrieved from TOVS it has an accuracy of the same order as the seasonal amplitude of atmospheric CO$_2$ (approximately 10 ppm) (Engelen & Stephens 2004) and the vertical sensitivity is largest in the mid-troposphere, making surface flux estimations have a large uncertainty. Additionally, regional biases cause the inferred fluxes to be not useful for biogeochemical analysis (Chevallier et al. 2005b).

The next generation of satellites that were capable of measuring CO$_2$ were launched in 2002. The SCanning Imaging Absoprtion spectroMeter for Atmospheric CHartography (SCIAMACHY) instrument, launched onboard the ENVISAT satellite in March 2002, is a passive ultraviolet (UV) - visible (VIS) - near infrared (NIR) hyper-spectral spectrometer designed to investigate tropospheric and stratospheric composition and processes with a nadir instantaneous field of view (IFOV) of 60 x 30 km$^2$ at the ground (Barkley et al. 2006, Bovensmann et al. 1999). Observations of CO$_2$ were retrieved from the 1.57 $\mu$m radiance channel (Barkley et al. 2006a, Buchwitz et al. 2005) and have been shown to follow the CO$_2$ seasonal cycle, but with a bias of approximately 4% and standard deviation of about 3% to in-situ observations (Barkley et al. 2006b). The relative accuracy and precision were both determined to be 1-2% for monthly averages for approximately a 7° x 7° footprint (Schneising et al. 2008). CO$_2$ retrieved from SCIAMACHY observations were unable to meet the 1% precision required for a 8° x 10° footprint to improve upon surface network measurements (Barkley et al. 2006). The differences between SCIAMACHY column average dry air mole fraction of CO$_2$ ($X_{CO2}$)
with ground based observations and model data were attributed to the low variability of global CO$_2$ distribution, and the inherent difficulties in retrieval algorithms to be able to account correctly for light scattering caused by clouds and aerosols (Houweling et al. 2005).

The SCIAMACHY instrument is most sensitive to CO$_2$ concentrations near the boundary layer due to it measuring in the near infrared (NIR) wavelength range. This sensitivity can be expressed by an averaging kernel, which indicates how sensitive a retrieval algorithm is to the true atmospheric profile for different altitude ranges. Figure 2.10 illustrates how the averaging kernel for SCIAMACHY peaks close to the boundary layer.

![SCIAMACHY NIR averaging kernels](image)

**Figure 2.10.** SCIAMACHY NIR averaging kernels from the Full Spectral Initiation (FSI) CO$_2$ retrieval algorithm for a range of SZA using a spectral window of 1561.03 to 1585.39 nm and albedo of 0.2 (left), with the mean averaging kernel given by the black dashed line. The normalised mean weighting function of AIRS which observes in the TIR spectral range is given (right) for comparison. Figure taken from Barkley et al. (2006).

Figure 2.10 also gives the averaging kernel of another instrument which is sensitive
broadly throughout the troposphere, peaking at 12 km in the tropics (Barkley et al. 2006). This instrument, the Atmospheric Infrared Sounder (AIRS) instrument, was launched in May 2002 onboard NASA's AQUA satellite and is a high spectral resolution spectrometer that measures radiances in the TIR wavelength range (3.7-15.4 $\mu$m) with an IFOV of roughly 13.5 x 18 km$^2$ at nadir. AIRS radiances have been assimilated directly into the European Centre for Medium-Range Weather Forecasts (ECMWF) four-dimensional variational (4D-Var) system, allowing tropospheric CO$_2$ to be estimated (Engelen et al. 2004). However, due to atmospheric mixing in the upper troposphere, the CO$_2$ concentrations estimated from AIRS are quite zonal and can only provide information about broad features of surface fluxes (Chevallier et al. 2005a).

Similarly, the Infrared Atmospheric Sounding Interferometer (IASI) and Advanced Microwave Sounding Unit (AMSU), launched together onboard the European MetOp platform in October 2006, can be used to retrieve CO$_2$ in the upper troposphere using the TIR wavelength range. Although these instruments provide accurate tropospheric CO$_2$ concentrations (0.5% monthly for a 5$^\circ$ x 5$^\circ$ footprint) they give little information on surface fluxes (Chevallier et al. 2005a, Crevoisier et al. 2009).

Studies have revealed that retrievals of CO$_2$ in the NIR exceed the performance of those in the TIR and provide more lower tropospheric information, provided the atmospheric scatterers are properly constrained (e.g. Christi & Stephens 2004). Furthermore, using the combination of the 1.61 and 2.06 $\mu$m NIR CO$_2$ bands with the 0.76 $\mu$m O$_2$ A band can provide precisions of 0.3-2.5 ppm when the scattering optical depth is less than 0.3 (Kuang et al. 2002). Incorporating the above, the latest generation of satellite instruments, launched in 2009, enable CO$_2$ to be estimated from observed radiances in these three bands. The NASA Orbiting Carbon Observatory (OCO) satellite consisted of a grating spectrometer designed to measure these bands (Connor et al. 2008). Unfortunately, during the launch of the OCO satellite, in February 2009, it failed to reach its desired orbit so no observations were possible. However, the Japanese Greenhouse gases Observing SATellite (GOSAT) was successfully launched into orbit in January 2009 to become the first satellite with the potential to allow CO$_2$ to be inferred within the required precision (JAXA 2009).
2.6 GOSAT

2.6.1 Overview

GOSAT was successfully launched on the 23rd January 2009 by JAXA’s H-IIA rocket and was the first dedicated greenhouse gas satellite (JAXA 2009). Onboard GOSAT is the Thermal And Near-infrared Sensor for carbon Observation (TANSO) that is comprised of two instruments; a Fourier Transform Spectrometer (FTS) that is used to measure greenhouse gases, and a Cloud and Aerosol Imager (CAI) that provides information about the scene, such as cloud and aerosol quantities. Both instruments are kept looking towards the Earth’s geocentric direction by a three-axis attitude control system. The power required for this as well as the sensor operation is gained from two solar panels that face the Sun. The satellite has a Sun synchronous orbit, optimising the power supply, and is quasi-recurrent with 14 + 2/3 revolutions per day (Kuze et al. 2009). GOSAT provides global measurements of total column CO$_2$ from its SWIR bands and of mid-tropospheric sub-columns from its TIR band.

The GOSAT project was developed jointly by the Japan Aerospace Exploration Agency (JAXA), the National Institute for Environmental Studies (NIES) and the Ministry of Environment (MOE). Its primary aim was to estimate emissions and absorptions of greenhouse gases with an increased accuracy on a sub-continental scale and to provide the environmental administration with an assessment of forest carbon balances, regional emissions and absorptions (JAXA et al. 2008). Research using GOSAT may provide an increased understanding of the global distribution and temporal variations of greenhouse gases. This could develop our knowledge of the carbon cycle on a global scale and its influence on the climate, which is essential for the prediction of climate change and its possible impacts. Additionally, it aims to lead to new developments in both Earth observation satellite measurement techniques and the approach of greenhouse gas measurements.

Studies have indicated (e.g. Kadygrov et al. 2009) that the assimilation of GOSAT observations into inverse models may provide an uncertainty reduction greater than that of surface networks alone and a larger improvement when both are assimilated (see Figure 2.11). Furthermore, the regional reduction in uncertainty has been assessed (see Figure 2.12 for example) and has shown large improvements over South America, Africa and Asia (Kadygrov et al. 2009).
Figure 2.11. Comparison of global CO$_2$ flux uncertainties and standard deviation of monthly mean observations of $X_{CO2}$ estimated for the assimilation of existing ground based measurement stations (dashdotted line), GOSAT (solid line), GOSAT and ground based stations (dashed line), and a 50% reduction level of existing total flux uncertainties. Figure taken from Kadygrov et al. (2009).
Figure 2.12. Annual CO$_2$ flux uncertainty reduction of GOSAT observations with 1.8 ppm precision for monthly mean and systematic error of 1 ppm. Figure taken from (Kadygrov et al. 2009).
2.6.2 Instruments

2.6.2.1 TANSO-FTS

TANSO-FTS comprises of three units; an electronic circuit unit, a control unit and an optical unit. The electronic unit is composed of a main and a redundant digital processing circuit that functions as a data processor, data-telemetries-commands-transmission interface with the satellite bus system and includes the temperature controller. The control unit acts as a controller for the cooling machine, the electronic cooling machine, the optical path switcher and solar irradiance diffusing board. It also functions as a driver and controller of the interferometer and the pointing mechanism. The optical unit contains the pointing mechanism (with redundancy), the blackbody diffusing board, the solar irradiance diffusing mechanism, the optical path switcher, monitoring camera, Fourier interferometer, relay and band splitting optics, the detector optics, analogue processing circuit and the cooling machine (Kuze et al. 2009).

The interferometer, based on the original Michelson interferometer (Michelson & Morley 1887), is a device that can split a beam of radiation into two separate paths and then recombine these after a path difference has been introduced. This creates a condition whereby interference between the two beams can occur. The intensity variations of the resulting beam is measured as a function of path difference by the detector.

The interferometer used by TANSO-FTS is of double pendulum type as shown in Figure 2.13. It uses corner cube mirrors which are statically aligned with each other and the rotary motor to maximise modulation efficiency with the minimum optical shear (Kuze et al. 2009). These are attached to a swing arm that is moved by the rotary motor with less than 1% speed stability, creating uniform frequency of the output modulated signal (Kuze et al. 2009). A rotary speed of four seconds is required in order to achieve a small enough sampling of the electrical bandwidth and the necessary signal to noise ratio (SNR). This is used as the nominal time for interferogram acquisition, although TANSO-FTS is also capable of faster scanning modes that allow denser observations but at the expense of SNR (Kuze et al. 2009).

Between the mirrors is a beamsplitter, where a beam of radiation from an external source can be partially reflected to one mirror and partially transmitted to the other mirror. Once reflected from the mirrors both beams return to the beam split and are partially reflected, transmitted and interfere. Due to the effect of interference, the intensity of the beam at the detector depends on the optical path difference (OPD) of the two beams. The OPD created are $\pm \, 2.5 \text{ cm}$, four times as long as the mechanical motion, and are

59
obtained to make both the real and imaginary spectra for suitable phase correction (Kuze et al. 2009).

The width of the beam splitter was selected to be larger than the maximum OPD to try to reduce the effects of channelling, which is caused by multiple reflections at the band separation optics (Kuze et al. 2009). The resulting light is reflected from a collecting mirror towards a circular slit, which due to its physical geometry forces the different spectral bands to be adequately aligned. The circular slit’s geometry was designed to provide the maximum efficiency while maintaining within the maximum IFOV divergence for band 2 for 0.2 cm$^{-1}$ spectral resolution (Kuze et al. 2009). This defines the IFOV to be 15.8 mrad, equivalent to 10.5 km projected onto the surface of the Earth (Kuze et al. 2009). The narrowed light passes to a collimating mirror and is then split through dichroic filters for each band in series. To minimize the background TIR, these filters transmit longer wavelength light and reflect shorter wavelength light. This min-
imises the polarization sensitivity of the shorter wavelengths which then travel though a narrow band-pass filter. These are divided by a beam splitter into two polarizations, parallel (P) and perpendicular (S), that are then measured by two detectors. The band-pass filters are fixed to reduce channelling, to decrease stray light and to discard shorter wavelength light to evade aliasing (indistinguishable signals). While band 1 is measured by two Silicon detectors, both bands 2 and 3 are measured using Indium Gallium Arsenide (InGaAs) detectors that are non-biased and cooled to -40 °C using thermoelectric coolers which minimises dark currents. The TIR light is observed by a Photo-Conductive (PC) Mercury Cadmium Telluride (HgCdTe) (MCT) detector that is cooled to 70 K using a pulse tube cooler with low vibration, which does not influence the interferogram (Kuze et al. 2009).

![Figure 2.14. Example interferogram data. Figure obtained from JAXA et al. (2008).](image)

The interferogram is essentially the interference pattern of the two beams recorded as the OPD changes, and an example of this is shown in Figure 2.14. The measured spectral radiances are obtained by inverse Fourier transforming the interferograms and provide information of absorption in the atmosphere. Figure 2.15 shows example radiances for each of the GOSAT spectral bands.
Figure 2.15. Example TANSO-FTS radiances for each spectral band with the main atmospheric gases that absorb radiation within those bands shown. Figure taken from JAXA et al. (2008).
2.6.2.2 TANSO-CAI

TANSO-CAI consists of an electronics unit and an optics unit with the aim to observe radiances which could allow different aerosol types and optical thicknesses to be derived. The electronics unit functions as a data processor and command interface with the satellite. The optics unit has three telescopes covering four spectral bands; 0.380 ± 0.005 µm (band 1), 0.674 ± 0.005 µm (band 2), 0.870 ± 0.005 µm (bands 3), and 1.600 ± 0.01 µm (band 4). These bands were chosen specifically to avoid water vapour and oxygen absorption, as well as to provide information on polarisation which can not be obtained by the observing angles due to TANSO-CAI being very close to nadir (Kuze et al. 2009). Additionally, each spectral band was selected to provide useful information; band 1 can observe aerosols over land where the albedo is small, bands 2 and 3 are shorter/longer than the red-edge where reflectance of vegetation changes, and band 4 gives measurements in the same spectral region as the CO₂ in TANSO-FTS band 2. TANSO-CAI has a wide swath of ± 36.1° with continuous spatial coverage and higher spatial resolution than TANSO-FTS providing sub-pixel information, such that it can detect clouds and aerosols outside of the TANSO-FTS field of view (Kuze et al. 2009).

2.6.3 Observation Modes

GOSAT TANSO-FTS nominally performs a cross-track scanning pattern (see Figure 2.16) where the number of lattice points can be varied from one to nine points per one cross-track scan, with five points being normal (Kuze et al. 2009). Additionally, it can measure in a specific observation mode which provides targeted observations for validation and experimental purposes. It can also measure over the ocean using the sun glint mode, aiming where the Sun light is reflected off of the waves of the ocean causing the radiance to be very high. However, this can only be measured at certain solar zenith angles which vary with latitude depending on the season. This overcome the difficulty of measuring SWIRs over the ocean that have very low albedo, hence a low radiance and low signal-to-noise ratio (SNR). TANSO-FTS also applies a signal amplification factor (gain) to the SWIR signals to amplify the signal to an appropriate voltage to compensate for high and low intensities. It has the option of three gain levels; low, medium and high. Normally the high gain is used due to most land surfaces having a low albedo, however the medium gain is applied over high albedo regions such as deserts. The fourth instrument channel measures the TIR region that allows vertical profiling over both land and ocean. SWIRs can only be measured in day time since it depends on reflected sun light, but TIR measurements are made throughout both day and night time, as the Earth will ra-
diate TIR from the surface. The instrument uses a solar irradiance (light directly from the Sun) calibration for the SWIR observations and a blackbody calibration for the TIR observations. It also performs deep-space and lunar calibrations for both SWIR and TIR bands (Kuze et al. 2009). Furthermore, the specific observation mode has been utilized for vicarious calibration of both TANSO-FTS and TANSO-CAI by aiming at the Railroad Valley Desert in Nevada, USA (Kuze et al. 2011).

**Figure 2.16.** Diagram illustrating the observation points of TANSO-FTS in 5-point mode for one scene over Australia during April 2009.
2.7 Current Operational Performance

GOSAT has generally performed as expected during its current lifetime, however a couple of issues have arisen. The solar diffuser is used to calibrate the spectral radiances observed by TANSO-FTS with the irradiance from the Sun. Unfortunately, degradation of the solar diffuser has been found, see Figure 2.17. Since all TANSO-FTS spectral radiances rely on the solar diffuser for signal calibration a correction would be required for this. Until recently, a preliminary radiometric factors were provided by NIES to try to compensate for the degradation. However, Yoshida et al. (2012) provides a more accurate method of correction, but this has not been applied to the results shown in Chapter 4.

![Solar Diffuser Degradation](image)

**Figure 2.17.** Solar diffuser degradation factor for each TANSO-FTS spectral band and polarisation.

Secondly, instability has been found in the pointing mirror of TANSO-FTS leading to the location of the observed points being incorrect. Microvibrations in the pointing mirror increase towards larger pointing angles causing the outermost observation points to become most affected with cross-track pointing being up to approximately 1 km incor-
rect in some cases (Kuze et al. 2011). This is further complicated as the pointing mirror also rotates along-track causing the outermost cross-track points to be at a larger angle. Additionally, the pointing mirror reverses in the cross-track direction at the outermost points, causing these points to be increasingly susceptible to unstable movements, with errors in the IFOV location of up to about 10 km (Yoshida 2011). The pointing error was preliminarily analysed by JAXA by comparing onboard camera (CAM) images with TANSO-FTS locations. The amplitude of the offset and the variability was found to be substantially reduced when using three cross-track points, rather than five points, likely due to the pointing mirror being moved by smaller angles. Therefore, after August 2009 the nominal observation mode was changed from five to three points. The pointing error has been investigated by NIES and a separate product (simplified L2 Input Data Sets) has only recently been made available in 2012 which includes corrected geolocation data.

Also, a non-linear response of the band 1 detector to the intensity of incident radiation was found recently (Frankenberg et al. 2011a) and is currently being characterised and calibrated for (Crisp et al. 2012, Suto et al. 2011). Additionally, Frankenberg et al. (2011a) found GOSAT observes fluorescence lines in the O$_2$ A band. The latest TANSO-FTS Level 1B datasets (130.130C, 141.141C, and 150.150C) released by NIES include corrections for the band 1 detector.
2.8 Summary

Although previous satellites have successfully provided global spatial and temporal trends in CO₂, their designs were not suited to the precision required to allow surface flux inversions to resolve sources and sinks of CO₂ on a sub-continental scale. Both OCO and GOSAT were specifically planned to meet this need, and with the successful launch of GOSAT there is currently a satellite that could potentially allow precise CO₂ measurements to be inferred. The next chapter describes a new retrieval algorithm developed to retrieve CO₂ from GOSAT observations.
Chapter 3

Retrieval of Atmospheric CO$_2$ from GOSAT

3.1 Algorithm Overview

The University of Leicester Full Physics (UoL-FP) retrieval algorithm was developed to infer column-averaged dry air mole fractions of CO$_2$ ($X_{CO2}$) from GOSAT SWIR measurements. UoL-FP retrieval utilisés the algorithm developed for the NASA Orbiting Carbon Observation (OCO) mission (Boesch et al. 2006, 2011, Connor et al. 2008, Crisp et al. 2012, Parker et al. 2011), which was lost due to a launch vehicle malfunction (Boesch et al. 2011). The UoL-FP algorithm and the algorithm used for the NASA Atmospheric CO$_2$ Observations from Space (ACOS) project and the NASA OCO-2 mission (Crisp et al. 2012, O’Dell et al. 2012), are two parallel developments based on the OCO algorithm and thus both algorithms follow a similar retrieval strategy. While the UoL-FP algorithm utilises the OCO algorithm, the ACOS algorithm is a re-development of it such that the implementation of both algorithms are independent of each other. Both retrieval algorithms differ in their definition of the state vector, a priori values, and a priori covariances, especially in the treatment of aerosols and cirrus clouds. There are also differences in spectroscopy, sounding selection methods, and post-screening criteria. All of these aspects can lead to differences in algorithm performance and $X_{CO2}$.

The UoL-FP retrieval algorithm comprises of a pre-processing algorithm, a cloud-screen, a post-processing algorithm and the OCO Level 2 Full Physics retrieval algorithm which has been modified to be used with GOSAT observations. This chapter provides details of the UoL-FP retrieval algorithm design and how $X_{CO2}$ can be accurately retrieved from GOSAT SWIR radiances.
3.2 Retrieval Algorithm

The key problem of retrieving CO$_2$ from measured SWIR spectra is that the discrete measurements are a function of many parameters (e.g. CO$_2$, H$_2$O, CH$_4$, albedo, temperature, pressure, and aerosols) that we may or may not accurately know, creating an underconstrained situation with no unique solution. It is therefore necessary to use additional prior information to constrain the solution to be realistic compared to real atmospheric observations. An explanation of the relevant aspects of retrieval theory are given in this section and a detailed description is given in Rodgers (2000). A brief overview of this module, illustrating the main aspects involved, is shown in Figure 3.1.

The UoL-FP algorithm has been designed to accurately retrieve X$_{CO2}$ from SWIR spectra by simultaneously fitting the 0.76 $\mu$m O$_2$ A band, the 1.61 $\mu$m and the 2.06 $\mu$m CO$_2$ bands (Boesch et al. 2006, 2011, Connor et al. 2008). The algorithm employs the inverse method, where an iterative retrieval system based on Bayesian optimal estimation (maximum likelihood estimation) fits the simulated spectral radiance to the measured spectral radiance in order to infer X$_{CO2}$ (Rodgers 2000). The OCO algorithm has been modified to perform retrievals of X$_{CO2}$ from each of the GOSAT SWIR bands using the wavelength ranges of 0.7575-0.7722 $\mu$m, 1.5880-1.6231 $\mu$m, and 2.0399-2.0833 $\mu$m.

The simulated spectral radiance is derived using a forward model that uses solar, radiative transfer, and instrument models to simulate the spectral radiance of a scene based on a set of parameters, known as the state vector, with initial a priori values. The low-streams interpolation functionality (O’Dell 2010) is used to accelerate the Linearized Discrete Ordinate Radiative Transfer model (LIDORT) (Spurr et al. 2001) which is combined with a fast 2-orders-of-scattering vector radiative transfer code (Natraj & Spurr 2007).
Figure 3.1. Overview of the retrieval module used in the UoL-FP retrieval algorithm.
3.3 The State Vector

The state vector gives the retrieved parameters and consists of a 20-level profile of CO$_2$ volume mixing ratio (VMR) and 20-level logarithmic extinction profiles of cirrus and two aerosol types. The number of profile levels chosen provides a compromise between retrieval accuracy and computational speed. In addition, the state vector includes multiplicative scaling factors for CH$_4$ VMR and H$_2$O VMR, an additive offset for a temperature profile, surface pressure, surface albedo, spectral albedo slope, and spectral shift/stretch. An additive O$_2$ A band intensity offset is also retrieved to mitigate the effects of fluorescence and the GOSAT Band 1 non-linear response to the intensity of incident radiation, that is currently being characterized and calibrated (Crisp et al. 2012, Frankenberg et al. 2011a, Suto et al. 2008). The state vector structure presently consists of 97 state vector elements as shown in Table 3.1.

<table>
<thead>
<tr>
<th>Description</th>
<th>Parameters</th>
<th>Number of Elements</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aerosols</td>
<td>3 x 20 levels</td>
<td>60</td>
</tr>
<tr>
<td>CO$_2$</td>
<td>20 levels</td>
<td>20</td>
</tr>
<tr>
<td>Albedo</td>
<td>2 x 3 bands</td>
<td>6</td>
</tr>
<tr>
<td>Dispersion</td>
<td>2 x 3 bands</td>
<td>6</td>
</tr>
<tr>
<td>Zero-Level Offset</td>
<td>Scalar</td>
<td>1</td>
</tr>
<tr>
<td>Surface Pressure</td>
<td>Scalar</td>
<td>1</td>
</tr>
<tr>
<td>Temperature</td>
<td>Scalar</td>
<td>1</td>
</tr>
<tr>
<td>Water Vapour</td>
<td>Scalar</td>
<td>1</td>
</tr>
<tr>
<td>CH$_4$</td>
<td>Scalar</td>
<td>1</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td>97</td>
</tr>
</tbody>
</table>

The UoL-FP retrieval algorithm includes a pre-processing stage that generates all the a priori values needed for retrievals and applies any necessary calibrations to the GOSAT measurements. These a priori values are used firstly in the cloud screen and then in the CO$_2$ retrievals. The cloud screen is based on an O$_2$ A band retrieval of surface pressure and is explained in detail in Section 3.21. The O$_2$ A Band retrieval has a state structure of 5 elements; surface pressure, albedo scaling, albedo slant, spectral shift and spectral stretch. The retrievals for CO$_2$ are then performed on only cloud screened exposures and use a state structure that represents the atmosphere, surface and instrument (see Table 3.1). The results from these CO$_2$ retrievals are gathered together and subjected
to a post-processing stage, which essentially applies a post-screen to the results to only keep retrievals which performed well, see Section 3.22. An overview of the algorithm is given by Figure 3.2, showing the main steps involved to infer $X_{CO_2}$ from GOSAT observations.

**Figure 3.2.** Overview of the UoL-FP retrieval algorithm, providing the flow of information passing between different steps of the algorithm. These steps are grouped into four main modules; the pre-processing, cloud-screen, $CO_2$ retrieval and post-processing modules.
3.4 Forward Model

The forward model simulates spectra based on the state vector, $\mathbf{x}$, that comprises of $n$ parameters, such as temperature, pressure and volume mixing ratios that are retrieved. The measurement vector, $\mathbf{y}$, represents a set of $m$ measurements and can be related to the state vector by

$$\mathbf{y} = \mathbf{F}(\mathbf{x}, \mathbf{b}) + \mathbf{e}$$  \hspace{1cm} (3.1)

where the vector $\mathbf{y}$ is the measurement containing GOSAT radiances with measurement error $\mathbf{e}$, the forward function $\mathbf{F}$ represents the relationship between the measurement vector and state vector, and the forward function parameter, $\mathbf{b}$, represents all non-retrieved parameters that are used in the forward model but are assumed to already be accurately known.

The forward model consists of three major modules; a radiative transfer (RT) model, a solar model, and an instrument mode. The solar model described in Boesch et al. (2011) was used, which provides a high-resolution empirical list of absorption lines for the full solar disk. It includes over 20,000 absorption lines with their corresponding frequency, strength and doppler width. The solar spectrum is obtained by calculating the Planck function for the Sun and multiplying with the listed solar absorption lines. The continuum radiance of the solar model is based on a polynomial fit to the NIR range of the extra-terrestrial solar spectrum acquired by the SOLSPEC instrument (Thuillier et al. 2003). The solar model is assumed time independent but allows the continuum radiance to be adjusted by the Earth-Sun distance. Figure 3.3 shows the solar continuum with solar absorption lines for each SWIR band retrieved.

![Figure 3.3](image)

**Figure 3.3.** The solar continuum and solar absorption lines of each SWIR band used in the UoL-FP retrieval algorithm.

The RT model calculates the transfer of radiation throughout the beam path based on a priori information for atmospheric and surface parameters. It uses the LIDORT model which simultanously derives backscatter intensities for a plane parallel, multi-layered,
multiple-scattering atmosphere at arbitrary elevation angles and a set of atmospheric variations. The computational speed of the RT model was increased by using two orders of scattering and the low-streams interpolation functionality (O’Dell 2010). This computes low accuracy line-by-line RT calculations which are corrected by an error curve derived from the computation of both low and high accuracy RT calculations for a number of wavelengths throughout the bands.

For absorption within the Earth’s atmosphere, the RT model utilises tabulated spectroscopic parameters for each trace gas. CO\textsubscript{2} and O\textsubscript{2} include line-mixing and are taken from v3.2 of the OCO linelists (Crisp et al. 2012), whereas CH\textsubscript{4} and H\textsubscript{2}O are from the TCCON linelists (Wunch et al. 2011a). These are based on HITRAN08 (Rothman et al. 2009) with updates to CO\textsubscript{2} (Toth 2005), H\textsubscript{2}O (Toth et al. 2008, Jenouvrier et al. 2007) and CH\textsubscript{4} (Frankenberg et al. 2011b). The absorption lines and strengths are applied to the solar spectrum using Voigt line shapes.

The instrument model is used to generate noise and intensity and different wavelengths with respect to the instrument line shape (further discussed in Section 3.19), using a priori information of the instrument parameters.
3.5 Jacobian

Measurements are typically non-linear, so linearisation of the forward model about some reference state, \( x_0 \), is required to be able to examine the information content within a measurement. Providing that the forward model is linear within the error bounds of the retrieval, Equation 3.1 can be linearised to get

\[
y - F(x_0) = \frac{\partial F(x)}{\partial x} (x - x_0) + e
\]

(3.2)

where the partial derivative \( \frac{\partial F}{\partial x} \) denotes the sensitivity of the Forward model to the change in the state vector, \( x \). This quantity is known as the Jacobian, \( K \), and is a matrix consisting of \( m \) forward model elements (number of measurements) with respect to \( n \) state vector elements (number of parameters retrieved). Thus if there are more elements in the measurement vector than unknowns in the state vector (\( m > n \)) then the retrieval can be described as over-constrained. Similarly if there are less elements in the measurement vector than unknowns in the state vector (\( m < n \)) then the retrieval is under-constrained. The state vector is described in detail in Section 3.3.

An example of the partial derivatives of each state vector element is given in Figures 3.4, 3.5 and 3.6, which were calculated during a CO\(_2\) retrieval from GOSAT for the 4th September 2009 over Lamont/USA. Figure 3.4 gives the Jacobians of CO\(_2\), aerosols and cirrus for each atmospheric retrieval level and wavelength retrieved. The Jacobian for CO\(_2\) is large in the 1.61 and 2.06 \( \mu \)m bands, specifically where CO\(_2\) absorption lines occur, and zero in the O\(_2\) A band where no CO\(_2\) absorption occurs. Additionally, the Jacobian for CO\(_2\) remains large throughout the atmospheric profile. The aerosol/cirrus Jacobians show some sensitivity to all absorption lines (O\(_2\), H\(_2\)O, CO\(_2\), etc) in all three wavelength bands. The aerosol Jacobian is larger towards lower atmospheric levels, whilst the cirrus Jacobian is only sensitive at about 200 hPa, reflecting the optical depth at each level.

Figure 3.5 shows the Jacobians of surface pressure, temperature, H\(_2\)O, CH\(_4\) and O\(_2\) A band zero level intensity offset for the same example retrieval. The Jacobian of surface pressure shows sensitivity to absorption lines, whilst the temperature scalar Jacobian appears inversely proportional to all absorption lines. The Jacobian of the H\(_2\)O scale factor relates to water vapour absorption lines, thus there is no sensitivity in the O\(_2\) A band. Similarly, there are no CH\(_4\) absorption lines in the O\(_2\) A band, so the Jacobian of the CH\(_4\) scale factor only corresponds to CH\(_4\) absorption lines in the 1.61 and 2.06 \( \mu \)m bands. The zero level offset Jacobian only relates to the O\(_2\) A band where this intensity
offset is retrieved.

The albedo, albedo slope and dispersion shift/stretch Jacobians for the example retrieval are shown in Figure 3.6. The albedo Jacobian is zero for all bands, showing no sensitivity of the forward model to a change in albedo, since albedo acts to change the overall intensity of each band and is not related to absorption. The albedo slope changes the intensity and absorption across each band, so the Jacobian therefore shows sensitivity to all absorption lines and varies in sensitivity across each band. The dispersion shift represents the wavelength shift needed for the Forward model to match the measured spectra. Since the pre-processing module adjusts the wavenumbers of the GOSAT measured spectra to match a well-known Solar absorption line, the dispersion shift will not change much during a retrieval. Thus, the Jacobian of the dispersion shift is zero in all bands. However, the dispersion stretch is also retrieved and the Forward model will be sensitive to any variations of this as it can cause corresponding modelled and measured absorption lines to differ in wavelength, as illustrated by the Jacobian.
FIGURE 3.4. Example of the Jacobians of CO$_2$, aerosols and cirrus profiles for each SWIR band retrieved, with the measured signal from GOSAT given in the top panels for reference. The profile levels are shown using different colours, where red represents the boundary layer and black represents the top of the atmosphere. This retrieval was from GOSAT SWIR radiances for the 4th September 2009 over Lamont/USA and passed all screens.
Figure 3.5. Example of the Jacobians of surface pressure, temperature, $\text{H}_2\text{O}$, $\text{CH}_4$ and $\text{O}_2$ A band zero level intensity offset. This retrieval was from GOSAT SWIR radiances for the 4th September 2009 over Lamont/USA.
Figure 3.6. Example of the Jacobians of albedo, albedo slope and dispersion shift/stretch. This retrieval was from GOSAT SWIR radiances for the 4th September 2009 over Lamont/USA.
3.6 Inverse Method

In an under-constrained retrieval scenario there are less measurements than unknown state vector elements. This can be constrained using a priori information.

In an over-constrained retrieval case there are more measurements than unknown state vector elements, but even in an ideal situation the extra measurement information does not provide any new information to the retrieval. It is more likely in reality that there would be no solution that agrees with all of the measurements, since each will have an error associated with it. Whether under or over-constrained, an optimal estimation approach may be used to find the best solution.

3.7 Optimal Estimation

Optimal estimation uses an a priori estimate of the state vector to constrain the retrieval to find reasonable solutions, which might not be possible from only the measurements. Since this does not obtain a unique solution, the optimal solution can be gained by two possibilities; the most probable state where \( \frac{dP(x,y)}{dx} = 0 \) or the expected state, \( \hat{x} \), which is the mean state averaged over its probability density function (PDF)

\[
\hat{x} = \int xP(x \mid y) \, dx \quad (3.3)
\]

Assuming the probability fits a Gaussian model, hence the PDF is symmetric about its maximum, the most probable state equals the expected state.

The most probable state is determined by minimising the cost function. The cost function represents the different minimum costs yielded by optimising the difference between the measured and modelled spectra as well as the difference between the state vector and the a priori. The cost function can be written as

\[
X^2 = (y - Kx)^T S_y^{-1}(y - Kx) + (x - x_a)^T S_a^{-1}(x - x_a) \quad (3.4)
\]

where \( x_a \) is the a priori estimate of the state vector. The most probable state can then be expressed as

\[
\hat{x} = x_a + (K^T S_y^{-1}K + S_a^{-1})^{-1}K^T S_y^{-1}(y - Kx_a) \quad (3.5)
\]
3.8 Iterative Approach

The most probable state found might be far from the true solution. To reach a state closer to the truth an iterative approach can be used, with each iteration re-running the forward model but with priori information gained from the previous iteration. This priori information is most likely to be closer to the true solution and would result in the forward model producing a state closer to the truth. However, that may not always be the case. Therefore, to attempt to obtain a solution closer to the truth the Gauss-Newton scheme (Rodgers 2000) is used to force the state to significantly change to find a state closer to the truth. This iterative approach minimises the cost function, such that for each iteration step the forward model is used to re-calculate the Jacobians and the modelled spectra. Assuming the retrieval is not too non-linear the Gauss-Newton iteration solution can be given as

\[ x_{i+1} = x_i + \left[ S^{-1} + K_i^T S e^{-1} K_i \right]^{-1} \left[ K_i^T S e^{-1} (y - F(x_i)) - S^{-1} (x_i - x_a) \right] \] (3.6)

where the first iteration, \( x_0 \), is set to the a priori state vector, \( x_a \). However, if the solution is too far from the true solution or if the retrieval is too non-linear then the Gauss-Newton method becomes inadequate. In this scenario the Levenberg-Marquardt method (Rodgers 2000) is included whereby an adjustable factor, \( \gamma \), is used to minimise the cost function. The iteration is then

\[ x_{i+1} = x_i + \left[ (1 + \gamma) S^{-1} + K_i^T S e^{-1} K_i \right]^{-1} \left[ K_i^T S e^{-1} (y - F(x_i)) - S^{-1} (x_i - x_a) \right] \] (3.7)

After each iteration \( \gamma \) is either increased or decreased by an arbitrary factor depending on the cost function. If the cost function increases then \( \gamma \) is increased and the same iteration is re-attempted. Whereas if the cost function decreases then \( \gamma \) is decreased, the state vector is updated, and the next iteration is performed. This process continues until the retrieval converges.

3.9 Convergence

At the end of an iteration three tests are performed. First, the updated cost function plus the previous cost function is compared to the cost function forecast plus the previous cost function to give a ratio, as described in Section 3.8. If this ratio is \(< 0.25\) then the model has diverged away from the measurement and the iteration is repeated with an increased
\( \gamma \) factor. If the ratio is between 0.25 and 0.75 then the model has become closer to the measurement but not close enough so another iteration is performed. If the ratio is > 0.75 then the model and measurement are said to have converged (Rodgers 2000). The second test of convergence compares the error variance derivative \((d\sigma^2)\) with the \(d\sigma^2\) of the previous iteration. If the difference is less than or equal to the number of state vector elements then convergence is achieved, otherwise iterations are continued. The error variance derivative is expressed as

\[
d\sigma_i^2 = d\mathbf{x}_{i+1}^T \hat{\mathbf{S}}^{-1} d\mathbf{x}_{i+1}
\]

(3.8)

where the covariance of the retrieved state is given by \(\hat{\mathbf{S}}\). Both of these tests must be passed for the retrieval to actually converge. The third test computes the \(\chi^2\) of the fit to check quantitively how good the model compares to the measurement. If the normalised \(\chi^2\) is less than 1 then it is a good fit, between 1 and 10 is a reasonable fit, and greater than 10 is a poor fit (Rodgers 2000). However, the normalised \(\chi^2\) is also dependent on the assumptions of noise.

Upon convergence the retrieved \(X_{CO2}\) can be resolved as

\[
X_{CO2} = \mathbf{h}^T \hat{\mathbf{x}}
\]

(3.9)

where the pressure weighting operator, \(\mathbf{h}\), has elements of zero for all non-CO\(_2\) elements.

The averaging kernel expresses the ability of the retrieval to constrain the true state, hence how sensitive the algorithm is to the truth throughout the atmosphere, and is defined as

\[
\mathbf{A} = \frac{d\hat{\mathbf{x}}}{d\mathbf{x}} = \mathbf{SK}^T \mathbf{S}_e \mathbf{K}
\]

(3.10)

where \(\mathbf{S}\) is the a posteriori error covariance, which provides a measure of the estimated accuracy of the solution and is given by

\[
\mathbf{S} = (\mathbf{K}^T \mathbf{S}_e \mathbf{K} + \mathbf{S}_a^{-1})^{-1}
\]

(3.11)

The column averaging kernel can then be calculated as

\[
\mathbf{a}^T = \mathbf{h}^T \mathbf{A}
\]

(3.12)

where an averaging kernel with a value of 1 is good and 0 is bad. Additionally, the \(X_{CO2}\) error variance is computed by

\[
\sigma^2_{X_{CO2}} = \mathbf{h}^T \mathbf{S} \mathbf{h}
\]

(3.13)
3.10 Covariance Matrix

The covariance matrix defines the correlation and constraint of a state vector element within the retrieval scheme. $S_y$ is the measurement error covariance matrix where the diagonal elements are the variances of the individual elements of $y$ and the off-diagonal elements give the covariance between the different elements of $y$. Similarly the covariance of the a priori data can be given by the covariance matrix $S_a$.

For all retrievals a single a priori covariance matrix has been used, with a column variability of 12 ppm for CO$_2$ (based on global estimates of Dufour & Breon (2003)) which decreases with altitude, by $\sim$10% at the boundary layer to $\sim$1% in the stratosphere (see Figure 3.7). For surface pressure a standard deviation of 4 hPa was used to allow for more difficult topographies. To account sufficiently for the expected large variability of aerosols and cirrus clouds, the profiles were given a large flexibility with a covariance of 50 times the 1-$\sigma$ uncertainty for each atmospheric level (see Figure 3.8).

![Figure 3.7. CO$_2$ covariance used for all retrievals of $X_{CO2}$. Zero represents the top of the atmosphere and 20 represents the boundary layer.](image-url)
Figure 3.8. Aerosol/cirrus covariance of 50 times the 1-σ uncertainty. Zero represents the top of the atmosphere and 20 represents the boundary layer.

3.11 Theoretical retrieval assessment of GOSAT and OCO

To assess the potential of retrieving $X_{CO2}$ from GOSAT SWIR bands, scenes were simulated and retrieved for different surface types and SZAs using the OCO Level 2 retrieval algorithm (Boesch et al. 2006), with the instrument model based on pre-flight information of GOSAT. A non-apodized sinc-function was used to simulate the ILS for an OPD of 2.5 cm and field of view (FOV) of 15.8 mrad. This relates to a full-width-half-maximum (FWHM) of 0.35 cm$^{-1}$ for band 1 and 0.27 cm$^{-1}$ for bands 2 and 3 of TANSO-FTS. The noise has been calculated for each band by using the SNR values given in Suto et al. (2008), for an albedo of 0.3 and SZA of 30°.

Figures 3.9 and 3.10 show the $X_{CO2}$ averaging kernels and retrieval errors, respectively, for different simulated scenarios. $X_{CO2}$ error gives the expected error of $X_{CO2}$ retrievals, which is found to be less than $\sim$1.5 ppm over all surfaces, except over snow and ocean in nadir where the error is in the order of 1-2%. However, GOSAT measures
over the ocean with the Sun-glint mode where the errors are low. Therefore, it is expected that GOSAT will not provide useful measurements of $X_{CO_2}$ over areas of inland water where nadir mode is used, and over regions with snow cover (such as mountains, poles, and seasonally high latitudes). The averaging kernels describe the sensitivity of the instrument and retrieval algorithm to retrieve $X_{CO_2}$ at different altitudes. The averaging kernels peak near the surface and remain high throughout the troposphere, hence the $X_{CO_2}$ near the boundary layer is retrieved which will provide important insight into the source and sinks of CO$_2$ over land.

Similarly, Figures 3.11 and 3.12 show simulated averaging kernels and expected $X_{CO_2}$ error for the OCO instrument for a variety of surface types. The averaging kernels of OCO are very similar to that of GOSAT, however OCO is more sensitive at higher SZAs over snow surfaces than GOSAT. The $X_{CO_2}$ error of GOSAT is in general higher than OCO (e.g. $\sim$0.5 ppm larger for conifer/vegetation), although ocean (Sun-glint) and desert surfaces show similar $X_{CO_2}$ errors as OCO. Also, GOSAT shows much higher $X_{CO_2}$ errors ($\sim$2 ppm larger) over snow and ocean (nadir) surfaces than OCO. Both instruments show that the $X_{CO_2}$ error depends on the surface type and increases with SZA and AOD.

\textbf{Figure 3.9.} Calculated averaging kernels (AK) for simulated scenes of different surface types and SZAs, all with a 0.1 total column optical depth and instrument model based on GOSAT pre-flight characteristics.
FIGURE 3.10. Calculated $X_{CO2}$ error for simulated scenes of different surface types, SZA, and total column optical depth. All simulations used an instrument model based on GOSAT pre-flight characteristics.
Figure 3.11. Calculated averaging kernels (AK) for simulated scenes of different surface types and solar zenith angle, all with a 0.1 total column optical depth and instrument model based on OCO pre-flight characteristics (courtesy of H. Boesch, 2009).
Figure 3.12. Calculated $X_{CO_2}$ error for simulated scenes of different surface types, solar zenith angle, and total column optical depth. All simulations used an instrument model based on OCO pre-flight characteristics (courtesy of H. Boesch, 2009).
3.12 Testing the spectral window setup

There are four CO$_2$ absorption bands within the SWIR spectral range of TANSO-FTS (see Figure 3.13); the 1.58 µm CO$_2$ band (1.558 - 1.592 µm range), the 1.61 µm CO$_2$ band (1.588 - 1.623 µm range), the 2.01 µm CO$_2$ band (1.980 - 2.024 µm range), and the 2.06 µm CO$_2$ band (2.040 - 2.083 µm range). Thus, providing a choice of wavelength ranges to retrieve CO$_2$ from. Additionally, there are two polarised signals available; parallel (P) and perpendicular (S) to the incident plane. Each of these absorption bands and polarisations may result in a different retrieved $X_{CO2}$ value, due to instrument and spectroscopy variations. To test for any differences and to determine which bands or polarisations are most suitable to use, each of these bands and polarisations were individually used to retrieve $X_{CO2}$.

![Figure 3.13](image.png)

**Figure 3.13.** The first TANSO-FTS band 2 and 3 radiances obtained, illustrating the CO$_2$ absorption regions within these bands. Figure taken from JAXA (2009).

The measured radiances were calibrated with the initial radiometric conversion coefficients and the retrievals used the official GOSAT ILS from JAXA (version 006) (Kuze et al. 2009) and tabulated absorption cross-sections of HITRAN08 (Rothman et al. 2009). The a priori were obtained from the pre-processing module. A single Gaussian profile of 2 km height and width with an AOD of 0.075 and scattering optical properties of Kahn type 4b (carbonaceous and dusty environments) were used for the a priori aerosol (Kahn et al. 2001). The state vector included a 20 level profile of CO$_2$, a scale factor for H$_2$O, albedo, albedo slope, and dispersion shift and stretch.
Figures 3.14 and 3.15 show the retrieved $X_{CO_2}$ for each polarisation and CO$_2$ absorption band for Park Falls/USA and Darwin/Australia, providing northern and southern hemisphere scenes. The $X_{CO_2}$ values retrieved range between $\sim$325 ppm and $\sim$410 ppm, with the 2.01 $\mu$m $X_{CO_2}$ being generally $\sim$20 ppm higher than that of the other CO$_2$ bands. The 3rd exposure of Park Falls shows a different distribution of $X_{CO_2}$ than the other two exposures. Using the official NIES TANSO-CAI L1B false RGB image of this scene, this exposure appears to coincide with a cloud, whereas the other two exposures appear cloud-free. Similarly, the 3rd and 8th exposures of Darwin show a different distribution of $X_{CO_2}$ than the other two exposures and also coincide with clouds. This highlights are few issues; some clouds remain even after the NIES cloud-screen has been applied, the retrieved $X_{CO_2}$ of a cloudy exposure can be biased high or low, and the amount that the $X_{CO_2}$ is biased for a cloudy exposure is different for each CO$_2$ absorption band and polarisation. Furthermore, the retrieved $X_{CO_2}$ is similar ($\sim$0.5 to 2 ppm different) for each polarisation, but for cloudy exposures there is a large difference between polarisations (up to $\sim$25 ppm). If the cloudy exposures and 2.01 $\mu$m band $X_{CO_2}$ retrievals are neglected, then all the other $X_{CO_2}$ values retrieved are roughly consistent (within $\sim$10 ppm of each other) and have a mean value of $\sim$389 ppm for Park Falls and $\sim$385 ppm for Darwin, which are very similar to the global levels of CO$_2$ observed by NOAA at Mauna Loa (Masarie & Tans 1995).

These preliminary results were all retrieved using a single spectral window, such as the 1.61 $\mu$m CO$_2$ band (1.588 - 1.623 $\mu$m range). However, Kuang et al. (2002) found that using a combination of the 0.76 $\mu$m O$_2$ A band, 1.58 $\mu$m and 2.06 $\mu$m CO$_2$ bands provided improved precisions in simulations.

O’Brien et al. (1998) demonstrated that the 0.76 $\mu$m O$_2$ A band can provide constraints on surface pressure, which can be uncertain due to topographic and local weather variations. Additionally, it can provide constraints on optical path length variations, where uncertainties due to cloud and aerosol scattering can exist. Whilst the 1.58 $\mu$m and 1.61 $\mu$m CO$_2$ bands are almost free of other absorbing atmospheric gases, variations in water vapour produce uncertainties in the retrieved $X_{CO_2}$ by broadening the CO$_2$ absorption lines slightly and effecting the dry-air fraction when calculating the $X_{CO_2}$ (Kuang et al. 2002). However, the 2.01 $\mu$m and 2.06 $\mu$m CO$_2$ bands contain weak water vapour absorption lines that can provide a constraint on the water vapour. The 2.01 $\mu$m and 2.06 $\mu$m CO$_2$ bands contain CO$_2$ absorption lines that are strong enough to be sensitive to scattering, so provide additional constraints on optical path length uncertainties from clouds and aerosols (Kuang et al. 2002). However, the 2.01 $\mu$m CO$_2$ band contains lots of water vapour absorption that may cause intereference. Furthermore, the 1.58 $\mu$m
Figure 3.14. X\textsubscript{CO\textsubscript{2}} retrieved from GOSAT between 23rd - 30th April 2009 within 500 km of the Park Falls/USA TCCON site. Each retrieval was designed to retrieve a single polarisation signal (P or S) and a single CO\textsubscript{2} absorption band (shown by different colours). The index represents different TANSO-FTS exposures.

CO\textsubscript{2} band has O\textsubscript{2} day glow emissions that may potentially perturb measurements in this band (Boesch et al. 2006). O\textsubscript{2} day glow emission is known to occur in the mesosphere and thermosphere due to three mechanisms; the excitation of O\textsubscript{2} from a collision with an O(\textsuperscript{1}D) atom produced in the photolysis of O\textsubscript{3} or O\textsubscript{3}, resonant absorption of oxygen bands (including the O\textsubscript{2} A band), or the chemical reacton between two oxygen atoms and either a N\textsubscript{2} or O\textsubscript{2} molecule (Sheese et al. 2010).

These preliminary results showed the X\textsubscript{CO\textsubscript{2}} retrieved from GOSAT to be similar between the different CO\textsubscript{2} bands, with the exception of the 2.01 \textmu m CO\textsubscript{2} band which gave much higher values (due to poor spectroscopy). Thus, to reduce spectroscopic biases the 2.01 \textmu m CO\textsubscript{2} band should not be used. Additionally, the 1.61 \textmu m CO\textsubscript{2} band gave X\textsubscript{CO\textsubscript{2}} values closer to the 2.06 \textmu m CO\textsubscript{2} band than that of the 1.58 \textmu m CO\textsubscript{2} band, and
$X_{CO2}$ retrieved from GOSAT between 23rd - 30th April 2009 within 500 km of the Darwin/Australia TCCON site. Each retrieval was designed to retrieve a single polarisation signal (P or S) and a single CO$_2$ absorption band (shown by different colours). The index represents different TANSO-FTS exposures. Since the 1.58 µm CO$_2$ band contains day glow it is not appropriate to use. Therefore, the combination of the 0.76 µm O$_2$ A band, 1.61 µm and 2.06 µm CO$_2$ bands was tested in the UoL-FP retrieval algorithm. The O$_2$ A band provides information on surface pressure and aerosols, which is highly beneficial to help constrain the retrieval of $X_{CO2}$.

$X_{CO2}$ was retrieved from GOSAT using different spectral bands in the UoL-FP retrieval algorithm for observations within ±5 degrees over the Lamont/USA TCCON site between April and September 2009. The exposures were pre-screened for a SNR > 50 and the NIES cloud-screen was used (Taylor et al. 2012). Figure 3.16 shows the difference in retrieved $X_{CO2}$ from the 1.61 µm CO$_2$ band with the retrieval setup to use either; the 0.76 µm O$_2$ A band with the 1.61 µm CO$_2$ band, the 2.06 µm CO$_2$ band, or all three bands together. On average, there is little difference in $X_{CO2}$ when including the O$_2$ A
band, but the scatter in $X_{CO_2}$ is of the order of $\sim$5 ppm. However, since the surface pressure was fixed to the a priori assumption any constraints provided by the inclusion of the $O_2$ A band would not effect the surface pressure. The $X_{CO_2}$ retrieved from the 2.06 $\mu$m CO$_2$ band is higher on average but the $X_{CO_2}$ values lower than about 370 ppm are much lower than that retrieved using the 1.61 $\mu$m CO$_2$ band. These low values are potentially influenced by aerosols which the 2.06 $\mu$m CO$_2$ band is sensitive to, due to aerosol optical properties have more effect in this wavelength range. When all three bands are retrieved simultaneously, it produces $X_{CO_2}$ that appears an amalgam of the effects already found. Although the precision of the data changes little between the different setups, it allows for the possibility of additional constraints and potentialy useful additional parameters to be retrieved such as surface pressure and clouds/aerosols.
Figure 3.16. $X_{CO_2}$ retrieved from GOSAT using different spectral bands in the UoL-FP retrieval algorithm for observations within $\pm 5^\circ$ of the Lamont/USA TC-CON site between April and September 2009. The data was pre-screened and cloud screened but no post-screening was applied. The top left panel shows the $X_{CO_2}$ retrieved using the $1.61 \mu m$ CO$_2$ band compared with that retrieved when using the $1.61 \mu m$ CO$_2$ band and the $0.76 \mu m$ O$_2$ A band. The top right panel shows a comparison of the $X_{CO_2}$ retrieved from the $1.61 \mu m$ and $2.06 \mu m$ CO$_2$ bands separately. The bottom panel shows the $X_{CO_2}$ retrieved using the $1.61 \mu m$ CO$_2$ band compared with that retrieved when using the $0.76 \mu m$ O$_2$ A band, $1.61 \mu m$ and $2.06 \mu m$ CO$_2$ bands.
3.13 Profile of Carbon Dioxide Concentration

The a priori CO$_2$ profiles were obtained from 3 hourly model 3.75° longitude by 2.5° latitude gridded 2009 fields of the Laboratory of Climate Sciences and Environment (LCSE) General Circulation Model of Laboratoire de Meteorologie Dynamique (LMDZ) (Pickett-Heaps et al. 2011). A correction of 18.25 ppm was applied to the 2009 CO$_2$ profiles that was derived from a comparison with National Oceanic and Atmospheric Administration (NOAA) observations over Mauna Loa, see Figure 3.17. Additionally, yearly increments of 1.63 ppm and 2.36 ppm were taken from global NOAA observations (Masarie & Tans 1995) and applied to the CO$_2$ profiles for 2010 and 2011 respectively, to account for increasing atmospheric CO$_2$ concentrations (see Figure 3.18).

![Figure 3.17](image)

Figure 3.17. $X_{CO2}$ calculated for LMDZ 2009 model CO$_2$ fields interpolated to the time and location of CO$_2$ observations of Mauna Loa/Hawaii, compared to $X_{CO2}$ calculated for Mauna Loa observations (left). LMDZ $X_{CO2}$ was found to be 18.25 ppm lower than Mauna Loa on average. After applying this offset to the LMDZ $X_{CO2}$ (right) a difference is still observed, with LMDZ estimating a smaller range of $X_{CO2}$ than observed by Mauna Loa.

However, the Mauna Loa comparison is only based on a small sample size, therefore the offset LMDZ $X_{CO2}$ was compared to $X_{CO2}$ retrieved from different TCCON sites (see Figure 3.19), including; Lamont/USA, Park Falls/USA, Orleans/France, Bremen/Germany, Garmisch/Germany, Bialystok/Poland, Tsukuba/Japan, Darwin/Australia, Wollongong/Australia, and Lauder/New Zealand. This analysis provides comparisons of $X_{CO2}$ for both northern and southern hemispheres at various locations, and shows the calculated offset to give LMDZ $X_{CO2}$ of similar value as TCCON for all locations. The LMDZ $X_{CO2}$ estimated for northern hemisphere sites shows a good comparison to TCCON with correlation coefficients ranging from 0.86 to 0.98. However, the LMDZ $X_{CO2}$
FIGURE 3.18. Mean yearly CO$_2$ from NOAA in-situ observations (top) and the yearly increment in CO$_2$ (bottom).

is found to estimate a lower range of X$_{CO2}$ values than observed by TCCON, as previously found with the Mauna Loa comparison. For the southern hemisphere sites, LMDZ poorly reproduces the X$_{CO2}$ retrieved from TCCON with correlation coefficients ranging from 0.48 to 0.64. Again, the offset applied to LMDZ provides X$_{CO2}$ values of a similar mean value as TCCON for southern hemisphere sites. Thus, applying this offset to the model CO$_2$ allows the retrievals to use a CO$_2$ a priori profile with values that are on average closer to the truth.
Figure 3.19. $X_{CO2}$ calculated for LMDZ 2009 model CO$_2$ fields interpolated to the time and location of TCCON observations for nine different sites, compared to the retrieved $X_{CO2}$ from TCCON. The one-to-one line is shown in black and each observation is shown by a black diamond. The correlation coefficient per site is given by $r$. 
3.14 Methane Concentration

The a priori profile of CH$_4$ is acquired from a TM3 model run (Houweling 2009) and is interpolated to the desired latitude, longitude and time. This is only retrieved as a scaling of the profile, assuming the profile shape to be true. The retrieval uses a standard deviation of 0.32 for the CH$_4$ scale factor.

3.15 Surface Pressure

The European Centre for Medium-Range Weather Forecasts (ECMWF) provides model data that includes assimilated observations from surface buoy and satellite measurements (ECMWF 2008). ECMWF provides atmospheric profiles of pressure, temperature and specific humidity on a 1.125 degree by 1.125 degree global grid with 90 layers every 6 hours. Given the latitude, longitude and altitude of a site of interest the surface pressure can be determined from these profiles. All 90 layer datasets can be converted onto 91 levels using coefficients provided by ECMWF (ECMWF 2007). ECMWF also provides potential data, $\phi$, for the lowest level of the same grid, which can be used to find the geopotential height, $Z_g$, of each grid point level as

$$Z_g = \frac{\phi}{g}$$

where gravity, $g$, is calculated as a function of latitude and approximate altitude. Taking the four surrounding grid points of the site of interest, the pressure, $P$, at the site altitude can be found for each grid point by using the hydrostatic equation

$$P = P_0 e^{-\frac{Z}{Z_0}}$$

where $P_0$ is the pressure of the grid point level lower than the site altitude, $Z$ is the difference in altitude between the grid point level and the site altitude and $Z_0$ is the scale height defined as

$$Z_0 = \frac{RT}{Mg}$$

where $R$ is the ideal gas constant, $T$ is the average temperature across the differential, $M$ is the Molar mass of wet air and $g$ is the gravitational acceleration as a function of latitude and altitude. The Molar mass of wet air can be calculated by

$$M = \rho_d(1 - SH) + \rho_wSH$$
where SH is the ECMWF specific humidity, $\rho_d$ is the dry air mass and $\rho_w$ is the mass of wet air. The site altitude can be obtained from a global digital elevation model with a horizontal grid spacing of 30 arc seconds named GTOPO30 provided by the U.S. Geological Survey (USGS 1996). Note, that in the case that the site altitude is lower than the lowest level of a grid point, the pressure is calculated with respect to the lowest level. Also, the temperature and molar mass are extrapolated downwards based on the lapse rate and gradient of the 5 lowest levels above, respectively. The surface pressure for the site can then be resolved by interpolating the pressures with latitude, longitude and time.

To validate the surface pressure two sites were chosen; Southern Great Plains (SGP) /USA, and Black Forest/Germany. Since pressure changes with altitude, highly variable topographies will also vary in pressure. These sites are flat and mountainous in topography respectively, providing an assessment of the extremes in topography where data from observations are available. The Atmospheric Radiation Measurement (ARM) program (Ackerman & Stokes 2003) has a central facility in SGP which takes in situ ground based measurements of surface pressure using the Temperature, Humidity, Wind and Pressure Sensors (THWAPS) (Peppler et al. 2008), which has a precision of 0.1 hPa and an uncertainty of 0.7 hPa (ARM 2011a). Figure 3.20 shows that the surface pressure is within 1 hPa of THWAPS observations. Surface Pressure observations were also made by the ARM mobile facility that visited Black Forest, Germany, in 2007 with the Surface Meteorological Instrument (MET) (Peppler et al. 2008), which has a precision of 0.1 hPa and an uncertainty of 0.7 hPa (ARM 2011b). Figure 3.20 illustrates that over a mountainous topography the surface pressure can still be resolved to within 1 hPa. Hence, the surface pressure a priori could be constrained to less than 0.1%. In the retrieval a standard deviation of 4 hPa is used to allow for more difficult topographies.

Initially, the a priori surface pressure was calculated for the centre of the TANSO-FTS IFOV location. However, regions of large topographic variations may cause this surface pressure to be unrealistic for the entire IFOV. Therefore, a new method was created that calculates the surface pressure for all known altitudes within the IFOV. The altitude information is obtained from GTOPO30 (USGS 1996) which has a 30-arc second resolution (approximately 1 km), allowing roughly 70 surface pressures to be calculated within the IFOV (∼10.5 km diameter) from which the mean and standard deviation can be computed.

The a priori surface pressure was calculated using each method for GOSAT observations within ±5° of the Lamont TCCON site between April and December 2009. Figure 3.21 gives a comparison of these surface pressures and shows differences up to ∼3.5 hPa, which is within the assumed a priori surface pressure 1-sigma uncertainty used
In the retrievals.

On average, a bias in retrieved surface pressure of $\sim 4$ hPa is observed (see Figure 3.22), which is lower than reported by Crisp et al. (2012). This bias has been suggested to be due to deficiencies in the O$_2$ A band spectroscopy and that scaling the line strength of O$_2$ could reduce it (Butz et al. 2011). For this retrieval, instead of scaling the line strength, the systematic overestimates of surface pressure are corrected by normalizing the retrieved X$_{CO_2}$ with the ratio of retrieved surface pressure and ECMWF surface pressure.

The effect of normalising the retrieved X$_{CO_2}$ with the surface pressure ratio was tested for retrievals from GOSAT. Figure 3.23 shows a one year comparison of the retrieved X$_{CO_2}$ globally with and without the surface pressure normalisation applied. In general, the X$_{CO_2}$ becomes higher in value when the surface pressure normalisation is applied, thus reducing the X$_{CO_2}$ bias compared to TCCON.
Figure 3.21. Comparison of surface pressure calculated for the GOSAT IFOV centre and for the mean across the IFOV, for GOSAT observations within ±5° of the Lamont TCCON site. All surface pressures were calculated from ECWMF and interpolated to the time and location of the observation.
FIGURE 3.22. Histogram of the difference between the retrieved surface pressure from GOSAT observations between July 2009 and June 2010 globally, and ECMWF surface pressure calculated for the times and locations of the GOSAT observations.
Figure 3.23. Comparison of $X_{CO_2}$ retrieved from GOSAT with and without surface pressure normalisation applied. Shown are pre-screened, cloud-screened and post-screened GOSAT observations globally between July 2009 and June 2010.
3.16 Temperature and Water Vapour

The ECMWF specific humidity data can be used to generate water vapour VMR profiles using the equation

\[ H_{2OVMR} = 10^6 \left( \frac{SH}{(R_d R_w) - (SH((R_d R_w) - 1))} \right) \]  

(3.18)

where \( R_d \) and \( R_w \) are the gas constants in dry and wet air respectively. ECMWF also provides temperature profiles which, along with the \( H_{2OVMR} \), are interpolated with latitude, longitude and time to the specific observation. A constant \( O_2 \) VMR profile of 0.2095 is also used as an a priori. Often the retrieved surface pressure may be larger than the a priori surface pressure such that the atmospheric profiles for temperature and water vapour may be extrapolated below the a priori surface pressure value. In some cases this results in spurious values in the lowest levels of the profiles. To resolve this a static 20 level pressure grid is always used, and the a priori profiles are extrapolated using the lapse rate for temperature and gradient for water vapour of the 5 atmospheric levels above the surface pressure, such that a reasonable extrapolation occurs as shown in Figure 3.24. The temperature and \( H_2O \) profiles are assumed correct but the retrieval is allowed to scale them. For the temperature offset a standard deviation of 3.2 K is used and for the \( H_2O \) scale factor a standard deviation of 0.32 is used.
Figure 3.24. Example of the extrapolation technique used for temperature and water vapour profiles. The black lines show the time interpolated profile with the lapse rate/gradient applied, whereas the other profiles show the profiles for each time frame which have been simply extrapolated below the surface pressure. These profiles are already interpolated with latitude and longitude from four ECMWF grid points.
Atmospheric aerosols can scatter light into or out of the measured light path causing the measured intensity to vary and the optical path length to change. Additionally, some aerosols can absorb some of the light being measured, reducing the signal intensity and effecting the absorption lines of trace gases. Initially, the retrieval algorithm was setup to retrieve a single aerosol profile with an a priori AOD equal to the Moderate Resolution Imaging Spectroradiometer (MODIS) monthly mean AOD inferred for that 1° x 1° grid cell location. MODIS is a remote sensing instrument that is onboard the Terra and Aqua satellites that were launched in 1999 and 2002, respectively. However, aerosols are known to be highly variable and using a monthly mean AOD could potentially be improved.

The ARM Climate Research Facility includes a network of ground-based instruments located around the world. The central facility at Lamont/USA and the Darwin facility in Darwin/Australia are both located near to the respective TCCON sites. AOD was obtained from these ARM sites within the observation period of GOSAT using a Cimel sunphotometer (CSPHOT) instrument. The CSPHOT measures direct solar irradiance and sky radiance from the ground at visible and near infrared wavelengths to calculate atmospheric absorption and scattering (Gregory 2011). It derives AOD with a high accuracy (between 0.01 and 0.02 (Gregory 2011)), providing a useful source of in-situ AOD.

\[ \tau_{0.76\mu m} = \tau_{0.87\mu m} \ast \left( \frac{\lambda_{0.76}}{\lambda_{0.87}} \right)^{-\alpha_{0.50\mu m} - 0.87\mu m} \]  \hspace{1cm} (3.19)

where \( \tau \) represents the AOD, \( \lambda \) the wavelength and \( \alpha \) the angstrom coefficient. Figures 3.25 and 3.26 show the retrieved \( X_{CO2} \) from GOSAT over Lamont and Darwin, respectively, when using ARM AOD as a priori. For comparison, the retrieved \( X_{CO2} \) from GOSAT when using MODIS AOD as a priori and \( X_{CO2} \) inferred from TCCON are shown. The \( X_{CO2} \) retrieved from GOSAT using ARM AOD is very similar in value to the \( X_{CO2} \) retrieved when using MODIS AOD and follow the same temporal pattern in general. Both sets of GOSAT retrievals show a bias of about -6 ppm on average compared to TCCON, with ARM AOD based retrievals being slightly lower than MODIS AOD based retrievals.

However, the ARM network is sparsely distributed globally, making it unsuitable to
Figure 3.25. $X_{CO_2}$ inferred from TCCON compared with $X_{CO_2}$ retrieved from GOSAT using the UoL-FP retrieval algorithm where the a priori aerosol AOD equals either MODIS monthly mean AOD or ARM temporally closest AOD, calculated for the $O_2$ A band. All TCCON $X_{CO_2}$ data between April and mid-July 2009 from the Lamont site is shown in green and only GOSAT observations within ±5° of the TCCON site and within 2 hours of an ARM AOD measurement are given. All retrieved GOSAT observations using MODIS AOD are given by cyan triangles and those using ARM AOD by orange triangles. The daily mean of the retrieved $X_{CO_2}$ from GOSAT is given in blue and red where MODIS and ARM AOD were used respectively (with error bars representing the $X_{CO_2}$ daily scatter). The retrieved observations from GOSAT were post-screened for low $\chi^2$ in bands 1 and 2 to filter out retrievals with poor fits of the simulated spectra to GOSAT spectra.

be used for global retrievals of $X_{CO_2}$ from GOSAT. Although satellite measurements of aerosols exist, they are limited to only provide measurements at the locations and times of their orbital paths, not coincident with GOSAT. Although GOSAT includes a cloud and aerosol imager (TANSO-CAI) no aerosol data product has been released yet. Currently, the only way to obtain temporally and spatially coincident aerosol information is to use model estimates.
Figure 3.26. $X_{CO2}$ inferred from TCCON compared with $X_{CO2}$ retrieved from GOSAT using the UoL-FP retrieval algorithm where the a priori aerosol AOD equals either MODIS monthly mean AOD or ARM temporally closest AOD, calculated for the O$_2$ A band. All TCCON $X_{CO2}$ data between April and mid-July 2009 from the Darwin site is shown in green and only GOSAT observations within $\pm 5^\circ$ of the TCCON site and within 2 hours of an ARM AOD measurement are given. All retrieved GOSAT observations using MODIS AOD are given by cyan triangles and those using ARM AOD by orange triangles. The daily mean of the retrieved $X_{CO2}$ from GOSAT is given in blue and red where MODIS and ARM AOD were used respectively (with error bars representing the $X_{CO2}$ daily scatter). The retrieved observations from GOSAT were post-screened for low $\chi^2$ in bands 1 and 2 to filter out retrievals with poor fits of the simulated spectra to GOSAT spectra.

The Global and regional Earth-system (Atmosphere) Monitoring using Satellite and in-situ data (GEMS) provides re-analysis model AOD data on 1.125° x 1.125° grid cells globally. GEMS includes two natural aerosol types, sea salt and desert dust, that are surface and weather dependent. These aerosols types both have 3 different sizes; small, medium and large. In addition to the natural aerosols, five anthropogenic aerosols are included. These comprise of sulphate, hydrophilic/hydrophobic organic matter and
hydrophilic/hydrophobic black carbon. The GEMS model obtains information about these aerosols from the Global Fire Emission Database (GFED), the Speciated Particulate Emission Wizard (SPEW), and climatologies of the Emission Database for Global Atmospheric Research (EDGAR) (Simmons 2010). Thus, the GEMS model incorporates emissions of aerosols from fires as well as from domestic, industrial, power generation, transport and shipping activities (Simmons 2010). To complement these aerosol sources, the GEMS model assimilates global measurements of AOD from MODIS. One disadvantage of using the GEMS model is that it does not include any aerosol emissions from volcanic activity. The GEMS model (experiment f1kd) provides the total AOD at multiple wavelengths every twelve hours at 00:00 and 12:00, with a model forecast every three hours proceeding an analysis.

$X_{CO_2}$ was retrieved from GOSAT using a single aerosol profile with an a priori AOD equal to the GEMS AOD interpolated to the time and location of the GOSAT observation. The a priori AOD at the $O_2$ A band was calculated from the GEMS 0.670 $\mu$m and 0.865 $\mu$m AOD data similar to Equation 3.19, where the angstrom coefficient was calculated as

$$a_{0.670\mu m - 0.865\mu m} = - \frac{\ln \left( \frac{\tau_{0.670\mu m}}{\tau_{0.865\mu m}} \right)}{\ln \left( \frac{\lambda_{0.670\mu m}}{\lambda_{0.865\mu m}} \right)}$$

Figure 3.27 shows the retrieved $X_{CO_2}$ from GOSAT when using GEMS AOD as the a priori AOD, compared to retrieved $X_{CO_2}$ when an a priori AOD of 0.15 was always used. The retrieved $X_{CO_2}$ is very similar between the two methods, with slightly lower values when GEMS AOD was used. This indicates that the a priori AOD, if not extreme in value, does not affect the retrieval of $X_{CO_2}$ significantly.

Instead of retrieving aerosol extinction for each atmospheric level, one option could be to retrieve the height, width and optical depth of a Gaussian shaped aerosol extinction profile. This test provides a different approach to aerosols, allowing the peak of the aerosol distribution to be retrieved, rather than letting the optical depth of each atmospheric layer to vary individually. The retrieval algorithm was modified for this test and $X_{CO_2}$ was retrieved from GOSAT between April 2009 and February 2010 for observations within $\pm 5^\circ$ of the Lamont TCCON site. Figure 3.28 shows this retrieved $X_{CO_2}$ compared to $X_{CO_2}$ inferred by TCCON. Between April and July 2009, the retrieved $X_{CO_2}$ shows similar values to that previously seen in Figure 3.25. On average, the retrieved $X_{CO_2}$ is biased low compared to TCCON by roughly 6 ppm with large daily mean scatters ranging from $\sim 2$ ppm to $\sim 20$ ppm. Thus, changing the way the aerosol profile shape is retrieved does not improve the retrievals of $X_{CO_2}$. 

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FIGURE 3.27. A comparison of $X_{CO_2}$ retrieved from GOSAT between April 2009 and April 2010 using the UoL-FP retrieval algorithm with an a priori optical depth either equal to 0.15 always or equal to GEMS AOD. The retrieved observations were pre-screened, cloud-screened, and then post-screened for low $\chi^2$ values.

In addition to the aerosol profile shape and optical depth, the optical properties assumed could be changed. Instead of using a mixture of carbonaceous and dust aerosols based on Kahn et al. (2001) for a single aerosol, different aerosol types could be given separate profiles. The retrieval algorithm was modified to retrieve four aerosol logarithmic extinction profiles, each a priori profile having a Gaussian shape with a height and width of 2 km and an optical depth of 0.05. Using a Lorenz-Mie code (Mishchenko et al. 2002) for spherical particles, these aerosols were given optical properties based on GEMS medium sized desert dust, sulphate, hydrophobic organic matter and hydrophobic black carbon properties. Figure 3.29 shows the retrieved $X_{CO_2}$ from GOSAT using this setup for observations within $\pm 5^\circ$ of the Lamont TCCON site, compared to TCCON $X_{CO_2}$. The retrieved $X_{CO_2}$ shows a reduced bias ($\sim$ 4 ppm) compared to TCCON than previously found and a much lower daily scatter, between 1 ppm and 8 ppm.
An alternative method could be to use multiple aerosol mixtures rather than separate aerosol types, this would allow a faster computational time of the retrievals by reducing the number of state vector elements. To test this, aerosol optical properties were computed using a Mie code for carbonaceous/dusty continental and carbonaceous/sooty continental aerosol mixtures described in Kahn et al. (2001). Both these mixtures were given the same a priori extinction profiles that were Gaussian shaped with a height and width of 2 km, and an optical depth of 0.05 each. Figure 3.30 shows the retrieved $X_{CO_2}$ from GOSAT for this retrieval setup, for observations between April 2009 and February 2010 within $\pm 5^\circ$ of the Lamont TCCON site. The retrieved $X_{CO_2}$ was found to have a mean bias of $\sim 6$ ppm and a daily scatter between 1 ppm and 10 ppm.

Furthermore, it is not just aerosols that can cause absorption and scattering of light in the atmosphere, but clouds too. Although the cloud-screen already removes most observations influenced by clouds, some scenes are falsely flagged as clear. These scenes most likely include thin cirrus clouds that are low absorbing, letting the light still be measured by GOSAT. Therefore, the inclusion of cirrus in the state vector may allow these observations to still be retrieved successfully. A Gaussian shaped profile with a 10 km height, 2 km width and optical depth of 0.05 was used as a priori to test this. Optical properties for an effective radius of 60 $\mu$m was assumed for a non-spherical ice particles obtained from the Baum model (Baum et al. 2005a,b). The aerosol was setup the same as the last test, where two aerosol profiles of the same profile shape were used with different aerosol mixtures were used for the optical properties, but this test also includes the cirrus profile described. The retrieved $X_{CO_2}$ is shown in Figure 3.31 for GOSAT observations between April 2009 and February 2010 within $\pm 5^\circ$ of the Lamont TCCON site. Compared to TCCON, the retrieved $X_{CO_2}$ has a daily mean bias of approximately 3 ppm and a daily mean scatter between 1 ppm and 10 ppm. Importantly, less exposures were filtered out by the post-screen, meaning that retrievals of exposures that previously poorly fitted the measured GOSAT spectra are now retrieved. This gives $X_{CO_2}$ values within the same range as the exposures whose retrievals previously fitted the measurements well. Furthermore, the retrieved $X_{CO_2}$ shows a large increase of 3 ppm compared to exactly the same retrievals but without the cirrus included. Thus, not only does this approach produce a larger $X_{CO_2}$ dataset but it provides a lower bias.

Eguchi et al. (2007) found that cirrus clouds vary in height depending on the latitude, with larger values towards the equator (as shown in Figure 3.32). Since the use of a cirrus profile had a large effect on the retrieved $X_{CO_2}$, it was important to follow it up by testing the use of profile shape that varies with latitude. Mean values of cirrus cloud center heights and widths were obtained from Eguchi et al. (2007) for the tropics and
Figure 3.28. $X_{CO2}$ retrieved from GOSAT using the UoL-FP retrieval algorithm with the aerosol height, width and optical depth retrieved. All pre-screened and cloud-screened GOSAT observations within $\pm 5^\circ$ of the Lamont TCCON site between April 2009 and February 2010 were retrieved and post-screened for low $\chi^2$ values. The $X_{CO2}$ retrieved from TCCON is given in green with the daily mean shown in orange. The retrieved GOSAT $X_{CO2}$ is given by cyan triangles and the daily mean by blue circles, with the daily scatter given as error bars.

Subtropics in both hemispheres, mid-latitudes in both hemispheres, and globally (see Table 3.2). These values were interpolated onto a $1^\circ$ latitudinal grid and for every grid point a Gaussian shaped cirrus profile was created with the corresponding height and width. Thus providing a latitudinal dependent cirrus profile that can be used with retrievals of $X_{CO2}$ from GOSAT.

Figure 3.33 shows a comparison of the retrieved $X_{CO2}$ from GOSAT when using a constant cirrus profile and when using a latitudinally dependent cirrus profile, for observations between April 2009 and February 2010 within $\pm 5^\circ$ of the Lamont TCCON site. On average, higher $X_{CO2}$ and a lower retrieved cirrus optical depth is retrieved when a latitude dependent cirrus profile is used. Thus, using a latitudinally dependent cirrus profile reduces the $X_{CO2}$ bias to TCCON, by providing the retrieval with a more realistic
Figure 3.29. $X_{CO2}$ retrieved from GOSAT using the UoL-FP retrieval algorithm with four different aerosol profiles retrieved. The a priori of each aerosol was given a Gaussian shaped profile with a height and width of 2 km and optical depth of 0.05, but different optical properties. The optical properties were computed using a Mie code for the GEMS medium sized desert dust, sulphate, hydrophobic organic matter and hydrophobic black carbon. All pre-screened and cloud-screened GOSAT observations within $\pm 5^\circ$ of the Lamont TCCON site between April 2009 and February 2010 were retrieved and post-screened for low $\chi^2$ values. The $X_{CO2}$ retrieved from TCCON is given in green with the daily mean shown in orange. The retrieved GOSAT $X_{CO2}$ is given by cyan triangles and the daily mean by blue circles, with the daily scatter given as error bars.

Based on these tests, three atmospheric extinction profiles are used. Two of these profiles are the same shape, Gaussian profile with a 2 km height and width, and have a total column aerosol optical depth of 0.05 each (as shown in Figure 3.34). The only difference between these profiles are the aerosol properties used for each. Based on Kahn et al. (2001) one uses classification 4b and the other uses classification 5b. Therefore providing one profile designed for carbonaceous and dusty environments and the other profile for carbonaceous and sooty environments. The third profile allows for the pos-
Figure 3.30. $X_{CO2}$ retrieved from GOSAT using the UoL-FP retrieval algorithm with 2 aerosol profiles, both of Gaussian shape with a height and width of 2 km and optical depth of 0.05 each. The optical properties were computed using a Mie code for carbonaceous/dusty continental and carbonaceous/sooty continental aerosol mixtures described in Kahn et al. (2001). All pre-screened and cloud-screened GOSAT observations within ±5° of the Lamont TCCON site between April 2009 and February 2010 were retrieved and post-screened for low $\chi^2$ values. The $X_{CO2}$ retrieved from TCCON is given in green with the daily mean shown in orange. The retrieved GOSAT $X_{CO2}$ is given by cyan triangles and the daily mean by blue circles, with the daily scatter given as error bars.

Possibility of a cirrus cloud and is based on the height and thickness described in Eguchi et al. (2007). A Gaussian profile of 0.1 optical depth was created for each of the latitude regions described using the height and thickness given and then a look up table based on every 1 degree of latitude was created globally by interpolating between these regions with respect to latitude. Each retrieval uses the cirrus profile defined in the look up table that is closest to the specific latitude of the retrieved scene. The cirrus properties are constructed from ice cloud models based on the Baum model (Baum et al. 2005a,b). In the O2 A Band the ratio of scattering to absorption for cirrus has very little dependence on the particle radius but a high dependence on optical depth. Conversely at the 1.61 μm
Fig. 3.31. $X_{CO2}$ retrieved from GOSAT using the UoL-FP retrieval algorithm with a cirrus profile and 2 aerosol profiles. The cirrus a priori profile had a Gaussian shape with a height of 10 km, width of 2 km and optical depth of 0.05. The cirrus optical properties were obtained from the Baum model (Baum et al. 2005a) and given an effective radius of 60 $\mu$m. Both aerosols had an a priori profile of Gaussian shape with a height and width of 2 km and optical depth of 0.05 each. The optical properties were computed using a Mie code for carbonaceous/dusty continental and carbonaceous/sooty continental aerosol mixtures described in Kahn et al. (2001). All pre-screened and cloud-screened GOSAT observations within $\pm 5^\circ$ of the Lamont TCCON site between April 2009 and February 2010 were retrieved and post-screened for low $\chi^2$ values. The $X_{CO2}$ retrieved from TCCON is given in green with the daily mean shown in orange. The retrieved GOSAT $X_{CO2}$ is given by cyan triangles and the daily mean by blue circles, with the daily scatter given as error bars.

and 2.03 $\mu$m bands, there is more absorption by cirrus and therefore a higher sensitivity to the particle radius (Zhang et al. 2010). A large ice particle of an effective radius of 60 $\mu$m is assumed for all retrievals, to provide the retrieval algorithm with a high variability of the ratio of scattering to absorption between bands.
FIGURE 3.32. Cirrus cloud characteristics as a function of latitude, calculated from the Geoscience Laser Altimeter System (GLAS) and the Ice, Cloud, and Land Elevation Satellite (ICESat) between 1st October and 18th November 2003. The altitude of the top of the cirrus cloud is given in blue, the altitude of the bottom of the cirrus cloud is given in red and the cirrus cloud thickness is given in purple. The cirrus cloud optical depth is given in orange and the cirrus cloud temperature is given in green. The average Outgoing Longwave Radiation (OLR) obtained from NOAA satellites during the same time period is shown in black (Eguchi et al. 2007).

Table 3.2. Mean cirrus cloud heights and widths calculated from GLAS and ICESat data for the tropics and subtropics in both hemispheres, mid-latitudes in both hemispheres, and globally (Eguchi et al. 2007).

<table>
<thead>
<tr>
<th>Latitude region</th>
<th>Latitude range</th>
<th>Height [km]</th>
<th>Width [km]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tropics</td>
<td>15°S - 15°N</td>
<td>13.6</td>
<td>1.6</td>
</tr>
<tr>
<td>Subtropics (northern hemisphere)</td>
<td>15°N - 30°N</td>
<td>12.0</td>
<td>1.6</td>
</tr>
<tr>
<td>Subtropics (southern hemisphere)</td>
<td>15°S - 30°S</td>
<td>10.7</td>
<td>1.4</td>
</tr>
<tr>
<td>Middle latitude (northern hemisphere)</td>
<td>30°N - 60°N</td>
<td>9.2</td>
<td>1.6</td>
</tr>
<tr>
<td>Middle latitude (southern hemisphere)</td>
<td>30°S - 60°S</td>
<td>8.7</td>
<td>1.6</td>
</tr>
<tr>
<td>Globe</td>
<td>86°S - 86°N</td>
<td>10.3</td>
<td>1.6</td>
</tr>
</tbody>
</table>
Figure 3.33. $X_{CO2}$ retrieved from GOSAT between April 2009 and February 2010 within ±5° of the Lamont TCCON site, when a constant cirrus profile of Gaussian shape with a height of 10 km and width of 2 km was assumed for all retrievals compared to when a latitude dependent cirrus profile based on Eguchi et al. (2007) was used. Both retrievals were otherwise setup the same and both were given an a priori optical depth of 0.05 and the same optical properties. A comparison of the retrieved $X_{CO2}$ is given by the left panel and a comparison of the retrieved cirrus optical depth is given by the right panel.
FIGURE 3.34. Gaussian aerosol profiles of 2 km height width and with 0.05 AOD.
Figure 3.35. Gaussian profiles based on the height and thickness described for various regions by Eguchi et al., (Eguchi et al. 2007).
3.18 Surface Properties

The albedo, $\alpha$, is calculated from the spectral continuum of the GOSAT L1B using the reflectivity

$$\alpha = \frac{\pi S_{GOSAT}}{S_{Solar} \cos(SZA)}$$  \hspace{1cm} (3.21)

where SZA is the solar zenith angle, and the Solar irradiance and observed GOSAT radiance are given by $S_{Solar}$ and $S_{GOSAT}$ respectively. The retrieval uses two albedo parameters for each spectral band, giving the albedo for the centre wavelength of the band and the slope of the albedo. The slope of the albedo is set to zero in the a priori. The covariance for albedo is completely open (standard deviation of 1) and the slope a priori error uses a standard deviation of 0.01 cm$^{-1}$ so that the band edges can vary by 50% to compensate for any apodization artifacts introduced during the FFT of the TANSO-FTS interferogram.

3.19 Instrument Properties

The recommended radiometric calibration and solar diffuser degradation correction (preliminary factors) were applied to the GOSAT Level 1B files (050.050C, 080.080C, 110.1-10C and 130.130C) which covered the years 2009 to 2011 and were acquired via the GOSAT User Interface Gateway.

The radiative transfer model produces synthetic spectra for a specified observation geometry and surface/atmospheric state. Before comparing these spectra to calibrated measured spectra they are corrected for instrument dispersion, and convolved with the instrument line shape (ILS) function and polarisation. For polarisation to be correctly used a Mueller matrix (Kuze et al. 2009) is required to combine the different Stokes vectors. However, the Mueller matrix is derived using the observation viewing angles which are known to have pointing issues due to microvibrations causing instability. Therefore, the intensity is approximated by using the mean of the polarised intensities:

$$I = \frac{P + S}{2}$$  \hspace{1cm} (3.22)

The noise is derived from the standard deviation of the out of band radiances for each band (see Figure 3.36), which in the absence of noise would centre on zero. Similarly to
the signal, the noise from each polarisation is combined as

\[ N = \sqrt{N_F^2 + N_S^2} \]  

(3.23)

**Figure 3.36.** Example raw radiance for each GOSAT SWIR band showing out of band characteristics used to derive the noise (grey).

Systematic structures in the residual of the spectral fit were found (typically solar lines), that are most likely driven by insufficient spectroscopy. As a consequence, the reduced \( \chi^2 \) of the spectral fit will increase with the theoretical SNR of the sounding. The \( \chi^2 \) could become reduced by masking these spectroscopic peaks, which can be achieved by increasing the noise.

Figure 3.37 shows the real mean squared (RMS)/Signal compared to 1/SNR for each band for \( X_{CO2} \) retrieved from GOSAT for one 3-day cycle in June 2009. The noise can be inflated by using an empirical relationship between the RMS/signal and 1/SNR for each band (Crisp et al. 2012) of

\[ \frac{\text{RMS}}{\text{Signal}} = A + \frac{B}{\text{SNR}} \]  

(3.24)

where A and B are the coefficients derived (see Table 3.3). Using these coefficients the noise was then modified by

\[ \text{Inflated Noise} = A \ast \text{Signal} + B \ast \text{Noise} \]  

(3.25)

The GOSAT 3-day cycle was re-retrieved using the inflated noise and the difference of retrieved \( \chi^2 \) of the fit for each band is shown in Figure 3.38. The \( \chi^2 \) was found to become much lower when the noise was increased, as expected.

With the release of new TANSO-FTS L1B data versions, the noise coefficients were re-calculated for each version. Figure 3.39 shows the fit of RMS/Signal and 1/SNR for each band with each available data version shown, where September 2009 is L1B version
Figure 3.37. Fit of relative RMS (RMS/Signal) and 1/SNR for each GOSAT SWIR band retrieved by the UoL-FP algorithm for global observations of a 3-day orbital cycle between 1st-3rd June 2009. Additionally, the difference between the retrieved and expected $\chi^2$ is highlighted for different ranges in value. A linear fit of the data is given by the dotted line for each band, with its equation provided in the top left of each panel.

V050, December 2009 is L1B version v080, and May 2010 is L1B version v100. These fits vary only slightly from the initial coefficients derived and are also similar to that reported by ACOS (Connor 2010). For simplicity, the mean of the coefficients from each L1B version was used.

Table 3.3. Derived coefficients used to inflate the noise.

<table>
<thead>
<tr>
<th>Test</th>
<th>A (Band 1)</th>
<th>B (Band 1)</th>
<th>A (Band 2)</th>
<th>B (Band 2)</th>
<th>A (Band 3)</th>
<th>B (Band 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial (007.007V)</td>
<td>0.00139045</td>
<td>0.765909</td>
<td>0.00222652</td>
<td>0.544227</td>
<td>0.00637773</td>
<td>0.561081</td>
</tr>
<tr>
<td>1st-3rd September 2009 (080.080C)</td>
<td>0.0038452676</td>
<td>0.65677958</td>
<td>0.0021347245</td>
<td>0.70029702</td>
<td>0.0039319740</td>
<td>0.65454255</td>
</tr>
<tr>
<td>1st-3rd December 2009 (050.050C)</td>
<td>0.004277149</td>
<td>0.57120579</td>
<td>0.0017439434</td>
<td>0.77762136</td>
<td>0.0039563910</td>
<td>0.63522226</td>
</tr>
<tr>
<td>1st-3rd May 2010 (100.100C)</td>
<td>0.0042883528</td>
<td>0.57437039</td>
<td>0.0023632774</td>
<td>0.51059092</td>
<td>0.0036055675</td>
<td>0.65775003</td>
</tr>
<tr>
<td>Mean</td>
<td>0.00422045</td>
<td>0.600785</td>
<td>0.00226265</td>
<td>0.662836</td>
<td>0.00383131</td>
<td>0.649172</td>
</tr>
<tr>
<td>ACOS (080.050C)</td>
<td>0.00234298</td>
<td>0.643008</td>
<td>0.00165767</td>
<td>0.634136</td>
<td>0.00287656</td>
<td>0.633723</td>
</tr>
</tbody>
</table>

However, recent updates in spectroscopy have reduced the systematic structures in
the residual of the spectral fit, giving the expected $\chi^2$ values. Therefore, the original noise is now sufficient to use.

The dispersion gives the pixel-wavelength mapping and consists of two parameters for each band; the start wavenumber $W$ and the wavenumber increment $D$ needed to reach the next pixels wavenumber. The wavenumber, $\nu$, for detector pixel, i, and spectrometer, k, is given by

$$\nu_i = W_k + D_k$$

(3.26)

The dispersion is given in the GOSAT L1B data but requires adjusting in all SWIR bands. This adjustment can be calculated by the difference of the spectra away from a well known single strong Fraunhofer line, otherwise known as a Solar line, (12985.163 cm$^{-1}$) in the O$_2$ A band. Additionally, the O$_2$ A band dispersion requires shifting one spectral point lower to become approximately correct. It should be understood that although all SWIR bands dispersions are adjusted they are only approximate and require the dispersion to be retrieved.

To describe the response of the instrument to light with wavelength an instrument line shape function (ILS) is used, which was obtained from the JAXA ILSF model (Kuze et al. 2009). Band 1 has the ILS provided for both polarisations due to band 1 being sensitive to displacement of the optical axis with its shorter wavelengths, whereas bands 2 and 3 have accurate alignment. The ILS model gives the ILS for 3 different wavelengths in band 1 and 4 different wavelengths in both bands 2 and 3.

To mitigate the effect of the GOSAT band 1 non-linear response to the intensity of incident radiation on the retrieval of $X_{CO2}$, an additive zero level intensity offset was retrieved for the O$_2$ A band. To check that the inclusion of this does not cause the retrieved $X_{CO2}$ to become worse, retrievals with and without the zero level offset applied were retrieved for GOSAT pre-screened and cloud-screened observations for September 2009 globally. Figure 3.41 shows the comparison of retrieved $X_{CO2}$ from retrievals with and without the zero level offset retrieved in the O$_2$ A band. For low $X_{CO2}$ values, slightly larger $X_{CO2}$ values are found when a zero level offset is included in the retrieval. However, in general, the $X_{CO2}$ remains similar. This method may also partially mitigate fluorescence observed by Frankenberg et al. (2011a).
Figure 3.38. Comparison of retrieved $\chi^2$ and SNR for each GOSAT SWIR band retrieved by the UoL-FP algorithm for global observations of a 3-day orbital cycle between 1st-3rd June 2009. The left column shows the data for each band when the original noise was used, and the right column shows each band when the inflated noise was used. Note that the y-axis scales are different, giving much lower $\chi^2$ values when the inflated noise was used.
FIGURE 3.39. Linear fits of RMS/Signal and 1/SNR for each GOSAT SWIR band retrieved by the UoL-FP algorithm for global observations of a 3-day orbital cycle in 3 different time frames, covering different TANSO-FTS L1B data versions. The initial/previous fit is shown in black and that reported similarly by ACOS in cyan.
FIGURE 3.40. ILS used in the retrievals for each GOSAT SWIR spectral band with different wavenumbers given.
Figure 3.41. Comparison of retrieved $X_{CO_2}$ from GOSAT using the UoL-FP retrieval algorithm with and without an O$_2$ A band zero level offset retrieved. These retrievals were performed on pre-screened and cloud-screened GOSAT observations for September 2009 globally. Details of the algorithm are provided in Chapter 3.
3.20 Pre-Processing Screen

A pre-screen is applied to all GOSAT observations during the pre-processing algorithm such that only land observations with a SNR > 50 in all bands are used. Although observations with a low SNR could be retrieved, it would result in a poor retrieval fit, since small changes in the signal can not be distinguished from the noise. Also, poor retrieval fits are expected over snow surfaces (see Section 3.11) so only scenes with a SZA < 70° are used to reduce the computational requirement of performing retrievals globally. Additionally, the pre-screen selects non-saturated measurements with an along-track angle error < 0.05, cross-track angle error < 0.007, and where the IFOV surface pressure standard deviation is < 10 hPa. The along-track and cross-track filters are used to keep only measurements where the pointing has not been adversely affected by microvibrations.

The IFOV of TANSO-FTS is ~10 km in diameter and within this area the topography can change greatly. Although the pre-processing uses the mean atmospheric conditions within the IFOV, this might not provide useful a priori in extreme scenes. Also, the surface can vary significantly over such a large region, leading to variations in albedo that are unaccounted for in the retrieval. Since surfaces with a higher albedo reflect more light the observed radiance will be biased towards that surface type and therefore that location and not the entire IFOV scene. In this instance, the retrieval would require the atmospheric conditions for that location and not the mean of the IFOV. Screening to remove scenes with a high surface pressure variation helps mitigate this issue.

It is possible that during a TANSO-FTS observation the interferogram can become saturated by light. If the interferogram becomes saturated, then when the FFT is performed the spectral radiance and absorption lines will be larger everywhere leading to a poor retrieval. Therefore, a new method for detecting GOSAT saturation was created to allow these observations to be filtered out. This detection method uses the mean out-of-band radiances, since the spectra should have higher values throughout the measured spectral ranges when saturation occurs. Initially all three SWIR bands were analysed but the saturation is best detected with this method in band 2, as shown in Figure 3.44. The band 2 saturation detection was compared with the \( \chi^2 \) of X\(_{CO2} \) retrievals for band 2 (see Figure 3.45) and a clear trend was found with larger \( \chi^2 \) values. TANSO-FTS is capable of measuring radiances using different gain modes, such that it can cope with high intensity signals by using the medium gain so that saturation should not arise. To attempt to determine why the saturation occurs, the maximum signal of band 2 was compared with the saturation detection test for each gain mode. This is given in Figure 3.47, which distinctly shows that saturation happens when the signal intensity is highest for
observations using high gain. This highlights that the medium gain mode should be used in additional measurement locations.

**Figure 3.42.** Mean out-of-band radiances of band 1 P (left) and S (right) compared with latitude and retrieved $X_{CO2}$ for 1st-7th September 2009 globally.

**Figure 3.43.** Mean out-of-band radiances of band 2 P (left) and S (right) compared with latitude and retrieved $X_{CO2}$ for 1st-7th September 2009 globally.

**Figure 3.44.** Mean out-of-band radiances of band 3 P (left) and S (right) compared with latitude and retrieved $X_{CO2}$ for 1st-7th September 2009 globally.

### 3.21 Cloud Screen

One major disadvantage of using SWIR observations is that the IFOV must be free of thick clouds as these will cause irradiation to be reflected towards the satellite from a higher altitude, that of the cloud, rather than the Earth’s surface where source/sink information can be gained. Therefore, a cloud screen is necessary to filter out exposures where thick clouds are within the IFOV. When irradiation gets reflected towards the satellite from a thick cloud the effective surface is that of the cloud rather than the Earth’s surface. This results in the surface pressure being that at the cloud altitude, which typically is significantly lower than that at the Earth’s surface pressure. An estimate of
the surface pressure can be obtained by performing an O$_2$ A band retrievals where the surface pressure is retrieved. A fast O$_2$ retrieval is implemented, by using a small spectral window (13056 to 13074.8 cm$^{-1}$) where the surface pressure information is obtained, to reduce the computational time as the cloud screen is performed on a large volume of data (all pre-processed exposures). If the retrieved surface pressure is compared to the a priori surface pressure then the difference in surface pressure can be determined. Since the a priori surface pressure is well constrained, a surface pressure difference of greater than 20 hPa would normally only occur if a thick cloud is present in the IFOV, thus the
exposures can be cloud screened using this method.

This cloud screen approach has been verified by using a simulator (see Chapter 5) to compare the O$_2$ A band retrievals with the simulated cloud optical depth. Spectral radiances for each GOSAT SWIR band were simulated using 61 level profiles of pressure, water vapour VMR, CO$_2$ VMR, O$_2$ VMR, temperature, extinction for 11 different aerosols, liquid water extinction and ice water extinction. Also included was surface pressure and for each band values for albedo shift and slope, and spectral shift and stretch. The cloud screening method was then applied to these radiances and the retrieved State Vector elements compared to the true values.

Figures 3.49 and 3.50 show histograms of the retrieved parameters for clear and cloudy scenes, where clear scenes have a total column optical depth of less than 0.3. This clearly shows how the difference in surface pressure can be used to screen for many cloudy scenes, with the statistics given in Table 3.4. The cloud screen is shown to remove $\sim$86% of cloudy scenes and keep $\sim$94% of clear scenes.

The statistics can be compared to those of alternative cloud screening methods by calculating two diagnostic quantities (Taylor et al. 2012); the accuracy of the cloud screen to correctly classify clear and cloudy scenes, and the positive predictive value that specifies the percentage of clear scenes classified as clear. The accuracy, ACC, can be given as

$$\text{ACC} = \frac{\text{True clear} + \text{True cloudy}}{\text{Total number of scenes}} = 87.90\%$$ (3.27)

and the positive predictive value, PPV, as

$$\text{PPV} = \frac{\text{True Clear}}{\text{True Clear} + \text{False Clear}} = 69.59\%$$ (3.28)

Taylor et al. (2012) compares two different cloud screening methods which have been applied to GOSAT; a full O$_2$ A band surface pressure retrieval method (ABO2 Cloud Screen) and the CLAUDIA-CAI cloud screen which utilises the TANSO-CAI. The ACC and PPV over land for the ABO2 cloud screen are 78.2% and 63.0%, and for the CLAUDIA-CAI cloud screen are 79.1% and 71.0% respectively. In that study the cloudiness truth of the scenes was determined by the MODIS cloud mask product (MYD35), rather than by simulated scenes. Although the values can not be directly compared to those of this simulation as the simulated scenes do not include any variation of cloudiness within the IFOV of the simulated observation, the values are in general similar.
Table 3.4. Cloud screen statistics from simulations.

<table>
<thead>
<tr>
<th>Description</th>
<th>Number of Elements</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>9923</td>
<td>100.00</td>
</tr>
<tr>
<td>Total clear truth</td>
<td>2544</td>
<td>25.64</td>
</tr>
<tr>
<td>Total cloudy truth</td>
<td>7379</td>
<td>74.36</td>
</tr>
<tr>
<td>True clear retrieved</td>
<td>2387</td>
<td>24.06</td>
</tr>
<tr>
<td>False clear retrieved</td>
<td>1043</td>
<td>10.51</td>
</tr>
<tr>
<td>True cloudy retrieved</td>
<td>6336</td>
<td>63.85</td>
</tr>
<tr>
<td>False cloudy retrieved</td>
<td>157</td>
<td>1.58</td>
</tr>
</tbody>
</table>

**Figure 3.49.** Histograms of $\chi^2$, surface pressure difference, albedo and dispersion with frequencies separated into true/false clear/cloudy, using the cloud screening method described in this section for simulated observations of GOSAT for two repeat cycles (6 days) globally.
Figure 3.50. This shows the same as Figure 3.49 but with the x-axis scale reduced such that more detail can be viewed where clear and cloudy scenes need distinguishing.
3.22 Post-Processing Screen

Some retrievals will inherently perform poorly due to incorrect prior assumptions. Additionally, by giving the aerosols a large flexibility it can increase the chances of retrievals not converging and obtaining the solution from a local minima, rather than the true optimal solution. Therefore some quality assurance should be used with the retrieval results. Post-screening criteria is applied to the successful retrievals to reduce the scatter and to remove problematic and potentially biased retrievals. An overview of the empirically derived post-screening criteria is shown in Table 3.5, with its application shown for a combination of seven TCCON sites (see Chapter 4) over a two year period in Figure 3.51. The post screen threshold values were inferred from correlations of various parameters with the X\textsubscript{CO2} bias between GOSAT and TCCON for coincident observations over Lamont/USA and Darwin/Australia (see Chapter 4) and by considering the distribution and scatter of global X\textsubscript{CO2} data retrieved for August 2009.

Table 3.5. UoL-FP v3G Post-Screen parameter threshold limits.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Lower Threshold</th>
<th>Upper Threshold</th>
</tr>
</thead>
<tbody>
<tr>
<td>Outcome</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Band 1 (\chi^2)</td>
<td>0.6</td>
<td>1.3</td>
</tr>
<tr>
<td>Band 2 (\chi^2)</td>
<td>0.6</td>
<td>1.5</td>
</tr>
<tr>
<td>Band 3 (\chi^2)</td>
<td>0.4</td>
<td>1.5</td>
</tr>
<tr>
<td>A Posterior Error</td>
<td></td>
<td>1.6</td>
</tr>
<tr>
<td>Number of Divergences</td>
<td></td>
<td>2</td>
</tr>
<tr>
<td>Retrieved - A priori Surface Pressure (hPa)</td>
<td>0</td>
<td>16</td>
</tr>
<tr>
<td>Retrieved Aerosol 1 Optical Depth</td>
<td>0.3</td>
<td></td>
</tr>
<tr>
<td>Retrieved Aerosol 2 Optical Depth</td>
<td>0.4</td>
<td></td>
</tr>
<tr>
<td>Retrieved Ice Optical Depth</td>
<td>0.05</td>
<td></td>
</tr>
<tr>
<td>Total Optical Depth</td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td>BTD 8-11</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Cirrus Test</td>
<td>0.98</td>
<td>1.05</td>
</tr>
<tr>
<td>Retrieved Albedo Band 1 / Band 3</td>
<td></td>
<td>4</td>
</tr>
<tr>
<td>Retrieved Temperature Scale (K)</td>
<td>-2</td>
<td></td>
</tr>
<tr>
<td>Retrieved H\textsubscript{2}O Scale Factor</td>
<td>0.7</td>
<td>1.3</td>
</tr>
<tr>
<td>Retrieved A priori Dispersion Band 2 Shift (cm(^{-1}))</td>
<td>-0.022</td>
<td>0.022</td>
</tr>
<tr>
<td>Retrieved Intensity Offset</td>
<td>-0.002</td>
<td>0.0014</td>
</tr>
</tbody>
</table>

The post-screen selects only converged retrievals with a good spectral fit (where the
FIGURE 3.51. Box-wisker plot of $X_{CO_2}$ retrieved from GOSAT for seven TCCON sites combined between April 2009 and May 2011. The effect of each post-screen parameter is shown sequentially along the x axis.

normalized $\chi^2$ of the fit residual is $\sim 1$) and a low posterior error of $X_{CO_2}$ (<1.6 ppm). In addition, we also filter for a number of parameters related to thin clouds and aerosols, which include: the difference between retrieved and ECMWF surface pressure (0-16 hPa), the retrieved cirrus optical depth (<0.05), the band 4 (TIR) brightness temperature difference (BTD) between 8 $\mu$m and 11 $\mu$m (where $>0$ indicates cloud/cirrus), a test for cirrus based on the radiance ratio of 2.1038 $\mu$m to 2.0906 $\mu$m (where cirrus absorption occurs below a value of 0.98), and the ratio of 0.76 $\mu$m to 2.06 $\mu$m albedo that indicates ice/snow if the value is $>4$. Scenes containing a large retrieved aerosol amount (AOD $>0.5$) are also removed. Figure 3.52 gives the correlation matrix of the state vectors used, showing many cross-correlations between statevectors. As most of the correlating state vectors are included in previously mentioned filters, water vapour and temperature which also show some correlation with $X_{CO_2}$ are used to remove cases where clear outliers exist (indicating a poor retrieval).

Thin cirrus clouds will likely not be filtered out by the cloud screen and are difficult to detect in the SWIR spectral range so three different techniques were designed to screen for cirrus clouds. The first method was to utilise the TANSO-FTS TIR band and
Figure 3.52. Example correlation matrix of the state vectors used in the UoL-FP retrieval algorithm, obtained from a $X_{CO_2}$ retrieval over Lamont/USA on April 22$^{nd}$ 2009. For the state vector profiles (e.g. CO$_2$, aerosols and ice) the top of the atmosphere is given by the 1$^{st}$ level. Note that only 19 levels are shown here due to the retrieved surface pressure being higher than the two lowest a priori pressure levels.

calculate the brightness temperature difference (BTD) between 8.3 and 11.15 $\mu$m, which has previously been used with MODIS observations to characterise scenes with clouds (Ackerman et al. 1998). This BTD can be calculated using Planck’s law as

$$BTD = \left[ \frac{h\nu_8}{k \ln \left( 1 + \frac{2h\nu_8^3}{\nu_8 c^2} \right)} \right] - \left[ \frac{h\nu_{11}}{k \ln \left( 1 + \frac{2h\nu_{11}^3}{\nu_{11} c^2} \right)} \right]$$

(3.29)

where $\nu_8$ is the 8.3 $\mu$m wavenumber, $\nu_{11}$ is the 11.15 $\mu$m wavenumber, h is the Planck
constant \((6.6262 \times 10^{-34} \text{ J s})\), \(c\) is the speed of light \((2.99792458 \times 10^8 \text{ m s}^{-1})\) and \(k\) is the Boltzmann constant \((1.38062 \times 10^{-23} \text{ J K}^{-1})\). This technique should indicate if cirrus clouds exist if the BTD is greater than \(\sim 0\) (Ackerman et al. 1998).

![Figure 3.53](image)

**Figure 3.53.** The retrieved \(X_{CO2}\) that has been filtered out due to the 8-11 \(\mu\text{m}\) BTD threshold, for August 2009 globally.

The second technique uses the ratio of the retrieved albedo of band 1 to band 3. In band 1 ice/snow has a very high albedo, whilst all other surface types have a lower albedo. In band 3 ice/snow has the lowest albedo, with all other surfaces having low albedos but higher than ice/snow. By using the ratio of band 1 to band 3 albedo, if the scene contains ice/snow then the ratio will be larger. Thus this method has the potential to detect cirrus (ice cloud). Additionally, it can detect surface ice/snow which would also lead to a reduced accuracy in the \(X_{CO2}\) retrieval due to a lower SNR, so would be preferable to screen out as well.

The third technique uses the fact that cirrus strongly absorbs radiances in the band 3 spectral range, with the absorption peaking towards the centre of the band. This cirrus test makes use of this absorption peak by taking the ratio of the continuum radiance at the edge of the band where the absorption is low \((2.1038 \mu\text{m})\) to the continuum radiance at the middle of the band where at the peak cirrus absorption \((2.0904 \mu\text{m})\). If no cirrus exists then the cirrus test would give a value of \(\sim 1\), but if cirrus does exist in the scene then the value ought to be slightly \(< 1\). This is clearly demonstrated in Figure 3.55, which shows the cirrus test values for the set of simulated scenes used in the cloud screen analysis, for details see Section 3.21. This indicates that a threshold of \(\sim 0.98\) would be suitable to remove scenes that contain cirrus.

Figure 3.56 shows the difference between with and without the post-screen applied
Figure 3.54. Ratio of band 1 retrieved albedo to band 3 retrieved albedo for August 2009 globally. The upper plot shows its correlation with retrieved $X_{CO_2}$ with the density of values given by colour and the red vertical line showing the threshold limit. The lower plot shows the retrieved $X_{CO_2}$ that has been filtered out due to this parameters threshold.

for parameters related to the quality of the retrieval performance. When no post-screen is applied, a $X_{CO_2}$ bias is found with the number of divergences and the number of iterations. There is also a large number of retrievals with poor $\chi^2$ values. Once post-screened, the $X_{CO_2}$ biases are significantly reduced and only retrievals with a good spectral fit remain.

An example of the spectral fit and the fit residual is given in Figure 3.57, and shows that in general the retrieval fits the measured radiance well with low residuals. However, some structures remain in the residual indicating that some spectral lines are not fitted well, with the larger residual peaks relating to solar absorption lines. Future updates in spectroscopy could improve the spectral fit.
Figure 3.55. Cirrus test for realistic simulated scenes, as described in Section 3.21. It is compared against the simulated ice optical depth with the density of values given by the colour scale.
FIGURE 3.56. Example application of the post-screen on different parameters for GOSAT pre-screened and cloud screened observations September 2009. The left hand panels show the results when no post-screen is applied, and the right hand panels show the same data but filtered using the post-screen. For each panel, the colour represents the number of results within that bin with reference to the colourbar, the pink line gives a linear fit of the data, and r gives the correlation of the parameter against $X_{CO2}$. 
Figure 3.57. Example spectral fits and fit residuals for each band retrieved, where the measured radiances are given in red, the simulated radiances in green and the residual in gray. This example exposure is located over Lamont/USA where the surface is vegetation and the retrieved total optical depth was $\sim 0.1$. 
3.23 Summary

This chapter described the UoL-FP retrieval algorithm and how it can be used to infer $X_{CO_2}$ from cloud screened GOSAT SWIR measured radiances. The vertical sensitivity and $X_{CO_2}$ error of GOSAT was assessed using pre-flight instrument information. This shows surfaces with a low albedo in the wavelength range of the GOSAT bands (such as ocean and snow) will result in errors of about 1%, whilst brighter surfaces give much lower errors and better sensitivity to the boundary layer. The next chapter shows the $X_{CO_2}$ inferred from GOSAT using the UoL-FP retrieval algorithm and compares with ground based measurements, global chemistry transport model calculations and alternative retrieval algorithms.
Chapter 4

Atmospheric CO₂ retrieved from GOSAT: Validation, model comparison and algorithm intercomparison

4.1 Introduction

To assess the ability of the UoL-FP retrieval algorithm (version 3G) to accurately and precisely estimate $X_{CO2}$ from GOSAT, it is essential to compare this with alternative observational data and independent model data. It is also useful to compare to alternative algorithms as this can potentially indicate retrieval algorithm dependent biases. This chapter provides an assessment of the UoL-FP retrieval algorithm through comparisons with:

1. In-situ ground based observations of $X_{CO2}$ derived from solar absorption spectra measured by TCCON FTS instruments.
2. The GEOS-Chem global 3-D chemistry transport model atmospheric CO₂ estimations (Feng et al. 2011).
3. Alternative algorithms that retrieve $X_{CO2}$ from GOSAT.

A total of 1,718,561 GOSAT observations globally over land between June 2009 and May 2010 have been processed using the UoL-FP retrieval algorithm (v3G), reducing to 764,290 scenes (44.47%) once pre-screened. These were then cloud-screened, leaving 372,479 scenes (21.67%) that were retrieved. Details of this are given in Table 4.1. Additionally, a total of 306,890 GOSAT observations over land for thirteen TCCON
sites between April 2009 and May 2011. Once pre-screened and cloud-screened 24,980 (8.11%) observations remained, details of which can be found in Table 4.2.

Table 4.1. UoL-FP v3G retrieval statistics for 1 year of daytime, land only global GOSAT observations.

<table>
<thead>
<tr>
<th>Year</th>
<th>Total from L1B</th>
<th>Passed Pre-Screen</th>
<th>O2 Completed</th>
<th>Cloudy Exposures</th>
<th>Clear Exposures</th>
<th>Retrieval Completed</th>
<th>Passed Post-Screen</th>
</tr>
</thead>
<tbody>
<tr>
<td>200906</td>
<td>126745</td>
<td>55303</td>
<td>51768</td>
<td>27863</td>
<td>23905</td>
<td>22881</td>
<td>5724</td>
</tr>
<tr>
<td>200907</td>
<td>139663</td>
<td>61514</td>
<td>57799</td>
<td>31098</td>
<td>26701</td>
<td>25660</td>
<td>7474</td>
</tr>
<tr>
<td>200908</td>
<td>139615</td>
<td>60201</td>
<td>56234</td>
<td>30330</td>
<td>25904</td>
<td>24928</td>
<td>7306</td>
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<tr>
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<td>62622</td>
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<td>29217</td>
<td>27771</td>
<td>27012</td>
<td>6761</td>
</tr>
<tr>
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<td>70907</td>
<td>63121</td>
<td>27433</td>
<td>35688</td>
<td>34135</td>
<td>6197</td>
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<td>72724</td>
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<td>4563</td>
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<td>58267</td>
<td>54104</td>
<td>29010</td>
<td>25094</td>
<td>23719</td>
<td>5015</td>
</tr>
<tr>
<td>Total</td>
<td>1718561</td>
<td>764290</td>
<td>697038</td>
<td>324559</td>
<td>372479</td>
<td>352937</td>
<td>62278</td>
</tr>
</tbody>
</table>

4.2 Comparisons with TCCON CO2 measurements

TCCON is a network of ground-based, solar absorption, near infrared, Fourier transform spectrometers that measure atmospheric columns of the gases CO2, CO, CH4, H2O and others with a precision of 0.25% for CO2 (Wunch et al. 2010, 2011a), making it an ideal dataset for validation of retrieved XCO2 from GOSAT. Since the TCCON instruments are direct solar-viewing, the effects of aerosol and high cirrus cloud are negligible. To maintain consistency between different TCCON observatories, all sites use the same instrumentation and the same software for data processing and analysis. This includes the conversion of raw interferograms into spectra, which are then spectrally fitted using the GFIT algorithm with subsequent quality controls applied to the column abundances (Wunch et al. 2011a). Wunch et al. (2010) compared the retrieved XCO2 from TCCON with aircraft observations using 14 coincident profiles and found that a single, global calibration factor of 0.989 accurately matches the data within the error. Subsequently, Messerschmidt et al. (2011) found additional European TCCON sites to be consistent with this calibration factor. Additionally, an airmass dependent correction factor was applied as described in Wunch et al. (2011a).
$X_{CO_2}$ retrievals from GOSAT were performed between April 2009 and May 2011 over thirteen TCCON sites: Lamont/USA, Park Falls/USA, Orleans/France, Bialystok/Poland, Bremen/Germany, Karlsruhe/Germany, Garmisch/Germany, Sodankyla/Finland, Tsukuba/Japan, Izana/Spain, Darwin/Australia, Wollongong/Australia, and Lauder/New Zealand. Figure 4.1 shows $X_{CO_2}$ retrieved from GOSAT compared with $X_{CO_2}$ inferred from TCCON for each of these thirteen TCCON sites, where GOSAT observations within a $5^\circ$ radius are used. The lower right panel of Figure 4.1 gives the mean bias, standard deviation and correlation of all daily mean coincident GOSAT $X_{CO_2}$ data compared to TCCON daily mean coincident $X_{CO_2}$ data. This gives an overall bias of -0.08 ppm, scatter of 2.52 ppm and correlation of 0.72 between GOSAT and TCCON $X_{CO_2}$.

Although there are few daily coincidences between GOSAT and TCCON over Izana, there is a good agreement between the single soundings for Izana, with GOSAT reproducing the seasonal cycle very well but with a mean bias of approximately -3.36 ppm compared to TCCON. Although the bias is small for Lauder (-0.44 ppm), there are very few coincident points with TCCON and those that are coincident have a large standard deviation of 4.11 ppm (using daily mean data). Similarly, Tsukuba has a low bias (-0.35 ppm), a large standard deviation (3.42 ppm) and only a small number of coincident points between GOSAT and TCCON observations. Unlike most other sites, Tsukuba TCCON data was not available for the duration of the GOSAT observations. Furthermore, there was a lack of TCCON data available for the Karlsruhe and Sodankyla TCCON sites, reducing the number of GOSAT/TCCON coincidences in those locations. Since few GOSAT observations exist over Sodankyla, due to clouds, this site does not provide statistically meaningful data. The other sites all provide at least 200 coincidences between GOSAT and TCCON.

Figure 4.2 gives the locations of each TCCON site and the locations of pre-screened and cloud-screened coincident GOSAT observations. No GOSAT observations are actually observed directly at Izana, instead the coincident observations are located over the Sahara desert. Only Lamont and Parkfalls have purely land within a $5^\circ$ radius of the site, all other sites include areas of ocean where observations either do not exist or are removed in the pre-screen. Furthermore, Izana, Lauder and Tsukuba are located on islands, thus reducing the number of GOSAT/TCCON coincidences in those locations. Since few GOSAT observations exist over Sodankyla, due to clouds, this site does not provide statistically meaningful data. The other sites all provide at least 200 coincidences between GOSAT and TCCON.

Bremen, Karlsruhe, and Garmisch TCCON sites are all located nearby each other, with overlapping coincidence regions. Many GOSAT observations are coincident with all three of these TCCON sites, hence making the overall GOSAT/TCCON comparison
statistics biased towards these observations. Some of the GOSAT observations coincident with these three sites are also coincident with the Orleans TCCON site. Bremen has the least number of coincident GOSAT observations that are also coincident with the Orleans site. For these reasons, Garmisch and Karlsruhe will no longer be used in this comparison.

Figure 4.3 shows the statistics of $X_{CO_2}$ retrieved from GOSAT compared to TCCON for different spatial coincidence limits. The number of coincident observations (or sample size) increases with larger spatial coincidence thresholds, simply due to GOSAT normally observing in a regular spatial pattern (such as the 5-point cross-track mode). Hence, the choice of coincidence criteria should not only be a compromise between the sample size and the spatial distance from a TCCON site, but must also take into account the statistics between GOSAT and TCCON coincident data.

In general, the $X_{CO_2}$ bias between GOSAT and TCCON observations improves with a smaller spatial coincidence (using daily mean coincident data). However, with a latitudinal coincidence less than about 2° the bias becomes worse. The bias remains almost constant below a longitudinal coincidence of 5°, whilst higher longitudinal coincidence limits produce larger biases. The bias calculated from coincident single soundings shows the same patterns but with lower values overall.

The $X_{CO_2}$ standard deviation of daily mean coincident points is almost homogeneous for all spatial coincidence thresholds considered, with a mean value of $\sim 2.45$ ppm. However, the $X_{CO_2}$ standard deviation of coincident single soundings increases with bigger spatial coincidences (from $\sim 2.2$ ppm to $\sim 2.7$ ppm), especially with larger latitudinal coincidence limits. Similarly, the correlation coefficient of daily mean coincident GOSAT and TCCON $X_{CO_2}$ is fairly homogeneous for all spatial coincidence limits with a value of $\sim 0.76$, whereas the correlation coefficient of $X_{CO_2}$ from coincident single soundings decreases with larger spatial coincidences (from $\sim 0.76$ to $\sim 0.68$).

Using the statistics from both daily mean and single sounding coincident data, a latitude threshold higher than 2° and a longitude threshold less than or equal to 5° would provide the best GOSAT/TCCON comparison. A spatial coincidence of 5° would therefore provide a compromise between larger spatial coincidence limits with more observations and smaller latitude limits with reduced $X_{CO_2}$ standard deviations and correlation coefficients.

Figure 4.4 shows the statistics of $X_{CO_2}$ retrieved from GOSAT compared to TCCON for different temporal coincidence limits. In general, reduced temporal coincidence gives in a small improvement in the $X_{CO_2}$ bias and standard deviation, but the number of
observations increases with higher temporal limits. There are between 1 and 3 GOSAT overpassing orbits of each TCCON site, providing all GOSAT observations within ~70 minutes. Whereas, TCCON measurements are taken throughout daytime, observing the variability of airmass transport and source/sink processes (Butz et al. 2011). Since the diurnal cycle of CO₂ changes rapidly, a large temporal coincidence for TCCON would not reflect the same airmass as observed by GOSAT. Thus, as a compromise between the number of observations and the diurnal cycle a temporal coincidence of ±2 hours would be suitable.

All pre-screened, cloud-screened and post-screened GOSAT measurements over land within a coincidence criteria of 5° of each TCCON site and ±2 hours of TCCON observations were used for only the seven TCCON sites selected. Figure 4.5 shows a comparison of the retrieved X_CO₂ from GOSAT with coincident TCCON X_CO₂ data for these seven TCCON sites (without averaging kernels applied). It also shows the correlation of coincident daily mean GOSAT X_CO₂ with the average of all TCCON X_CO₂ within ±2 hours of each coincident GOSAT data. GOSAT X_CO₂ is observed to have values consistent with TCCON retrievals for all sites for the entire time period and a very good agreement between coincident data is found. In particular, the seasonal cycle observed for northern hemispheric sites is well reproduced by GOSAT with correlation coefficients between 0.56 and 0.85, similar to that observed by (Butz et al. 2011, Oshchepkov et al. 2012a, Wunch et al. 2011b). Consistent to the TCCON measurements, the two southern hemisphere sites, Darwin and Wollongong, show weaker seasonal cycles than the northern hemisphere sites. However, the retrieved X_CO₂ from GOSAT over Wollongong is found to have a slightly enhanced seasonal cycle.

Gaps exist in the GOSAT and TCCON time-series due to clouds and instrumental issues. The number of points per site varies mainly due to seasonal cloud cover and the number of overpassing orbits. The number of soundings of each station varies from 260 for Bremen to 2445 for Lamont, with the number of coincident days with TCCON between 19 and 261, respectively. The Lamont site has the largest sample size due to multiple orbit overpasses within the coincidence criteria and less clouds than the other sites.

Using the data for all sites (lower right in Figure 4.5), the average bias of GOSAT compared to TCCON is calculated as -0.20 ppm with a standard deviation of 2.26 ppm (using daily means), which is similar to that found by Butz et al. (2011) and lower than reported by other algorithms Crisp et al. (e.g. 2012), Morino et al. (e.g. 2011), Oshchepkov et al. (e.g. 2012a). Using single soundings instead of daily means, a standard deviation of 2.46 ppm is found. The average of the bias per station is 0.10 ppm with a range of
-0.87 to 0.77 ppm and a standard deviation of 0.56 ppm (using daily mean data). The mean correlation coefficient between GOSAT and TCCON observations is found to be 0.75. The mean bias and scatter are largely influenced by the large number of soundings over the Lamont site. The highest scatter is observed for Bremen, which has the lowest number of scenes (see Table 4.3 for details).

The standard deviation of the retrieved $X_{CO2}$ inferred from comparisons to coincident TCCON data is substantially (factor of ~2 on average) larger than that predicted from the a posteriori error, which is found on average to be 1.14 ppm. The additional scatter observed is likely introduced by cloud perturbations, variations of aerosols, and other geophysical parameters. A similar value is reported by O’Dell et al. (2012) from simulations.

There are some outliers between GOSAT and TCCON retrievals that have passed the screening and are likely due to scattering caused by undetected clouds or aerosols. Thus, the effect of applying a stricter post-screen was tested, where reduced upper threshold limits were applied for the retrieved ice optical depth (0.03 limit), the retrieved total optical depth (0.3 limit) and the ratio of retrieved band 1 albedo to band 3 albedo (2 limit). When applied, this caused many of the outliers to be removed, reducing the mean bias to -0.08 ppm and the scatter to 1.92 ppm (of daily means), but at the cost of the number of exposures (see Table 4.4 for details).

This comparison has not taken into account the effect of the different averaging kernels $A$ and a priori, which describe the sensitivity of a retrieval algorithm to the true state throughout the atmosphere (Rodgers 2000, Rodgers & Connor 2003). The retrieved $X_{CO2}$ for GOSAT and TCCON depends on the averaging kernel and the a priori used in the retrieval algorithm according to:

$$X_{CO2} = h^T x_a + h^T A (x - x_a)$$

(4.1)

where $h^T$ is the transpose of the pressure weighting function (O’Dell 2010), $x$ is the true VMR profile, and $x_a$ is the a priori VMR profile. A consequence of Equation 4.1 is that the true and the retrieved $X_{CO2}$ will differ if $A$ is different from the Unity matrix. This difference is referred to as the smoothing error and it will be different for the GOSAT and TCCON retrievals due to their different averaging kernels and a priori values.

An assessment of this effect on the comparison of $X_{CO2}$ retrievals from GOSAT and TCCON was performed by calculating the $X_{CO2}$ retrieved from GOSAT and TCCON using GEOS-Chem model (described in Section 4.3) calculations as the true CO$_2$ profile. Since the GEOS-Chem model calculations will not necessarily represent the true atmo-
spheric CO$_2$ profiles these values for smoothing error differences will only represent a rough estimate.

For each of the seven TCCON sites considered, GEOS-Chem was interpolated temporally to the observation time of TCCON and GOSAT using the GEOS-Chem grid cell that includes the TCCON site. The averaging kernels were applied to GEOS-Chem for the period of April 2009 to December 2010 where both model, TCCON and GOSAT data exist. Only TCCON data that was temporally closest to GOSAT data, which was coincident within ±2 hours and 5° of each TCCON site was used.

The mean smoothing errors of TCCON and GOSAT were calculated as 0.12 ppm and 0.27 ppm respectively. The mean smoothing error difference between the model X$_{CO2}$ with GOSAT averaging kernels applied and the model X$_{CO2}$ with TCCON averaging kernels applied was found to be -0.14 ppm. The average of the mean smoothing error difference per station is -0.22 ppm with Lamont having the smallest mean smoothing error difference of -0.05 ppm and Orleans having the largest mean smoothing error difference of -0.50 ppm. The standard deviation of the mean smoothing error difference per site is 0.18 ppm. The mean of the scatter of the smoothing error difference per station is 0.19 ppm, with a standard deviation of 0.09 ppm. The smoothing error difference is observed to have a small seasonal cycle dependence, with northern hemispheric sites varying from approximately -1.5 ppm to 0.5 ppm and a lower amplitude for southern hemisphere stations where the seasonal cycle is reduced. Hence, for southern hemispheric sites the difference of the smoothing errors has a very small impact on the comparison of GOSAT with TCCON. However, for northern hemispheric sites there is a seasonal cycle dependence between GOSAT and TCCON X$_{CO2}$ due to the smoothing error difference (see Figure 4.9). If the point-by-point smoothing error difference is applied to the coincident TCCON data, the mean bias of GOSAT compared to TCCON is found to increase by 0.14 ppm but the standard deviation and correlation coefficient remain the same (see Table 4.5 for further details). It cannot be expected that the estimates are necessarily representing the correct value for each individual sounding as this would require that the model reproduces the shape of the CO$_2$ profile at each time step and location. Indeed, it is found that including the smoothing error estimates from GEOS-Chem to the GOSAT-TCCON comparisons worsens the comparisons for some sites while it improves it for others.
Table 4.2. Statistics for the UoL-FP v3G comparison to all 13 TCCON stations.

<table>
<thead>
<tr>
<th>TCCON site</th>
<th>Total from L1B</th>
<th>Passed pre-screen</th>
<th>Percent cloudy</th>
<th>Passed post-screen</th>
<th>Coincident days</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodankyla (67.368° N)</td>
<td>35864</td>
<td>2858</td>
<td>74.25</td>
<td>83</td>
<td>19</td>
</tr>
<tr>
<td>Bialystok (53.230° N)</td>
<td>29376</td>
<td>8321</td>
<td>61.16</td>
<td>459</td>
<td>54</td>
</tr>
<tr>
<td>Bremen (53.100° N)</td>
<td>24054</td>
<td>5484</td>
<td>66.36</td>
<td>260</td>
<td>19</td>
</tr>
<tr>
<td>Karlsruhe (49.100° N)</td>
<td>30371</td>
<td>8706</td>
<td>65.62</td>
<td>383</td>
<td>47</td>
</tr>
<tr>
<td>Orleans (47.970° N)</td>
<td>24047</td>
<td>9511</td>
<td>61.32</td>
<td>715</td>
<td>47</td>
</tr>
<tr>
<td>Garmisch (47.476° N)</td>
<td>25505</td>
<td>8479</td>
<td>65.02</td>
<td>328</td>
<td>40</td>
</tr>
<tr>
<td>Park Falls (45.945° N)</td>
<td>33277</td>
<td>5785</td>
<td>55.52</td>
<td>421</td>
<td>86</td>
</tr>
<tr>
<td>Lamont (36.604° N)</td>
<td>30984</td>
<td>9888</td>
<td>39.44</td>
<td>2445</td>
<td>261</td>
</tr>
<tr>
<td>Tsukuba (36.051° N)</td>
<td>21636</td>
<td>3320</td>
<td>55.18</td>
<td>129</td>
<td>19</td>
</tr>
<tr>
<td>Izana (28.300° N)</td>
<td>11614</td>
<td>1094</td>
<td>11.52</td>
<td>230</td>
<td>8</td>
</tr>
<tr>
<td>Darwin (12.424° S)</td>
<td>17273</td>
<td>4511</td>
<td>39.64</td>
<td>990</td>
<td>54</td>
</tr>
<tr>
<td>Wollongong (34.406° S)</td>
<td>13182</td>
<td>4971</td>
<td>50.45</td>
<td>570</td>
<td>56</td>
</tr>
<tr>
<td>Lauder (45.038° S)</td>
<td>9707</td>
<td>1069</td>
<td>49.49</td>
<td>28</td>
<td>16</td>
</tr>
<tr>
<td>Total</td>
<td>306890</td>
<td>73997</td>
<td>56.49</td>
<td>7041</td>
<td>726</td>
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</tbody>
</table>
FIGURE 4.1. Comparison of retrieved $X_{CO_2}$ from GOSAT with retrieved $X_{CO_2}$ from thirteen TCCON site for coincident cloud-screened observations within 2 hours and 5° between April 2009 and May 2011. All TCCON data shown in light green, coincident daily mean TCCON shown in dark green, all GOSAT data shown in light red and daily mean GOSAT shown in dark red (with daily mean points with only one value per day are represented by open circles). The average difference between the daily means is given as the bias (ranging from -3.36 to 2.01 ppm), the standard deviation of daily means by the Std (between 1.89 and 4.11 ppm) and the correlation coefficient of daily means by $r$ (-0.10 at Darwin to 0.85 at Park Falls). The lower right panel gives the correlation of daily mean coincident retrieved GOSAT $X_{CO_2}$ with daily mean TCCON $X_{CO_2}$ within ±2 hours for the thirteen sites. This gives an overall bias of -0.08 ppm, scatter of 2.52 ppm and correlation of 0.72 between GOSAT and TCCON $X_{CO_2}$. There are no daily mean coincident TCCON points for Izana, so no statistics exist for this location and therefore had no effect on the overall bias, standard deviation or correlation coefficient.
**Figure 4.2.** Locations of pre-screened and cloud-screened TANSO-FTS observations between April 2009 and May 2011 within 5° of thirteen TCCON sites. The locations of the TCCON sites are represented by a blue circle and GOSAT observation locations are given by red circles. Additionally, a green surface represents land and white shows an ocean surface. The dotted line represents a 2° grid.
Figure 4.3. Statistics for a range of latitude and longitude coincidence limits for the comparison of GOSAT and TCCON $X_{CO2}$. The left column shows the mean bias (top left), standard deviation (left second row), correlation coefficient (left third row), and sample size of daily mean GOSAT $X_{CO2}$ coincident with TCCON. Similar statistics are shown in the right column where GOSAT single soundings are coincident with TCCON individual observations. Additionally, a temporal coincidence of 2 hours between GOSAT and TCCON observations was used.
Figure 4.4. Statistics for a range of latitude, longitude, and temporal coincidence limits for the comparison of GOSAT and TCCON X$_{CO2}$. The sample size of coincident daily mean data (left column), mean bias of coincident daily mean data (middle column), and standard deviation of coincident single soundings (right column) is shown for three different temporal coincidence limits of GOSAT and TCCON X$_{CO2}$. The top row shows the statistics for a temporal coincidence of 30 minutes between GOSAT and TCCON observations. Similarly, the middle and bottom row shows the statistics for a temporal coincidence of 2 hours and 6 hours, respectively.
FIGURE 4.5. Comparison of retrieved $X_{CO_2}$ from GOSAT with retrieved $X_{CO_2}$ from seven TCCON site for coincident cloud-screened observations within 2 hours and 5° between April 2009 and May 2011. All TCCON data shown in light green, coincident daily mean TCCON shown in dark green, all GOSAT data shown in light red and daily mean GOSAT shown in dark red (with daily mean points with only one value per day are represented by open circles). The average difference between the daily means is given as the bias (ranging from -0.87 to 0.77 ppm), the standard deviation of daily means by the Std (between 2.04 and 2.96 ppm) and the correlation coefficient of daily means by r (-0.10 at Darwin to 0.85 at Park Falls). The lower right panel gives the correlation of daily mean coincident retrieved GOSAT $X_{CO_2}$ with daily mean TCCON $X_{CO_2}$ within ±2 hours for the seven sites. This gives an overall bias of -0.20 ppm, scatter of 2.26 ppm and correlation of 0.75 between GOSAT and TCCON $X_{CO_2}$.

FIGURE 4.6. The Lamont/USA TCCON averaging kernel as a function of atmospheric pressure for different SZA (left). The right panel shows these averaging kernels for different SZA when interpolated onto a finer grid of 0.25°.
Table 4.3. Statistics for the UoL-FP v3G comparison to TCCON, where the bias, $\sigma$ and $r$ were calculated using coincident daily mean data and other parameters calculated from individual coincident points.

<table>
<thead>
<tr>
<th>TCCON site</th>
<th>Total from L1B</th>
<th>Passed pre-screen</th>
<th>Passed cloudy</th>
<th>Passed post-screen</th>
<th>Coincident days</th>
<th>Bias (ppm)</th>
<th>$\sigma$ (ppm)</th>
<th>$r$ (Pearson)</th>
<th>Single sounding $\sigma$ (ppm)</th>
<th>A Posterior error (ppm)</th>
<th>Mean smoothing error difference (ppm)</th>
<th>$\sigma$ smoothing error difference (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bialystok (53.230° N)</td>
<td>29376</td>
<td>8321</td>
<td>61.16</td>
<td>459</td>
<td>54</td>
<td>0.09</td>
<td>2.28</td>
<td>0.83</td>
<td>2.20</td>
<td>1.24</td>
<td>-0.21</td>
<td>0.32</td>
</tr>
<tr>
<td>Bremen (53.100° N)</td>
<td>24054</td>
<td>5484</td>
<td>66.36</td>
<td>260</td>
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<td>0.50</td>
<td>2.96</td>
<td>0.56</td>
<td>2.00</td>
<td>1.20</td>
<td>-0.06</td>
<td>0.22</td>
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<td>1.20</td>
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<td>0.26</td>
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<tr>
<td>Darwin (12.424° S)</td>
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<td>4511</td>
<td>39.64</td>
<td>990</td>
<td>54</td>
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<td>-0.10</td>
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<td>0.77</td>
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<td>2.90</td>
<td>1.09</td>
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<tr>
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<td>0.75</td>
<td>2.46</td>
<td>1.14</td>
<td>-0.14</td>
<td>0.26</td>
</tr>
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Table 4.4. Statistics for the UoL-FP v3G comparison to TCCON with stricter post-screen, where the bias, $\sigma$ and $r$ were calculated using coincident daily mean data and other parameters calculated from individual coincident points.

<table>
<thead>
<tr>
<th>TCCON site</th>
<th>Total from L1B</th>
<th>Passed pre-screen</th>
<th>Percent cloudy</th>
<th>Passed post-screen</th>
<th>Coincident days</th>
<th>Bias (ppm)</th>
<th>$\sigma$ (ppm)</th>
<th>$r$ (Pearson)</th>
<th>Single sounding $\sigma$ (ppm)</th>
<th>A Posterior error (ppm)</th>
<th>Mean smoothing error difference (ppm)</th>
<th>$\sigma$ smoothing error difference (ppm)</th>
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<tr>
<td>Bialystok (53.230° N)</td>
<td>29376</td>
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<td>Bremen (53.100° N)</td>
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<td>0.84</td>
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<tr>
<td>Lamont (36.604° N)</td>
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<td>9888</td>
<td>39.44</td>
<td>1448</td>
<td>194</td>
<td>-0.17</td>
<td>1.70</td>
<td>0.83</td>
<td>2.23</td>
<td>1.13</td>
<td>-0.05</td>
<td>0.24</td>
</tr>
<tr>
<td>Darwin (12.424° S)</td>
<td>17273</td>
<td>4511</td>
<td>39.64</td>
<td>711</td>
<td>47</td>
<td>-0.45</td>
<td>2.15</td>
<td>-0.06</td>
<td>2.04</td>
<td>0.85</td>
<td>-0.08</td>
<td>0.07</td>
</tr>
<tr>
<td>Wollongong (34.406° S)</td>
<td>13182</td>
<td>4971</td>
<td>50.45</td>
<td>363</td>
<td>48</td>
<td>0.70</td>
<td>2.14</td>
<td>0.48</td>
<td>1.96</td>
<td>1.04</td>
<td>-0.26</td>
<td>0.10</td>
</tr>
<tr>
<td>Total</td>
<td>172193</td>
<td>48471</td>
<td>53.57</td>
<td>3355</td>
<td>413</td>
<td>-0.08</td>
<td>1.84</td>
<td>0.81</td>
<td>1.92</td>
<td>1.11</td>
<td>-0.14</td>
<td>0.26</td>
</tr>
</tbody>
</table>
Figure 4.7. Example averaging kernels for TCCON and GOSAT over Lamont/USA (left) and the corresponding CO$_2$ VMR vertical profile (right). The estimated smoothing error for TCCON and GOSAT with respect to GEOS-Chem is given, and the smoothing error difference between TCCON and GOSAT is stated.
FIGURE 4.8. Correlation of GEOS-Chem with and without TCCON averaging kernels applied for data within 5° of seven TCCON sites between April 2009 and December 2010 (left). Similarly, a correlation between GEOS-Chem with and without GOSAT averaging kernels applied is shown in the centre. The mean difference (Diff) or smoothing error between with and without averaging kernels applied is given for TCCON and GOSAT. The smoothing error difference is estimated from a comparison between GEOS-Chem with TCCON and GOSAT averaging kernels applied (right).
Figure 4.9. Calculated smoothing error differences for seven TCCON sites, for coincident cloud-screened observations within 2 hours and $5^\circ$ between April 2009 and December 2010. The standard deviation of the smoothing error difference values are given by the Std (between 0.07 and 0.32 ppm).
Table 4.5. Statistics for the UoL-FP v3G comparison to TCCON with and without TCCON data being corrected by the smoothing error difference (SED) between GOSAT and TCCON. The bias, $\sigma$ and $r$ were calculated using coincident daily mean data and other parameters calculated from individual coincident points, between April 2009 and December 2010.

<table>
<thead>
<tr>
<th>TCCON site</th>
<th>Bias without SED (ppm)</th>
<th>$\sigma$ without SED (ppm)</th>
<th>$r$ without SED (Pearson)</th>
<th>Bias with SED (ppm)</th>
<th>$\sigma$ with SED (ppm)</th>
<th>$r$ with SED (Pearson)</th>
<th>Bias difference (ppm)</th>
<th>$\sigma$ difference (ppm)</th>
<th>$r$ difference (Pearson)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bialystok (53.230° N)</td>
<td>-0.13</td>
<td>2.34</td>
<td>0.73</td>
<td>-0.34</td>
<td>2.47</td>
<td>0.7</td>
<td>-0.21</td>
<td>0.13</td>
<td>-0.03</td>
</tr>
<tr>
<td>Bremen (53.100° N)</td>
<td>0.43</td>
<td>2.73</td>
<td>0.64</td>
<td>0.39</td>
<td>2.77</td>
<td>0.64</td>
<td>-0.04</td>
<td>0.04</td>
<td>0.00</td>
</tr>
<tr>
<td>Orleans (47.970° N)</td>
<td>-0.44</td>
<td>2.48</td>
<td>0.73</td>
<td>-0.91</td>
<td>2.44</td>
<td>0.74</td>
<td>-0.47</td>
<td>-0.04</td>
<td>0.00</td>
</tr>
<tr>
<td>Park Falls (45.945° N)</td>
<td>-0.37</td>
<td>2.84</td>
<td>0.77</td>
<td>-0.71</td>
<td>2.85</td>
<td>0.77</td>
<td>-0.34</td>
<td>0.01</td>
<td>0.00</td>
</tr>
<tr>
<td>Lamont (36.604° N)</td>
<td>-0.08</td>
<td>2.23</td>
<td>0.66</td>
<td>-0.09</td>
<td>2.23</td>
<td>0.66</td>
<td>-0.01</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Darwin (12.424° S)</td>
<td>-0.96</td>
<td>2.36</td>
<td>0.09</td>
<td>-1.06</td>
<td>2.35</td>
<td>0.10</td>
<td>-0.10</td>
<td>-0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Wollongong (34.406° S)</td>
<td>0.91</td>
<td>2.47</td>
<td>0.24</td>
<td>0.67</td>
<td>2.47</td>
<td>0.24</td>
<td>-0.24</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Total</td>
<td>-0.15</td>
<td>2.44</td>
<td>0.66</td>
<td>-0.29</td>
<td>2.44</td>
<td>0.66</td>
<td>-0.14</td>
<td>0.00</td>
<td>0.00</td>
</tr>
</tbody>
</table>
4.3 Comparisons to the GEOS-Chem model

One year of GOSAT $X_{CO_2}$ retrievals was compared with CO$_2$ atmospheric concentrations from the GEOS-Chem global 3-D chemistry transport model (v8-02-01) (Feng et al. 2011). The GEOS-Chem simulations use assimilated GEOS-5 meteorology from the Global Modeling and Assimilation Office based at NASA Goddard to drive the model. For the surface CO$_2$ fluxes, GEOS-Chem assimilates annual fossil fuel emissions estimated from CDIAC, biofuel emissions obtained from Yevich & Logan (2003) climatology, monthly biomass burning emissions extracted from the third version of the Global Fire Emission Database (GFEDv3) that uses observations of the land surface by ground-based and satellite instruments, monthly ocean fluxes that are derived from sea-surface pCO$_2$ observations (Takahashi et al. 2009), and biospheric fluxes that are computed using the CASA biosphere model (Randerson et al. 1997). The CASA model is three hourly and is constrained by GEOS meteorology output and Normalized Difference Vegetation Index (NDVI). The mixing depths and surface fields are updated every three hours and the GEOS-5 meteorology is updated every six hours. The 4D fields include 47 vertical levels and a horizontal grid resolution of 2$^\circ$ latitude by 2.5$^\circ$ longitude.

The GEOS-Chem simulations are forced by posterior fluxes inferred from the GLOBALVIEW CO$_2$ product, that includes 2009 and 2010 data (Maserie & Tans 1995). The GLOBALVIEW data from 78 surface sites are assimilated to estimate monthly surface fluxes over 144 global regions by using an ensemble Kalman Filter (EnKF) (Feng et al. 2009, 2011). Feng et al. (2011) reported that the GEOS-Chem model is within 1.5 ppm of free and upper troposphere aircraft vertical profile measurements and CO$_2$ retrieved from AIRS observations. Additionally, GEOS-Chem is on average within 0.5 to 1.0 ppm of observed partial CO$_2$ columns from the HIAPER HIPPO project (James Barlow, personal communication, 2012). The GEOS-Chem model vertical transport errors are estimated to be less than $\sim$2 ppm. This is supported by complementary model evaluation studies of CH$_4$, CH$_3$CCl$_3$, and SF$_6$ (e.g., Fraser et al. 2011, Patra et al. 2011). TCCON $X_{CO_2}$ was compared with that of GEOS-Chem (convolved with the TCCON averaging kernels) for each of the seven sites considered in the previous section and GEOS-Chem was found to have a mean bias of -0.09 ppm, standard deviation of 1.23 ppm and correlation coefficient of 0.93, comparing better than GOSAT with TCCON (see Figure 4.10). Based on the current understanding of model performance it can be concluded that the model should well reproduce large scale features of the atmospheric CO$_2$ distribution and that differences between GOSAT and GEOS-Chem are likely due to GOSAT retrieval biases. Although, model errors could also contribute to any observed differences.
Figure 4.10. Comparison of $X_{CO_2}$ calculated from GEOS-Chem (with TCCON averaging kernels applied) with retrieved $X_{CO_2}$ from seven TCCON site for cloudscreened observations within 5° between April 2009 and May 2011. All TCCON data shown in light green, coincident daily mean TCCON shown in dark green, all GEOS-Chem data shown in light red and daily mean GEOS-Chem shown in dark red (with daily mean points with only one value per day are represented by open circles). The average difference between the daily means is given as the bias (ranging from -0.56 to 0.79 ppm), the standard deviation of daily means by the Std (between 0.80 and 1.49 ppm) and the correlation coefficient of daily means by $r$ (0.80 at Darwin to 0.97 at Orleans). The lower right panel gives the correlation of daily mean retrieved GEOS-Chem $X_{CO_2}$ with daily mean TCCON $X_{CO_2}$ for the seven sites. This gives an overall bias of -0.09 ppm, scatter of 1.23 ppm and correlation of 0.93 between GEOS-Chem and TCCON $X_{CO_2}$.

Figure 4.11 shows the comparison between $X_{CO_2}$ from GOSAT and GEOS-Chem for each season, where the GEOS-Chem CO$_2$ profiles were interpolated to the locations and times of the GOSAT observations, and convolved with the averaging kernel (Equation 4.1) to obtain $X_{CO_2}$ as measured by GOSAT. Overall, the spatial and temporal distribution of $X_{CO_2}$ agrees well between GOSAT and GEOS-Chem. The mean difference observed between GOSAT and GEOS-Chem $X_{CO_2}$ varies from -1.43 to -0.82 ppm seasonally, with the standard deviation ranging from 2.41 to 2.71 ppm. High correlations of the seasonal latitudinal gradient are found with correlation coefficients ranging between 0.73 and 0.96, but slightly poorer point-by-point correlation coefficients with values be-
Figure 4.11. Comparison of $X_{CO_2}$ retrieved from GOSAT and $X_{CO_2}$ calculated from GEOS-Chem CO$_2$ profiles convolved with the scene-specific GOSAT averaging kernel. For each season, GOSAT is shown at the top, GEOS-Chem in the middle, and a zonal mean comparison of them at the bottom. The global maps indicate the largest differences are observed over desert regions, such as the Sahara and central Asia. The zonal mean comparison shows a very good agreement between GOSAT and GEOS-Chem. It also includes the average TCCON $X_{CO_2}$ from each site which in general agrees, with observed differences mostly due to zonal averaging of GOSAT/GEOS-Chem data.

Over desert regions such as the Sahara, Saudi Arabia and Australia, differences are found between GOSAT and GEOS-Chem of up to 3 ppm. The medium gain mode is predominantly used over desert areas where surface reflectance causes the signal intensity to be high. Suto et al. (2008) found that TANSO-FTS instrument micro-vibrations produced a larger effect on medium gain observations and $X_{CO_2}$ retrievals of medium gain have been observed to be $\sim$1% higher than that of high gain (Crisp et al. 2012). However, $X_{CO_2}$ retrieved from the UoL-FP v3G algorithm for desert regions observed with the high gain mode, such as central Asia, are found to also give similar differences in $X_{CO_2}$ compared to GEOS-Chem. Since all these regions include high albedo in the
1.61 and 2.06 $\mu$m CO$_2$ bands and high levels dust, the $X_{CO2}$ difference is not solely due to gain issues (as suggested by Crisp et al. (2012)) but also potentially an aerosol or other instrumental related issue.

Figure 4.12. Comparison of $X_{CO2}$ retrieved from GOSAT and $X_{CO2}$ calculated from GEOS-Chem CO$_2$ profiles convolved with the scene-specific GOSAT averaging kernel, for a number of different regions between June 2009 and May 2010. The locations of these regions are indicated on the map by the black boxes (top left). The top centre and top right panels give the time series of GOSAT and GEOS-Chem data for the northern and southern hemispheres, showing the seasonal cycle to be well matched but with GOSAT data offset lower by -1.50 ppm in the northern hemisphere and -0.63 ppm in the southern hemisphere. The second row of plots show a very good agreement between GOSAT and GEOS-Chem both in value and seasonality for regions containing urban environments. Below these, the time series of forested regions are shown, with GOSAT reproducing the GEOS-Chem data very well but with a larger scatter and fewer data points. The bottom panels show the time series for 3 different desert regions, with GOSAT matching the seasonal cycle of GEOS-Chem very well but offset lower in value from -0.86 to -2.99 ppm.

The time series of $X_{CO2}$ from GOSAT and GEOS-Chem for a number of different regions are shown in Figure 4.12. The seasonal cycle is found to match very well between
GOSAT and GEOS-Chem for both northern and southern hemispheres, but a difference of 1.5 ppm is observed in the northern hemisphere throughout the year and a smaller difference of 0.63 ppm in the southern hemisphere. GOSAT and GEOS-Chem $X_{CO2}$ show a high consistency for both Europe and the USA, with a $X_{CO2}$ difference of 0.20 ppm and 0.56 ppm and correlation coefficients of 0.73 and 0.70, respectively. On average, GOSAT and GEOS-Chem agree for South Asia with a mean difference of 0.53 ppm and correlation coefficient of 0.68, but differences are viewed in the seasonal cycle with GEOS-Chem over-estimating the $X_{CO2}$ in autumn 2009 and under-estimating during spring 2010.

Few GOSAT soundings are observed over the Amazon due to tropical clouds, and those measurements that are retrieved have a large standard deviation (3.56 ppm) that is potentially due to partial cloud contamination. Similarly, a larger scatter is seen in tropical Africa during autumn and spring for GOSAT, whereas winter and summer have a reduced scatter and have similar values as GEOS-Chem. A very high agreement is observed for NW Russia with a correlation coefficient of 0.81 and a mean $X_{CO2}$ difference of 0.43 ppm.

In central Asia where a desert region is observed with the high gain mode, the seasonal cycle of GOSAT and GEOS-Chem is observed to agree well ($r=0.76$), but consistently with a $\sim 2$ ppm difference in value. Also, a $\sim 3$ ppm mean difference is found over the Sahara between GOSAT and GEOS-Chem $X_{CO2}$ where a desert region is observed with the medium gain mode, but the seasonal cycle agrees well with a correlation coefficient of 0.80. Australia, which comprises partially of desert and is observed partially with medium and high gain modes, also shows a difference in $X_{CO2}$ between GOSAT and GEOS-Chem with a value of 0.86 ppm.
4.4 Bias Correction Scheme

A bias correction method was developed to help identify and reduce the observed biases of retrieved $X_{CO2}$ from GOSAT, especially over desert areas. Similar to Wunch et al. (2011b), this bias correction method is based on a multivariate linear regression of the difference between pseudo observations and retrieved $X_{CO2}$ from GOSAT between July 2009 to June 2010 over the region south of 25°S where the variability of CO$_2$ is low. The pseudo-observations were generated using GEOS-Chem calculations to provide information on vertical profile and spatial variations while the temporal variations and absolute values were scaled to match Wollongong/Australia (34.406°S) and Lauder/New Zealand (45.038°S) TCCON measurements. Initially all retrieval parameters and observation related parameters were used. Similar to Wunch et al. (2011b), the four parameters with the highest correlation were identified and used to obtain the following equation:

$$X_{CO2}^{\text{bias-corrected}} \text{[ppm]} = X_{CO2}^{\text{retrieved}} \text{[ppm]} + 4.19 - 0.564 \times \text{CO}_2\_\text{Signal\_Ratio} - 0.193 \times (\Delta P \text{[hPa]}) + 56.8 \times \text{Ice\_Optical\_Depth} + 0.256 \times 10^{-5} \times \text{O}_2\_\text{Albedo\_Slope} \quad (4.2)$$

These parameters have been identified by the regression which does not provide a means of identifying the physical link. A correlation ($r=0.37$) of the $X_{CO2}$ difference is observed with the ratio of the mean signal of the 1.61 $\mu$m CO$_2$ band to the 2.06 $\mu$m CO$_2$ band ($\text{CO}_2\_\text{Signal\_Ratio}$), which may be a consequence of the spectroscopic differences of the different CO$_2$ bands, but might also relate to aerosols or albedo. Spectroscopic errors may also result in systematic effects in the retrieved surface pressure as shown by Wunch et al. (2011b) and the $X_{CO2}$ difference is found to correlate ($r=0.30$) with delta surface pressure ($\Delta P$), even though the $X_{CO2}$ is already normalised with the surface pressure ratio. An anti-correlation ($r=-0.22$) of the $X_{CO2}$ difference with retrieved cirrus optical depth ($\text{Ice\_Optical\_Depth}$) is observed which indicates the path length was altered. Based on the geographical location of the bias correction for this parameter it can be inferred that the ice optical depth refers to cirrus clouds. An anti-correlation ($r=-0.29$) of the $X_{CO2}$ difference is also found with the slope of the band 1 albedo ($\text{O}_2\_\text{Albedo\_Slope}$) which may be due to errors in the O$_2$ A band spectroscopy, zero-level offsets, or variations in signal intensity over vegetation regions that could be caused by not accounting...
Figure 4.13. Illustration of $X_{CO_2}$ calculated from GEOS-Chem within 5° of Wollongong/Australia and Lauder/New Zealand combined between April 2009 and December 2010 with TCCON averaging kernels applied (dark green) compared to $X_{CO_2}$ inferred from TCCON (light green). When the daily mean difference between GEOS-Chem and TCCON is applied as a ±5 day running mean to the GEOS-Chem data, the GEOS-Chem $X_{CO_2}$ becomes closer in value to TCCON, as shown in Red.

For the known fluorescence (Frankenberg et al. 2011a).

Whilst Wunch et al. (2011b) found biases partially due to airmass and the $O_2$ A band signal, here no correlation is found between the airmass and the $X_{CO_2}$ difference ($r = 0.01$) nor the $O_2$ A band signal ($r=0.01$). Additionally, a zero level offset in the $O_2$ A band is already retrieved to mitigate the effects of the GOSAT Band 1 non-linearity and partially fluorescence (Butz et al. 2011, Frankenberg et al. 2011a, 2012, Suto et al. 2008).

This bias correction was applied to one year (June 2009 to May 2010) of $X_{CO_2}$ retrieved from GOSAT globally and compared to $X_{CO_2}$ calculated from the GEOS-Chem model with GOSAT averaging kernels applied. The annual mean global difference is found to be reduced from -1.22 ppm to -0.68 ppm and the correlation to increase from 0.61 to 0.74.

Figure 4.15 shows a comparison between GOSAT and GEOS-Chem $X_{CO_2}$ latitudinal gradients for each season, comparing with and without the bias correction. The seasonal latitudinal gradients show improvements with increases of the correlation between 0.01 and 0.18 depending on the season, for DJF the improvement in correlation coefficient is marginal. The majority of the improvement occurs in the northern hemisphere, especially
**Figure 4.14.** Correlation of the $X_{CO2}$ bias with the ratio of band 2 to band 3 mean radiance (top left), the retrieved cirrus optical depth (top right), the $\Delta P$ (bottom left), and the retrieved band 1 albedo slope (bottom right). Where the bias is between GOSAT and the GEOS-Chem 3D chemistry transport model with GOSAT averaging kernels applied and corrected with a fit to Wollongong and Lauder TC-CON measurements. The colour gives the density of values within the bin range.

Over the Sahara region.

The time series of bias corrected $X_{CO2}$ retrieved from GOSAT compared to GEOS-Chem data for different regions is shown in Figure 4.16. For most regions it is found that the bias, standard deviation and correlation coefficient have improved. In particular, the bias correction has considerably reduced the larger differences previously seen over desert regions; the difference over the Sahara has reduced from -2.99 ppm to -1.20 ppm, the difference over Australia has reduced to about half, and the difference over central Asia improved from -2.08 ppm to -0.94 ppm. However, for the Sahara region, the standard deviation and correlation coefficient becomes worse. Also, for some regions (Tropical Africa and NW Russia) it is found that the bias increases when applying the bias correction.
Figure 4.15. The top panels show zonal mean comparisons for each season of $X_{CO2}$ retrieved from GOSAT and $X_{CO2}$ calculated from GEOS-Chem CO$_2$ profiles convolved with the scene-specific GOSAT averaging kernel. The bottom panel gives revised comparisons where the GOSAT $X_{CO2}$ was bias corrected. The latitudinal gradients of the bias corrected GOSAT $X_{CO2}$ match very well to GEOS-Chem with correlations between 0.78 and 0.97. The bias correction leads to increases of the correlation between 0.01 and 0.18, and shows a large improvement over the latitudes that contain the Sahara.
Figure 4.16. Comparison of bias corrected $X_{CO_2}$ retrieved from GOSAT and $X_{CO_2}$ calculated from GEOS-Chem CO$_2$ profiles convolved with the scene-specific GOSAT averaging kernel, for a number of different regions between June 2009 and May 2010. The locations of these regions are indicated on the map by the black boxes (top left). GOSAT agrees very well with GEOS-Chem for both hemispheres and all regions. GOSAT shows a good reproduction of the seasonal cycle of GEOS-Chem data, with the exception of south Asia where GOSAT is higher in the summer period. Minor offsets exist over the desert regions, such as Australia, the Sahara, and central Asia, which is a large improvement compared to the non-bias corrected $X_{CO_2}$ shown in Figure 4.12. Additionally, the scatter of all regions is less than that of the non-bias corrected $X_{CO_2}$. Furthermore, the bias correction has brought the $X_{CO_2}$ over the Amazon to be of the same magnitude as GEOS-Chem.
4.5 Comparisons of GOSAT with Land Surface Types

The $X_{CO_2}$ difference between GOSAT and GEOS-Chem was compared against the land surface type for each observation. The surface type was determined by using the most common land cover type within a coincidence of $\pm 0.1^\circ$ of the Moderate Resolution Imaging Spectroradiometer (MODIS) Level 3 yearly 0.05$^\circ$ land cover type (MCD12C1) for 2007 (assuming the land cover has not significantly altered between 2007 and 2010). Figures 4.17 and 4.18 show this comparison for each of the GOSAT gain modes separately.

For retrievals of GOSAT high gain data, GOSAT and GEOS-Chem correlate well with $X_{CO_2}$ differences less than 0.23 ppm for surfaces that consist of forest, cropland, savanna, wetland, or urban environments. However, larger $X_{CO_2}$ differences (up to $\sim 3.2$ ppm) are found for surfaces that consist of desert, shrubland, grassland or snow.

Where medium gain data has been retrieved, $X_{CO_2}$ differences are found to be similar to that of high gain data. The retrieved $X_{CO_2}$ from GOSAT is found on average to be $\sim 1.5$ ppm lower than GEOS-Chem over both shrubland and grassland surfaces. The largest $X_{CO_2}$ difference of 2.75 ppm is observed over desert surfaces. The lowest difference (0.35 ppm) is found over savanna surfaces. No observations over other surface types exist due to the specific observation locations where medium gain was used. The $X_{CO_2}$ differences are found to be larger with certain land surface types, specifically deserts, irrespective of the gain setting used.
FIGURE 4.17. Comparison of $X_{CO_2}$ retrieved from high gain GOSAT data with GEOS-Chem $X_{CO_2}$ calculations for different MODIS land cover types. The red dashed line shows the one-to-one line, B represents the mean $X_{CO_2}$ difference, and $\sigma$ gives the standard deviation of $X_{CO_2}$. GOSAT and GEOS-Chem correlate well with $X_{CO_2}$ differences less than 0.23 ppm for surfaces that consist of forest, cropland, savanna, wetland, or urban environments. However, larger $X_{CO_2}$ differences (up to $\sim$3.2 ppm) are found for surfaces that consist of desert, shrubland, grassland or snow.
Figure 4.18. Comparison of $X_{CO_2}$ retrieved from medium gain GOSAT data with GEOS-Chem $X_{CO_2}$ calculations for different MODIS land cover types. The red dashed line shows the one-to-one line, B represents the mean $X_{CO_2}$ difference, and $\sigma$ gives the standard deviation of $X_{CO_2}$. The retrieved $X_{CO_2}$ from GOSAT is found on average to be $\sim$1.5 ppm lower than GEOS-Chem over both shrubland and grassland surfaces. The largest $X_{CO_2}$ difference of 2.75 ppm is observed over desert surfaces. The lowest difference (0.35 ppm) is found over savanna surfaces.
4.6 Comparisons to alternative retrieval algorithms

The retrieved $X_{\text{CO}_2}$ from GOSAT using the UoL-FP algorithm was compared with five alternative algorithms for two years of data over 11 TCCON sites; Lamont/USA, Park Falls/USA, Sodankyla/Finland, Bialystok/Poland, Bremen/Germany, Orleans/ France, Garmisch/Germany, Tsukuba/Japan, Darwin/Australia, Wollongong/Australia and Lauder/ New Zealand. The following sections give a brief description of each algorithm (further details are given in Oshchepkov et al. (2012b)).

4.6.1 NIES PPDF D Retrieval Algorithm

The National Institute for Environmental Studies (NIES) Photon Path Length Probability Density Function (PPDF) Differential Optical Absorption Spectroscopy (DOAS) retrieval algorithm (NIES PPDF D) uses the maximum a posteriori method of optimal estimation (Rodgers 2000) with the DOAS technique (Oshchepkov et al. 2012a). It includes the HSTAR, high resolution RSTAR, radiative transfer model (Kokhanovsky et al. 2010), based on a discrete ordinate algorithm (Duan et al. 2005) and uses tabulated absorption cross-sections of HITRAN08 (Rothman et al. 2009). The solar model is based on a high resolution solar irradiance database provided by Dr. R. Kurucz (http://kurucz.harvard.edu/sun/irradiance2008/). The instrument model uses the official GOSAT ILS from JAXA (version 006) (Kuze et al. 2009). It applies a pre-screen that uses a PPDF based on the radiative transfer model (Bril et al. 2007, S. Oshchepkov & Yokota 2008, Oshchepkov et al. 2009, 2011, 2012a) and the NIES cloud-screen (Taylor et al. 2012). Only the 1.61 $\mu$m band was retrieved within the wavenumber range of 6180-6380 cm$^{-1}$.

The a priori CO$_2$ and CH$_4$ were obtained from daily profiles of the NIES TM global atmospheric transport model (Maksyutov et al. 2008). The a priori H$_2$O, temperature and surface pressure were extracted from 6 hourly model outputs from the Japan Meteorological Agency (JMA) on 0.5 degree horizontal resolution with 21 vertical levels. The albedo a priori was taken from the 16-daily MODIS land surface albedo product (MOD43B3) (Schaaf et al. 2002). The main differences between the retrieval algorithms include the; a priori, pre-screen, radiative transfer model, and solar model.
4.6.2 NIES L2 Retrieval Algorithm

The NIES L2 algorithm uses the maximum a posteriori method of optimal estimation with the Levenberg-Marquardt method (Rodgers 2000) and an accurate single scattering and approximate multiple scattering radiative transfer model of Duan et al. (2005). It uses the same absorption cross-sections, solar model, ILS and a priori as the NIES PPDF D retrieval algorithm. The polarisations were combined using a preliminary Mueller Matrix providing a single intensity to be retrieved. It pre-screens for saturated H_2O absorption lines as a test for cirrus clouds and uses the NIES cloud-screen (Taylor et al. 2012). The state vector consists of CO_2, CH_4 and H_2O 15-level VMR profiles, uniform 0-2 km AOD, surface pressure, albedo, dispersion and a temperature scaling factor. Only the 1.61 µm band was retrieved with the wavenumber ranges of 6180-6380 cm^{-1} for CO_2 and 5900-6150 cm^{-1} for CH_4. Additionally, all solar absorption lines were omitted from the retrieval.

The a priori aerosol optical properties and AOD were obtained from semi-real time daily outputs of the three-dimensional aerosol transport model, the Spectral Radiation-Transport Model for Aerosol Species (SPRINTARS) (Takemura et al. 2003), for soil dust, carbonaceous, sulfate, and sea-salt aerosols uniformly distributed within a 2-km layer from the surface (Yoshida et al. 2010). The key differences between the retrieval algorithms include the; a priori, pre-screen, radiative transfer model, and solar model, scattering approach, and the intensity used.

4.6.3 ACOS Retrieval Algorithm

The ACOS retrieval algorithm used for the NASA Atmospheric CO_2 Observations from Space (ACOS) project and the NASA OCO-2 mission is also based on the OCO algorithm (like the UoL-FP retrieval algorithm) but has been re-developed (O’Dell et al. 2012, Crisp et al. 2012). Here, it used HITRAN08 absorption cross-sections (Rothman et al. 2009) with additional line mixing in the O_2 A band. It simultaneously retrieved the O_2 A band, the 1.61 µm and 2.06 µm CO_2 bands with the average intensity of each polarisation. The state vector consisted of a 20-level CO_2 VMR profile and 20-level logarithmic extinction profiles of two aerosols, cloud and cirrus. Additionally, surface pressure was retrieved as well as a temperature offset, a scaling factor for H_2O, albedo, albedo slope and dispersion shift/stretch.

It obtained its a priori surface pressure, temperature, and water vapour from ECMWF. The a priori CO_2 VMR was extracted from latitudinal zonal land/ocean LMDZ 2004
model fields. The a priori albedo was computed from GOSAT reflectance. The a priori aerosol, cloud and cirrus profiles were constructed to provide a low altitude aerosol, a middle altitude aerosol, a middle high altitude cloud, and a high cirrus cloud. The optical properties of the aerosols used were types 2b and 3b from Kahn et al. (2001), representing a mixture of dusty maritime and coarse dust and a mixture of carbonaceous and black carbon maritime, respectively. The cloud and cirrus optical properties were taken from the Baum model (Baum et al. 2005a,b). It used a pre-screen of SNR $>70$, a cloud-screen based on the difference between ECMWF surface pressure and the surface pressure retrieved from a full O$_2$ A band retrieval (Taylor et al. 2012), and a post-screen based on $\chi^2$ of each band, a posterior error, retrieved AOD, surface pressure and degrees of freedom.

4.6.4 RemoTeC Retrieval Algorithm

The Netherlands Institute for Space Research (SRON) retrieval algorithm (Butz et al. 2011) uses the Phillips-Tikhonov regularisation method of the CO$_2$ profile in addition to the least squares difference minimisation method with Gauss-Newton method applied (Rodgers 2000). These are balanced by a regularisation parameter. It uses a vector radiative transfer model based on the Gauss-Seidel solution of the radiative transfer equation and forward adjoint perturbation theory for the derivatives. It uses the tabulated absorption cross-sections of HITRAN08 (Rothman et al. 2009) and assumes AOD to be negligible by forcing retrievals into vanishing aerosol and cirrus optical thickness. It simultaneously retrieves the 1.61 $\mu$m CO$_2$ band, CO$_2$ concentrations and microphysical properties of aerosol and cirrus particles. Filtered null-space components of the retrieved CO$_2$ profile are filled by 2005 model fields of CarbonTracker (Peters et al. 2007).

The a priori surface pressure, temperature and water vapour were obtained from ECMWF, a priori CO$_2$ from CarbonTracker 2005 model fields, and albedo from GOSAT reflectance. The NIES cloud-screen (Taylor et al. 2012), a pre-screen of SNR $>70$ and a post-screen based on $\chi^2$ and AOD was used.

4.6.5 Intercomparison Results

Only retrieved $X_{CO2}$ within a spatial coincidence of 5° radius of a TCCON site was used. All coincident statistics use the mean $X_{CO2}$ within $\pm2$ hours ($\pm1$ hour for NIES 01.XX and NIES PPDF D algorithms) for temporal coincidence to TCCON data and
these values are daily averaged. These statistics are shown in Figure 4.19, which shows the correlation of each algorithm with TCCON for all sites combined.

The NIES 01.XX algorithm shows a large offset to TCCON which is known to be due to the existence of CO₂ absorption lines within the solar irradiance database used (Yoshida 2011). The NIES 01.XX results also have a large scatter which is most likely due to the assumption of a fixed aerosol profile between 0 - 2 km altitude globally. The ACOS B2.8 algorithm also has a large offset which is known to be mainly due to a surface pressure bias of 11 hPa (Crisp et al. 2012). However, the ACOS B2.9 algorithm attempts to correct for this surface pressure bias by scaling the O₂ A band absorption lines by a factor of 1.025 and also includes an O₂ A band zero level offset to help alleviate the effects of the band 1 nonlinearity. With these updates the ACOS B2.9 algorithm correlates well (r= 0.77) with TCCON. The NIES PPDF D algorithm has the highest correlation (r= 0.79) but the least amount of data. Currently it removes a lot of useful data that are only partially contaminated by optical path modification. The RemoTeC 1.0 algorithm also gives a good correlation (r= 0.76) with TCCON.

The UoL-FP algorithm (no bias correction applied) has the highest amount of data and shows a good but slightly lower correlation (r= 0.72) with TCCON but has a number of outliers included which will affect this correlation. As shown in Section 4.2 if a stricter post-screen is applied then the TCCON comparison statistics improve substantially but at the cost of the amount of data. Most algorithms give a similar correlation to TCCON (UoL-FP 3G, ACOS B2.9, Remotec 1.0 and NIES PPDF D) but the small differences in the statistics should not be over-interpreted as the different screening applied to each algorithm can easily account for these differences, and the screening can easily be changed slightly to improve the statistics or allow more data.
FIGURE 4.19. Correlation diagrams between GOSAT and ground based FTS measurements of $X_{\text{CO}_2}$ for six algorithms. The GOSAT data were selected within a circle of 5° latitude/longitude radius centered at each FTS station. The $X_{\text{CO}_2}$ from ground based FTS data were the mean values measured within ±2 hours (±1 hour for NIES 01.XX and PPDF D) of GOSAT overpass time and both GOSAT and TCCON data were daily averaged. Red lines correspond to the best fit for all sites and green line is one-to-one correspondence. The number of coincident scans and characteristics of statistical relationships between ground based FTS and GOSAT $X_{\text{CO}_2}$ are listed in the inserts of each panel.
4.7 Conclusions

$X_{CO2}$ retrievals from GOSAT using the UoL-FP v3G algorithm were performed and show from comparison to ground-based TCCON data that the retrievals have a random error of $\sim 2.5$ ppm, twice the a posteriori error estimate. With stricter screening this precision reduces to $\sim 1.9$ ppm, which should be sufficient to allow improved surface flux estimates (e.g., Chevallier 2007b, Chevallier et al. 2009, Miller et al. 2007).

The mean bias inferred from TCCON comparisons is 0.2 ppm with a station-to-station variation of 0.56 ppm (standard deviation). Thus, over TCCON sites the $X_{CO2}$ retrieval is very accurate with biases approaching only a few tenths of a ppm, which is close to typical retrieval requirements of systematic bias for $X_{CO2}$ (e.g. Chevallier 2007b, Chevallier et al. 2009, Miller et al. 2007). The latest results from other algorithms (Bril et al. 2007, Yokota et al. 2009, Butz et al. 2011, Morino et al. 2011, Wunch et al. 2011b, Yoshida et al. 2010, Crisp et al. 2012, Oshchepkov et al. 2012a) find similar results with a variation in the number of soundings potentially due to the screening criteria applied.

However, over desert regions, which are not observed by TCCON, it is found that the retrieved $X_{CO2}$ from GOSAT shows significant (up to $\sim 3$ ppm) differences when compared to GEOS-Chem model calculations which are believed to be biases in the GOSAT retrieval. Similar findings have been reported in Crisp et al. (2012) who suggested that these biases are related to instrument gain. Although here, biases are observed for desert regions observed with both settings of the gain, leading to the speculation that these biases could be introduced by desert dust or related to the high surface albedo in the CO$_2$ bands.

This study highlights the need for further algorithm improvements, especially over the deserts, but improvements in instrument calibration or spectroscopy might also lead to reduced biases for these regions. Until sufficient algorithmic improvements have been achieved, one would suggest to either omit $X_{CO2}$ retrievals over deserts or apply the described bias correction method which significantly reduces the biases in our $X_{CO2}$ retrieved over deserts. However, except for the desert regions, the $X_{CO2}$ retrievals approach the point where they should be useful for the inversion of CO$_2$ surface fluxes with data assimilation methods (e.g. Baker et al. 2010, Chevallier et al. 2009, Feng et al. 2009).
Chapter 5

Developing an Improved Scattering Approach

5.1 Scattering Issue

Clouds and aerosols can cause light to become scattered. This can have a number of effects on a satellite measurement; light can be scattered into and contribute to the observed radiance, light can scatter out of and reduce the observed radiance, and the photon path length of the observed light can be changed. This can lead to biases in the retrieved $X_{CO_2}$ and therefore any improvement on the current retrieval scattering approach used would be highly beneficial. This chapter describes the development of a simulator and its use to test a new scattering approach and its comparison to the $X_{CO_2}$ retrievals shown in the previous chapter.

5.2 Simulator

5.2.1 Simulator Overview

The UoL-FP Simulator was developed to allow multiple SWIR retrieval approaches to be assessed. Although the simulator has primarily been used to investigate alternative aerosol methods for use in SWIR CO$_2$ retrievals from GOSAT, it can be applied to many different parameter issues (such as the assessment of the cloud screen, given in Section 3.21). A flexible scheme enables parameters to be selected from various sources, allowing the simulator to be easily modified for other applications. The simulator uses
the UoL-FP retrieval algorithm to simulate spectral radiances based on a priori information of various atmospheric/surface/instrumental parameters that are described in the following sections. An overview of the simulator is given in Figure 5.1, showing the main aspects involved to get from various source data to simulated radiances.

**Figure 5.1.** Diagram showing an overview of the simulator design, with arrows showing the flow of information. Grey boxes represent source data, yellow boxes indicate calculations being performed, purple boxes represent calculated selections of source data, blue boxes show input data for the Forward model, and orange boxes highlight the main computations of the Forward model.
5.2.2 Instrument Inputs

The simulator was designed to allow multiple orbit schemes to be used. This has been utilised to simulate spectral radiances for GOSAT orbits for two 3-day repeat cycles as well as Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations (CALIPSO) orbits for a six day period, although the flexibility could allow alternative observations to be simulated. The parameters for the orbits of GOSAT are obtained from pre-processed GOSAT L1B data, see Chapter 3. This includes the satellite position, solar and observing angles, relative velocity, altitude and observed IFOV location.

To simulate a CALIPSO orbit, these parameters are obtained directly from CALIPSO Level 1 data, but the retrieval algorithm remains setup for GOSAT observations such that GOSAT observations are simulated but for a CALIPSO orbit path. Similarly, the spectral dispersion is currently designed for the GOSAT SWIR bands with the GOSAT spectral resolution of $0.2 \text{ cm}^{-1}$. The same dispersion for each band is used for every simulation to reduce any uncertainties this could potentially introduce.

The noise is derived as a function of the spectral signal using a modelled function derived from real GOSAT high gain measurements. Equation 5.1 defines the function and Table 5.1 gives the coefficients derived by comparing noise, $N$, with signal, $S$, from pre-processed GOSAT L1B data for August 2009, as shown in Figure 5.2.

$$N = AS + B$$  \hspace{1cm} (5.1)

<table>
<thead>
<tr>
<th>GOSAT Band</th>
<th>A</th>
<th>B</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.0046960347</td>
<td>1.1245479E-09</td>
</tr>
<tr>
<td>2</td>
<td>0.0025181001</td>
<td>6.1902212E-10</td>
</tr>
<tr>
<td>3</td>
<td>0.0045985506</td>
<td>3.8230411E-10</td>
</tr>
</tbody>
</table>
Figure 5.2. Correlation of real GOSAT noise and signal for August 2009 with the post-screen applied. A linear fit of the correlation is shown in blue with the equation of the fit giving the coefficients to be used in the GOSAT noise model.
5.2.3 Radiative Transfer Inputs

5.2.3.1 Atmospheric Parameters

The atmospheric temperature, pressure and water vapour are derived from ECMWF operational analysis data. The atmospheric CO$_2$ is obtained from LMDZ 2009 data and CH$_4$ from the TM3 model. All these parameters are derived using the same method as described in Chapter 3.

5.2.3.2 Surface Parameters

The albedo is generated from MODIS Level 3 16-day average global albedo data at 0.8585 µm, 1.6400 µm and 2.1300 µm. These wavelengths were selected as they are reasonably close in wavelength to the three GOSAT SWIR wavelengths; 0.765 µm, 1.61 µm and 2.06 µm. The albedo for these three GOSAT bands are derived using the corresponding wavelength MODIS albedos as well as spectral dependence data from the Airborne Visible InfraRed Imaging Spectrometer (AVIRIS) and the Advanced Spaceborne Thermal Emission and Reflection radiometer (ASTER) for different surface types, as defined in Table 5.2. MODIS Level 3 land cover data provides a global dataset of surface/biome types, allowing each MODIS grid cell (0.05° x 0.05°) to have its albedo derived using the corresponding spectral dependance from either AVIRIS or ASTER. The spectral dependence for each surface type is shown in Figure 5.3. The spectral dependence, $\beta$, is interpolated to the wavelength of MODIS and that of the desired GOSAT wavelength, $\lambda$. This provides a wavelength dependent ratio that is applied to the MODIS albedo to get the albedo at the desired wavelength, as given by Equation 5.2. For simplicity, the albedo slope was assumed flat within each band. Figures 5.4, 5.5 and 5.6 show the derived albedo globally for each GOSAT SWIR band using this method for two 3-day repeat orbit cycles of GOSAT.

\[
Albedo_{\lambda} = \frac{Albedo_{\text{MODIS}}\beta_{\lambda}}{\beta_{\text{MODIS}}} \tag{5.2}
\]
Table 5.2. Spectral dependence source for each MODIS surface type.

<table>
<thead>
<tr>
<th>MODIS Surface Type</th>
<th>Spectral Dependency Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>Assumed constant albedo</td>
</tr>
<tr>
<td>Evergreen needleleaf forest</td>
<td>AVIRIS: Evergreen needleleaf forest</td>
</tr>
<tr>
<td>Evergreen broadleaf forest</td>
<td>AVIRIS: Evergreen broadleaf forest</td>
</tr>
<tr>
<td>Deciduous needleleaf forest</td>
<td>AVIRIS: Deciduous needleleaf forest</td>
</tr>
<tr>
<td>Deciduous needleleaf forest</td>
<td>AVIRIS: Deciduous broadleaf forest</td>
</tr>
<tr>
<td>Mixed forests</td>
<td>AVIRIS: Mixed forest</td>
</tr>
<tr>
<td>Closed shrubland</td>
<td>AVIRIS: Closed shrubs</td>
</tr>
<tr>
<td>Open shrubland</td>
<td>AVIRIS: Open shrubs</td>
</tr>
<tr>
<td>Woody savannas</td>
<td>AVIRIS: Woody savanna</td>
</tr>
<tr>
<td>Savannas</td>
<td>AVIRIS: Savanna</td>
</tr>
<tr>
<td>Grasslands</td>
<td>AVIRIS: Grassland</td>
</tr>
<tr>
<td>Permanent wetlands</td>
<td>AVIRIS: Wetlands</td>
</tr>
<tr>
<td>Croplands</td>
<td>AVIRIS: Croplands</td>
</tr>
<tr>
<td>Urban and built-up</td>
<td>ASTER: Urban Scenario</td>
</tr>
<tr>
<td>Cropland/natural vegetated mosaic</td>
<td>AVIRIS: Croplands</td>
</tr>
<tr>
<td>Snow and ice</td>
<td>ASTER: Medium Snow</td>
</tr>
<tr>
<td>Barren or sparsely vegetated</td>
<td>AVIRIS: Bare</td>
</tr>
<tr>
<td>Unknown</td>
<td>Assumed constant albedo</td>
</tr>
</tbody>
</table>
Figure 5.3. Spectral dependence for the different surface types used from AVIRIS and ASTER with vertical lines giving MODIS and GOSAT wavelengths used in the albedo method.
Figure 5.4. Derived Albedo for GOSAT band 1 for two 3-day GOSAT repeat orbit cycles averaged into 5°x5° grid cells.

Figure 5.5. Derived Albedo for GOSAT band 2 for two 3-day GOSAT repeat orbit cycles averaged into 5°x5° grid cells.
FIGURE 5.6. Derived Albedo for GOSAT band 3 for two 3-day GOSAT repeat orbit cycles averaged into 5°x5° grid cells.
5.2.3.3 GEMS Scattering Parameters

The GEMS model (experiment f1kd until October 2009 and f93i afterwards) provides mass mixing ratio profiles (MMR) for eleven different aerosol types, ice water content (IWC) MMR profiles and liquid water content (LWC) MMR profiles on 60 sigma levels every twelve hours at 00:00 and 12:00, with a model forecast every three hours proceeding an analysis. However, to reduce computational time only data for every six hours was used. These sigma levels are converted to pressure layers, $P_{\text{Layer}}$, using coefficients $A$ and $B$ (provided by GEMS) and the surface pressure, $P$, as

$$P_{\text{Layer}} = A_{\text{Layer}} + B_{\text{Layer}} P$$  \hspace{1cm} (5.3)

The pressure levels are determined by taking the middle pressure between layer pressures. The MMR data are interpolated to each simulated observation location and time using the four closest grid points for the two closest six hour intervals. The extinction for each aerosol can then be derived from the MMR (Knappett 2012, Simmons 2010) by

$$E = \frac{\mu \zeta}{g}$$  \hspace{1cm} (5.4)

where $E$ is the extinction, $\zeta$ is the MMR, $g$ (gravity) is defined as 9.80665 and $\mu$ is the mass extinction coefficient (MEC) for GEMS aerosols interpolated to the relative humidity and $O_2$ A band wavelength, see Figure 5.7. The relative humidity was derived from the ECMWF specific humidity, $S$, by

$$RH = \frac{p_{H_2O}}{p_{\text{Liquid}}} \times 100$$  \hspace{1cm} (5.5)

where RH is the relative humidity, $p_{\text{Liquid}}$ is the saturated vapour pressure and $p_{H_2O}$ is the partial pressure of water vapour which can be derived to be

$$p_{H_2O} = \frac{SPM_{AIR}}{M_{H_2O}}$$  \hspace{1cm} (5.6)

The molecular mass of air, $M_{AIR}$, and of $H_2O$, $M_{H_2O}$, is assumed to be 28.97 g mol$^{-1}$ and 18.015422 g mol$^{-1}$ respectively. There are numerous ways to estimate the saturated vapour pressure (Murphy & Koop 2005) but the method described by Murphy & Koop (2005) currently provides the most suitable function throughout normal atmospheric tem-
Figure 5.7. Mass extinction coefficients (MEC) for each of the GEMS aerosol types (Benedetti 2010). Black carbon (BC1) and desert dust aerosols (DD1, DD2 and DD3) are assumed to be independent of humidity (top left), with hydrophilic and hydrophobic black carbon having the same MEC. Organic matter (bottom left), sea salt (top right) and Sulphate (bottom right) (OM1, SS1/SS2/SS3, and SO4) are assumed to be humidity dependent.

\[
\ln(p_{\text{liquid}}) \approx 54842763 - \frac{6763.22}{T} - 4.210 \ln(T) + 0.000367T + \tanh[0.0415(T - 218.8)] \\
\left(53.878 - \frac{1331.22}{T} - 9.44523 \ln(T) + 0.014025T\right)
\]  

(5.7)

The average extinction between layers is assumed as the extinction for each level with the top and bottom levels assumed to have half of the closest layers extinction. To verify that this method of calculating the extinction is approximately correct, it was computed for a wavelength of 0.55 \(\mu\)m and compared to the GEMS AOD product at 0.55 \(\mu\)m. This is shown in Figure 5.8 and proves to correlate well with some slight scatter. Note that
the AOD, $\tau$, was calculated by

$$\tau = \frac{E_i + E_{i+1}}{2} |P_i - P_{i+1}|$$

(5.8)

where $E$ is the extinction, $P$ is the pressure and $i$ represents the atmospheric level. The extinction profiles for ice and liquid clouds are calculated using the method by Slingo & Schrecker (1982),

$$E = \frac{3\zeta}{2R_{eff}\rho} \frac{RT}{MgP}$$

(5.9)

where the effective radius is given by $R_{eff}$, $\rho$ is the density (assumed 0.92 for ice and 1 for liquid), $R$ is the ideal gas constant, and the second term on the right hand side is added to convert from altitude layers to pressure layers. The effective radius of liquid cloud particles was calculated using the Liu & Daum (2000) method. This is given as

$$R_{eff} = 0.13 \left( \frac{3RT}{4\pi MP} \right)^{\frac{1}{3}} \left( \frac{\zeta}{N} \right)^{\frac{1}{3}-0.11}$$

(5.10)
where $N$ is $600 \, \text{cm}^{-1}$ over land. The method for the ice effective radius is taken from Sun & Rikus (1999) and is given as

$$R_{\text{eff}} = \frac{1}{2} \left[ \frac{3\sqrt{3}}{8} \left( 45.8966\zeta^{0.2214} + (0.7957\zeta^{0.2535}T - 83.15) \right) \right. \\
\left. (1.2351 + (0.0105T - 273.15)) \right]$$ (5.11)

where $T$ is the layer temperature. Similarly, the extinction is converted from layers to levels. Not only are the effective radii used within the extinction calculation, but the effective radii at the pressure level where the maximum ice/liquid cloud extinction exists, hence cloud height, is used to define the ice/liquid cloud scattering properties within the retrieval. Ice and liquid cloud scattering properties were obtained from the Baum model (Baum et al. 2005a,b) for different effective radii; every $5 \, \mu\text{m}$ between 10 to 90 $\mu\text{m}$ for ice cloud particles and every $1 \, \mu\text{m}$ between 4 and 64 $\mu\text{m}$ for liquid cloud particles. For each simulation the Baum model scattering properties with the effective radii closest to that calculated from GEMS was used.

To check the simulated optical depths and effective radii provide an approximation of the real world they have been compared to the MODIS TERRA Level 3 daily product. A comparison of the AOD, given in Figure 5.9, shows that both the magnitude and spatial distribution are very similar, which was expected since GEMS assimilates MODIS AOD. Some difference are, however, expected due to the humidity being different. The calculated GEMS cloud and cirrus optical depths show a similar spatial pattern, see Figure 5.10. Although, MODIS gives much higher values over the poles, few GOSAT soundings will be observed at these latitudes and those that are will get screened out. The magnitude of the cloud/cirrus optical depths are on average lower in GEMS than MODIS. Figure 5.11 shows the effective radii of cloud/cirrus for GEMS and MODIS. The GEMS water cloud effective radii have a similar spatial distribution and magnitude as MODIS, although some differences can be seen over Siberia, the Himalayas, and the south of the Indian Ocean. The GEMS cirrus cloud effective radii have a similar magnitude on average, but are clearly lower than MODIS around the equator. Although some differences exist between GEMS and MODIS the parameters still provide a range of reasonable values. Since these differences are specific to cloud/cirrus, the effect is small as most cloudy scenes will be filtered out by a cloud screen.

Although the ice and liquid cloud properties were obtained from the Baum model, the aerosol properties were derived from refractive index (RI) values and assumptions of the GEMS aerosol sizes and modes (Benedetti 2010). However, the GEMS model is
known to include spin-down effects mainly in the dust but also in all other aerosol types (Morcrette 2012). This leads to the model overestimating the amount of small desert dust particles and underestimating the amount of large desert dust particles (Morcrette 2012). To compensate for this issue, the small and medium GEMS desert dust aerosols sizes were replaced with accumulation mode desert dust particle sizes and the large GEMS desert dust replaced by the coarse desert dust particle sizes from the climatology of
Figure 5.11. Comparison of calculated mean GEMS liquid cloud (top left) and ice cloud (top right) effective radius for all 6 hour time frames and MODIS mean daily liquid cloud (bottom left) and ice cloud (bottom right) effective radius between 08/09/2009 and 14/09/2009. The effective radii at the atmospheric level where the highest extinction exists is used for GEMS. Similarly, the MODIS retrieved effective radii is most sensitive within the cloud/cirrus. Areas where no valid data exists is given by white.

Kahn et al. (2001), such that the desert dust aerosols have larger sizes. The organic matter, black carbon and sulphate aerosols were replaced by the equivalent aerosol sizes from the Kahn et al. (2001) climatology which have less extreme sizes. The sea salt aerosols were kept as the GEMS sizes as the quantity of these aerosols over land is much smaller.

The RI of sulphate and sea salt were calculated by interpolating RI values from Hale & Querry (1973) between 0.2 and 0.7 mm, Palmer & Williams (1974) between 0.7 and 2.0 mm and Downing & Williams (1975) between 2.0 and 1000 mm. Additionally, Tang & Munkelwitz (1991) RI of 1.330 at 0.633 mm was used as the reference standard and Kou et al. (1993) data was used for the imaginary RI between 0.7 and 2.7 mm. These RI were computed for a humidity of 50% by spectrally interpolating between the RI of the dry aerosols and the RI of water using formulas which were derived from accurate laboratory measurements (Tang & Munkelwitz 1991, 1994, 1996) of the change in RI with relative humidity of those aerosols (Lacis 2010).

The RI for hydrophobic and hydrophilic organic matter were obtained from Dick...
et al. (2007) for a humidity of $4 - 10\%$ and $44 - 76\%$, respectively. The RI for dust and soot were taken from Shettle & Fenn (1979) and are not humidity dependant. Using these RI values (shown in Figures 5.12 and 5.13) and the assumed sizes and modes given in Table 5.3, the phase function moments, geometric cross-section, extinction and scattering cross-sections and asymmetry parameter for each aerosol were computed using the Mie code of Mishchenko et al. (2002). This computes the scattering of light by polydisperse homogeneous spherical particles using Lorenz-Mie scattering (Mishchenko et al. 1999). The radius standard deviation gives a measure of the width of the size distribution around the effective radius, with the minimum and maximum radius providing the limits. All GEMS aerosols are approximated with a log normal size distribution (Benedetti 2010) that can be expressed as

$$n (r) = \text{constant} \times r^{-1} \exp \left[ - \frac{\left( \ln r - \ln r_g \right)^2}{2 \ln^2 \sigma_g} \right]$$  \hspace{1cm} (5.12)$$

where $r$ is the radius, $r_g$ is the geometric radius and $\sigma_g$ is the standard deviation of the radius. Using a range of sizes provides a polydisperse distribution which is more realistic than a monodisperse sample since aerosols are highly variable in size (Williams et al. 2002). However, the desert dust types based on Kahn et al. (2001) are obtained (courtesy of V. Natraj) from the OCO orbit simulator (O’Brien et al. 2009) which computes these aerosols as polydisperse non-spherical homogenous particles.

The properties for the two log-normal size distribution modes (a and b) were combined for each sea salt size using the method by Abdou et al. (1997). The two modes are assumed equally weighted, such that the mean of the modes is used for the extinction cross-section, scattering cross-section, geometric cross-section and asymmetry parameter. The phase function, $P$, can be given by

$$P (\Omega, \lambda) = \frac{P_a \omega_a}{\omega_a + \omega_b} + \frac{P_b \omega_b}{\omega_a + \omega_b}$$  \hspace{1cm} (5.13)$$

where $\lambda$ is the minimum and maximum wavelengths for each GOSAT SWIR band, $\Omega$ is the scattering angle and $\omega$ is the single scattering albedo. Note that the asymmetry parameter gives an indication of the scattering direction, where a value of -1 gives a strong backwards directed peak and 1 gives a strong forward directed peak. Figure 5.14 shows the key derived properties for the eleven aerosols. It gives the asymmetry parameter for each of the eleven aerosols, showing that all aerosols scatter strongly forward at the $O_2$ A band wavelength and smaller aerosols have a weaker forward scattering peak with increasing wavelength. The angstrom coefficient was calculated from the extinction cross
Figure 5.12. Assumed refractive index real values for the GEMS aerosol types.

section, \( C_{\text{ext}} \), at different wavelengths by

\[
\alpha = \frac{\ln \left( \frac{C_{\text{ext}\lambda}}{C_{\text{ext}\lambda_r}} \right)}{\ln \left( \frac{\lambda}{\lambda_r} \right)}
\]  

(5.14)

where \( \lambda_r \) is the reference wavelength (in this case at the \( \text{O}_2 \) A band) and \( \lambda \) is the wavelength of the other bound that the angstrom coefficient is to be calculated between. Note that the angstrom coefficients are lower for larger particles. The ability of each aerosol to absorb or scatter light is shown by the single scattering albedo, which is simply the ratio:

\[
\omega = \frac{C_{\text{sca}}}{C_{\text{ext}}}
\]  

(5.15)

where \( C_{\text{sca}} \) represents the scattering cross section. Black carbon has a low single scattering albedo, so is much more absorbing than the other aerosol types.
The phase function of each aerosol type is given at each GOSAT band in Figures 5.15, 5.16, and 5.17, showing the scattering amount from a forward to backward direction. Both dust types scatter strongly in the forward direction with a slight backwards peak. Accumulation dust differs from coarse dust in the CO$_2$ bands with a reduced backwards peak. Sea-salt mostly scatters light in a forward direction with a minor backwards peak and changes little between bands. The other aerosols all give similar scattering directions of a weaker forward peak that reduces with increasing wavelength.
Table 5.3. Aerosol sizes corresponding to the 11 GEMS aerosols (Benedetti 2010).

<table>
<thead>
<tr>
<th>Aerosol</th>
<th>Abbreviation</th>
<th>Effective Radius</th>
<th>Radius Standard Deviation</th>
<th>Minimum Radius</th>
<th>Maximum Radius</th>
</tr>
</thead>
<tbody>
<tr>
<td>Small Sea Salt (mode a)</td>
<td>SS1a</td>
<td>0.1992</td>
<td>1.90</td>
<td>0.0300</td>
<td>0.50</td>
</tr>
<tr>
<td>Medium Sea Salt (mode a)</td>
<td>SS2a</td>
<td>0.1992</td>
<td>1.90</td>
<td>0.5000</td>
<td>5.00</td>
</tr>
<tr>
<td>Large Sea Salt (mode a)</td>
<td>SS3a</td>
<td>0.1992</td>
<td>1.90</td>
<td>5.0000</td>
<td>20.00</td>
</tr>
<tr>
<td>Small Sea Salt (mode b)</td>
<td>SS1b</td>
<td>0.1992</td>
<td>2.00</td>
<td>0.0300</td>
<td>0.50</td>
</tr>
<tr>
<td>Medium Sea Salt (mode b)</td>
<td>SS2b</td>
<td>0.1992</td>
<td>2.00</td>
<td>0.5000</td>
<td>5.00</td>
</tr>
<tr>
<td>Large Sea Salt (mode b)</td>
<td>SS3b</td>
<td>0.1992</td>
<td>2.00</td>
<td>5.0000</td>
<td>20.00</td>
</tr>
<tr>
<td>Small Desert Dust †‡</td>
<td>DD1</td>
<td>0.47</td>
<td>2.60</td>
<td>0.0500</td>
<td>2.00</td>
</tr>
<tr>
<td>Medium Desert Dust †‡</td>
<td>DD2</td>
<td>0.47</td>
<td>2.60</td>
<td>0.0500</td>
<td>2.00</td>
</tr>
<tr>
<td>Large Desert Dust †‡</td>
<td>DD3</td>
<td>1.90</td>
<td>2.60</td>
<td>0.5000</td>
<td>15.00</td>
</tr>
<tr>
<td>Hydrophobic Organic Matter †</td>
<td>OM1</td>
<td>0.13</td>
<td>1.80</td>
<td>0.0070</td>
<td>2.00</td>
</tr>
<tr>
<td>Hydrophilic Organic Matter †</td>
<td>OM2</td>
<td>0.124</td>
<td>1.80</td>
<td>0.0067</td>
<td>1.90</td>
</tr>
<tr>
<td>Hydrophobic Black Carbon †</td>
<td>BC1</td>
<td>0.1180</td>
<td>2.00</td>
<td>0.0050</td>
<td>0.50</td>
</tr>
<tr>
<td>Hydrophilic Black Carbon †</td>
<td>BC2</td>
<td>0.14396</td>
<td>2.00</td>
<td>0.0061</td>
<td>0.61</td>
</tr>
<tr>
<td>Sulphate</td>
<td>SO4</td>
<td>0.08</td>
<td>1.88</td>
<td>0.0070</td>
<td>0.81</td>
</tr>
</tbody>
</table>

† Kahn et al. (2001) aerosol sizes used.
‡ Non-spherical aerosol (other aerosols are spherical).
* Hydrophilic black carbon was assumed a growth factor of 1.22 (Popovicheva et al. 2008).
§ Hydrophilic organic matter was assumed a growth factor of 1.05 (Malm et al. 2005), but since the Kahn et al. (2001) organic matter is given as hydrophilic, the sizes were scaled down by one over the growth factor to get the corresponding hydrophobic organic matter sizes.
FIGURE 5.14. Derived aerosol properties for each GEMS aerosol type for the start wavelength of each GOSAT SWIR band. The angstrom coefficient between bands 1 and 2 is given by the green stars and the angstrom coefficient between bands 2 and 3 is given by the red diamonds. For both the single scattering albedo and the asymmetry parameter band 1 is given by the purple triangles, band 2 by the green stars and band 3 by the red diamonds.
FIGURE 5.15. Derived phase function moments for the start wavelength of GOSAT band 1 as a function of scattering angle for each of the eleven GEMS aerosol types. An angle of zero points directly in the forward direction.
Figure 5.16. Derived phase function moments for the start wavelength of GOSAT band 2 as a function of scattering angle for each of the eleven GEMS aerosol types. An angle of zero points directly in the forward direction.
Figure 5.17. Derived phase function moments for the start wavelength of GOSAT band 3 as a function of scattering angle for each of the eleven GEMS aerosol types. An angle of zero points directly in the forward direction.
5.2.3.4 CALIPSO Scattering Parameters

CALIPSO was launched in April 2006 and contains three instruments; the Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP), the Imaging Infrared Radiometer (IIR), and the Wide Field Camera (WFC). Here, data products of the CALIOP instrument have been used to derive atmospheric profiles of aerosols, clouds and cirrus clouds. This can be used instead of the GEMS aerosols/clouds.

The CALIPSO Lidar Level 2 aerosol profile data provides atmospheric profiles of total extinction per 40 km ground track at a number of different wavelengths, as well as altitude, pressure, temperature and relative humidity. The extinction profiles exist on altitude levels but are needed on pressure levels for input into the UoL-FP retrieval algorithm. To convert between altitude and pressure levels the hydrostatic equation can be used

\[
\frac{dP}{dz} = -\rho g
\]  

where \( P \) is pressure, \( z \) is altitude, \( g \) is the acceleration due to gravity (assumed constant and equal to a mean sea level value of 9.80665 m s\(^{-1}\)), and \( \rho \) is the density. The density can be substituted for as

\[
\frac{dP}{dz} = -\frac{g}{1000} \left[ \left( \frac{P_{dry}}{(R_{dry}T)} \right) + \left( \frac{P_{wet}}{(R_{wet}T)} \right) \right]
\]

where temperature is \( T \), the ideal gas constant for dry air is \( R_{dry} \) (which is the ideal gas constant divided by the dry air mass, 0.02897 kg mol\(^{-1}\)), the ideal gas constant for wet air is \( R_{wet} \) (which is the ideal gas constant divided by the wet air mass, 0.018015422135 kg mol\(^{-1}\)). The partial pressure of wet air \( P_{wet} \) can be derived from the relative humidity, RH, and the saturated vapour pressure, \( p_{Liquid} \), using the method of Murphy & Koop (2005) previous described in Section 5.2.3.3 by

\[
P_{wet} = RH \frac{p_{Liquid}}{100}
\]

The partial pressure of dry air is then obtained by

\[
P_{dry} = P - P_{wet}
\]

To simulate GOSAT radiances the aerosol/cloud/cirrus extinctions need to be for the \( \text{O}_2 \) A band, therefore CALIOP extinction profiles at 532 and 1064 nm are used. The total
optical depth, \( \tau \), at each of these wavelengths can be calculated by

\[
\tau = \sum \frac{dP_i}{dzi} \frac{(E_i + E_{i+1})}{2} dP_{i,i+1}
\]  

(5.20)

where \( E \) is extinction and and \( i \) denotes the atmospheric level. These optical depths can be used to compute an angstrom coefficient that relates optical depth with wavelength. The angstrom coefficient can be calculated as

\[
\alpha = \frac{\ln \left( \frac{\tau_{532}}{\tau_{1064}} \right)}{\ln \left( \frac{\lambda_{1064}}{\lambda_{532}} \right)}
\]  

(5.21)

The optical depth at the \( \text{O}_2 \) A band (765 nm) is then approximated as

\[
\tau_{765} = \tau_{532} \left( \frac{\lambda_{765}}{\lambda_{532}} \right)^{-\alpha}
\]  

(5.22)

The extinction profile at the \( \text{O}_2 \) A band can be computed from the 532 nm extinction profile by multiplying it by the ratio \( \tau_{765}/\tau_{532} \).

The CALIPSO Lidar Level 2 Vertical Feature Mask (VFM) data separates the atmospheric profile into many altitude layers and classifies what features exist in each layer, such as clear air, cloud or aerosol. These features are also sub-categorized for each aerosol/cloud type, as shown in Figure 5.18. Any VFM data that was categorized as Invalid, Surface, Sub-Surface or No Signal was excluded from all computations. Stratospheric features were assumed to be cloud. Clouds were separated into liquid and ice clouds using the cloud phase, where horizontally and randomly orientated ice was taken to be ice cloud, and unknown and water taken to be liquid cloud. Any aerosol that was sub-categorized as Non-determined or Other was assumed as desert dust.

The VFM data is separated into three altitude ranges with numerous discrete layers within each range: -0.5 to 8.2 km, 8.2 to 20.2 km, and 20.2 to 30.1 km. The lowest altitude range gives VFM data for 15 discrete increments for every 5 km of the ground track. However, the VFM data for the other altitude ranges are averaged over larger ground track lengths. To obtain the VFM data on a regular grid the upper altitude data is re-binned onto the same grid as the lowest altitude range.

The fractional amount of aerosol within an extinction profile layer, is calculated as the number of discrete VFM increment layers that contain aerosol divided by the total number of increment layers within the 40 km ground track. Similarly, the fractional amount of cloud and sub-categorized clouds/aerosols are calculated in the same way.
FIGURE 5.18. CALIOP VFM for one orbit track with features given by different colours. This example notably shows water clouds over the majority of the orbit track, polluted dust and smoke over Southern Africa, and a large amount of desert dust over the Sahara.

To get separate extinction profiles for aerosol, liquid cloud and ice cloud the extinction profile is split up into those three profiles for each layer according to the ratio of their fractional amounts.

The particle properties for ice and liquid clouds are taken from the Baum model (Baum et al. 2005a,b) as described in Section 5.2.3.3, using half of the effective particle size obtained from CALIPSO IIR level 2 data as the effective radius for both liquid and ice cloud particles.

Omar et al. (2004) estimates the aerosol properties of the CALIPSO model by comparisons to observations by AERONET. Each aerosol type is assumed to be a bi-modal aerosol of fine and coarse modes (Mielonen et al. 2009, Omar et al. 2004, Simmons 2010), as shown in Table 5.4. Based on these parameters, the aerosol scattering properties for each aerosol type and mode were computed using the Mie code explained in Section 5.2.3.3, where the RIs were interpolated linearly to the wavelength of each GOSAT band. The volume fraction of each mode was used to weight the mixing of the
modes using the method by Abdou et al. (1997), which states that the combined single scattering albedo can be given as

\[ \omega_{M,\lambda} = \sum_{k=1}^{k} f_k \omega_{k,\lambda} \]  \hspace{1cm} (5.23)

where M indicates the combined mixture, k represents each aerosol type/mode, and f is the fraction used to weight the aerosols contribution to the mixture. Also, Abdou et al. (1997) gives the combination of phase functions as:

\[ P(\Omega, \lambda) = \sum_{k=1}^{k} \frac{f_k P_{k,\lambda} \omega_{k,\lambda}}{\omega_{M,\lambda}} \]  \hspace{1cm} (5.24)

However, the UoL-FP retrieval algorithm was designed to calculate all scattering values relative to the O\textsubscript{2} A band. Therefore, Equation 5.23 and 5.24 use the relative single scattering albedo of each aerosol. The relative extinction cross section for an aerosol can be given as

\[ C_{ext,k,\lambda,rel} = \frac{C_{ext,k,\lambda}}{C_{ext,k,\lambda,r}} \]  \hspace{1cm} (5.25)

The combined relative extinction cross section is then given by:

\[ C_{ext,M,\lambda,rel} = \sum_{k=1}^{k} f_k C_{ext,k,\lambda} \]  \hspace{1cm} (5.26)

Similarly, the relative scattering cross section of the mixture is expressed by:

\[ C_{sca,M,\lambda,rel} = \sum_{k=1}^{k} f_k C_{sca,k,\lambda} \]  \hspace{1cm} (5.27)

The six aerosol types were then combined, weighted by the total column fraction of each type, for each atmospheric aerosol extinction profile. The fraction of each aerosol type that is used to create the aerosol property mixtures is shown in Figure 5.19. This, for example, shows that aerosols over the Sahara are almost completely weighted to desert dust.

The properties of each aerosol type are given in Figure 5.20, which shows a low angstrom coefficient for all aerosol types. The single scattering albedo is highest for clean marine aerosol, showing it is highly scattering and absorbs little light. For clean continental, polluted dust and smoke aerosols about half gets scattered and half absorbed. The asymmetry parameter shows all aerosol types to have a strong forward scattering
Figure 5.19. Optical depth fractions of each of the CALIPSO aerosol types for simulated scenes globally between 08/09/2009 to 14/09/2009.

peak at the O_2 A band, with clean marine aerosol strongly forward scattering at all SWIR bands of GOSAT. Dust, polluted dust, polluted continental and smoke aerosols all have a reduced forward scattering at the CO_2 bands compared to that of the O_2 A band. The phase functions (Figures 5.21, 5.22 and 5.23) show that the scattering directions of the coarse modes changes little with wavelength, but the fine modes become much less forward scattering, almost isotropic, at the wavelengths of the CO_2 bands. This becomes important as when the modes are combined the scattering direction at the CO_2 bands highly depends on the volume fractions used to weight the each modes contribution to the aerosol type. For example, this is demonstrated by the polluted continental aerosol that comprises approximately half the fine mode and half the coarse mode, so the forward scattering reduces for the CO_2 bands as shown by the asymmetry parameter in Figure 5.20.

To check the simulated scenes are realistic, the calculated aerosol/cloud/cirrus parameters were compared MODIS. Although the calculated effective radii for liquid and ice clouds are designated the same for CALIPSO, the values have a reasonably large
FIGURE 5.20. Derived aerosol properties for each CALIPSO aerosol type for the start wavelength of each GOSAT SWIR band. The angstrom coefficient between bands 1 and 2 is given by the green stars and the angstrom coefficient between bands 2 and 3 is given by the red diamonds. For both the single scattering albedo and the asymmetry parameter band 1 is given by the purple triangles, band 2 by the green stars and band 3 by the red diamonds. The aerosol types are represented as; CM is Clean Marine, DD is Desert Dust, PC is Polluted Continental, CC is Clean Continental, PD is Polluted Dust, and SM is smoke.

variation (between 0 and \(\sim 60 \, \mu m\)) with the bulk of the data covering the ranges of both liquid and ice effective radii from MODIS, as shown in Figure 5.24. Thus, the calculated effective radii from CALIPSO provides a realistic basis for the cloud/cirrus properties used in the simulations. Comparing the optical depth with MODIS, Figure 5.25, reveals that both the liquid and ice cloud optical depths are highly under-estimated. However, CALIPSO measurements can not penetrate through thick clouds at these wavelengths.
Therefore, any extinction profile that should include a thick cloud will contain invalid data and not be used. This leaves CALIPSO soundings that contain either thin or no clouds, hence biasing the optical depths to lower values compared to MODIS. Additionally, this effectively acts as an initial cloud screen, since any scene containing thick cloud would be filtered out. The optical depth calculated for aerosol has a similar range as MODIS but with the values on average to be much lower than MODIS. Figure 5.26 shows the spatial distribution of AOD which follows the same patterns as both MODIS and GEMS. Some differences in AOD are expected as both GEMS and MODIS provide AOD data for coarse grids of \( \geq 110 \times 110 \) km (1° x 1°), whereas CALIPSO AOD has been calculated for a much smaller area of 0.07 x 40 km.
FIGURE 5.22. Derived phase function moments for the start wavelength of GOSAT band 2 as a function of scattering angle for each of the CALIPSO aerosol types and modes. An angle of zero points directly in the forward direction.
<table>
<thead>
<tr>
<th>Aerosol Type</th>
<th>Mode</th>
<th>Real RI at 532 nm</th>
<th>Imaginary RI at 532 nm</th>
<th>Real RI at 1064 nm</th>
<th>Imaginary RI at 1064 nm</th>
<th>Minimum Radius (µm)</th>
<th>Maximum Radius (µm)</th>
<th>Effective Radius (µm)</th>
<th>Radius Standard Deviation (µm)</th>
<th>Volume Fraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clean Marine</td>
<td>Fine</td>
<td>1.400</td>
<td>0.005</td>
<td>1.400</td>
<td>0.005</td>
<td>0.05</td>
<td>0.30</td>
<td>0.15</td>
<td>1.60</td>
<td>0.025</td>
</tr>
<tr>
<td>Clean Marine</td>
<td>Coarse</td>
<td>1.390</td>
<td>0.0005</td>
<td>1.390</td>
<td>0.0005</td>
<td>0.30</td>
<td>10.00</td>
<td>1.22</td>
<td>1.60</td>
<td>0.975</td>
</tr>
<tr>
<td>Desert Dust</td>
<td>Fine</td>
<td>1.414</td>
<td>0.0036</td>
<td>1.495</td>
<td>0.0043</td>
<td>0.05</td>
<td>0.30</td>
<td>0.12</td>
<td>1.48</td>
<td>0.223</td>
</tr>
<tr>
<td>Desert Dust</td>
<td>Coarse</td>
<td>1.414</td>
<td>0.0036</td>
<td>1.495</td>
<td>0.0043</td>
<td>0.30</td>
<td>10.00</td>
<td>2.84</td>
<td>1.91</td>
<td>0.777</td>
</tr>
<tr>
<td>Polluted Continental</td>
<td>Fine</td>
<td>1.404</td>
<td>0.0063</td>
<td>1.439</td>
<td>0.0073</td>
<td>0.05</td>
<td>0.30</td>
<td>0.16</td>
<td>1.53</td>
<td>0.531</td>
</tr>
<tr>
<td>Polluted Continental</td>
<td>Coarse</td>
<td>1.404</td>
<td>0.0063</td>
<td>1.439</td>
<td>0.0073</td>
<td>0.30</td>
<td>10.00</td>
<td>3.55</td>
<td>2.07</td>
<td>0.469</td>
</tr>
<tr>
<td>Clean Continental</td>
<td>Fine</td>
<td>1.380</td>
<td>0.0010</td>
<td>1.380</td>
<td>0.0010</td>
<td>0.05</td>
<td>0.30</td>
<td>0.21</td>
<td>1.61</td>
<td>0.050</td>
</tr>
<tr>
<td>Clean Continental</td>
<td>Coarse</td>
<td>1.455</td>
<td>0.0340</td>
<td>1.455</td>
<td>0.0340</td>
<td>0.30</td>
<td>10.00</td>
<td>2.63</td>
<td>1.90</td>
<td>0.950</td>
</tr>
<tr>
<td>Polluted Dust</td>
<td>Fine</td>
<td>1.452</td>
<td>0.0109</td>
<td>1.512</td>
<td>0.0109</td>
<td>0.05</td>
<td>0.30</td>
<td>0.13</td>
<td>1.51</td>
<td>0.241</td>
</tr>
<tr>
<td>Polluted Dust</td>
<td>Coarse</td>
<td>1.452</td>
<td>0.1370</td>
<td>1.512</td>
<td>0.1370</td>
<td>0.30</td>
<td>10.00</td>
<td>3.16</td>
<td>1.99</td>
<td>0.759</td>
</tr>
<tr>
<td>Smoke</td>
<td>Fine</td>
<td>1.517</td>
<td>0.0234</td>
<td>1.541</td>
<td>0.0234</td>
<td>0.05</td>
<td>0.30</td>
<td>0.14</td>
<td>1.56</td>
<td>0.329</td>
</tr>
<tr>
<td>Smoke</td>
<td>Coarse</td>
<td>1.517</td>
<td>0.0298</td>
<td>1.541</td>
<td>0.0298</td>
<td>0.30</td>
<td>10.00</td>
<td>3.73</td>
<td>2.14</td>
<td>0.671</td>
</tr>
</tbody>
</table>
Figure 5.23. Derived phase function moments for the start wavelength of GOSAT band 3 as a function of scattering angle for each of the CALIPSO aerosol types and modes. An angle of zero points directly in the forward direction.

Figure 5.24. Correlation of the liquid (left) and ice (right) effective radius calculated from CALIPSO with that retrieved from MODIS between 08/09/2009 to 14/09/2009 for CALIPSO soundings. The calipso soundings have much lower values for cloud/cirrus optical depths since the CALIOP measured radiance will often not penetrate through clouds, leaving incomplete extinction profiles that get screened out, hence removing most cloud scenes.
Figure 5.25. Correlation of the aerosol (left), liquid cloud (centre) and ice cloud (right) optical depth calculated from CALIPSO with that retrieved from MODIS between 08/09/2009 to 14/09/2009 for CALIPSO soundings.

Figure 5.26. AOD calculated from CALIPSO between 08/09/2009 to 14/09/2009 for CALIPSO soundings.
5.3 Initial Tests

To aid the construction of a new approach to atmospheric scattering it was first neccessary to determine which aspects are important in the retrieval of \( X_{CO2} \). Although there is a high spatial/temporal variation of aerosols and clouds, this can essentially be described by three components; the total column optical depth, the vertical distribution, and the particle properties.

The first test was to investigate how the AOD affects the retrieved \( X_{CO2} \). Here the simulator (with GEMS aerosols/clouds) was used to simulate GOSAT spectra for different amounts of AOD, with each AOD scenario having 100 realistic scenes simulated. All of these spectra were retrieved with the same aerosol profile (Gaussian shape peaking at a height of 2 km with a width of 2 km), the same aerosol properties, the same a priori AOD of 0.2, and all other a priori and statevectors the same. Figure 5.27 shows how the results of this test demonstrate that the UoL-FP retrieval algorithm can accurately retrieve \( X_{CO2} \) within \(~0.1\) AOD of the truth and remains within a \( X_{CO2} \) error of 0.03 ppm between a true AOD of 0.025 to 0.4 range. The performance of the retrieval to accurately retrieve \( X_{CO2} \) reduces for large differences (\( >0.2\)) between the a priori and true AOD, hence it is important to provide the retrieval with an a priori AOD that is not too small or too large as the retrieval will find it difficult to reach the true optimal solution.

Secondly, using the simulator in a similar fashion, the effect of the vertical distribution of aerosol on retrieved \( X_{CO2} \) was assessed. Here, spectra were simulated with a Gaussian shaped aerosol profile peaking at 2 km. The simulations used 100 realistic scenes and all were retrieved using different scenarios of the a priori aerosol profile peak height. Figure 5.28 shows that the mean resulting \( X_{CO2} \) error varies very little with the peak height of the aerosol profile. However the range of \( X_{CO2} \) values increases with the difference in height, with large differences between lower and upper altitudes.

To evaluate the impact of the aerosol properties on the retrieval of \( X_{CO2} \), spectra were simulated for 100 realistic scenes using the Kahn aerosol mixture 4b. Retrievals of \( X_{CO2} \) were performed on these simulated spectra for 11 different scenarios, where each scenario used only the properties of a single GEMS aerosol type. The results are given in Figure 5.29 and show that there is a large variation of \( X_{CO2} \) error. There are smaller \( X_{CO2} \) errors for larger sized aerosols and larger \( X_{CO2} \) errors for smaller sized aerosols, which is likely due to the Kahn 4b mixture being a combination of medium and large sized particles. The aerosol properties cause the \( X_{CO2} \) error to be significantly higher than that of AOD or vertical distribution differences. Hence it is by far the most crucial aerosol component to get closest to truth for accurate \( X_{CO2} \) retrievals.
Likewise, the same components were investigated for water clouds and cirrus clouds in exactly the same manner, with the components replaced with cloud/cirrus respectively. For the particle properties of both water clouds and cirrus clouds a range of effective radii obtained from the Baum model (Baum et al. 2005a,b) were used.

Figure 5.30 shows that the retrieval achieves a low X\(_{CO2}\) error when the assumed a priori cloud optical depth is within 0.15 of the truth, but increases rapidly with higher optical depth differences. For cirrus, the retrieval only performs accurately where the a priori optical depth is within 0.025 of the truth. The cirrus retrieval is extremely poor at capturing the true optical depth which clearly has an impact on the X\(_{CO2}\) error above a 0.05 difference in optical depth. Since a cloud screen is applied to real GOSAT observations, scenes with a high optical depth will already be omitted from X\(_{CO2}\) retrievals. Therefore, predominantly scenes with a low optical depth or cirrus will remain, revealing that it is more important to get the a priori cirrus optical depth close to truth than cloud
optical depth.

The effect of cloud height is shown in Figure 5.32 and shows a clear relationship with $X_{CO2}$ error of $\sim0.02$ ppm/km. Figure 5.33 shows the effect of cirrus height and gives a similar trend but with a larger gradient of $\sim0.075$ ppm/km and an anti-correlation with retrieved optical depth. Again, cirrus has a far larger affect on $X_{CO2}$ than water clouds and it is important to have an accurate a priori assumption of cirrus vertical distribution.

Similarly, there is a trend of effective radius with $X_{CO2}$ error (see Figures 5.34 and 5.35). The gradient is higher with cirrus than cloud, with values of $\sim0.35$ ppm/10$\mu$m and $\sim0.10$ ppm/10$\mu$m respectively. The $X_{CO2}$ error is approximately double that created by incorrect optical depth or cirrus height, making it more vital to get closer to the truth for retrievals of $X_{CO2}$.

**Figure 5.28.** Mean retrieved $X_{CO2}$ of 100 simulated GOSAT spectra for different peak heights of a Gaussian shaped aerosol profile. The blue line gives the $X_{CO2}$ error and the green line shows the AOD error, with the standard deviation of the 100 results for each height scenario shown by the errorbars.
Figure 5.29. Mean retrieved $X_{CO_2}$ from retrievals using different aerosol types with 100 simulated GOSAT spectra. The blue line gives the $X_{CO_2}$ error and the green line shows the AOD error, with the standard deviation of the 100 results for each height scenario shown by the errorbars. The dust properties used were based purely on GEMS, giving DD1, DD2 and DD3 as small, medium and large dust particles, respectively.
Figure 5.30. Mean retrieved $X_{CO_2}$ of 100 simulated GOSAT spectra for different cloud optical depths. The blue line gives the $X_{CO_2}$ error and the green line shows the optical depth error, with the standard deviation of the 100 results for each optical depth scenario shown by the errorbars.

Figure 5.31. Mean retrieved $X_{CO_2}$ of 100 simulated GOSAT spectra for different cirrus optical depths. The blue line gives the $X_{CO_2}$ error and the green line shows the optical depth error, with the standard deviation of the 100 results for each optical depth scenario shown by the errorbars.

Figure 5.32. Mean retrieved $X_{CO_2}$ of 100 simulated GOSAT spectra for different peak heights of a Gaussian shaped cloud profile. The blue line gives the $X_{CO_2}$ error and the green line shows the optical depth error, with the standard deviation of the 100 results for each height scenario shown by the errorbars.

Figure 5.33. Mean retrieved $X_{CO_2}$ of 100 simulated GOSAT spectra for different peak heights of a Gaussian shaped cirrus profile. The blue line gives the $X_{CO_2}$ error and the green line shows the optical depth error, with the standard deviation of the 100 results for each height scenario shown by the errorbars.
Figure 5.34. Mean retrieved $X_{CO_2}$ from retrievals using different cloud effective radii with 100 simulated GOSAT spectra. The blue line gives the $X_{CO_2}$ error and the green line shows the optical depth error, with the standard deviation of the 100 results for each height scenario shown by the errorbars.

Figure 5.35. Mean retrieved $X_{CO_2}$ from retrievals using different cirrus effective radii with 100 simulated GOSAT spectra. The blue line gives the $X_{CO_2}$ error and the green line shows the optical depth error, with the standard deviation of the 100 results for each height scenario shown by the errorbars.
5.4 New Scattering Approach Design Basis

Whilst clouds have a large effect on the retrieval accuracy, a cloud screen could be used to remove most cloud contaminated scenes in real retrievals of $X_{CO_2}$ from GOSAT. For this reason the new scattering approach was designed to provide initially a better approach to aerosols. The current retrieval method (UoL-FP v3G) gives the retrieval a large flexibility to allow more information from the measured radiance to be used to gain aerosol information. However, this method relies upon the same two aerosol mixtures globally. One way this method could be improved is to use aerosol mixtures that are specific to the observed location and time. At present there are a number of observation methods that provide aerosol information (e.g. satellite observations, ground based measurements and aircraft observations), however due to the fact that aerosols are highly variable both spatially and temporally none of these sources can provide adequate information for this approach. Thus, the new approach was designed to use model aerosol information obtained from the GEMS model.

The aerosol extinction profiles for each of the eleven GEMS aerosol types were derived using the method explained in Section 5.2.3.3. To keep the retrieval computational time low and for direct comparison to the current retrieval method, the eleven aerosol profiles were reduced to two profiles. For each GEMS latitude/longitude grid cell the aerosol profiles are temporally interpolated to the local mid-day time, as GOSAT observes day-time SWIR radiances. Since the model only provides estimates and not the true known aerosol information, some spatial smoothing is applied by taking the eight closest surrounding grid cells. The mean profiles of each aerosol type are calculated for this enlarged region and the two aerosol types which have the largest AOD are separated into two profile groups. The profile shape of all other aerosol profiles are correlated with both of these two profiles and then added to the group of which each aerosol correlates highest with. The aerosol profiles within each group are then added together to maintain the total AOD. This selection method of grouping the aerosols ideally contains at least one aerosol which will significantly contribute to the scene, allowing the retrieval to utilise both profiles for flexibility. Clearly the case can exist where only one aerosol type significantly contributes to a scene, in which case the retrieval would likely be forced to only use that groups profile. To overcome this possibility and allow more adaptability to the retrieval a lower AOD limit of 0.025 per group profile was enforced. There are cases, such as desert dust over the Sahara, where the AOD can be too large for the retrieval algorithm to accurately perform retrievals for (Kuang et al. 2002). To compensate for this, where the combined total AOD is greater than 0.275 in value, both group profiles
are reduced such that their ratio remains the same (with exception to the lower limit) and their combined total AOD equals this upper limit. The aerosol properties of each group were also designed to be specific to the scene by combining the individual aerosol types of each group, using the method described in Section 5.2.3.3 where the AOD fractions as the weighting of aerosol types contributing to the combination.

5.5 Comparison of Scattering Approach Tests

Although the new scattering approach design includes all three main aerosol components, it is possible that certain aspects might not provide any benefit over the flexibility of the current retrieval method. Therefore, in addition to the current retrieval method and the new scattering design being compared though retrievals of simulated scenes, each aerosol component of the new design is tested individually too. The next section explains each of these tests and makes a comparison between them.

5.5.1 Model Based Simulations: Aerosols Only

Spectral radiances of two 3-day cycles of GOSAT were simulated with scenes containing aerosols but not cirrus/clouds, using the GEMS model in the simulator. This provided a total of 9858 simulated exposures that were distributed globally using the GOSAT 5-point observation mode. The simulated spectra were retrieved using the same instrument, surface and atmospheric variables with the exception of the aerosol components which were different for each test;

- Use the current retrieval method (UoL-FP v3G) aerosol approach for reference to compare new tests against. Here the cirrus profile is omitted leaving only two aerosol profiles with a total AOD of 0.1.
- Use the current retrieval method but replace the AOD with that of the new scattering approach design.
- Use the current retrieval method but replace the aerosol profiles with that of the new scattering approach design, where the profile shape is retained but its total AOD is reduced to 0.1 for all scenes.
- Use the current retrieval method but replace the aerosol optical properties with that of the new scattering approach design.
Use all three aspects of the new scattering approach design.

The retrievals of each test were quality filtered to remove any poor retrievals, similar to the post-screening applied to the real GOSAT retrievals. However, many of the post-screen filter parameters do not exist in the simulations, so a more basic screen was applied that filters for only converged retrievals with <3 divergences, a total retrieved AOD of <0.3 and a difference between true and retrieved surface pressure of >-1 hPa. Not only are these parameters included in the post-screen, the parameters and filter thresholds were obtained empirically from comparisons with the $X_{CO_2}$ bias of the current retrieval approach that was used to retrieve the simulated scenes. Figure 5.36 shows the $X_{CO_2}$ bias and number of exposures for each of the filter parameters, as well as the threshold values that provide an appropriate compromise between the number of exposures and maintaining a low $X_{CO_2}$ bias and scatter. Although stricter screening could be applied, it would reduce the number of exposures and bias the screening to this retrieval approach setup.

The statistics on the number of exposures passing the quality filter, the resulting $X_{CO_2}$ bias and standard deviation of each test is given by Table 5.5. In addition to the five retrieval tests stated above, the simulated scenes were retrieved using exactly the same a priori values as used in the simulations, but one aerosol was used instead of all 11 aerosols. The one aerosol profile was the sum of all 11 aerosol profiles, so as to keep the profile shape and AOD the same, and the optical properties of the 11 aerosols were mixed together (weighted with the total column AOD). The only difference between the simulations and retrievals in this test was that the simulations mixed the aerosol optical properties at each atmospheric layer, whereas the retrievals used the aerosol optical properties mixed for the total atmospheric column. This test results in a $X_{CO_2}$ bias of -0.02 ppm with standard deviation of 0.63 ppm, showing some small differences between the retrieval and simulation that are due to the difference in mixing of optical properties and other retrieval related artefacts.

The current retrieval method (UoL-FP v3G) has a $X_{CO_2}$ bias of 0.06 ppm and a standard deviation of 0.82 ppm. The new scattering approach method shows an improvement upon this, with a lower $X_{CO_2}$ bias of 0.03 ppm and a reduced standard deviation of 0.69 ppm, such that it almost approaches the precision and accuracy of the retrievals using the simulation a priori. Looking at each new scattering approach aspect separately, using the AOD produces a slightly higher bias of 0.07 ppm and a worse standard deviation of 0.92 ppm. Using the new profiles results in a slightly lower bias of 0.05 ppm but higher standard deviation of 0.87 ppm (compared to the current retrieval method). When the
new aerosol properties are used the standard deviation improves (0.76 ppm) but the bias becomes slightly larger (0.07 ppm). In general, the difference in bias and standard deviation between the retrieval tests is small and all tests perform well on average (biases <0.07 ppm). Furthermore, the number of exposures left once quality screened is similar between retrieval tests, ranging between 50.3% and 54.1%.

Table 5.5. Retrieval statistics for each of the aerosol retrieval tests of aerosol only loaded simulated scenes.

<table>
<thead>
<tr>
<th>Aerosol test</th>
<th>Number of screened retrievals</th>
<th>$\Delta X_{CO2}$ mean (ppm)</th>
<th>$\Delta X_{CO2}$ σ (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Retrieval using simulation a priori</td>
<td>5249</td>
<td>-0.02</td>
<td>0.63</td>
</tr>
<tr>
<td>Current retrieval method (UoL-FP v3G)</td>
<td>4962</td>
<td>0.06</td>
<td>0.82</td>
</tr>
<tr>
<td>Current retrieval method with the new scattering approach AOD</td>
<td>4988</td>
<td>0.07</td>
<td>0.92</td>
</tr>
<tr>
<td>Current retrieval method with the new scattering approach aerosol profiles</td>
<td>5002</td>
<td>0.05</td>
<td>0.87</td>
</tr>
<tr>
<td>Current retrieval method with the new scattering approach aerosol properties</td>
<td>5332</td>
<td>0.07</td>
<td>0.76</td>
</tr>
<tr>
<td>New scattering approach method AOD, aerosol profiles and aerosol properties</td>
<td>5297</td>
<td>0.03</td>
<td>0.69</td>
</tr>
</tbody>
</table>

Figure 5.37 shows the difference between true and retrieved $X_{CO2}$ of each retrieval test to assess potential geographical biases. Large $X_{CO2}$ differences are found over desert regions in the current retrieval method (UoL-FP v3G), as already observed in comparisons between real GOSAT observations and GEOS-Chem model estimations (see Section 4.3. Additionally, other biases can be seen in Europe, North East Asia, central and southern Africa, and East USA. The retrieval using simulation a priori provides a basis for what could best be achieved and shows no desert biases and much lower biases elsewhere in the world. Some small biases are found though, mainly located over East
Similarly, the new scattering approach method also shows low biases throughout the world, but the aerosol is quite similar to the aerosol used in the simulations so is expected to perform well. Using each aspect of the new scattering approach method with the current retrieval method provides tests that are quite different to the simulations. When either the new AOD or new profiles are used the geographical biases remain similar to that of the current retrieval method. However, when the new optical properties are used the biases over the desert regions significantly reduce, as well as biases in other areas such as Europe and central/southern Africa. This leaves only small geographical biases similar to that produced by the new scattering approach method. Thus, changing the aerosol optical depth and profile shape has little or no benefit to the retrieval of $X_{CO2}$ but using aerosol optical properties that are scene dependent and closer to the truth provides a large improvement in $X_{CO2}$. These results agree with the conclusion of the initial aerosol tests. Hence, if scene dependent aerosol optical properties were used in retrievals of $X_{CO2}$ from real GOSAT measurements then better results would be expected.
Figure 5.36. Comparison of $X_{CO2}$ bias with each quality filter parameter for retrievals of aerosol only simulations using the current retrieval algorithm (right column). The screened data is given in blue and the data filtered out is shown in red. The left column shows the corresponding number of exposures when different filter thresholds are applied to each parameter.
Figure 5.37. Difference between true and retrieved $X_{CO2}$ of each retrieval test, where a is the retrieval using the simulation a priori, b is the current retrieval method with the new AOD, c is the current retrieval method, d is the current retrieval method with the new aerosol profile shapes, e is the new scattering approach method, and f is the current retrieval method with the new aerosol optical properties. Grey areas show where no data exists.
5.5.2 Model Based Simulations: Aerosols, Clouds and Cirrus

In real GOSAT observations the scenes would not just contain aerosols but also possibly cirrus and water clouds, therefore it is neccessary to test how these would interact with the aerosol methods. Here, the simulations are setup in exactly the same way as in the previous section, but cirrus and water clouds are included in the scenes according to what the GEMS model estimates. Before any retrievals were performed the simulated scenes were cloud screened, where scenes were determined as clear if the $O_2$ A band retrieved surface pressure was within $<20$ hPa of the truth. The five different retrieval tests were performed on 3281 cloud screen exposures and quality filtered. The resulting statistics are given in Table 5.6. The new AOD method shows the lowest $X_{CO2}$ bias and standard deviation, but the smallest number of exposures. The differences between the statistics of each test are quite small and all tests perform well on average ($X_{CO2}$ bias $<0.56$ ppm).

Table 5.6. Retrieval statistics for each of the retrieval tests of aerosol/cirrus/cloud loaded simulated scenes.

<table>
<thead>
<tr>
<th>Aerosol test</th>
<th>Number of screened retrievals</th>
<th>$\Delta X_{CO2}$ mean (ppm)</th>
<th>$\Delta X_{CO2}$ $\sigma$ (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current retrieval method (UoL-FP v3G)</td>
<td>1362</td>
<td>-0.49</td>
<td>2.02</td>
</tr>
<tr>
<td>Current retrieval method with the new scattering approach AOD</td>
<td>1331</td>
<td>-0.43</td>
<td>1.89</td>
</tr>
<tr>
<td>Current retrieval method with the new scattering approach aerosol profiles</td>
<td>1331</td>
<td>-0.55</td>
<td>2.09</td>
</tr>
<tr>
<td>Current retrieval method with the new scattering approach aerosol properties</td>
<td>1403</td>
<td>-0.51</td>
<td>2.01</td>
</tr>
<tr>
<td>New scattering approach method AOD, aerosol profiles and aerosol properties</td>
<td>1393</td>
<td>-0.56</td>
<td>1.97</td>
</tr>
</tbody>
</table>

Figure 5.38 shows the difference between true and retrieved $X_{CO2}$ of each retrieval test to assess potential geographical biases. Some negative biases are found in all test results, suggesting there to be a small issue with the simulations. Large regional biases
are found over desert regions, such as the Sahara, in the current retrieval method (UoL-FP v3G), the new AOD method, and the new aerosol profile shapes method. This bias is significantly reduced when the new aerosol optical properties method is used or the new scattering approach method is used. Changing the aerosol optical depth and profile shape has little benefit to the simulated retrieval of $X_{CO_2}$ but using scene dependent aerosol optical properties provides a large improvement in $X_{CO_2}$, as found in the previous section. However, this may not be true for real retrievals due to GOSAT instrument calibrations coupling with spectroscopy and aerosols.

**Figure 5.38.** Difference between true and retrieved $X_{CO_2}$ of each retrieval test, where a is the current retrieval method with the new AOD, b is the current retrieval method, c is the current retrieval method with the new aerosol profile shapes, d is the new scattering approach method, and e is the current retrieval method with the new aerosol optical properties.
5.5.3 Verification with Satellite Based Simulations

Since GEMS data is used in both the simulations and the new scattering approach design it is a partially circular test, such that one would expect the new scattering approach to give high correlations. To break this circularity, completely independent aerosol information from CALIPSO was used in the simulations instead of GEMS. The calipso based simulations were cloud screened and then retrieved with each of the five tests.

A total of 4505 cloud screened exposures were retrieved and quality filtered (see Section for details) with the statistics given in Table 5.7. There is very little difference between the statistics of each test; $X_{CO_2}$ biases range between -0.08 to -0.01 ppm, the standard deviation of $X_{CO_2}$ ranges from 1.24 to 1.39 ppm, and the number of exposures range between 2997 and 3221. The current retrieval method (UoL-FP v3G) has the lowest $X_{CO_2}$ bias but the largest standard deviation. The new aerosol optical properties provide the largest number of exposures and the new scattering approach method gives the smallest standard deviation. However, the differences between tests are so small that a single test can not be justified as the best.

Table 5.7. Retrieval statistics for each of the retrieval tests of CALIPSO based aerosol/cloud/cirrus loaded simulated scenes.

<table>
<thead>
<tr>
<th>Aerosol test</th>
<th>Number of screened retrievals</th>
<th>$\Delta X_{CO_2}$ mean (ppm)</th>
<th>$\Delta X_{CO_2}$ $\sigma$ (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current retrieval method (UoL-FP v3G)</td>
<td>3193</td>
<td>-0.01</td>
<td>1.39</td>
</tr>
<tr>
<td>Current retrieval method with the new scattering approach AOD</td>
<td>2998</td>
<td>-0.03</td>
<td>1.29</td>
</tr>
<tr>
<td>Current retrieval method with the new scattering approach aerosol profiles</td>
<td>2997</td>
<td>-0.08</td>
<td>1.30</td>
</tr>
<tr>
<td>Current retrieval method with the new scattering approach aerosol properties</td>
<td>3221</td>
<td>-0.03</td>
<td>1.29</td>
</tr>
<tr>
<td>New scattering approach method AOD, aerosol profiles and aerosol properties</td>
<td>3116</td>
<td>-0.05</td>
<td>1.24</td>
</tr>
</tbody>
</table>

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Figure 5.39 shows the difference between the true and retrieved $X_{CO2}$ of each retrieval test to assess potential geographical biases. The current retrieval method (UoL-FP v3G) retrieves lower $X_{CO2}$ values over most of the world, except for over desert regions, such as the Sahara, where much higher values than the truth are retrieved. Similar biases are found in retrievals using the new AOD or new profiles. However, when the new aerosol optical properties are used the biases over the desert regions reduce, as well as the negative biases in other areas. Similarly, the new scattering approach method also gives reduced biases, suggesting that it is influenced mostly by the aerosol optical properties and less so by the profile shape and AOD. Hence, using scene dependent aerosol optical properties reduces geographical biases and thus gives an improvement in $X_{CO2}$. This concurs with the findings from the GEMS based simulations discussed in Sections 5.5.1 and 5.5.2.
Figure 5.39. Difference between true and retrieved $X_{CO2}$ of each retrieval test, where a is the current retrieval method with the new AOD, b is the current retrieval method, c is the current retrieval method with the new aerosol profile shapes, d is the new scattering approach method, and e is the current retrieval method with the new aerosol optical properties.
5.6 Application to GOSAT

Although the simulations show that using spatially dependent aerosol optical properties improves the current retrieval approach, the true outcome can only be found by testing it with real measurements. $X_{CO_2}$ was retrieved using the spatially dependent aerosol optical properties method (to be further known as UoL-FP v3.4G) for the same GOSAT data that was used in the UoL-FP v3G retrievals so that a direct comparison of the retrieved results was possible. This includes $X_{CO_2}$ retrieved over 7 different TCCON sites for a 2 year period and one year globally with a comparison to the GEOS-Chem model, as described in Chapter 4.

Figure 5.40 shows a comparison of $X_{CO_2}$ retrieved from GOSAT using the UoL-FP v3.4G method with $X_{CO_2}$ retrieved by the TCCON. The same TCCON sites and data have been used as that shown in Chapter 4. The mean GOSAT-TCCON bias of the UoL-FP v3.4G retrieval algorithm is found to be -0.18 ppm with a standard deviation of 2.45 ppm and correlation coefficient of 0.70. Compared to the results of UoL-FP v3G (see Chapter 4), this method shows a slightly lower bias but a slightly larger standard deviation and slightly worse correlation coefficient. Table 5.8 gives the statistics for the UoL-FP v3.4G retrieval method, showing that the number of retrieved exposures passing the post-screen is 4494, which is lower than that found for UoL-FP v3G. Similarly, the number of coincident exposures with TCCON is slightly less than that found for UoL-FP v3G. However, it should be noted that the post-screen was optimised for UoL-FP v3G and not for UoL-FP v3.4G, and could result in good retrievals being incorrectly filtered out. To avoid this potential issue, the GOSAT-TCCON bias retrieved for UoL-FP v3G and UoL-FP v3.4G were compared for only exposures that exist after post-screening in both datasets. This is shown in Figure 5.41, and it is found that the UoL-FP v3.4G method reduces the bias by -0.05 ppm. However, the bias is less important than geographical biases which can be assessed with a comparison to the GEOS-Chem 3-D chemistry transport model.

One year of global GOSAT data was retrieved using the UoL-FP v3.4G method. Figure 5.42 shows a comparison of each season between GOSAT and GEOS-Chem, with large scale features consistent with that previously found using UoL-FP v3G (see Chapter 4). However, in general, there are slightly more geographical biases than that found with UoL-FP v3G. Additionally, the zonal mean comparisons show slightly worse correlation coefficients, ranging between 0.62 and 0.94, with the majority of the differences being towards the poles.

The time series of UoL-FP v3.4G $X_{CO_2}$ from GOSAT and GEOS-Chem for a number
of different regions are shown in Figure 5.43. The seasonal cycle is found to match very well between GOSAT and GEOS-Chem for both northern and southern hemispheres, but a difference of 1.59 ppm is observed in the northern hemisphere throughout the year and a smaller difference of 0.79 ppm in the southern hemisphere, which are both larger than that found with UoL-FP v3G. GOSAT and GEOS-Chem $X_{CO_2}$ show a high consistency for both Europe and the USA, with a $X_{CO_2}$ difference of -0.43 ppm and -0.16 ppm and correlation coefficients of 0.69 and 0.72, respectively. These values are similar to that observed with UoL-FP v3G, as is that of South Asia, which on average, GOSAT and GEOS-Chem agree with a mean difference of -0.27 ppm and correlation coefficient of 0.70, but differences are viewed in the seasonal cycle with GEOS-Chem over-estimating the $X_{CO_2}$ in autumn 2009 and under-estimating during spring 2010.

The retrieved $X_{CO_2}$ over the Amazon shows a small improvement (compared to UoL-FP v3G) with the bias reducing to 1.17 ppm and standard deviation reducing to 3.31 ppm. Similarly, the bias and standard deviation over NW Russia is improved with values of -0.38 ppm and 3.58 ppm, respectively. Tropical Africa remains very similar to that of UoL-FP v3G.

In central Asia where a desert region is observed with the high gain mode, the seasonal cycle of GOSAT and GEOS-Chem is observed to agree well ($r=0.73$), but consistently with a $\sim2$ ppm difference in value, similar to that of UoL-FP v3G. Also, a $\sim3$ ppm mean difference is found over the Sahara between GOSAT and GEOS-Chem.
Figure 5.40. Comparison of UoL-FP v3.4G retrieved $X_{\text{CO}_2}$ from GOSAT with retrieved $X_{\text{CO}_2}$ from seven TCCON site for coincident cloud-screened observations within 2 hours and $5^\circ$ between April 2009 and May 2011. All TCCON data shown in light green, coincident daily mean TCCON shown in dark green, all GOSAT data shown in light red and daily mean GOSAT shown in dark red (with daily mean points with only one value per day are represented by open circles). The average difference between the daily means is given as the bias (ranging from -1.21 to 0.94 ppm), the standard deviation of daily means by the Std (between 1.84 and 2.87 ppm) and the correlation coefficient of daily means by r (-0.11 at Darwin to 0.84 at Park Falls). The lower right panel gives the correlation of daily mean coincident retrieved GOSAT $X_{\text{CO}_2}$ with daily mean TCCON $X_{\text{CO}_2}$ within ±2 hours for the seven sites. This gives an overall bias of -0.18 ppm, scatter of 2.45 ppm and correlation of 0.70 between GOSAT and TCCON $X_{\text{CO}_2}$.

$X_{\text{CO}_2}$ where a desert region is observed with the medium gain mode, but the seasonal cycle agrees well with a correlation coefficient of 0.76. Australia, which comprises partially of desert and is observed partially with medium and high gain modes, also shows a difference in $X_{\text{CO}_2}$ between GOSAT and GEOS-Chem with a value of -1.14 ppm. No improvement is observed in any of these desert regions, compared to UoL-FP v3G.

In the UoL-FP v3G retrievals, $X_{\text{CO}_2}$ biases were found with most parameters (see Chapter 4). Figure 5.44 shows a comparison of UoL-FP v3G and UoL-FP v3.4G results for correlations of $X_{\text{CO}_2}$ with various retrieval parameters. The UoL-FP v3.4G method significantly reduces all of these parameter biases, indicating that the retrieval performs
Figure 5.41. Comparison of the point-by-point biases of UoL-FP v3G and UoL-FP v3.4G retrieved $X_{CO2}$ from GOSAT with retrieved $X_{CO2}$ from seven TCCON site for coincident cloud-screened observations within 2 hours and 5° between April 2009 and May 2011. UoL-FP v3G daily mean GOSAT-TCCON bias is shown in red (with daily mean points with only one value per day are represented by open circles). Similarly, the UoL-FP v3.4G daily mean GOSAT-TCCON bias is shown in blue. The average bias difference ranges from -0.01 to 0.40 ppm, the standard deviation (Std) ranges between 0.30 and 0.54 ppm, and the correlation coefficient (r) has a station-to-station range of 0.98 to 0.99. The lower right panel gives the correlation of the daily mean biases and shows an overall bias difference of -0.05 ppm, scatter of 2.34 ppm and correlation of 0.71 between UoL-FP v3G and UoL-FP v3.4G. Hence, this shows UoL-FP v3.4G improves the $X_{CO2}$ slightly.

much better. In Chapter 4, a bias correction scheme was introduced that attempts to correct the retrieved $X_{CO2}$ using four parameters that correlate most with the $X_{CO2}$ bias. Figure 5.44 shows the correlation of these parameters with $X_{CO2}$ for the UoL-FP v3G method and the UoL-FP v3.4G method. This shows that the UoL-FP v3.4G method virtually removes the band 1 albedo slope bias and the band 2 / band 3 mean signal bias, suggesting that these biases were not spectroscopy/signal related but actually aerosol related biases and are corrected using this method. The biases with ice optical depth and surface pressure difference still exist, indicating that these may not be related to aerosols. The ice optical depth bias may potentially be due to the retrieval algorithm not accounting sufficiently for cirrus/clouds, which has been shown to create $X_{CO2}$ biases...
Figure 5.42. Comparison of UoL-FP v3.4G $X_{CO_2}$ retrieved from GOSAT and $X_{CO_2}$ calculated from GEOS-Chem CO$_2$ profiles convolved with the scene-specific GOSAT averaging kernel. For each season, GOSAT is shown at the top, the difference between GOSAT and GEOS-Chem in the middle, and a zonal mean comparison of them at the bottom. The global maps indicate the largest differences are observed over desert regions, such as the Sahara and central Asia. The zonal mean comparison shows a good agreement between GOSAT and GEOS-Chem, but worse than that observed with UoL-FP v3G (see Figure 4.11. It also includes the average TCCON $X_{CO_2}$ from each site which in general agrees, with observed differences mostly due to zonal averaging of GOSAT/GEOS-Chem data.

(see Section 5.3).
Figure 5.43. Comparison of UoL-FP v3.4G $X_{CO_2}$ retrieved from GOSAT and $X_{CO_2}$ calculated from GEOS-Chem CO$_2$ profiles convolved with the scene-specific GOSAT averaging kernel, for a number of different regions between June 2009 and May 2010. The locations of these regions are indicated on the map by the black boxes (top left). The top centre and top right panels give the time series of GOSAT and GEOS-Chem data for the northern and southern hemispheres, showing the seasonal cycle to be well matched but with GOSAT data offset lower by -1.59 ppm in the northern hemisphere and -0.79 ppm in the southern hemisphere. The second row of plots show a very good agreement between GOSAT and GEOS-Chem both in value and seasonality for regions containing urban environments. Below these, the time series of forested regions are shown, with GOSAT reproducing the GEOS-Chem data very well but with a larger scatter and fewer data points. The bottom panels show the time series for 3 different desert regions, with GOSAT matching the seasonal cycle of GEOS-Chem very well but offset lower in value from -1.14 to -3.38 ppm.
Figure 5.44. Comparison of UoL-FP v3G and UoL-FP v3.4G results for correlations of $X_{CO_2}$ with various retrieval parameters. The blue diamonds represent the data, the red line gives a linear fit of the data with the equation of the fit given in red, and the correlation coefficient is given by $r$. 

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Figure 5.45. Comparison of UoL-FP v3G and UoL-FP v3.4G results for correlations of $X_{CO_2}$ with each of the parameters previously used for the bias correction. The blue diamonds represent the data, the red line gives a linear fit of the data with the equation of the fit given in red, and the correlation coefficient is given by $r$. 
5.7 Scattering Conclusion

A flexible simulation algorithm was designed and used to test how various aerosol and cloud components effect retrievals of $X_{CO2}$. The aerosol properties were found to have the largest effect on $X_{CO2}$. The simulator was also used to test different aerosol setups in the retrieval of $X_{CO2}$, finding that spatially dependent AOD and profile shapes do not reduce the $X_{CO2}$ bias. However, spatially dependent aerosol optical properties show an improvement in the $X_{CO2}$ retrieved, with the bias and scatter reduced. Therefore, a new approach was applied to real retrievals of $X_{CO2}$ from GOSAT where the aerosol optical properties were replaced with spatially dependent aerosol optical properties. A comparison with seven different TCCON sites showed that the new approach reduces the bias slightly. A comparison with GEOS-Chem showed that the geographical biases are not improved. This may be due to potential problems with instrument calibration and spectroscopy that are coupled with aerosols in the retrieval of $X_{CO2}$. However, the biases observed in different retrieval parameters, such as the band 1 albedo slope and the ratio of band 2 to band 3 mean signal, have been significantly reduced and thus suggests the new method removes some of the biases, but not all.
Chapter 6

Conclusions

6.1 Assessment of GOSAT retrievals

CO₂ retrievals with high accuracy and precision from satellites is a major challenge but can be of great benefit for the understanding of the carbon cycle. Here, an existing optimal estimation retrieval algorithm developed for the OCO mission was modified and applied to retrieved CO₂ from SWIR radiances observed by the Japanese GOSAT satellite. The main developments of the algorithm included the setup of the instrument and retrieval strategy as well as creation of the pre-processing, cloud screen and post-processing aspects. These advancements set the algorithm apart from the original and was therefore named as the UoL-FP retrieval algorithm.

The algorithm has successfully retrieved X\textsubscript{CO₂} from GOSAT observations for over two years. Comparisons were performed to observations by the TCCON ground based network with a mean smoothing error difference of -0.14 ppm. The single sounding precision was found to be between 2.04 and 2.96 ppm and the bias between -0.87 and 0.77 ppm for seven TCCON sites distributed in the USA, Europe and Australia. Using a stricter post-screen improves the comparison by reducing the single sounding precision to 1.50 to 2.15 ppm and the bias to -0.6 to 0.7 ppm, but at the cost of the number of exposures. The mean bias inferred from TCCON comparisons is very accurate with mean biases approaching only a few tenths of a ppm, which is close to typical retrieval requirements of systematic bias for X\textsubscript{CO₂} (e.g. Chevallier 2007b, Chevallier et al. 2009, Miller et al. 2007). Future improvements in spectroscopy, instrument calibration and atmospheric scattering are expected to provide improved results.

Global seasonal comparisons of retrieved X\textsubscript{CO₂} from GOSAT to model calculations of the GEOS-Chem 3D chemistry transport model are demonstrated. A correlation of
0.61 was determined between GOSAT and GEOS-Chem, with a correlation of 0.76 for the latitudinal gradient. A high level of consistency between GOSAT and GEOS-Chem was observed over expected source regions for both the seasonal cycles and amplitudes of $X_{CO_2}$. Large biases were found over central Asia and the Sahara, but with the development of a bias correction scheme these were significantly reduced.

To further improve and test the retrieval algorithm, a simulator was built to allow multiple retrieval approaches to be tested and was initially used to analyse the effects of aerosols and clouds on the retrieval of $X_{CO_2}$. It was further used to develop a new scattering method for use in the $X_{CO_2}$ retrievals based on aerosol information from the GEMS model. This new method was applied to real GOSAT $X_{CO_2}$ retrievals and a comparison to TCCON gave an improvement of 0.02 ppm for the bias. Although a global comparison with the GEOS-Chem model showed the geographical biases to become slightly worse towards the poles, the biases of some retrieval parameters largely improved. This simulator has provided a framework for future retrieval algorithm developments.

6.2 Outlook

6.2.1 Future Implications

The $X_{CO_2}$ retrievals approach the point where they should be useful for the inversion of CO$_2$ surface fluxes with data assimilation methods (e.g. Baker et al. 2010, Chevallier et al. 2009, Feng et al. 2009). If assimilated into chemistry transport models, the high density of globally inferred $X_{CO_2}$ from GOSAT has the potential to reduce model uncertainties of surface fluxes, especially over regions poorly sampled by ground-based measurements. This could lead to an improved quantification of terrestrial CO$_2$ sources and sinks, and therefore a better understanding of the carbon cycle. This would allow more accurate predictions of future climate change and improved methods of adaptation. Furthermore, as government legislations, both national and international, aim to reduce future emissions of CO$_2$, accurate predictions of global CO$_2$ fluxes using satellite data could allow improved policies and global monitoring.

6.2.2 Retrieval algorithm development

Improving the retrieval algorithm is an ongoing task and there are many ways that it could be done. Any enhancement upon the existing spectroscopy would clearly be beneficial.
The a priori could also be improved, for example having the CO\textsubscript{2} profiles for years surpassing 2009 would provide better spatial estimates as it would include the biomass burning for the actual years.

Although the aerosol approach has already been further developed, there is still other options that could be explored that might prove to be superior. Such options might include; deriving aerosol information from TANSO-CAI as this would provide it exactly for the TANSO-FTS observation location and time, using an alternative aerosol model (e.g. SPRINTARS (Takemura et al. 2003)), or using generic small, medium and large sized aerosol particles for the scattering properties as this could give a more flexible scheme. The use of TANSO-CAI observations has been limited so far by data storage constraints. However, with increased storage capabilities this data could potentially provide information including; aerosol optical depth, aerosol type, cloud optical depth, cloud type, and aerosol/cloud distribution within the IFOV. TANSO-CAI could then also be used as an alternative cloud screening method as demonstrated by Taylor et al. (2012), allowing the possibility of comparing cloud fraction with X\textsubscript{CO2}.

The simulator could be used to test many different retrieval designs, including new approaches of aerosols, cirrus clouds and water clouds as well as retrieval algorithm variations, such as using alternative spectral bands or different state vector parameters. For this reason, the simulator could also be further developed to provide more realistic simulations of scenes. This could involve albedo, topographic, aerosol, cirrus, and cloud variations across the IFOV.

Possible ways to improve the present X\textsubscript{CO2} could be to apply a new bias correction method or to advance the pre/cloud/post-screens. Finally, performing more detailed inter-comparisons with alternative retrieval algorithms could indicate possible retrieval algorithm dependent issues that could be improved.

6.2.3 Future missions

Whilst GOSAT was the first greenhouse gas dedicated satellite, there are many more future missions currently planned. The next satellite that will allow CO\textsubscript{2} to be retrieved will be the Orbiting Carbon Observatory 2 (OCO-2) satellite through the NASA Earth System Science (ESSP) program, for launch in 2014. OCO-2 will include one instrument consisting of three high resolution grating spectrometers that will observe O\textsubscript{2} and CO\textsubscript{2} absorption bands with a spatial resolution of 1 x 1.5 km\textsuperscript{2} and swath width of 10 km. This high spatial resolution instrument has the potential to make observations in the small gaps between clouds, thus providing more measurements globally.
It is proposed that in 2015/2016 the Tropical Carbon Mission (TCM) may launch, which is a bilateral Earth Observation mission between the UK and US space agencies. It will contain a NASA Orbiting Carbon Observatory 3 (OCO) instrument and a UK Carbon Monoxide and Methane Spectrometer (CMS). It will measure between $\pm 35^\circ$ latitude and will reduce sampling biases from clouds and the diurnal CO$_2$ cycle by using a precessing orbit at 430 km altitude and 35$^\circ$ inclination. This is proposed to complement the OCO-2 mission by providing more measurements over the tropical regions.

There will be a GOSAT-2 satellite launching in 2016. Its design specifications will be finalised this year, but it will likely be of similar design to GOSAT. This would have the benefit of globally repeating measurements over the same locations, but with the disadvantage of limited coverage.

The Carbon Monitoring Satellite (CarbonSat) is planned to be launched in 2018 as part of the ESA Earth Explorer Opportunity Missions (EE-8). CarbonSat will comprise of two instruments; a Cloud and Aerosol Imager (CAI) and an Imaging Spectrometer (IS) that will observe radiances in the O$_2$, CH$_4$ and CO$_2$ absorption bands. It will have a spatial resolution of 2 x 2 km$^2$ and swath of 500 km, providing a very large coverage and highly dense dataset of observations globally.

For possible launch in 2018 is the MicroCarb satellite being developed as part of the CNES microsatellite series. The instrument design will either be a scattering grating spectrometer or a static Fourier Transform interferometer and will measure radiances in the O$_2$ and CO$_2$ absorption bands. If this mission proves beneficial it could lead to a constellation of microsatellites providing increased sampling.

The last mission currently planned is the Active Sensing of CO$_2$ Emissions over Night, Days and Seasons (ASCENDS) satellite for launch in 2021 or later as a NASA Earth science decadal survey study. ASCENDS is currently planned to use laser-based differential absorption of O$_2$ and CO$_2$. This active method would allow observations to be made at night time, at high latitudes and through all seasons irrespective of how much cloud exists. For improved flux interpretation, it is planned to also contain a passive CO sensor.

The present retrieval algorithm and simulator were designed with a flexible scheme to allow alternative instruments to be used. Therefore, these could potentially be used with any of the future missions, allowing the development of a long-term dataset of $X_{CO2}$. 

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6.2.4 Final comments

This thesis describes a new retrieval algorithm capable of inferring $X_{CO2}$ from remotely sensed SWIR satellite observations. A comparison with coincident ground-based TCCON measurements shows that with strict filtering the retrieval algorithm should be sufficient to allow improved surface flux estimates (e.g., Chevallier 2007b, Chevallier et al. 2009, Miller et al. 2007). Furthermore, over TCCON sites the $X_{CO2}$ retrieval is very accurate with biases approaching only a few tenth ppm, which is close to typical retrieval requirements of systematic bias for $X_{CO2}$ (e.g. Chevallier 2007b, Chevallier et al. 2009, Miller et al. 2007).

The latest results from other algorithms (Bril et al. 2007, Yokota et al. 2009, Butz et al. 2011, Morino et al. 2011, Wunch et al. 2011b, Yoshida et al. 2010, Crisp et al. 2012, Oshchepkov et al. 2012a) find similar results with a variation in the number of soundings potentially due to the screening criteria applied. However, over desert regions, which are not observed by TCCON, it is found that the retrieved $X_{CO2}$ from GOSAT shows significant (up to $\sim$3 ppm) differences when compared to GEOS-Chem model calculations which are believed to be biases in the GOSAT retrieval. Similar findings have been reported in Crisp et al. (2012) who suggested that these biases are related to instrument gain. Although here, biases are observed for desert regions observed with both settings of the instrument gain, showing that these biases are only partially due to gain and this leads to the speculation that these biases could be introduced by desert dust or related to the high surface albedo in the CO$_2$ bands. This study highlights the need for further algorithm improvements, especially over the deserts. However, except for the desert regions, the $X_{CO2}$ retrievals approach the point where they should be useful for the inversion of CO$_2$ surface fluxes with data assimilation methods (e.g. Baker et al. 2010, Chevallier et al. 2009, Feng et al. 2009), which could then lead to a better understanding of the carbon cycle and improvements in future climate predictions.

Finally, this thesis also describes new simulator that can be used for the development of SWIR retrieval algorithms. Its use has been demonstrated for testing new scattering methods for retrievals of $X_{CO2}$ from GOSAT, but the flexibility of the simulator design is such that it could be applied to future missions.
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