MEASUREMENTS OF AMMONIA AND NITROUS OXIDE EMISSIONS FROM POTATO FIELDS IN CENTRAL WASHINGTON USING DIFFERENTIAL OPTICAL ABSORPTION SPECTROSCOPY (DOAS), TRACER DISPERSION, AND STATIC CHAMBER METHODS

By

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MEASUREMENTS OF ATMOSPHERIC AMMONIA AND NITROUS OXIDE EMISSIONS FROM POTATO FIELDS IN CENTRAL WASHINGTON USING DIFFERENTIAL OPTICAL ABSORPTION SPECTROSCOPY (DOAS), TRACER DISPERSION, AND STATIC CHAMBER METHODS

Abstract

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There are limited studies that have measured the magnitude of Nitrogen (N) emissions from intensively managed croplands. In this research, concentrations of ammonia (NH₃) and nitrous oxide (N₂O) from center-pivot irrigated potato fields in central Washington were measured during the 2007 and 2008 growing seasons (June – August). A short-path differential optical absorption spectroscopy (DOAS) instrument that operates in the 200-240 nm wavelength spectral range was used to measure NH₃ concentrations. During some periods, sulfur hexafluoride (SF₆) tracer gas was released from the center of the fields and concentrations were measured along a crosswind sample line downwind of the fields to help characterize local dispersion conditions. A Gaussian plume dispersion model (SIMFLUX) was used, along with the SF₆ tracer data in an inverse approach to estimate NH₃ emission rates. A static chamber flux method was used to measure N₂O emissions with sample analysis via electron capture gas chromatography (ECD-GC). Nitrous oxide flux rates generally peaked between 2.5 and 3.5 hours after irrigation or fertigation. The amount of N lost as N₂O ranged from zero to 12% of the applied N fertilizer which is comparable to documented rates of 0.1 to 8% (Dobbie and Smith,
Measured NH₃ concentrations varied with application status, sprinkler arm location, and wind speed. The highest NH₃ concentrations measured in the summers of 2007 and 2008 were ~34 ppbv and ~112 ppbv, respectively. Both peaks occurred during fertigation, when the sprinkler arm was spraying almost directly towards the DOAS measurement path and winds greater than 1 m s⁻¹ were blowing towards the measurement path. Estimated NH₃ flux rates ranged from 9 μg N m⁻² s⁻¹ to 460 μg N m⁻² s⁻¹ for fertigation periods and from 0.6 μg N m⁻² s⁻¹ to 2 μg N m⁻² s⁻¹ for periods of no fertigation or irrigation, which indicates that a significant fraction of NH₃ loss occurs as a result of direct volatilization of NH₃ during fertigation. Overall, the average NH₃ flux during fertigation periods was 10% of N applied as fertilizer.
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Dedication

This thesis is dedicated to my mother and father for their unconditional love and support.
CHAPTER 1

INTRODUCTION

1.1 Motivation and Research Objectives

The increasing use of synthetic Nitrogen (N) fertilizers has led to an altered nitrogen cycle and a doubled amount of fixed N entering the biosphere (Vitousek et al., 2000). There are uncertainties in the magnitude of N emissions from various sources, including intensively managed croplands. Up to 50% of the applied N to croplands can be lost, with gaseous emissions being the dominant N loss mechanism (IFA, 2001).

Excess emissions of NH₃ and N₂O can have negative effects on human health and the environment. Ammonia reacts rapidly with both sulfuric and nitric acids in the atmosphere to form fine particles, which cause health problems, reduce visibility, cause acid deposition, and perturb the earth’s radiation balance (Pandis, 1995). Nitrous oxide is a potent greenhouse gas which ranks just below carbon dioxide (CO₂) and methane (CH₄) in global warming concerns (EPA, 2006). Due to the impacts on the environment and human health, it is important to quantify and understand the factors that control NH₃ and N₂O emissions.

In this research, concentrations of NH₃ and N₂O from potato fields in central Washington (WA) were measured during the 2007 and 2008 growing seasons (June – August). A short-path differential optical absorption spectroscopy (DOAS) instrument that operates in the 200-240 nm wavelength spectral range was used to measure NH₃ concentrations. During some periods, sulfur hexafluoride (SF₆) tracer gas was released from the center of the fields and concentrations were measured along a crosswind sample line downwind of the fields to help characterize local dispersion conditions. A Gaussian plume dispersion model (SIMFLUX) was used, along with the SF₆ tracer data in an inverse approach to estimate NH₃ emission rates. Nitrous oxide fluxes
were measured using a static chamber method and samples were analyzed using a gas chromato
graph equipped with an electron capture detector. The overall goals for this work are to improve our understanding of the loss of N gases from croplands and to account for the impact of these emissions upon regional air quality and global climate change.

This thesis is presented in six chapters. In the remainder of Chapter 1, the research location is described. In Chapter 2, literature describing the sources, controlling factors, and measured rate of N$_2$O and NH$_3$ emissions are reviewed. In Chapter 3, the measurement techniques used for measuring N$_2$O and NH$_3$ concentrations and the methods for estimating the flux rates are presented. Chapter 4 is a manuscript which summarizes the results in terms of NH$_3$ and N$_2$O concentration and emission patterns. Chapter 5 provides an analysis of errors. Finally, Chapter 6 provides conclusions from this research and possible future work.

1.2 Research Location Description

In central WA, approximately 62,000 hectares of cropland are used to grow more than 4.33 million tons of potatoes (data from 2005) (Liu et al., 2007). Information on NH$_3$ and N$_2$O emissions from the potato fields in the area are scarce making this a viable area of study. Potato crops are managed with high inputs of water and N fertilizers. The soil properties, climate, and management practices in the area promote N$_2$O and NH$_3$ losses from croplands. In section 1.2.1, the location of the measurement site will be discussed. The soil properties and climate as well as the management practices for the research location will be presented in sections 1.2.2 and 1.2.3, respectively.
1.2.1 Location Background Information

The potato fields used for this project are located approximately 16 miles east of Othello, WA, US. Summer 2007 measurements were conducted at potato field 19-7, shown in Figure 1.1 (N 46° 46.166’, W 118° 58.099’). Summer 2008 measurements were conducted at potato field 25-1, shown in Figure 1.2 (N 46° 45.659’, W 118° 52.357’). These fields were preferred over others because they were relatively flat and they were far from other potentially interfering sources of NH\textsubscript{3} and N\textsubscript{2}O. Fields of grass seed, alfalfa, and wheat, which are not typically managed with high inputs of N fertilizers, surrounded the two potato fields. Fields 19-7 and 25-1 were planted with ranger russet potatoes and with premier russet potatoes, respectively. The crop rotation for both fields is grass seed - wheat - potatoes.

![Figure 1.1: Aerial view of the potato field in central WA used for measurements in summer 2007 (Figure adapted from GoogleEarth). Surrounding the field were grass seed (A) and alfalfa (B) fields which are not typically managed with high amounts of N fertilizers.](image)
1.2.2 Soil Properties and Climate

Quincy fine sand soils are dominant in the region, where anaerobic conditions are easily maintained for long periods of time. The surface soil has a bulk density of 1.24 Mg per m$^3$. The soil porosity and pH are 0.53 and 6.6, respectively.

The growing period is characterized by lack of cloud cover, high daytime temperatures, and cool nights. The average annual rainfall is 150 mm so the crops are highly dependent on irrigation. As a result, crops are generally irrigated five out of seven days per week and often above the rate necessary for crop growth.
1.2.3 Management Practices

In winter, field preparation included deep ripping and disk plowing of the soil and ground rig fumigation. Deep ripping and disk plowing were conducted to break up traffic-induced or naturally occurring compaction layers and to increase soil moisture. In addition, potassium (K) and phosphorous (P) were applied.

In the spring, a chisel chopper was used to loosen and plant (seed) the soil. A rod weeder was used to kill weeds before plant emergence. Between two to six weeks after planting, the potato plants emerged from the ground and irrigation (application of water only) began. The plants began to grow tubers a few weeks after emergence. A dense, green canopy from approximately June to August was maintained. Heavy N fertilizer application began in early July. The potatoes were harvested during the first weeks of October, depending on the type of potato planted.

Water and N fertilizer were applied to the potato fields using a center-pivot irrigation system. In the center-pivot irrigation system, sprinklers were positioned over the crop canopy and along the length of connected pipe segments. The pipe segments were supported by trusses and mounted on wheeled towers. The sprinkler system rotated in a circular pattern and was fed with water or water with N fertilizer from the pivot point at the center of the field.

The timing and rate of irrigation and fertigation (application of water with N fertilizer) varied from day to day and from each field, under the discretion of the farmer. The type of fertilizer used for both fields was UAN solution 32, which contained 25% nitrate-N, 25% ammonium-N, and 50% urea-N. There were two rates for sprinkler rotation; 12 degrees clockwise (cw) per hour and 18 degrees cw per hour. For days of NH₃ measurement, 2.7 kg of N
was applied per acre per full 360° rotation. When pesticides were being applied, the sprinkler arm rotated 30 degrees cw per hour.
2.1 The Nitrogen Cycle

The N cycle describes the movement of different forms of N species between the atmosphere, biosphere, and geosphere. Nitrogen is essential for all life forms on Earth since it is an important component of DNA, RNA, and proteins, the building blocks of life. About 20% of the global N is in sedimentary rocks, and is not readily available to organisms. The Earth's atmosphere is the major reservoir of N and is composed of 78% diatomic nitrogen (N\textsubscript{2}) (Galloway et al., 2004). Most organisms cannot use N in the form of N\textsubscript{2} due to the strong triple bond that binds the two N atoms together.

The form usable by most organisms is reactive N (Nr), which includes all biologically active, photochemically reactive, and radiatively active N compounds. Reactive N compounds include inorganic reduced forms of N, such as NH\textsubscript{3} and ammonium (NH\textsubscript{4}\textsuperscript{+}). Inorganic oxidized forms of N, such as nitrogen oxides (NO + NO\textsubscript{2} = NO\textsubscript{x}), nitric acid (HNO\textsubscript{3}), nitrous oxide (N\textsubscript{2}O), nitrate (NO\textsubscript{3}\textsuperscript{-}) and organic compounds, such as urea, amines, proteins, and nucleic acids, are also included.

Diatomic N (N\textsubscript{2}) can be naturally converted to Nr in two ways: biological fixation and high-energy natural events. In biological fixation, symbiotic bacteria and free-living bacteria convert N\textsubscript{2} into Nr. Only about 10% of the Nr produced through this process is emitted into the atmosphere (Galloway et al., 1998). High-energy natural events, such as lightning strikes and forest fires, can break the strong triple bond that binds the atoms of N\textsubscript{2}. The resulting individual N atoms combine with molecular oxygen to form NO. Production of Nr through high-energy
natural events occurs only at the global rate of 3 to 10 Tg N year\(^{-1}\) (Prather et al., 2001 as cited in Galloway et al., 2001).

Human activities, mainly the production and use of synthetic N fertilizers and burning of fossil fuels, during the past century have altered the N cycle. The amount of fixed N that enters the biosphere has doubled (Vitousek et al., 2000). In 1913, the Haber-Bosch process, which describes the formation of NH\(_3\) by combining N\(_2\) and hydrogen in high temperature and pressure, was discovered. The NH\(_3\) formed is used to make fertilizer, which has led to increased agricultural productivity. It is predicted that by 2020, the current global production of synthetic N fertilizer of 80 Tg year\(^{-1}\) (1 Tg = 10\(^{12}\) g) will increase to 134 Tg year\(^{-1}\) (Vitousek et al., 2000).

The second main cause of human-fixed N is the burning of fossil fuels in automobiles, power generation plants, and industries. During the combustion of fossil fuels, N is emitted to the atmosphere as a waste product (NO) from either the oxidation of atmospheric N\(_2\) or organic N in the fuel. One study predicts that production of NO\(_x\) from fossil fuels will more than double in the next 25 years, from about 20 Tg year\(^{-1}\) to 46 Tg year\(^{-1}\) (Vitousek et al., 2000).

2.2 Human Health and Environmental Impacts

The human alteration of the N cycle has consequences. Excess emissions of NH\(_3\) and N\(_2\)O negatively impact the environment and human health. Ammonia reacts rapidly with both sulfuric and nitric acids in the atmosphere to form fine particles or aerosols (Pandis, 1995). Human exposure to aerosols negatively impact human health. Long-term exposures (months to years) cause deterioration of lung function and development of chronic bronchitis (EPA, 2003). Short-term exposures (hours to days) aggravate lung diseases and increases susceptibility to respiratory infections (EPA, 2003). Aerosols also reduce visibility in regional and urban areas.
and perturb the earth’s radiation balance (Pandis, 1995). In addition, deposition of NH₃ can lead to eutrophication and acidification of ecosystems (EPA, 2007).

Nitrous oxide is a potent greenhouse gas. The global warming potential (GWP) is a measure of how much a given mass of greenhouse gas contributes to global warming relative to the same mass of carbon dioxide (IPCC, 2007). The Intergovernmental Panel on Climate Change (IPCC) reports that the GWP of N₂O over a 100 year period is 310 (IPCC, 2007). Despite its low atmospheric concentration, N₂O ranks just below carbon dioxide and methane in importance. Due to their impacts to the environment and human health, it is important to quantify and understand the factors that control NH₃ and N₂O emissions.

### 2.3 Sources of Nitrous Oxide (N₂O)

#### 2.3.1 Natural Sources

Natural sources of N₂O are primarily from bacterial breakdown of N in soils and oceans through the microbial nitrification and denitrification. Tropical soils emit approximately 6.3 Tg of N₂O annually (IPCC, 1995). Of this, about 75% comes from wet forest soils and 25% comes from dry savannas. Temperate soils, including forest soils and grasslands, emit approximately 3.1 Tg of N₂O annually (IPCC, 1995). Tropical and temperate soils generally have different levels of nutrients, with tropical soils often being phosphorus limited while temperate soils are generally N limited. As a result, added N emissions tend to result in greater N₂O emissions in tropical soils than in temperate soils. Oceans produce approximately 4.7 Tg of N₂O year⁻¹ (EPA, 2006).
2.3.2 Anthropogenic Sources

According to the U.S. Greenhouse Gas Emissions Inventory, the estimated total N\textsubscript{2}O emission in the U.S. in 2006 was 367.9 Tg CO\textsubscript{2} Equivalent (Eq.), with an uncertainty range from 352.2 to 449.3 Tg CO\textsubscript{2} Eq. (EPA, 2008). Table 2.1 lists sources of N\textsubscript{2}O emissions in the U.S. and their contributions.

<table>
<thead>
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Agricultural soils account for a large portion of N\textsubscript{2}O emissions (265.0 Tg CO\textsubscript{2} Eq. in 2006) (EPA, 2008). The application of synthetic N fertilizer and livestock manure to croplands and pasture directly increase the amount of N in the soil, which can be converted into N\textsubscript{2}O. Indirect addition of N\textsubscript{2}O to soils occurs when applied fertilizer or manure volatilizes to NH\textsubscript{3} and is re-deposited onto the soil as particulate NH\textsubscript{4}\textsuperscript{+}, nitric acid, and NO\textsubscript{x}. Also, runoff and leaching of applied N to ground water and surface waters indirectly adds N\textsubscript{2}O to soils.
Nitrous oxide emissions from livestock manure and urine also occurs through nitrification and denitrification. The amount of N\textsubscript{2}O produced depends on the composition of the manure and urine, the type of bacteria present, and the amount of oxygen and liquid present. Emissions of N\textsubscript{2}O from manure management were estimated at 14.3 Tg CO\textsubscript{2} Eq. for the year 2006.

Nitrous oxide emissions also result from the reaction that occurs between N and oxygen during fossil fuel combustion. The type of fuel and pollution control device used as well as the maintenance and operating practices affect the amount of N\textsubscript{2}O emitted. Catalytic converters on vehicles can promote the formation of N\textsubscript{2}O, although the latest technical modifications to converters are addressing this problem. From 1990 to 2006, N\textsubscript{2}O emissions from mobile combustion decreased by 24 percent. The total N\textsubscript{2}O emission from mobile combustion was 33.1 Tg CO\textsubscript{2} Eq. in 2006 (approximately 9 percent of total U.S. N\textsubscript{2}O emissions).

Nitric acid production accounts for 15.6 Tg Eq. CO\textsubscript{2} for N\textsubscript{2}O emissions in 2006 while adipic acid production accounts for 5.9 Tg CO\textsubscript{2} Eq. Nitric acid is used for production of synthetic fertilizers, adipic acid, and explosives.

Wastewater treatment plants are another anthropogenic source of N\textsubscript{2}O emissions, where human sewage and other household wastewater end up. Nitrous oxide emissions result from both nitrification and denitrification of the N present in the wastewater, usually in the form of urea, ammonia, and proteins. Wastewater treatment plants account for 8.1 Tg Eq. CO\textsubscript{2} of N\textsubscript{2}O emissions.
2.4 Factors that Affect N₂O Emissions

There are many factors regulating the loss of N₂O from croplands to the atmosphere. Section 2.4.1 will discuss the role of nitrification and denitrification in N₂O emissions. In section 2.4.2, the role of crop management will be discussed.

2.4.1 Nitrification and Denitrification Rates

The microbial process of denitrification and nitrification are the major sources of N₂O emission from the soil. Nitrification is the biological oxidation of ammonium (NH₄⁺) to nitrite (NO₂⁻) or nitrate (NO₃⁻) under aerobic conditions and N₂O is a byproduct of nitrification (Seinfeld and Pandis, 1998). In denitrification, which is the bacterial process of converting nitrate (NO₃⁻) to dinitrogen (N₂) and N₂O under anaerobic conditions, N₂O is an intermediate (Seinfeld and Pandis, 1998). Nitrification rates are relatively uniform throughout a field while denitrification rates vary temporally and spatially (IFA, 2001). Various factors control whether nitrification or denitrification dominates in the production of N₂O.

Haerkort and MacKerron (2000) and the International Fertilizer Industry Association (IFA) (2001) have summarized the factors that affect nitrification and denitrification in crop production. They concluded that in most soils, nitrification depends on the amount of NH₄⁺ and oxygen in the soil as well as the soil pH and temperature. The amount of NH₄⁺ in the soil is derived from either mineralization of organic matter or from applied N fertilizer. Nitrification activity is generally suitable for pH levels between 4 to 8, and at high pH values (pH > 8) it may be inhibited due to toxic effects to the bacteria. When the soil temperature is above 4 °C, nitrification activity is favorable and is optimized when the soil temperature is between 20-40 °C. The factors that control nitrification are not the same for denitrification.
Denitrification depends on soil components such as the availability of organic carbon (C), oxygen, and NO₃⁻. Organic compounds, such as organic C, serve as electron donors and are sources of bacterial cellular material for denitrification activity. As soil oxygen content decreases, denitrification activity increases since anaerobic conditions are required for both synthesis and activity of the enzymes involved in the denitrification process. Denitrification rates generally increase linearly with NO₃⁻ concentrations up to 40 mg L⁻¹, while at relatively high NO₃⁻ concentrations denitrification is independent of NO₃⁻ concentration.

Similar to nitrification, the soil pH controls bacterial denitrification activity. Under acidic conditions (pH < 7), denitrification rates are slower than in alkaline conditions (IFA, 2001, Haverkort and MacKerron, 2000). In contrast to nitrification, soil temperature does not control denitrification activity because soil denitrifiers are adapted to and are capable of growing under a wide range of temperatures (Haverkort and MacKerron, 2000).

Other controlling factors for denitrification include the presence of decomposing organic matter and the magnitude of NH₃ volatilization. The presence of decomposing soil organic matter and plant litter produces anaerobic areas where high denitrification rates occur. NH₃ volatilized from applied fertilizer can be re-deposited onto the soil, indirectly affecting the availability of N for nitrification or denitrification (Haverkort and MacKerron, 2000).

2.4.2 Management Practices

Various agricultural management activities affect N₂O emissions from soils. Such activities include the application of nitrogen and excessive water and soil tillage. Nitrogen inputs to the soil, such as N fertilizer, animal manure, and crop residues, increase the amount of N available for nitrification and denitrification. Thus, N₂O emissions may increase with the N application rate. The production of N₂O is also related to the timing of fertilizer application.
When NH$_4^+$-based fertilizers are applied to the soil under conditions suitable for nitrification activity without any competition from plant uptake, N$_2$O emissions are likely to increase (IFA, 2001). This is also true when NO$_3^-$-based fertilizers are applied to soil under conditions suitable for denitrification activity.

Soil moisture is another factor since the soil moisture content affects the diffusion of oxygen through the soil. The rate of N$_2$O emission increases as the water-filled pore space (WFPS) in the soil increases. High N$_2$O emissions have been observed when the WFPS is between 60 and 90%, when the production of N$_2$O is dominated by denitrification (Dobbie et al., 1999, IFA, 2001, Flessa et al., 2002). High water content may result from heavy rain or excessive rates of irrigation. Below 60% WFPS nitrification, an aerobic process, is the dominant process for the production of N$_2$O (IFA, 2001).

Also, localized spatial variations in micro-relief (such as ridges and furrows or raised beds) or in water infiltration rates due to compaction, can result in localized wet spots. Usually, greater N$_2$O emission rates occur from tractor-compacted furrows than from ridges due to a greater reduction of air-filled pore space (Flessa et al., 2002, Smith et al., 1997). Flessa et al. (2002) measured mean WFPS values of 61% for tractor compacted inter-rows, 49% for uncompacted inter-rows, and only 30% for ridges for the growing period of potato fields in Germany.

Soil tillage may induce enhanced organic N mineralization (IFA, 2001). Soil tillage is typically conducted during planting and harvesting of root/tuber crops, and is the process of digging up the soil for the purpose of incorporating plant residues and/or fertilizer, seedbed preparation, weed control, and soil and water conservation. Soil tillage affects the gas diffusivity within the soil and the water and air holding capacity of the soil pores.
2.4.3 Other Factors

Soil properties, such as soil texture, affect N$_2$O emissions. Fine-textured soils hold water more tightly because they have more capillary pores within aggregates than sandy soils. Thus, in fine-textured soils, anaerobic conditions are easily maintained for longer periods of time (IFA, 2001).

Gas diffusion also has an influence on emissions of N$_2$O since before being emitted to the atmosphere, the N$_2$O gas must first diffuse through the soil pores. High soil water content, impeded drainage, soil compaction, fine soil texture, and soil surface sealing can all limit gas diffusion through the soil. Haverkort and MacKerron (2002) state that high denitrification activity may be present in soils that are close to saturation, but the emission of N$_2$O from the soil can be low. Thus, anaerobic conditions promote denitrification but limits gas diffusion.

2.5 Emission Rates of N$_2$O

The increase in atmospheric N$_2$O concentration of two to three percent every year is well documented but the sources of the increase are not (Vitousek et al., 2000). Nitrous oxide emissions from agriculture are typically reported as an emission factor (EF), which is the amount of N$_2$O emitted as a percentage of the N fertilizer applied. A default mean EF of 1.25 ± 1% of the N fertilizer applied was adopted by the Intergovernmental Panel on Climate Change (IPPC) in 1997 (Bouwman, 1996). This default EF is based on experiments that were at least 1 year in duration. The default EF is a general approximation and is independent of the specific crop type, region, environment, and management practices. Documented EF values range from 0.1 to 7.8% (Dobbie and Smith, 2003). The use of the default mean EF value is highly uncertain since there are extremely high spatial, temporal, inter-annual, and crop type variability in N$_2$O emissions.
Dobbie et al. (1999) measured N$_2$O emissions from intensively managed fields of ryegrass, winter wheat, potatoes, broccoli, and oilseed rape in Scotland. The fields of grassland, winter wheat, potato, and oilseed rape were treated with ammonium nitrate fertilizer. An ammonium nitrate/ammonium/phosphate/urea fertilizer mix was applied to the broccoli fields. Nitrous oxide fluxes varied widely throughout the year at all sites and also between sites. Emissions from the winter wheat, barley, and oilseed rape fields (0.2 to 0.7 kg N$_2$O per 100 kg applied N; EF: 0.2 to 0.7%) were consistently lower than emissions from the fields of grassland, potato, and broccoli (0.3 to 5.8 kg N$_2$O per 100 kg applied N; EF: 0.3 to 5.8%). Daily fluxes were as high as 1.2 kg N$_2$O hectare$^{-1}$ day$^{-1}$ and annual emissions ranged from 0.3 to 18.4 kg N$_2$O hectare$^{-1}$ year$^{-1}$. At all sites, fluxes generally increased sharply after N-fertilizer additions and gradually decreased with time. All of the reported emission rates were obtained from flux measurements using chambers.

Flessa et al. (2002) reported mean N$_2$O emissions of 1.6 and 2.0 kg hectare$^{-1}$ for the 1997 and 1998 growing periods (end of May to September) of two adjacent potato fields in Germany. The fields were applied with 75 and 37 kg N hectare$^{-1}$ for 1997 and 1998, respectively, resulting in EF values of 2.1% and 5.4%. Flessa et al. obtained these emission rates from flux measurements made by automated gas sampling chambers.

Haile-Mariam et al. (2008) reported N$_2$O losses of 0.5% (0.55 kg N hectare$^{-1}$) of applied fertilizer (112 kg N hectare$^{-1}$) in corn fields and 0.3% (0.59 kg N hectare$^{-1}$) of 224 kg N hectare$^{-1}$ applied fertilizer in potato fields in central Washington during the 2005 and 2006 growing seasons. The fields were applied with urea-ammonium-nitrate (UAN) solution (32% N) through a center pivot irrigation system. The fields were under sweet corn–potato rotation. Aerobic soil
conditions dominated, with soil WFPS ranging from 0.50–0.63 m³ m⁻³. Gas samples were collected using static chambers.

2.6 Sources of Ammonia (NH₃)

Global sources of NH₃ emissions and their estimated contributions and uncertainty range are listed in Table 2.2. Oceans are the major natural source of NH₃ emissions. Ammonia will volatilize from seawater at a rate that depends on the atmospheric NH₃ concentration, water salinity, and temperature (Bouwman et al., 1997). Another natural source of NH₃ is soils under natural vegetation. Ammonia is emitted by organisms within the soil that breakdown organic matter.

<table>
<thead>
<tr>
<th>Source</th>
<th>NH₃-N emission (Tg year⁻¹)</th>
<th>Uncertainty Range (Tg N year⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oceans</td>
<td>8.2</td>
<td>3-16</td>
</tr>
<tr>
<td>Soils under natural vegetation</td>
<td>2.4</td>
<td>0-10</td>
</tr>
<tr>
<td>Animal excreta</td>
<td>21.7</td>
<td>10-30</td>
</tr>
<tr>
<td>Synthetic fertilizer use</td>
<td>9.0</td>
<td>4.5-13.5</td>
</tr>
<tr>
<td>Crops and decomposition of crops</td>
<td>3.6</td>
<td>1.4-5.0</td>
</tr>
<tr>
<td>Biomass burning</td>
<td>5.9</td>
<td>3.9-7.7</td>
</tr>
<tr>
<td>Human excreta</td>
<td>2.6</td>
<td>1.3-3.9</td>
</tr>
<tr>
<td>Fossil fuel combustion, including aircraft</td>
<td>0.1</td>
<td>0.0-0.2</td>
</tr>
<tr>
<td>Industrial processes</td>
<td>0.2</td>
<td>0.1-0.3</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>53.6</strong></td>
<td><strong>40-70</strong></td>
</tr>
</tbody>
</table>

Agriculture contributes between 55% and 95% of the anthropogenic NH₃ emissions (McGinn and Janzen, 1997). Animal excreta are a major source, and there are two reaction pathways for NH₃ formation. One reaction pathway is the reaction of urea in urine with urease in the manure or soil to form liquid ammonia-N (NH₃ and NH₄⁺):

\[
\text{CO(NH}_2\text{)}_2 + 2\text{H}_2\text{O} \Leftrightarrow (\text{NH}_4\text{)}_2\text{CO}_3 \Leftrightarrow 2\text{NH}_4^+ + \text{CO}_3^{2-}
\]  

2.1
Ammonia emissions from urine deposited in natural areas are lower than from high density areas where animals are domesticated. The second reaction pathway is the microbiological anaerobic breakdown of organic N. Increasing human population, and therefore increasing food demand, has led to more domesticated animals and increased NH₃ emissions (Vitousek et al., 1997). Ammonia emissions also result from sprinkler application of dairy lagoon slurry from milking cows (Rumburg et al., 2006).

Application of synthetic fertilizer to croplands is another anthropogenic source of NH₃ emissions. Up to 50% of the applied N to croplands can be lost through gaseous emissions, leaching, runoff, or erosion. Generally, the dominant N loss processes are through gaseous emissions, mainly from volatilization of NH₃ and denitrification (IFA, 2001). Ammonia emissions can come from both soil and crop foliage. Ammonia emissions from soil depend on factors such as soil properties (i.e. soil pH, water content, and porosity). From crop foliage, the emissions depend on the plant developmental stage and a compensation point of atmospheric NH₃, below which vegetation acts as a source and above which vegetation acts as a sink (Lemon and van Houtte, 1980 as cited in McGinn and Janzen, 1997). Other factors such as the type and quantity of fertilizer applied and meteorological factors (i.e. wind speed, temperature, and precipitation) affect NH₃ emissions from soil and crop foliage.

Another source of NH₃ is from human breath, sweat, and waste. Emissions of NH₃ from treatment systems of human waste result from anaerobic degradation and are not well studied (Bouwman et al., 1997).

Incomplete combustion of fossil fuels also produces NH₃ emissions. Information on NH₃ emissions from fuel combustion is scarce and associated with large uncertainties (Bouwman et al., 1997). The use of emission controls in car engines to reduce NO and NO₂ emissions has led
to increased NH₃ emissions. Vehicles with three-way catalytic converters emit more NH₃ than cars with no emission controls (Bouwman et al., 1997). The combustion of coal and natural gas and industrial processes (i.e. production of explosives) also contribute to NH₃ emissions.

2.7 Regulating Factors of NH₃ Emission

There are many factors regulating the loss of NH₃ from soil-plant systems to the atmosphere. Section 2.7.1 will discuss factors that affect NH₃ emissions from the soil. In section 2.7.2, the factors that affect NH₃ emissions from the crop foliage will be discussed.

2.7.1 NH₃ Emissions from Soil

Emissions of NH₃ from soils result from biological degradation of organic compounds and are dependent on the following factors: the transformations of the total ammonical N (TAN) in the soil, which include NH₃-N + NH₄⁺-N, the soil properties, climatic factors, management practices, and the partial pressure between the atmosphere and the soil.

The NH₄⁺ in the fertilizer applied can be bacterially nitrified to nitrate (NO₃⁻), converted to NH₃, or remain as NH₄⁺ (Bouwman and Bouman, 2002). Ammonium will be in equilibrium with NH₃ depending upon the pH of the solution, as shown in the following reaction:

\[ \text{NH}_3 + \text{H}_2\text{O} \leftrightarrow \text{NH}_4^+ + \text{OH}^- \]

Soil properties such as soil pH, moisture content, infiltration rate, and cation exchange capacity (CEC) affect the rate of NH₃ volatilization from the soil to the atmosphere. Ammonia volatilization increases with increasing pH; the greatest NH₃ volatilization occurs when the soil pH is greater than 7.5 (Liu et al., 2007). From 0 to 30 % soil water content, the amount of NH₃ emission increases with increasing moisture due to the inability of the NH₃ molecules to displace water molecules (IFA, 2001). When the soil water content increases from 30 % to 100 %, NH₃
emissions decrease due to dissolution of NH$_3$ in the soil (IFA, 2001). The CEC is the capacity of a soil for ion exchange of positively charged ions between the soil and the soil solution. The lower the CEC, the more likely it is for NH$_3$ volatilization to occur (Bouwman and Bouman, 2002; Haverkort and MacKerron, 2000).

Climatic factors, such as wind speed and temperature, also affect NH$_3$ volatilization from the soil. Increasing wind speed promotes more rapid transport of NH$_3$ away from the water or soil to the atmosphere (Bouwman and Bouman, 2002; Haverkort and MacKerron, 2000). As soil temperature increases, the relative proportion of NH$_3$ to NH$_4^+$ increases while the solubility of NH$_3$ in water decreases (Bouwman and Bouman, 2002). Ammonia emissions may also increase as conditions of stress caused by crop disease, pests, and/or adverse weather conditions, such as drought, occur (Heckathorn and DeLucia, 1995 as cited in Bouwman et al., 1997).

The method and rate of fertilizer application can have a significant effect on the amount of NH$_3$ loss from fertilizer applications. Generally, applications that effectively bury the fertilizer (i.e. banding) result in lower NH$_3$ loss (IFA, 2001, Potter et al., 2001). While the effect of fertilizer application type is fairly well understood, studies conducted on the effect of N fertilizer application rate and the amount of NH$_3$ volatilization show conflicting results (IFA, 2001). Some studies reveal higher NH$_3$ volatilization rates at high N application rates while other studies show the opposite result.

The type of fertilizer applied influences the emissions of NH$_3$ since different types of fertilizer contain various percentages of N. When urea fertilizer is applied to very moist soils, the urea is hydrolyzed by the enzyme urease to produce a mixture of NH$_3$, NH$_4^+$, and bicarbonate (HCO$_3^-$) by the following reaction (Sommer et al., 2004):

\[
\text{CO(NH}_3\text{)}_2 + 2\text{H}_2\text{O} \leftrightarrow \text{NH}_3 + \text{NH}_4^+ + \text{HCO}_3^-
\]
The rate at which urea is hydrolyzed is related to urease activity, the availability of water, the pH of the solution, and temperature. The rate increases as water content and pH increases. Soil urease activity is optimized when the soil pH is between 8 and 9. The rate of urea hydrolysis also increases as temperature increases.

Ammonia volatilization is also driven by the atmospheric NH₃ concentration, and occurs when atmospheric NH₃ concentration is lower than the NH₃ concentration in the soil, liquid, or intercellular air space of the plant leaves (Bouwman and Bouman, 2002).

2.7.2 NH₃ Emissions from Crop Foliage

Plant leaves are another source of NH₃ emissions. Dissolved NH₃ and NH₄⁺ can be found in the water film, also known as the apoplastic solution, in the mesophyll cell walls of plant leaves. The amount of NH₃ and NH₄⁺ dissolved depends on the plant developmental stage, climate, fertilization, and the difference between the atmospheric NH₃ concentration and the NH₃ concentration in the leaves (Sommer et al., 2004). Ammonia will be emitted from a leaf if the leaf NH₃ concentration is greater than the atmospheric NH₃ concentration and NH₃ will be absorbed by the leaf in the reverse situation. Another source of NH₃ emission is decomposing potato tops and other plant residues. Generally, NH₃ volatilization increases with moisture, temperature, and N content of plant residues.

2.8 Emission Rates of NH₃

Agriculture is the largest source of global NH₃ emissions. Up to 50% of the applied N to croplands can be lost with gaseous emissions being the dominant N loss mechanism (IFA, 2001). There is a great deal of uncertainty in the magnitude of NH₃ emissions from various croplands. There are many factors that affect the emission rates and only a limited number of studies on the
emissions are available. A standard emission rate of 2.5 kg N hectare\(^{-1}\) year\(^{-1}\) has been used for all agricultural crops when compiling global emissions inventories, which contains a high degree of inaccuracy (Bouwman \textit{et al.}, 1997). Current U.S. emission estimates are based upon European measurements, which are not representative of actual farming practices in the U.S.

The IFA (2001) reports that there is a clear difference between measurements carried out in the fields and those performed inside laboratories. In addition, the measurement method used and the location where the measurements are made significantly affects the emission rates while the crop type is less significant (IFA, 2001). The average NH\(_3\) emission rates from grass are generally 20 and 10\% lower than from upland crops (croplands with no standing water throughout the growing season; i.e. corn and potato fields) and flooded systems, respectively (IFA, 2001).

The IFA (2001) report has grouped NH\(_3\) volatilization rates from upland crops by fertilizer type applied. Using micrometeorological techniques, the reported NH\(_3\) loss from broadcast applied UAN fertilizer ranges from 8\% to 18\%, with an average of 15\%. Broadcast application typically involves spreading dry fertilizer over the soil surface. Using dynamic flow chambers, the reported NH\(_3\) loss varies from negligible to 45\%. For the potato fields used for this study, UAN fertilizer was applied using a center-pivot irrigation system. From upland crops applied with ammonium bicarbonate, an NH\(_3\) loss rate of 21\% was measured in a field study using static chambers. Higher NH\(_3\) loss rates (as high as 30 to 70\%) were measured in a laboratory study using dynamic flow chambers. From calcium ammonium nitrate applications, NH\(_3\) loss rates of up to 6\% were reported from laboratory studies and much lower rates from field studies. Both the laboratory and field studies used dynamic flow chambers. Ammonia losses from upland crops applied with urea range from 15 to 20\%. This range comes from
various measurement techniques including chambers with and without forced flow, micrometeorological techniques, and wind tunnel measurements.

Studies have shown that the magnitude of NH$_3$ emissions resulting from decomposition of crop residue depends on the N content in the residue (McGinn and Janzen, 1997). In a study on perennial ryegrass herbage, 10% of the NH$_3$ applied was emitted to the atmosphere from herbage with 29.8 mg N g$^{-1}$ of leaf biomass, and no losses from herbage with 9.2 mg N g$^{-1}$ leaf biomass. In another study, 14% of the N applied was emitted into the atmosphere for a legume green manure (McGinn and Janzen, 1991). In the U.K., it is estimated that volatilized N losses by grassland decomposition accounts for 2.7% of the total N emissions in agriculture (Whitehead and Lockyer, 1989 as cited in McGinn and Janzen, 1997). Measured NH$_3$ emissions from decomposing sugar beet leave, potato tops, and field beam straw range from 0.9 to 3.7% of the N content (Sommer et al., 2004).

Little information is known on NH$_3$ volatilization losses in potato fields. The potato plant’s N uptake efficiency under current best management practices is approximately 65% (Lang et al., 1999) with a mean of 40% (Balasubramanian et al., 2004). Measured NH$_3$ emission rates from barley, wheat, and oilseed rape canopies range from 1 to 15 kg hectare$^{-1}$ year$^{-1}$ while measured NH$_3$ emissions from sugar beet leaves and potato shoots ranged from 8.6 to 12.6 kg N hectare$^{-1}$ year$^{-1}$ (Sommer et al., 2004).
CHAPTER 3

METHODOLOGY

This chapter describes the methods used for measuring N$_2$O and NH$_3$ concentrations and for determining flux rates from the potato fields. Mixing ratios of N$_2$O were measured using gas chromatography (GC) and flux rates of N$_2$O were estimated using a static chamber method. A short-path differential optical absorption spectroscopy (DOAS) instrument was used to measure concentrations of NH$_3$ along a path downwind of the fields. Sulfur hexafluoride (SF$_6$) was released from a point source from the center pivot of the fields to simulate NH$_3$ plume dispersion and to calibrate a Gaussian plume model (SIMFLUX). SIMFLUX was then used in an inverse modeling approach to estimate flux rates of NH$_3$ from the fields. Data were collected at potato fields in central WA, near Othello.

3.1 Measurement of N$_2$O

A disjunct eddy accumulation (DEA) system was used in summer 2006 for measurement of N$_2$O fluxes. In operation, air samples are partitioned into updraft and downdraft reservoirs over some averaging period based on the sign of the vertical velocity (Turnipseed et al., 2008). The total flux is related to the concentrations in the updraft and downdraft reservoirs. It was found that N$_2$O flux measurements with the DEA were below detection limits and so a static chamber method was then used for the summers of 2007 and 2008. For more information on the DEA system, refer to Turnipseed et al. (2008).
3.1.1 Static Closed Chamber Method and Flux Rate Calculation

Fluxes of N$_2$O were measured using a static closed chamber method (Hutchinson and Mosier, 1981). Chambers were installed between potato plant rows and sampled according to USDA-ARS GRACEnet (Greenhouse gas Reduction through Agricultural Carbon Enhancement network) protocols (USDA-ARS, 2003; www.GRACEnet.usda.gov). Each chamber used consisted of a base and a lid. Chamber bases were made from 30.5 cm inner diameter x 15 cm high polyvinyl chloride (PVC) cylinders. The bases were inserted 10 cm into the soil and at least 24 hours prior to sampling to allow for stabilization. The chamber lids (30.5-cm inner diameter by 7.5 cm high) were made from PVC caps and included a 2.54 cm sampling port, a built-in vent tube, and an internal 12-V DC 60 mm fan. The built-in vent tube prevented pressure perturbations by maintaining atmospheric pressure inside the chamber. The fan ensured sufficient gas mixing within the chamber for homogeneous gas concentrations and better simulated mass transfer from the soil to the atmosphere.

The approximate locations of the chambers are shown in Figure 3.1. The dashed blue lines and the solid red lines represent the road that lead to the center of the field used in summer 2007 and summer 2008, respectively. The wheel tracks of the sprinkler system are represented by the dashed gray concentric circles. The numbers in blue and red are chambers used in summer 2007 and summer 2008, respectively. Chambers 1, 2, 10, 11, 12, and 13 were placed near the dirt road that led to the center of the fields. Chambers 3, 4, 7, 8, and 14 were placed along the south edge of the fields. Chambers 5 and 6 were placed along the 4th wheel track, near the dirt road. Chamber 9 was placed approximately 10 m south of the dirt road, between the edge of the field and the 1st wheel track.
Air samples (about 30 ml) from within the chambers were collected every 15 minutes over a period of 1 hour after covering the chambers. After the first 1 hour of sampling, the chamber was uncovered for 30 minutes before the second hour of sampling began. Typically, a total of 4 hours of sampling was conducted for each chamber. To calculate the flux of N₂O, linear regression and the following calculation was used:

\[
\text{rate of change of } N_2O \text{ concentration inside chamber (ppm/min)} \times \frac{\text{chamber volume (L)}}{\text{chamber surface area (m²)}} = 10^{-6} \frac{L \text{ N}_2O/}{(m²*min)}
\]

The gas flux values were converted from a volumetric basis to a mass basis.
3.1.2 Advantages and Disadvantages of using the Chamber Method

The advantages of using the chamber method is that chambers are relatively inexpensive, easy to build, install, and remove, and have low detection limits. The disadvantages of using chambers include the inability to account for environmental conditions (i.e. winds) and disturbance to the soil which can easily occur when installing chamber bases. In addition, the method is labor intensive. Five air samples are needed to calculate one flux rate value. Chamber bases need to be installed at least 24-hours prior to sampling to minimize soil disturbance and root damage. Careful planning of sampling time is necessary to ensure that the measurements are made immediately after spraying of water or water with N fertilizer by the sprinkler system. Finally, numerous chambers are needed to capture the variability of N2O flux rates through the field due to heterogeneity of the soil.

3.1.3 Sample Analysis

Gas sampling was performed 0 to 6.5 hours after fertigation and irrigation events. Analysis of air samples for N2O concentration was conducted using a Hewlett-Packard 5890 Gas Chromatograph (GC) with an Electron Capture Detector (ECD). An ECD is best for analysis of compounds containing halogens and nitro groups (Poole and Poole, 1991). An Alltech Porapak Q (80/100) column was used and kept isothermal at 40 °C. The carrier gas was 95% Argon (Ar) and 5% methane (CH4). The GC flow rate ranged from 24.5 to 25.5 standard cubic centimeters per minute (sccm) and was maintained using an Alltech Digital Flow Check Flow Meter. A 777 ppbv N2O ± 10% (Scott-Marrin) standard was analyzed multiple times during each sampling day. All samples were analyzed within one day of collection since tests showed decreased concentrations (~25% reduced) from syringes analyzed after 24 hours.
3.2 Measurement of NH₃

3.2.1 Differential Optical Absorption Spectroscopy (DOAS)

Spectroscopy is the study of the interaction between radiation and matter as a function of wavelength \( (\lambda) \). Gas molecules absorb radiation if the radiation can electronically excite or change the vibrational or rotational energy of the molecule. A short-path differential optical absorption spectroscopy (DOAS) technique was used to measure NH₃ emissions from the potato fields. The DOAS technique avoids the NH₃ adhesion problem (Mount et al., 2002) that is common with in-situ methods using inlets and sampling tubes.

The DOAS technique measures the spectroscopic molecular absorption of trace gases over an open atmospheric path. Using Beer’s Law, which relates the amount of light absorbed to the number of molecules in the measurement path length, the concentration of NH₃ can be calculated. Beer’s Law states that there is a logarithmic dependence between the transmissivity of light through a substance and the product of the absorption coefficient of the molecule and the distance the light travels through the material. Beer’s Law can be expressed as:

\[
T = \frac{I(\lambda)}{I_o(\lambda)} = \exp^{-\sum \sigma_i(\lambda) L N_i} \tag{3.2}
\]

\( T \) - transmissivity;
\( \lambda \) - wavelength [nanometers];
\( I(\lambda) \) - light intensity in the presence of absorbers [counts];
\( I_o(\lambda) \) - light intensity without absorbers [counts];
\( \sigma_i(\lambda) \) - absorption cross-section of the \( i^{th} \) absorber expressed in [cm²];
\( N_i \) - concentration of the \( i^{th} \) absorber [molecules cm⁻³];
\( L \) - distance that the light travels through the material or the path length [cm].
The subscript “i” in the summation refers to any particular atmospheric specie that absorbs at the wavelength region of measurement and the sum occurs over all species that can absorb at wavelength $\lambda$. As a result, it is important to know all of the atmospheric species that absorb over the wavelength region measured and to account for them in the analysis. The DOAS instrument used for this project operates in the 200-240 nm wavelength range. In this wavelength range, atmospheric species NH$_3$, nitric oxide (NO), and sulfur dioxide (SO$_2$) absorb (Mount et al., 2002). The absorbances of NO and SO$_2$ are very low (below the detection limit) in the potato fields so they were not accounted for in the data analysis (Mount et al., 2002).

In addition to other gas species that absorb in the wavelength region measured, other atmospheric processes contribute to the measured spectrum and must be accounted for. Rayleigh scattering is wavelength dependent and describes the scattering of light by small molecules such as dinitrogen (N$_2$) and oxygen (O$_2$) which makeup about 98% of the atmosphere. As the wavelength decreases, the amount of scattering increases. For Rayleigh scattering, the ratio of the diameter of the scattering particle to the wavelength of measurement is much less than 1 and is computed using the following equation:

$$x = \frac{2\pi r}{\lambda}$$  \hspace{1cm} (3.3)

$x$ - size parameter

$r$ - characteristic dimension of the scattering particle;

$\lambda$ - wavelength of the light.

Rayleigh scattering occurs when $x << 1$. The Rayleigh scattering cross-section is computed from the Pendorf equation (Pendorf, 1957):

$$\sigma_R(\lambda) \approx \sigma_{RO} \cdot \lambda^{-4}$$  \hspace{1cm} (3.4)

where $\sigma_{RO} \approx 4.4 \times 10^{-16}$ cm$^2$ nm$^4$ for air.
Mie scattering (clouds and aerosols) occurs when the particles causing the scattering are larger than the wavelengths of radiation in contact with them. Similarly, Mie scattering can be modeled as an absorption cross-section. The cross-section for Mie scattering depends on the particles causing the scatter, but is typically given by $\lambda^{-1.6}$. Since NH$_3$ absorption is being measured in the mid-ultraviolet range (below the atmospheric cutoff of 290 nm) the solar spectrum does not need to be accounted for, a major benefit of measuring in this spectral region. Accounting for Rayleigh and Mie scattering, Beer’s law becomes the following:

$$\frac{I(\lambda)}{I_0(\lambda)} = \exp\left[\left(-\sum \sigma_i(\lambda)LN_i\right) + \varepsilon R(\lambda) + \varepsilon M(\lambda)\right] \quad 3.5$$

3.2.2 Instrument Description

Two versions of the instrument were used for NH$_3$ measurements. Details of the instrument used in summer 2007 are described in Mount et al. (2002). In this section, a brief description of the original instrument will be presented, followed by a description of the revised instrument.

A schematic diagram of the original DOAS instrument used for summer 2007 measurements is shown in Figure 3.2 and a picture of the instrument housing, which was a small trailer, is shown in Figure 3.3. A 150 W xenon (Xe) high-pressure emission lamp (source region of 0.5 mm, Hamamatsu Corp. model L2273) is used as a light source and is located at the side of the optical path of the primary mirror. The light from the Xe lamp is directed onto the 1$^{st}$ folding mirror, which then directs the light onto a f/5 parabolic primary mirror. The primary mirror is 40 cm in diameter, coated with aluminum (Al) and magnesium fluoride, and was made by Colorado Precision Optics. Only an annulus of 6 cm of light is illuminated onto the primary mirror since there is a second folding mirror that is located between the primary mirror and the 1$^{st}$ folding mirror which blocks part of the beam.
Figure 3.2: Schematic drawing of the original DOAS instrument showing the optical light path from the Xe lamp to the Reticon detector (Figure adapted from Mount et al., 2002).

Figure 3.3: The original DOAS instrument housing (cover removed), which was a trailer (Figure adapted from Mount et al., 2002).
From the primary mirror, the annulus of light is directed towards a set of four large retro-reflectors or a set of thirty-seven small retro-reflectors mounted on a tripod. Each of the four big retro-reflectors is 12.7 cm in diameter, and each of the small retro-reflectors is 9 cm in diameter. All are coated with Al and magnesium fluoride for protection. The retro-reflectors were obtained from Melles Griot Inc. The light from the retro-reflectors is displaced and returned to the primary mirror. The light is then intercepted by the second folding mirror, which directs the light towards the slit of a double crossed Czerny–Turner spectrograph. The focal length of the telescope optical system is 200 cm.

Inside the spectrograph, the light hits the 1st collimating mirror to make the rays parallel before it is reflected to a diffraction grating. The grating breaks the light up into its constituent colors. From the grating, the light travels to the 1st focusing mirror and then enters another entrance aperture. Once again, the light passes through a 2nd set of collimating mirror, grating, and focusing mirror. The spectrographs were double-crossed to reduce scattered light. The detector used for the original DOAS instrument was a homebuilt Reticon silicon photodiode array with 1024 pixels with each pixel 25 μm wide and 2.5 mm high. Spectral coverage was about 60 nm and spectral resolution was about 0.6 nm.

The original DOAS instrument had several limitations. The computer used to control the system was a Macintosh IIfx, which has been long discontinued. In addition, transporting the trailer to and into an agricultural field was laborious. Pulling the trailer had to be done carefully because the optics, detector, and other parts of the instrument are delicate and there were insufficient mountings to absorb the shock. In the field, the slightest movements of the trailer (e.g. heating and expansion of the tires) would result in loss of signal due to misalignment.
During the undertaking of this project, several parts of the DOAS instrument were revised from the prototype version used at the WSU dairy including the instrument housing, telescope and its light feed, and the detector. The revised DOAS instrument is more compact than the original instrument. The optics setup of the revised instrument is similar to the setup of the original instrument but the spectrograph is placed below instead of to the side of the 2\textsuperscript{nd} folding mirror and the primary mirror. The primary mirror (made by Stevens Optical) is only 32 cm in diameter instead of 40 cm, which allows for a more compact design. The light reflected from the retro-reflectors is returned to the primary mirror and then is intercepted by the 2\textsuperscript{nd} folding mirror, which directs the light downward to a 3\textsuperscript{rd} folding mirror. The 3\textsuperscript{rd} folding mirror directs the light into the entrance slit of the spectrograph. The angle of this third folding mirror is manually tilted in two axes by small motors so that peak light intensity can be directed into the spectrograph entrance slit. The f/number of the optical system is the same as the one used in the original instrument. A schematic diagram and a picture of the revised DOAS instrument are shown in Figure 3.4 and 3.5, respectively.
Figure 3.4: Schematic drawing of the revised DOAS instrument showing the optical light path from the Xe lamp to the CCD detector. The top figures illustrate a view from the top of the instrument, the bottom figure shows the view from the side.

Figure 3.5: The revised DOAS instrument with the cover removed.
The detector used in the revised instrument is a Princeton Instruments charge coupled device (CCD) detector system (spec-10:400B) with 400 pixels x 1340 pixels, with each pixel 19.4 µm wide and 19.4 µm high, and the spectral direction in the long direction. More details regarding the CCD detector can be found on the Princeton Instruments website under products-spectroscopy cameras-Spec-10-CCD. The WinSpec Spectroscopy Software version 2.5.16.5 was used for data collection. This software is compatible with Windows. A PC laptop was used for control of the system.

The main optics of the revised version are assembled on a 1.82 m by 0.85 m bench which is loaded in the back of a pickup truck. A special canopy was designed to house all of the necessary equipment (instrument, retro-reflectors, generator, etc.) for transport purposes. Two bicycle tires are attached to the sides of the instrument in order to maneuver the system in the field. A winch and two removable ramps are used to load the instrument into the back of the pickup truck. The instrument canopy rides on a set of six inflatable tires to increase stability during transportation and help absorb shock. Figure 3.6 shows a picture of the revised instrument housing.
To set up, the instrument is winched down the ramps. Four air jacks are attached to the corners of the instrument bench. The air jacks lift the instrument off the ground and the instrument is placed on three tripods. Unloading and set up of the instrument can be accomplished by three or more people and typically takes several hours. Aligning and fine tuning of the instrument to the retro-reflectors takes an additional several hours. Figure 3.7 is a picture of the revised DOAS instrument when it is setup for taking measurements.
3.2.3 Advantages and Disadvantages of the DOAS Technique

Use of the DOAS technique for measuring NH$_3$ concentrations has many advantages over traditional in-situ measurements (Mount et al., 2002). Ammonia adheres to surfaces, such as inlet tubes and sampling cells, and the use of the DOAS technique eliminates this problem. Results depend only on the quality of the independently measured cross-sections of the species and the algorithm used for data analysis (Mount et al., 2002). Finally, the DOAS technique can measure several gas species simultaneously (i.e. NO, SO$_2$, NH$_3$ for the wavelength region used).

The DOAS technique also has disadvantages. First, it is limited to molecular species that have spectrally structured absorption cross-sections in the wavelength region of the measurement. In addition, the DOAS technique is limited to times of good visibility. Rain and snow make measurements impossible due to the strong attenuation in the UV-Visible region. Lastly, the DOAS technique is labor intensive because the instrument must be monitored on an hourly basis to keep the instrument properly aligned.
There are many uncertainties associated with solving the Beer’s Law equation (eq. 3.3). \( I(\lambda) \) and \( I_o(\lambda) \) are generally not measured over the same path since the \( I_o \) measurement should not have any of the measured species in it. This negatively affects the spectral residual analysis. Changes in the instrument, such as lamp output, may occur due to changes in temperature and pressure that occur between measurement of \( I(\lambda) \) and \( I_o(\lambda) \). Generally these changes are very low frequency in spectral occurrence and so do not affect the analysis.

<table>
<thead>
<tr>
<th>Table 3.1: DOAS instrument characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Reticon Detector</strong></td>
</tr>
<tr>
<td>pixel size</td>
</tr>
<tr>
<td>25 μm wide x 2.5 mm high</td>
</tr>
<tr>
<td><strong>CCD Detector</strong></td>
</tr>
<tr>
<td>pixel size</td>
</tr>
<tr>
<td>20 μm x 20 μm</td>
</tr>
<tr>
<td><strong>Spectrograph</strong></td>
</tr>
<tr>
<td>focal length</td>
</tr>
<tr>
<td>375 cm</td>
</tr>
<tr>
<td><strong>Telescope</strong></td>
</tr>
<tr>
<td>focal length (original system)</td>
</tr>
<tr>
<td>200 cm</td>
</tr>
<tr>
<td>focal length (revised system)</td>
</tr>
<tr>
<td>160 cm</td>
</tr>
<tr>
<td><strong>Xe Lamp</strong></td>
</tr>
<tr>
<td>output power</td>
</tr>
<tr>
<td>150 W</td>
</tr>
<tr>
<td>arc length spot size</td>
</tr>
<tr>
<td>0.5 mm</td>
</tr>
<tr>
<td>wavelength range</td>
</tr>
<tr>
<td>185-2000 nm</td>
</tr>
<tr>
<td>average life</td>
</tr>
<tr>
<td>3000 hrs</td>
</tr>
<tr>
<td>output fluctuation</td>
</tr>
<tr>
<td>1%</td>
</tr>
<tr>
<td><strong>Other</strong></td>
</tr>
<tr>
<td>path length</td>
</tr>
<tr>
<td>~250 m</td>
</tr>
<tr>
<td>detection limit:</td>
</tr>
<tr>
<td>2007 data (original system)</td>
</tr>
<tr>
<td>5 ppbv</td>
</tr>
<tr>
<td>2008 data (revised system)</td>
</tr>
<tr>
<td>10 ppbv</td>
</tr>
<tr>
<td>spectral coverage</td>
</tr>
<tr>
<td>200-240 nm</td>
</tr>
</tbody>
</table>
3.2.4 Data Collection

Data was collected using the Reticon detector in the summer of 2007 and using the new CCD detector in the summer of 2008. The instrument was aligned towards the retro-reflectors so that the return signal counts were maximized. The integration time, which depends on the path length, was varied so that the maximum count rate was about 30,000 to 40,000 counts per integration time. Spectra were co-added so that data was averaged over five minute intervals.

Before taking actual measurements the electronic noise (dark spectra) of the instrument system was measured by placing a black slide in front of the entrance slit of the spectrograph and using the same settings as for the actual spectra. Changes in detector temperature caused changes in dark spectra. As a result, dark spectra measurements were also taken about every hour and were co-added and averaged. The average dark spectra were subtracted from the actual spectra measured (when the slide was removed). A zinc (Zn) emission lamp was placed in front of the entrance slit every hour to provide a strong spectral line fiducial at 213.9 nm. The Zn fiducial line was centered at a chosen pixel.

3.2.5 Limitations on Data Collection

Time was a major limitation for collection of data. Measurements had to be taken during the short potato growing season. The farmers who owned the fields would often make sudden changes to the fertigation and irrigation schedules. This would be due to uncontrollable changes in the weather, the crop's reaction to the fertigation and/or irrigation, and the amount of fertilizer and/or water available. Data collection was prohibited during times of chemigation (i.e. herbicides and/or pesticides) application because of health concerns.
Problems with the instrument were encountered in the field. Changes to the DOAS instrument were implemented without being fully tested (e.g. the replacement of the detector) before being used for data collection. High outside air temperature caused a significant problem with the CCD detector. The internal cooler of the CCD detector could not keep the detector cold enough when the outside air temperature was above ~24 °C. This problem was anticipated but not enough testing was performed prior to going to the fields and so a solution was initiated in the field. At the potato field near Othello, WA, the average outside air temperature during the afternoon was always above ~24 °C and even occasionally reached above 37 °C. An external radiator was used with cooling lines running through an ice bath in order to keep the detector running. However, variations in the dark current were sufficiently uncontrollable that the dark subtractions could not always be performed correctly.

3.2.6 Limitations on Data Quality

Analysis of the optical arrangement in January 2009 showed the focal position of the spectrograph was incorrect, which resulted in loss of light (signal). Lower signal to noise ratio (s/n) resulted in lower precision. However, this should not have affected the actual determinations of molecular concentration when sufficient NH₃ was present. It was also discovered that the mirrors within the spectrograph were not properly aligned and this resulted in a degraded spectral resolution. The crude temperature regulation of the CCD detector caused major problems with dark subtraction and overall had a significant negative effect on the measurement.

Figure 3.8 is a plot of the CCD Detector quantum efficiency (QE) with respect to wavelength. The QE, which is a measure of the detector’s electrical sensitivity to light, in the wavelength region used (200 to 240 nm) was below 10%, which resulted in lower s/n ratio.
The detection limit of the instrument depends on the magnitude of the spectral residuals of the data, which is the remainder of the spectral ratio of $I$ and $I_0$ after removal of the known absorbers (NH$_3$) as well as the spectrum of Mie and Rayleigh scattering. The residual spectrum should be random noise, but this level of precision is never reached due to systematic errors in the removal of the spectral signatures, detector temperature problems, and changes in the spectral structure of the Xe lamp. The original DOAS instrument was sensitive to a few ppbv of NH$_3$ while the revised version is only sensitive to at best 10 ppbv of NH$_3$.

The revised instrument can clearly be improved with more testing and modifications. Problems for our field work were accentuated by time constraints due to the late completion of the modified instrument, resulting in insufficient testing before using at the fields. The optical problems with the spectrograph resulted in degraded performance, but it was decided not to change this once the data collection was started. The main effects caused a degradation of s/n
and precision. Problems in the design of the telescope and feed optics were corrected part way through the experiment to increase the collection of light and significantly reduced the integration time.

3.2.7 Data Analysis

To convert the absorbance of NH$_3$ given by Beer’s law to concentration, a nonlinear least squares fitting method was used. The method fits the pattern of the cross-section of each gas molecule that absorbs in the wavelength range used to the natural logarithm of the ratio of $I/I_0$. In this case, only the cross-section of NH$_3$ was used since the absorbances of NO and SO$_2$ were found to be very low.

Figure 3.9 shows the cross-section spectra of NH$_3$ in the mid UV spectral region. This shows the ability of NH$_3$ to absorb light as a function of wavelength. For summer 2007 measurements, three absorption lines were used to reduce the data. For the summer 2008 data, only two absorption lines were used (wavelength range: 199nm to 207nm).
In addition to the cross-section spectra of NH$_3$, the nonlinear least squares algorithm accounts for the spectral pattern of Rayleigh and Mie scattering as well as the shift and stretch of the two spectra being ratioed. Shift and stretch in the spectra can result from displacement of the grating and detector as well as temperature changes experienced by the spectrograph. The method minimizes the spectral residual of patterns in the data after removal of the molecular spectra signatures.

Data analysis is conducted in two steps. First, a relative $I_0$, containing NH$_3$ features, was chosen from the measurement data (same-day) and used for initial data reduction. The NH$_3$ concentration for each measurement period was then calculated relative to the $I_0$ chosen. The actual NH$_3$ concentration was then determined by reducing the relative $I_0$ with an absolute $I_0$, a spectra with no NH$_3$ features taken in the laboratory.
3.2.8 Characterization of Dispersion Conditions using Tracer Studies

During some measurement periods, SF$_6$ tracer gas was released from the center of the fields and concentrations measured along a crosswind sample line downwind of the fields to help characterize local dispersion conditions. To conduct tracer studies, SF$_6$ was released as a continuous point source from ground-level at the center pivot of the field. The SF$_6$ release rates ranged from ~0.6 to ~0.7 L min$^{-1}$. Air flow was maintained using a 1.00 SLPM Tylan mass flow controller (S/N AA9605027) and a Tylan RO-32 control box. A BIOS International DryCal DC-Lite Flow Meter was used check flow rates. Sulfur hexafluoride is ideal for tracer studies because it is inert, concentrations can be measured with satisfactory accuracy at very low concentrations, and ambient concentrations of SF$_6$ are negligible. Even though SF$_6$ has a higher molecular weight (146.06 g mol$^{-1}$) than NH$_3$ (17.03 g mol$^{-1}$), the effect of this is negligible since turbulent diffusion is the dominant transport mechanism (Kaharabata et al., 2000).

Positioned downwind of the source were Trace Gas Automated Profile System (TGAPS) sample lines and syringe samplers. TGAPS is an automated sampling and analysis system, which continuously measures 5-minute averaged concentrations of SF$_6$ at seven different locations. The system draws air through 7 long inlet lines positioned along the length of the DOAS measurement path and air from these inlets is pumped into a set of seven 10-L Tedlar bags. While one set of 7 bags collects air samples, a second set of 7 bags is analyzed. During analysis, air from each bag is drawn to a continuous SF$_6$ analyzer (Flaherty et al., 2007). Between the sample analysis and collection, the Tedlar bags are evacuated completely to ensure that the bags are “clean” for the next sampling period.

Air samples were also collected using programmable, battery-operated syringe samplers. Each syringe sampler contains 12 syringes, which sequentially collect 30-minute averaged air
samples. Each syringe sampler is enclosed in an aluminum case, which serves to protect the 12-syringe unit inside. A sealed compartment inside the syringe sampler houses the electronics, stepper motors, and clock. Krasnec et al. (1984) provides a complete description of the syringe samplers used. The syringe samplers were also deployed along and beyond the DOAS measurement path downwind of the field and tracer release to provide a measure of plume width.

A GC equipped with an ECD and coupled with an electronic integrator was used to measure SF$_6$ concentrations. The air samples were combined with hydrogen (H$_2$) and then passed through a packed bed of palladium catalyst, which converted the oxygen (O$_2$) in the air to water (H$_2$O). Then, the air passed through a nafion tube with a countercurrent flow of N$_2$ to remove the H$_2$O from the sample stream. The sample stream was then passed to the ECD, where voltage changes due to the SF$_6$ in the stream provided the signal for concentration. See Benner and Lamb (1985) and Flaherty et al. (2007) for additional information about this type of detector system. Calibration was performed using 1005 pptv, 1625 pptv, and 1462 pptv Scott-Marrin Inc. calibration gases (certified accuracy ±5%). All samples were analyzed within a day of collection.

3.2.9 Field Setup

Figure 3.10 is a diagram of the field setup. Southwest winds are predominant in the central WA region. Thus, the DOAS instrument measurement path (path from the spectrograph to the retro-reflectors) was setup outside the edge on the northeast quadrants of the potato fields to capture the NH$_3$ plume as it traveled along the wind trajectory. Path lengths varied from ~150 to ~250 m.
Tracer sample lines and syringe samplers were set up along the DOAS measurement path and extended over a distance of 800 m so that ideally the entire plume of SF$_6$ would be captured. More details on how the dispersion studies were conducted will be provided in section 3.3.2.

![Diagram of the field setup.](image)

The only source of electrical power was at the center of the fields. As a result, instruments used in the fields (i.e. DOAS instrument) were powered using portable generators. A storage building with power located nearby was used to power other instruments such as the N$_2$O GC.

### 3.3 Measurement of Meteorological Variables

Meteorological information such as wind velocity, wind direction, and air temperature were measured using a sonic anemometer at 10 Hertz frequency. Soil moisture and temperature were measured using CS515s Water Content Reflectometers and copper/constantan
thermocouples (Campbell Scientific Inc.), respectively. Campbell Scientific CR10 data loggers were used to record data. Leaf stomatal conductance was measured using Decagon Devices Hand Held Leaf Porometer.

3.4 Determining Emission Rates of NH₃

A Gaussian plume dispersion model (SIMFLUX) was used with an inverse approach to estimate NH₃ emission rates. The first part of this section will present the basis of the Gaussian plume dispersion equation (section 3.4.1) and in section 3.4.2, the details of SIMFLUX will be presented.

3.4.1 The Gaussian Plume Dispersion Equation

The Gaussian plume dispersion model is useful for measuring the impacts of air pollutant sources on air quality. After a release, individual particles move according to velocity fluctuations which are random, resulting in a random distribution in each coordinate system. The result of this random distribution is that the plume contains more particles near the center of mass while fewer particles are present at increasing distances from the center of mass. It is reasonable to assume a Gaussian concentration distribution based on measurements of typical plumes and for ease of mathematical computation.

For a continuously emitting point source strength (Q) in grams per second (g s⁻¹) at an effective height (h) in meters above the ground, the concentration C at a receptor location x, y, and z in grams per cubic meter is given by the equation:

\[
C(x, y, z) = \frac{Q}{2\pi\sigma_y\sigma_z u} e^{-\frac{y^2}{2\sigma_y^2}} e^{-\frac{(z-h)^2}{2\sigma_z^2}} \left[ e^{-\frac{(z+h)^2}{2\sigma_z^2}} + e^{-\frac{(z+h)^2}{2\sigma_z^2}} \right]
\]

3.6
where x is the downwind distance of the receptor from the source in meters (m), y is the
crosswind distance in meters, z is the height of the receptor above the ground in meters, u is the
wind speed in meters per second, and $\sigma_y$ and $\sigma_z$ are the horizontal and vertical diffusion
coefficients, respectively, in meters. A complete general overview of the Gaussian plume
equation is discussed by Hanna et al. (1982).

Requirements for the Gaussian dispersion models are homogeneous turbulence, uniform
flow, no loss of pollutants, and steady state conditions (Arya, 1999). In addition, mean wind
speed is required to be larger than the standard deviations of turbulent velocity fluctuations, so
that the upstream or longitudinal diffusion can be neglected. Thus, significant wind shears,
inhomogeneous turbulence, and weak winds make the theoretical basis of Gaussian dispersion
modeling invalid. Finally, the methods for obtaining certain input parameters used in the
Gaussian plume dispersion equation are also subject to many assumptions and constraints. Such
critical parameters are the diffusion coefficients, $\sigma_y$ and $\sigma_z$, since they describe the shape of the
plume as it travels along in the mean wind direction. Various methods (Martin, Briggs, Taylor,
and Draxler) for determining $\sigma_y$ and $\sigma_z$ were compared with actual data obtained from a series of
SF$_6$ tracer dispersion studies conducted in the potato fields. A comparison of the various
methods is included in Appendix A.

3.4.2 SIMFLUX

A Gaussian plume model (SIMFLUX) was used in an inverse method to estimate NH$_3$
emission rates. SIMFLUX uses a dense array of point sources to calculate the dispersion of
emissions from point, line, and area sources. It is a versatile model because it is applicable for a
wide range of area source configurations. Each potato field was modeled as an array of
individual point sources at 1 m spacing. Howard et al. (1996) evaluated the effect of the length
of the spacing between each point source and found that the best model results occurred when the spacing between source points are less than one standard deviation of the lateral dispersion at any receptor location for the appropriate stability class.

Ammonia emission rates were back-calculated using observed NH$_3$ concentrations. The DOAS measurement path was modeled as a line of point receptors, approximately 400 m downwind from the center of the fields and at 1 m spacing. All measured variables, including wind speed and wind direction, were measured using a sonic anemometer at 10 Hz frequency and were input into the Gaussian plume equation. Analytical equations that represent the Pasquill-Gifford diffusion curves were employed for estimating the vertical and horizontal dispersion (Martin, 1976). For the results on the comparison of various sigma methods to be used in SIMFLUX, see Appendix A. The contribution from each point source was summed to estimate the concentration at each downwind receptor. An emission rate (Q) of 1 g s$^{-1}$ point$^{-1}$ was first used for all point sources and then adjusted until predicted NH$_3$ concentrations at the point receptors match observed concentrations.

Howard et al. (1996) discovered that significant errors result if a receptor is placed too far crosswind from the plume centerline. As a result, only flux rates calculated from periods that captured the shape of the Gaussian plume were considered to be valid. The percent of plume captured for each 30-min sampling period was calculated by determining the ratio of the area of the plume measured and the area of the entire plume predicted.
CHAPTER 4

Measuring Gaseous Losses of Nitrogen from Potato Fields in central Washington

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to be submitted to
Atmospheric Environment
4.1 Abstract

There are limited studies that have measured the magnitude of Nitrogen (N) emissions from intensively managed croplands. In this research, concentrations of ammonia (NH₃) and nitrous oxide (N₂O) from potato fields in central Washington were measured during the 2007 and 2008 growing seasons (June – August). A short-path differential optical absorption spectroscopy (DOAS) instrument that operates in the 200-240 nm wavelength spectral range was used to measure NH₃ concentrations. During some periods, sulfur hexafluoride (SF₆) tracer gas was released from the center of the fields and concentrations were measured along a crosswind sample line downwind of the fields to help characterize local dispersion conditions. A Gaussian plume dispersion model (SIMFLUX) was used, along with the SF₆ tracer data in an inverse approach to estimate NH₃ emission rates. A static chamber flux method was used to measure N₂O emissions with sample analysis via electron capture gas chromatography (ECD-GC). Nitrous oxide flux rates generally peaked between 2.5 and 3.5 hours after irrigation or fertigation. The amount of N lost as N₂O ranged from zero to 12% of the applied N fertilizer which is comparable to documented emission factor values of 0.1 to 8% (Dobbie and Smith, 2003). Measured NH₃ concentrations varied with application status, sprinkler arm location, and wind speed. The highest NH₃ concentrations measured in the summers of 2007 and 2008 were ~34 ppbv and ~112 ppbv, respectively. Both peaks occurred during fertigation, when the sprinkler arm was spraying almost directly towards the DOAS measurement path and winds greater than 1 m s⁻¹ were blowing towards the measurement path. Estimated NH₃ flux rates ranged from 9 μg N m⁻² s⁻¹ to 460 μg N m⁻² s⁻¹ for fertigation periods and from 0.6 μg N m⁻² s⁻¹ to 2 μg N m⁻² s⁻¹ for periods of no fertigation or irrigation, which indicates that a significant fraction
of NH$_3$ loss occurs as a result of direct volatilization of NH$_3$ during fertigation. Overall, the average NH$_3$ flux during fertigation periods was 10% of N applied as fertilizer.
4.2 Introduction

The increasing use of synthetic nitrogen (N) fertilizers has led to an altered N cycle and doubled amount of fixed N entering the biosphere (Vitousek et al., 2000). There are uncertainties in the magnitude of N emissions from various sources, including intensively managed croplands. Up to 50% of the applied N to croplands can be lost, with gaseous emissions being the dominant N loss mechanism (IFA, 2001). It is important to quantify and understand the factors that control ammonia (NH$_3$) and nitrous oxide (N$_2$O) emissions because they pose negative effects to human health and the environment. Ammonia reacts rapidly with both sulfuric and nitric acids in the atmosphere to form fine particles, which cause health problems, reduce visibility, cause acid deposition, and perturb the earth’s radiation balance (Pandis, 1995). Nitrous oxide is a potent greenhouse gas which ranks just below carbon dioxide and methane in global warming concerns.

In this research, emissions of NH$_3$ and N$_2$O from potato fields in central Washington (WA) were measured during the 2007 and 2008 growing seasons (June – August). A short-path differential optical absorption spectroscopy (DOAS) instrument (Mount et al., 2002) that operates in the 200-240 nm wavelength range was used to measure NH$_3$ concentrations. During some measurements, sulfur hexafluoride (SF$_6$) tracer gas was released from the center of the fields and concentrations were measured along a crosswind sample line downwind of the fields to help characterize local dispersion conditions. A Gaussian plume dispersion model (SIMFLUX) was used, along with the SF$_6$ tracer data, in an inverse approach to estimate NH$_3$ emission rates. A static chamber flux method was used to measure N$_2$O emissions with sample analysis via electron capture gas chromatography (ECD-GC).
The data collected will help improve CropSyst, a one-dimensional process-oriented model developed to simulate gaseous N emissions from various crop fields (Stockle, 1994). CropSyst is aimed to assist farmers in better management practices. The data will also improve NH$_3$ emission estimates in the AIRPACT-3 regional air quality forecast system (Chen et al., 2008; http://www.lar.wsu.edu/airpact-3/).

4.3 Materials and Methods

4.3.1 Description of Research Location

Field studies were conducted in 2007 and 2008 to characterize trace gas fluxes from potato fields near Othello, Washington, US. Two fields were chosen based on their flatness (for ease of setting up the DOAS instrument) and to maximize the distance from other interfering sources of NH$_3$ and N$_2$O. Fields of grass seed and wheat, which are not typically managed with high inputs of N fertilizers, surrounded the two potato fields. The crop rotation for both fields was grass seed - wheat - potatoes. The field used in summer 2007 was planted with ranger russet potatoes and the field used in summer 2008 was planted with premier russet potatoes.

Field preparation included deep ripping and disk plowing of the soil and ground rig fumigation during winter. Deep ripping and disk plowing were conducted to break up traffic-induced or naturally occurring compaction layers and to increase soil moisture. In addition, potassium (K) and phosphorous (P) were applied. In the spring, a chisel chopper was used to loosen and plant (seed) the soil. A rod weeder was used to kill weeds before plant emergence. Between two to six weeks after planting, the potato plants emerged from the ground and irrigation (application of water only) began. The plants began to grow tubers a few weeks after emergence. A dense, green canopy from approximately June to August was maintained. Heavy
N fertilizer application began in early July. The potatoes were harvested during the first weeks of October, depending on the type of potato planted.

A center-pivot irrigation system that rotated in a clockwise fashion was used for irrigation and fertigation (application of water with N fertilizer). The timing and rate of irrigation and fertigation varied from day to day and between the two fields, under the discretion of the farmer. The type of fertilizer used was UAN solution 32, which contains 25% nitrate-N, 25% ammonium-N, and 50% urea-N. During days of N$_2$O and NH$_3$ measurements, 2.7 kg of N was applied per acre per full 360° rotation. There were two rates for sprinkler rotation: 12 deg hr$^{-1}$ and 18 deg hr$^{-1}$. For calculations of the N fertilizer application rates, see Appendix C (section C.1). Table 4.1 summarizes the schedule and rate of irrigation, fertigation, and chemigation (application of pesticides) for all days of NH$_3$ measurement.
Table 4.1: Schedule and rate of application for days of NH$_3$ measurement

<table>
<thead>
<tr>
<th>Date/Time (PDT)</th>
<th>Application Status</th>
<th>Rate of Sprinkler Rotation (deg hr$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 am on 7/07/07 to 6 am on 7/10/07</td>
<td>Fertigation</td>
<td>12</td>
</tr>
<tr>
<td>6 am on 07/10/07 to 6 pm on 07/10/07</td>
<td>Nothing</td>
<td>--</td>
</tr>
<tr>
<td>6 pm on 7/10/07 to 3 pm on 7/17/07</td>
<td>Fertigation</td>
<td>12</td>
</tr>
<tr>
<td>3 pm on 7/17/07 to 3 am on 07/18/07</td>
<td>Chemigation</td>
<td>30</td>
</tr>
<tr>
<td>3 am on 07/18/07 to 5 am on 07/19/07</td>
<td>Fertigation</td>
<td>12</td>
</tr>
<tr>
<td>5 am on 07/19/07 to 4 pm on 07/19/07</td>
<td>Nothing</td>
<td>--</td>
</tr>
<tr>
<td>5 am on 7/22/08 to 5 am on 7/24/08</td>
<td>Fertigation</td>
<td>18</td>
</tr>
<tr>
<td>5 am on 7/24/08 to 5 am on 7/25/08</td>
<td>Nothing</td>
<td>--</td>
</tr>
<tr>
<td>5 am on 7/25/08 to 5 am on 7/27/08</td>
<td>Fertigation</td>
<td>18</td>
</tr>
<tr>
<td>5 am on 7/27/08 to 5 am on 7/28/08</td>
<td>Nothing</td>
<td>--</td>
</tr>
<tr>
<td>5 am to 7:30 am on 7/28/08</td>
<td>Fertigation</td>
<td>18</td>
</tr>
<tr>
<td>7:30 am to 4 pm on 7/28/08</td>
<td>Irrigation</td>
<td>18</td>
</tr>
<tr>
<td>4 pm on 7/28/08 to 4 am on 7/29/08</td>
<td>Irrigation</td>
<td>12</td>
</tr>
<tr>
<td>after 4 am on 7/29/08</td>
<td>Nothing</td>
<td>--</td>
</tr>
</tbody>
</table>

The growing period was characterized by clear skies, high daytime temperatures, and cool nights. The average rainfall in the area was 150 mm so the potato plants were highly dependent on irrigation. Quincy fine sand soils are dominant in the fields, where anaerobic conditions are easily maintained for long periods of time. The measured bulk density of the surface soil (0–10 cm) was 1.24 Mg m$^{-3}$. The soil porosity and pH were 0.53 and 6.6, respectively.
4.3.2 Field Setup

Figure 4.1 is a diagram of the field setup. Southwest winds were dominant. Thus, the DOAS measurement path (path from the spectrograph to the retro-reflectors) was setup along the northeast edge of the potato fields to capture the NH₃ plume as it traveled along the wind trajectory. Path lengths varied from ~150 to ~250 m. Dispersion studies were conducted to characterize local dispersion conditions. Sulfur hexafluoride was used as a tracer gas and was released as a point source at the center of the fields. The trace gas analytical profile system (TGAPS) and syringe samplers were set up along the DOAS measurement path and extended over a distance of 800 m so that ideally the entire plume of SF₆ would be captured.

![Diagram of the field setup](image-url)
4.3.3 Measurement of Nitrous Oxide Concentrations and Flux Rates

Fluxes of N$_2$O were measured using the static closed chamber method (Hutchinson and Mosier, 1981). Chambers were installed between potato plant rows and sampled according to USDA-ARS GRACEnet (Greenhouse gas Reduction through Agricultural Carbon Enhancement network) protocols (USDA-ARS, 2003; www.GRACEnet.usda.gov). Chamber bases were made from 30.5 cm inner diameter x 15 cm high polyvinyl chloride (PVC) cylinders. The bases were inserted 10 cm into the soil and at least 24 hours prior to sampling to allow for stabilization. Chamber lids (30.5 cm inner diameter and 7.5 cm high) each contained a built-in vent tube, an internal fan, and a sampling port.

Air samples (about 30 ml each) from within the chambers were collected every 15 minutes over a period of 1 hour after covering the chambers. Gas sampling was performed 0 to 6.5 hours after fertigation and irrigation events. A Hewlett-Packard 5890 GC (Hewlett Packard, Palo Alto, CA) with an isothermal (40 °C) Alltech Porapak Q (80/100) column and an Electron Capture Detector (ECD) was used for analysis of air samples for N$_2$O concentrations. The carrier gas was 95% Argon (Ar) and 5% methane (CH$_4$). The GC flow rate ranged from 24.5 to 25.5 standard cubic centimeters per minute (sccm) and was maintained using an Alltech Digital Check Flow Meter. A 777 ppbv ± 10% (Scott-Marrin) N$_2$O standard was analyzed multiple times each day to determine analytical uncertainties. All samples were analyzed within a day of collection and approximately 10% of all the samples were reanalyzed during each analytical period for QA/QC purposes. Nitrous oxide flux rates were calculated using the slope of the gas concentration over time within the chamber as described by Hutchinson and Mosier (1981).
Table 4.2 summarizes the dates, time periods, and application status for all days of N$_2$O flux measurement. Nitrous oxide flux rates measured on days of no irrigation or fertigation were found negligible and are not included in the table.

<table>
<thead>
<tr>
<th>Date</th>
<th>Time (PDT)</th>
<th>Application Status</th>
<th>Chamber No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>7/09/07</td>
<td>1 pm to 2 pm</td>
<td>water w/ N fertilizer</td>
<td>1, 2</td>
</tr>
<tr>
<td>7/11/07</td>
<td>8:45 am to 9:45 am</td>
<td>water w/ N fertilizer</td>
<td>4</td>
</tr>
<tr>
<td>7/11/07</td>
<td>1:45 pm to 2:45 pm</td>
<td>water w/ N fertilizer</td>
<td>5, 6, 7, 8</td>
</tr>
<tr>
<td>7/17/07</td>
<td>9:15 am to 2:45 pm</td>
<td>water w/ N fertilizer</td>
<td>9</td>
</tr>
<tr>
<td>6/20/08</td>
<td>12:10 pm to 5:10 pm</td>
<td>water only</td>
<td>10</td>
</tr>
<tr>
<td>6/21/08</td>
<td>9:45 am to 3:15 pm</td>
<td>water only</td>
<td>11, 12, 13</td>
</tr>
<tr>
<td>7/22/08</td>
<td>9:45 am to 3:15 pm</td>
<td>water w/ N fertilizer</td>
<td>14</td>
</tr>
</tbody>
</table>

4.3.4 Measurement of Ammonia

Atmospheric NH$_3$ concentrations were measured using an open path DOAS instrument. The instrument measures the photo-absorption of NH$_3$ in the mid-ultraviolet from 200 to 240 nm, and consists of the following parts: a 150 W Xenon lamp as a (UV) light source, a telescope to beam the UV light into the measurement path, retro-reflectors to direct the light back towards the telescope, optics to focus the light onto a spectrograph, a double crossed Czerny-Turner spectrograph, and a detector (Mount et al., 2002). For measurements made in summer 2007, the instrument was housed in a trailer and contained a multiplexing detector (Reticon). Details are described in Mount et al. (2002). A revised configuration of the instrument was used in the summer 2008 measurements, which is more compact, easier to maneuver in the fields, utilizes a charge coupled device (CCD) detector, and employs a PC laptop for control of the system (instead of a Macintosh IIfx used for the original version). A schematic and picture of the revised DOAS instrument are shown in Figures 4.2 and 4.3, respectively.
The optics setup is similar to the setup of the original instrument but the spectrograph is placed below instead of to the side of the second folding mirror. In addition, the primary mirror is only 32 cm in diameter (instead of 40 cm) which allows for a more compact design. The revised system uses a Princeton Instruments CCD detector system (spec-10:400B) with 1340 x 400 array (pixel size of 19.4 µm square).

Figure 4.2: Schematic drawing of the revised DOAS instrument showing the optical light path from the Xe lamp to the CCD detector.
During data collection, the instrument was aligned towards the retro-reflectors so that the return signal counts were maximized. The integration time, which depends on the path length, was varied so that the maximum count rate was about 30,000 to 40,000 counts per integration time. Spectra were co-added so that data were averaged over five minute intervals. Electronic noise (dark spectra) of the instrument system was measured every hour, co-added and averaged, and subtracted from the actual spectra measured. A zinc (Zn) emission lamp was placed in front of the entrance slit every hour to provide a strong spectral fiducial at 213.9 nm. A nonlinear least squares fitting method, which accounts for Rayleigh and Mie scattering as well as the shift and stretch of the spectra, was used for data analysis.

4.3.5 Sulfur Hexafluoride Tracer Studies

During some NH₃ measurements, SF₆ tracer studies were conducted to help characterize local dispersion conditions. Sulfur hexafluoride is ideal for tracer studies because it is inert, its concentration can be measured with satisfactory accuracy at very low concentrations, and ambient concentrations of SF₆ are negligible. Even though SF₆ has a higher molecular weight (146.06 g mol⁻¹) than NH₃ (17.03 g mol⁻¹), the effect of this is negligible since turbulent
diffusion is the dominant transport mechanism (Kaharabata et al., 2000). Pure SF$_6$ was released from a single point at the center of the fields at rates ranging from ~0.6 to ~0.7 L min$^{-1}$. Sulfur hexafluoride concentrations were measured at several points along and beyond the DOAS integration path using a Trace Gas Automated Profiling System (TGAPS) and portable syringe samplers. The total length covered by the TGAPS sampling locations and syringe samplers was greater than 800 m, which was typically long enough to capture background concentrations outside of the plume. Continuous 5-minute average concentrations of SF$_6$ were measured through TGAPS sample lines at 7 locations. For more information on the TGAPS system, refer to Flaherty et al. (2007). Programmable and battery-operated syringe samplers measured 30-minute average SF$_6$ concentrations. All samples were analyzed using a HP 5890 GC with an ECD detector coupled with an electronic integrator (Benner and Lamb, 1985). Scott-Marrin (Riverside, CA) SF$_6$ standards with ±5% certified accuracy were used for calibration of the GC.

4.3.6 Inverse Gaussian Plume Model (SIMFLUX) for Estimating Ammonia Flux Rates

A Gaussian plume dispersion model (SIMFLUX) (Howard et al., 1996) was used in an inverse approach to estimate NH$_3$ emission rates. For fertigation periods, the sprinkler arm was modeled as an array of point sources in a rectangular area (10 m by 400 m). For periods of no fertigation or irrigation, the entire potato field was modeled as an array of point sources in a circle (400 m radius). The DOAS measurement path was modeled as a line of point receptors. All point sources and receptors were spaced 1 m apart. Measured variables, including wind speed and wind direction, were input into the Gaussian plume equation. Analytical equations (Martin, 1976) that represent the Pasquill-Gifford diffusion curves provided the best match for the tracer studies conducted and, thus, were employed for estimating the vertical and horizontal dispersion. The contributions from each point source were summed to estimate the concentration
at each downwind receptor. Ammonia flux rates were calculated from the average predicted NH$_3$ concentration based upon a point source release of 1 g s$^{-1}$ point$^{-1}$ and the measured DOAS NH$_3$ concentration.

A set of criteria was developed to determine which periods of NH$_3$ measurements were suitable for estimating NH$_3$ flux rates using SIMFLUX. Only periods associated with wind speeds greater than 1 m s$^{-1}$, mean winds pointed towards the DOAS measurement path, wind direction standard deviations less than 30 degrees, and measured NH$_3$ concentrations greater than the detection limit of 10 ppbv for 2008 and 5 ppbv for 2007 were used. For periods of low wind speeds and high variation in wind direction, the theoretical basis of the Gaussian plume dispersion model becomes invalid (see Figure C.16 in Appendix C for determination of cutoff values).

Howard et al. (1996) discovered that significant errors result if a receptor is placed too far crosswind from the plume centerline. As a result, only periods that captured a significant portion of the NH$_3$ plume were used for estimation of flux rates. For each measurement period, SIMFLUX was used to determine the plume width along the extended DOAS measurement position. The portion of the NH$_3$ plume captured was calculated by determining the ratio of the area of the plume measured and the area of the entire plume predicted.

4.3.7 Measuring Meteorological Variables

Meteorological information such as wind velocity and air temperature were measured using a sonic anemometer at 10 Hz frequency located near the DOAS instrument. Soil moisture and temperature were measured using Campbell Scientific Inc. CS515s Water Content Reflectometers and copper/constantan thermocouples, respectively. Campbell Scientific CR10 data loggers were used to record data. Leaf stomatal conductance was measured using Decagon
Devices Hand Held Leaf Porometer. The water filled pore space (WFPS) was calculated as the ratio of the volumetric water content and soil porosity.

4.4 RESULTS

4.4.1 Nitrous Oxide Flux Rates

The key factors affecting N$_2$O emissions from the potato fields were soil WFPS, soil temperature, and time after fertilizer application. Figure 4.4 is a plot of all measured N$_2$O flux rates as a function of time after the sprinkler has passed for irrigation and fertigation periods. Individual plots for 7/17/07, 6/20/08, 6/21/08, and 7/22/08 (days when at least 4 hours of sampling were conducted per chamber) can be found in Figures B.1, B.2, B.3, and B.4 in Appendix B. As shown in Figure 4.4, N$_2$O flux rates ranged from zero to 120 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$ and from zero to 290 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$ immediately after irrigation and fertigation periods, respectively. The average N$_2$O flux rate for all measurements was 74 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$ with a standard deviation of 73 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$ and a median of 55 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$. The average N$_2$O flux rate after fertigation was 97 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$ (standard deviation of 92 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$, median of 61 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$). The average N$_2$O flux rate after irrigation was 53 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$ (standard deviation of 42 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$, median of 45 $\mu$g N$_2$O m$^{-2}$ s$^{-1}$). Generally N$_2$O flux rates peaked between 2.5 to 3.5 hours after passing of the sprinkler arm. As expected, the maximum measured N$_2$O flux rate occurred during fertigation since the N inputs to the soil increased the amount of N available for nitrification and denitrification.
Nitrous oxide flux rates as a function of soil surface temperature and soil WFPS are plotted in Figures 4.5 and 4.6, respectively. The soil WFPS ranged from 68% to 87%, and soil surface temperatures ranged from 14 °C to 29 °C. The greatest N₂O flux rate occurred when the soil surface temperature was 19 °C and the soil WFPS was 73%. Fertigation and irrigation events combined with the ability of the fine-textured soil to hold water tightly resulted in anaerobic soil conditions, thus allowing denitrification activity to occur. There was no clear trend in N₂O flux rates as a function of soil surface temperature since soil denitrifiers are adapted to and capable of growing under a wide range of temperatures (Haverkort and MacKerron, 2000).
Figure 4.5: Nitrous oxide flux rates as a function of soil surface temperature. The gray and red data points represent periods of irrigation and fertigation, respectively.

Figure 4.6: Nitrous oxide flux rates as a function of soil WFPS. The black and red data points represent periods of irrigation and fertigation, respectively.
Table 4.3 contains N₂O flux rates for fertigation periods with the corresponding fertilizer application rate and the amount of N lost. The amount of N lost as a function of amount applied ranged from zero to 12%, with a mean of 4%, which is comparable to documented emission factor (EF) values of 0.1 to 8% (Dobbie and Smith, 2003). The standard deviation was 3.9% and the median was 2.2%. Although denitrification activity was the main driver for observed N₂O fluxes, the bulk soil may not have been entirely anaerobic. Aerobic conditions may occur at the base of the plants where it is less compacted.

Table 4.3: Percent of N lost for days of N fertilizer application

<table>
<thead>
<tr>
<th>Date</th>
<th>Chamber #</th>
<th>Fertilizer App. Rate (μg N m⁻² s⁻¹)</th>
<th>N₂O Flux Rate (μg N m⁻² s⁻¹)</th>
<th>% N Lost</th>
</tr>
</thead>
<tbody>
<tr>
<td>7/9/2007</td>
<td>1</td>
<td>25</td>
<td>3.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>24</td>
<td>3.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>92</td>
<td>11.8</td>
<td></td>
</tr>
<tr>
<td>7/11/2007</td>
<td>5</td>
<td>68</td>
<td>8.8</td>
<td></td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>45</td>
<td>5.8</td>
<td></td>
</tr>
<tr>
<td></td>
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<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>12</td>
<td>1.6</td>
<td></td>
</tr>
<tr>
<td>7/17/2007</td>
<td>9</td>
<td>780</td>
<td>17</td>
<td>2.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>87</td>
<td>11.2</td>
</tr>
<tr>
<td></td>
<td></td>
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<td>3.4</td>
<td>0.44</td>
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<td></td>
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<td>3.4</td>
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<tr>
<td>7/22/2008</td>
<td>14</td>
<td>1170</td>
<td>16</td>
<td>1.3</td>
</tr>
<tr>
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<td></td>
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<td>19</td>
<td>1.7</td>
</tr>
<tr>
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<td></td>
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</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>20</td>
<td>1.7</td>
</tr>
</tbody>
</table>

4.4.2 Ammonia Concentrations

Measurements from 12:30 pm on 7/26/08 to 4:00 am on 7/27/08 (PDT) provide a good example of measured NH₃ concentrations. Figure 4.7 is a time-series plot of measured NH₃ concentrations, wind speed, wind direction, air temperature, and soil temperature (measured 8 inches below the surface). Figures 4.8 to 4.9 are polar plots illustrating the measured NH₃ concentrations as a function of sprinkler arm rotation angle (marked with circles) and wind
direction (marked with squares). Each polar plot shows the relationship between the location of the sprinkler arm relative to the DOAS measurement path and the magnitude of the measured NH$_3$ concentrations. Additional time-series plots and polar plots for the rest of the data can be found in Appendix C (Figures C.1 to C.15). Only NH$_3$ concentrations above the DOAS instrument detection limit (5 ppbv for 2007 and 10 ppbv for 2008) are shown in the polar plots.

From 12:30 pm to 5:30 pm on 7/26/08, the sprinkler arm was fertigating in the northwest quadrant of the field and west/southwest winds were dominant (Figure 4.8). The average wind speed was $\sim$2.6 m s$^{-1}$. Since the sprinkler arm was far from the DOAS measurement path, lower

![Figure 4.7: Time series plot of measured 30-minute average NH$_3$ concentration, wind speed, wind direction, air temperature, and soil surface temperature from 07/26/08 12:30 pm to 07/27/08 4 am (PDT). The red line signifies the NH$_3$ concentration detection limit of 10 ppbv. The bars on the wind direction data represent the 30-minute variation in wind direction (wind direction sigma).](image-url)
NH₃ concentrations were expected for this period. Measured NH₃ concentrations were only a few ppbv above the detection limit.

Figure 4.8: Polar plot of NH₃ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) from 12:30 pm to 5:30 pm on 07/26/08 (PDT). All NH₃ concentrations below the detection limit (10 ppbv) have been omitted.

Measured NH₃ concentrations dramatically increased as the sprinkler arm got closer to the DOAS measurement path (Figure 4.9). The sprinkler arm was in the NE quadrant of the potato field from 6:30 pm to 10:30 pm on 7/26/08 (PDT). The maximum NH₃ concentration measured was ~115 ppbv and occurred around 7:30 pm on 7/26/08 (PDT). At this time, the sprinkler arm was pointed directly towards the DOAS measurement path and winds (~2 m s⁻¹) were coming from the west/southwest.
Results show that NH$_3$ concentrations (measured downwind of the potato field) depend on sprinkler arm location, wind speed, and wind direction. The highest NH$_3$ concentrations measured in the summer of 2007 and the summer of 2008 were \(~34\) ppbv and \(~112\) ppbv, respectively. Both peaks occurred during periods of fertigation with the sprinkler arm pointed directly towards the DOAS measurement path and winds greater than \(1\) m s$^{-1}$ were blowing towards the DOAS measurement path. The field conditions for both days of peak NH$_3$ measurements are summarized in Table 4.4. The large difference between the two maximum values can be attributed to the greater rate of fertilizer application on 7/26/08 than on 7/18/07. In addition, the measured peak on 7/26/08 occurred at night whereas the measured peak on 7/18/07
occurred during the day. During the day, unstable conditions dominate whereas during the night, conditions are more stable and less mixing occurs.

Table 4.4: Field conditions for peak NH$_3$ concentrations

<table>
<thead>
<tr>
<th>Date/Time (PDT)</th>
<th>NH$_3$ Conc. (ppbv)</th>
<th>N Fertilizer App. Rate (μg NH$_3$ m$^{-2}$ s$^{-1}$)</th>
<th>WS (m s$^{-1}$)</th>
<th>WD (deg)</th>
<th>DOAS path location</th>
<th>Air Temp. (°C)</th>
<th>Soil Temp. (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>7/18/07, 12 pm</td>
<td>~34</td>
<td>950</td>
<td>~3</td>
<td>240</td>
<td>NE quadrant</td>
<td>22.2</td>
<td>21.7</td>
</tr>
<tr>
<td>7/26/08, 7:30 pm</td>
<td>~110</td>
<td>1425</td>
<td>~2</td>
<td>251</td>
<td>NE quadrant</td>
<td>27.2</td>
<td>20.4</td>
</tr>
</tbody>
</table>

The relationship between measured NH$_3$ concentrations normalized by wind speed and the sprinkler arm rotation angle during fertigation periods are shown in Figure 4.10. Only measurements that were above the DOAS instrument detection limit and associated with wind direction variations less than 30 degrees and wind speeds greater than 1.0 m s$^{-1}$ are reported. As expected, when the DOAS measurement path was positioned along the SE quadrant of the field, the highest NH$_3$ concentration was measured when the sprinkler arm rotation angle was 180°, or due south. When the DOAS instrument was positioned along the NE quadrant of the field, the highest NH$_3$ concentration was measured when the sprinkler arm rotation angle was north-northeast (25°).
Figure 4.10: Measured NH$_3$ concentrations (ppbv) normalized by wind speed (m s$^{-1}$) vs. sprinkler arm rotation angle (degrees) during fertigation periods. The green and blue data points are periods when the DOAS measurement path was positioned along the SE and NE quadrants of the potato field, respectively.

4.4.3 Ammonia Flux Rates

Ammonia measurements were first investigated to determine which periods were suitable for use in estimation of flux rates. Data were separated into periods of fertigation and periods of no irrigation nor fertigation (made 1 day after the sprinkler arm stopped irrigating or fertigating). Fertigation periods were further separated by the position of the sprinkler arm relative to the DOAS measurement path. “Far” and “close” fertigation periods are associated with the sprinkler arm being far or pointed away from and close to or pointed towards the DOAS measurement path, respectively.

Figure 4.11 is a polar plot of 30-minute averaged NH$_3$ concentrations normalized by wind speed versus sprinkler arm rotation angle. Since wind speeds greatly influenced measurements (section 4.4.2), NH$_3$ concentrations were normalized by wind speed to remove the effect of the
variable. Ammonia concentrations normalized by wind speed were generally greater for “close” fertigation periods (4 ppbv m\(^{-1}\) s\(^{-1}\) to 11 ppbv m\(^{-1}\) s\(^{-1}\)) than for “far” fertigation periods (1.7 ppbv m\(^{-1}\) s\(^{-1}\) to 3.7 ppbv m\(^{-1}\) s\(^{-1}\)). The difference between the two groups was expected since point sources that were “close” to the DOAS measurement path dispersed less than “far” point sources.

The magnitudes of measured NH\(_3\) concentrations normalized by wind speed for periods of no irrigation/fertigation were comparable to measurements made during “far” fertigation periods.

Figure 4.11: Polar plot of 30-min. averaged NH\(_3\) concentrations (ppbv) normalized by wind speed (m s\(^{-1}\)) versus sprinkler arm rotation angle (degrees). Measurements made during “far” and “close” fertigation periods are in blue and red, respectively. Measurements made when the DOAS path was along the SE quadrant are marked as circles while measurements made when the DOAS path was along the NE quadrant are marked as x’s.
Ammonia flux rates and percent NH₃ loss as a function of wind speed during fertigation periods and periods of fertigation/irrigation are plotted in Figures 4.12 and 4.13, respectively. Similar to NH₃ concentrations (section 4.4.2), NH₃ flux rates and NH₃ loss (ratio of NH₃ emission rate to NH₃ application rate) increased with increasing wind speed.

Figure 4.12: NH₃ flux rates (µg m⁻² s⁻¹) and NH₃ loss (%) as a function of wind speed (m s⁻¹) measured during fertigation periods.
The relationship between NH$_3$ flux rates and the portion of plume captured was investigated. The portion of plume captured for each 30-min sampling period was calculated by determining the ratio of the area of the plume measured to the area of the entire plume predicted, assuming both were Gaussian shaped plumes. Figures 4.14 and 4.15 are plots of NH$_3$ flux rates as a function of portion of the NH$_3$ plume captured by the DOAS measurement path for all fertigation periods and periods of no irrigation/fertigation, respectively. For “far” fertigation periods, 35% to 80% of the NH$_3$ plume was captured by the DOAS measurement path while 53% to 99% was captured for “close” fertigation periods. For periods when no irrigation/fertigation was applied, only 34% to 43% of the NH$_3$ plume was captured. There is no significant dependence of NH$_3$ flux rates with the percentage of plume captured for all of the periods, however, for fertigation periods, the flux rates tend to decrease with increasing percentage of plume captured. There is currently no plausible explanation for this.
Figure 4.14: NH₃ flux rates (µg m⁻² s⁻¹) as a function of the portion of plume captured (%) for fertigation periods.

Figure 4.15: NH₃ flux rates (µg m⁻² s⁻¹) as a function of the portion of plume captured (%) for periods of no irrigation/fertigation.
Other variables were investigated to see if they impacted NH$_3$ emission rates. The relationship between NH$_3$ flux rates and solar radiation, soil temperature, air temperature, and the time after the sprinkler arm passed the DOAS measurement path were investigated and no obvious trends were found. Periods used for estimation of NH$_3$ flux rates occurred between 10 am and 6 pm (PDT), when fluctuations in air and soil temperatures were small. Due to the limited range in air and soil temperatures, there was no obvious relationship between NH$_3$ concentrations and emissions and soil and air temperatures (see Figures C.21 to C.24 in Appendix C). For “far” fertigation periods, there was no dependence of NH$_3$ flux rates with the time after the sprinkler passed the DOAS measurement path (Figure 4.16).

![Figure 4.16: NH$_3$ flux rates and percent loss as a function of time after the sprinkler passed the DOAS measurement path for “far” fertigation periods.](image-url)
Table 4.5 shows the range of measured NH₃ concentrations, estimated flux rates, and percent of NH₃ lost grouped by application status. The percent of NH₃ lost was calculated by dividing the estimated NH₃ flux rate by the fertilizer application rate. There was no irrigation period that met the criteria parameters and, thus, there are no calculations of flux rates and percent of NH₃ lost for irrigation periods. Estimated NH₃ flux rates ranged from 34 μg NH₃ m⁻² s⁻¹ to 250 μg NH₃ m⁻² s⁻¹ and from 9.1 μg NH₃ m⁻² s⁻¹ to 320 μg NH₃ m⁻² s⁻¹ during fertigation periods when the sprinkler arm was “far” and “close” to the DOAS measurement path, respectively. Higher normalized NH₃ concentrations were observed when the sprinkler arm was closer to the DOAS path. However, in estimating the flux rates, dispersion is taken into account and the variation in calculated flux rates is smaller. When there was no irrigation/fertigation, NH₃ flux rates varied from 0.6 μg NH₃ m⁻² s⁻¹ to 2.1 μg NH₃ m⁻² s⁻¹.

<table>
<thead>
<tr>
<th>Application Status</th>
<th>Measured NH₃ Conc. (ppbv)</th>
<th>Q (μg NH₃ m⁻² s⁻¹)</th>
<th>NH₃ Loss (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Min</td>
<td>Average</td>
</tr>
<tr>
<td>No Irrigation or</td>
<td>8.4 to 18.5</td>
<td>0.6</td>
<td>1.2</td>
</tr>
<tr>
<td>Fertigation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(spray close to</td>
<td>9.9 to 34.0</td>
<td>9.1</td>
<td>80</td>
</tr>
<tr>
<td>DOAS)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fertigation</td>
<td>5.5 to 13.6</td>
<td>34</td>
<td>140</td>
</tr>
<tr>
<td>(spray far from</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DOAS)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Ammonia loss during fertigation ranged from 0.95% to 48.4% (average of ~8% for “far” periods and ~12% for “close” periods). The frequency distribution of NH₃ flux rates during fertigation are shown in Figure 4.17. The average NH₃ flux rate during all fertigation periods was 100 μg m⁻² s⁻¹. The standard deviation was ~84 μg m⁻² s⁻¹ and the median was ~75 μg m⁻² s⁻¹.
A total N loss of 47% was estimated for the entire period of heavy N fertigation (month of July). A total of 38% was lost through NH₃ emissions. For N lost as NH₃, the following assumptions were used:

- an average of 2.75 Mg of N (6050 lbs) total was applied to the potato fields
- the average NH₃ flux during irrigation was assumed to be 50% of the average NH₃ flux during fertigation
- the average NH₃ flux during periods of no irrigation nor fertigation was 1 μg m⁻² s⁻¹
- daytime and nighttime flux rates are the same, even though flux rates were only calculated from daytime measurements
- the emissions during irrigation and fertigation were based on the 400 m by 10 m area underneath the sprinkler arm
- the emissions during periods with no irrigation nor fertigation were constant throughout the month (the decrease of emissions with time after fertigation was not accounted for).
Of the 38% N lost through NH$_3$ emissions, 27% of the applied N was lost during periods of no fertigation/irrigation while 8% was lost during fertigation and 3% was lost during irrigation. More was lost during no irrigation/fertigation periods than fertigation or irrigation periods because of the greater time period and area of emissions used for that calculation.

Table 4.6: Total N Loss through NH$_3$ Emissions for the month of July.

<table>
<thead>
<tr>
<th>Application Status</th>
<th># Hours</th>
<th>Avg. Emission Rate (Q) (μg-NH$_3$ m$^{-2}$ s$^{-1}$)</th>
<th>Avg. Q (μg-N m$^{-2}$ s$^{-1}$)</th>
<th>July N loss (kg-N)</th>
<th>% of Lost N (NH$_3$ Emissions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Fertigation nor Irrigation</td>
<td>414</td>
<td>1.2</td>
<td>1</td>
<td>749</td>
<td>27%</td>
</tr>
<tr>
<td>Fertigation</td>
<td>180</td>
<td>100</td>
<td>84</td>
<td>218</td>
<td>8%</td>
</tr>
<tr>
<td>Irrigation</td>
<td>150</td>
<td>50</td>
<td>42</td>
<td>91</td>
<td>3%</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>549</strong></td>
<td><strong>165</strong></td>
<td><strong>123</strong></td>
<td><strong>1056</strong></td>
<td><strong>38%</strong></td>
</tr>
</tbody>
</table>

A total of 9% was lost through N$_2$O emissions. For N lost as N$_2$O the following assumptions were used:

- an average of 2.75 Mg of N (6050 lbs) total was applied to the potato fields
- the average N$_2$O flux after irrigation was assumed to be 100 μg N$_2$O m$^{-2}$ s$^{-1}$
- the average N$_2$O flux after fertigation was assumed to be 50 μg N$_2$O m$^{-2}$ s$^{-1}$
- N$_2$O flux rates on days of no irrigation nor fertigation were assumed negligible
- emission rates were assumed constant (no decrease with time)

Of the 9% N lost through N$_2$O emissions, 6% of the applied N was lost during fertigation and 3% was lost during irrigation.
Table 4.7: Total N Loss through N\textsubscript{2}O Emissions for the month of July.

<table>
<thead>
<tr>
<th>Application Status</th>
<th># Hours</th>
<th>Avg. Emission Rate (Q) (μg-N\textsubscript{2}O m\textsuperscript{-2}s\textsuperscript{-1})</th>
<th>Avg. Q (μg-N m\textsuperscript{-2}s\textsuperscript{-1})</th>
<th>July N loss (kg-N)</th>
<th>% of Lost N (N\textsubscript{2}O Emissions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Fertigation nor Irrigation</td>
<td>414</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0%</td>
</tr>
<tr>
<td>Fertigation</td>
<td>180</td>
<td>100</td>
<td>32</td>
<td>174</td>
<td>6%</td>
</tr>
<tr>
<td>Irrigation</td>
<td>150</td>
<td>50</td>
<td>16</td>
<td>72</td>
<td>3%</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>9%</td>
</tr>
</tbody>
</table>

4.5 Conclusions

Ammonia results show that as wind speeds increased, so did the measured NH\textsubscript{3} concentrations as well as estimated NH\textsubscript{3} flux rates and percent of NH\textsubscript{3} loss. Wind direction influenced measured NH\textsubscript{3} concentrations and, as a result, calculated flux rates for periods when winds blew towards the DOAS measurement path more accurately represent emission rates from the potato fields. There were no clear relationships between NH\textsubscript{3} emissions and solar radiation, soil and air temperature, and time after the sprinkler arm passed the DOAS measurement path.

Estimated NH\textsubscript{3} flux rates ranged from 9.1 μg NH\textsubscript{3} m\textsuperscript{-2}s\textsuperscript{-1} to 460 μg NH\textsubscript{3} m\textsuperscript{-2}s\textsuperscript{-1} for fertigation periods (average 100 μg NH\textsubscript{3} m\textsuperscript{-2}s\textsuperscript{-1}) and from 0.6 μg NH\textsubscript{3} m\textsuperscript{-2}s\textsuperscript{-1} to 2.1 μg NH\textsubscript{3} m\textsuperscript{-2}s\textsuperscript{-1} (average 1.2 μg NH\textsubscript{3} m\textsuperscript{-2}s\textsuperscript{-1}) for periods of no irrigation/fertigation which indicates that a significant fraction of NH\textsubscript{3} loss from these fields occurs as a result of direct volatilization of NH\textsubscript{3} during fertigation. The estimated NH\textsubscript{3} flux rates from this study are much higher than the standard rate of 0.01 μg NH\textsubscript{3} m\textsuperscript{-2}s\textsuperscript{-1} typically used for all crops when compiling global emission inventories (Bouwman et al., 1997), with the flux rate obviously dependent upon quantity of applied N. The amount of NH\textsubscript{3} loss ranged from 0.95% to 48% for fertigation periods (average 10%), which is comparable to reported NH\textsubscript{3} loss from other studies (range from 8 to 18% using micrometeorological methods and from zero to 45% using dynamic flow chambers) (IFA, 2001).
A total N loss of 47% was estimated for the entire period of heavy N fertigation (month of July). Of the 47% N lost, 38% was lost through NH₃ emissions and 9% was lost through N₂O emissions. Previous studies available in literature report similar total N loss in potato plants. Reported N loss under current best management practices has been reported to be ~35% of N applied (Lang et al., 1999) with a mean of 60% (Balasubramanian et al., 2004).

Nitrous oxide flux rates generally peaked between 2.5 and 3.5 hours after irrigation or fertigation and appeared to be a result of denitrification activity (soil WFPS ranged from 68% to 87%). Nitrous oxide flux rates ranged from negligible to 280 μg N₂O m⁻² s⁻¹ and are much higher than reported emission rates by Flessa et al. (2002), who reported mean N₂O emissions of 0.02 and 0.03 μg N₂O m⁻² s⁻¹ during the 1997 and 1998 growing periods of two adjacent potato fields in Germany using automated gas sampling chambers.

The average N loss as N₂O ranged from zero to 12% (average ~4%) of the applied N fertilizer which is comparable to documented EF values of 0.1 to 8% (Dobbie and Smith, 2003). The average N loss is much higher than that reported by Haile-Mariam et al. (2008), who reported N₂O losses of 0.3% of applied fertilizer in potato fields in central Washington during the 2005 and 2006 growing seasons using static chambers. Similar to the fields used in this study, the fields used by Haile-Mariam et al. (2008) were applied with UAN solution 32 through a center pivot irrigation system and aerobic soil conditions dominated in the fields.
5.1 Uncertainties in NH$_3$ Measurements

There are various errors associated with the measurements of NH$_3$. Uncertainty in NH$_3$ measurements result from errors in the absolute NH$_3$ absorption cross section of $\pm 20\%$ (Mount et al., 2002). The error in residual analysis was approximately $\pm 25\%$ and was due to uncontrollable variations in the dark current (electronic noise of the system) caused by the external radiator used for temperature regulation of the CCD detector, which resulted in difficult subtraction of darks during data analysis. Other errors include changes in the spectral output of the Xe lamp which could not be monitored at high frequency and illumination problems on the system optics from atmospheric turbulence. The overall error in the NH$_3$ concentration data combining all of these uncertainties in quadrature is estimated to be $\pm 30\% \pm 3$ppbv.

5.2 Uncertainties in NH$_3$ Flux Rates

The uncertainty in estimated NH$_3$ emission rates result from errors associated with the estimation of NH$_3$ concentrations as well as errors in SIMFLUX. SIMFLUX assumes uniform emission rates from all of the point sources within the field. Numerous factors (i.e. environmental factors, soil properties) regulate NH$_3$ loss from the soil and potato plants and these factors may not have been uniform throughout the field. During fertigation periods, only the area covered by the sprinkler arm was modeled (400 m by 10 m) as an NH$_3$ source. While other parts of the field contributed to NH$_3$ emissions, it was difficult to differentiate between direct volatilization from the spray of the sprinkler arm and the entire field.
There are also errors associated with the SF$_6$ tracer data used to calibrate the dispersion algorithm in SIMFLUX. During SF$_6$ tracer studies, the sources of error include changes in the SF$_6$ release rate, GC analysis (uncertainty in the gas standards ±5%), and error in duplicates. When compared with the SF$_6$ tracer studies, the equations by Martin (1976) for determining dispersion coefficients resulted in an average mean normalized gross error (MNGE) of 41% between predicted and observed SF$_6$ concentrations (Appendix A). The error in the NH$_3$ flux estimates is approximately ± 52%.

5.3 Uncertainties in N$_2$O Measurements

The uncertainty in N$_2$O measurements are derived from errors in chamber sampling and GC analysis. Errors associated with the GC analysis include the uncertainty in the gas standard utilized for the calibration of the voltage signal from the ECD (±10%) and the error in duplicate measurements (maximum error was ±18%). The total error ranged from ±11% to ±21%.

The errors associated with the chamber sampling cannot be quantified and include the uncertainty in the accuracy of volume measurements inside the chambers since soil surfaces were irregular. In addition, the concentration with time data was noisy and determination of flux rates required disregarding outliers based on judgment. Also, the number of measurements collected was not enough to capture the heterogeneity of N$_2$O emissions throughout the field. Other errors associated with chamber sampling should be minimal since installation and sampling of chambers were conducted according to the prudent guidelines of the USDA-ARS GRACEnet protocols (USDA-ARS, 2003). All samples were analyzed within a day of collection to prevent error due to leakage and approximately 10% of all the samples were reanalyzed during each analytical period to ensure data quality.
CHAPTER 6

CONCLUSIONS AND FUTURE WORK

There are limited studies that have measured the magnitude of Nitrogen (N) emissions from intensively managed croplands. It is important to understand the magnitude and factors controlling NH$_3$ and N$_2$O emissions because they pose negative effects to human health and the environment. In this research, concentrations and fluxes of ammonia (NH$_3$) and (N$_2$O) from potato fields in central Washington were measured during the 2007 and 2008 growing seasons (June – August).

Nitrous oxide fluxes were measured using a static chamber method and samples were analyzed using a GC-ECD. The chamber method was useful for investigating the processes controlling N$_2$O fluxes. However, the method made it difficult to extrapolate measured fluxes to field-scale estimates since there is a wide range of environmental conditions throughout the fields as well as a large variation in the soil properties.

This study is the first field experiment using the DOAS technique to measure NH$_3$ fluxes from potato fields. The DOAS technique proved to be an appropriate method for measuring NH$_3$ flux rates from the fields because it did not disturb the soil-plant-atmosphere pathway and it provided measurements integrated over a large area. However, NH$_3$ measurements were limited by the problems associated with the revised instrument (i.e. misalignment of mirrors, inability of the internal cooler within the CCD detector to regulate temperature), which disrupted data collection and decreased data quality. Correction of those problems will significantly enhance future measurements made by DOAS.
Suggested Future Work:

- Further improvement of the DOAS instrument (i.e. re-aligning of the mirrors, addition of an appropriate cooler to regulate temperature for the CCD detector) will improve future measurement quality and quantity.

- Measurements of NH₃ and other factors influencing emissions need to be made over a sufficiently longer time period so that the relationships between NH₃ emissions and influencing factors can be determined.

- The use of a relaxed-eddy accumulation technique (REA) concurrent with the DOAS technique would allow for continuous NH₃ flux measurements and provide complementary data to the current data set. (Zhu et al. (2000) have shown that the REA technique is successful in measuring NH₃ fluxes over a corn field.)

- Measurement of emissions from other potato fields under a different management practice. (The fields used in this study were continuously applied with small amounts of N fertilizer throughout the growing season. Other farmers apply large amounts of N fertilizer at one time only (i.e. at the beginning of the season)).

- Use of an automated system of chambers, which would allow for continuous, long-term measurements of N₂O and provide a better understanding of spatial and temporal variations in N₂O fluxes throughout the fields.
REFERENCES


APPENDIX A:

COMPARISON OF VARIOUS METHODS
FOR DETERMINING DISPERSION COEFFICIENTS
In this appendix, a brief overview of the various methods for determining the dispersion coefficients and the comparison results are presented.

**A.1 Description of Sigma Methods**

Critical input parameters to the Gaussian diffusion models are the dispersion coefficients ($\sigma_y$ and $\sigma_z$), which express the spread of the plume in the horizontal and vertical crosswind directions. These coefficients can be determined empirically or theoretically. The methods compared were:

1) Empirical methods:
   i) Martin’s equations based on Pasquill-Gifford diffusion curves (Martin, 1976)
   ii) Briggs’ equations for open-country conditions based on several field studies (Briggs, 1973)

2) Theoretical method: Taylor’s near-field dispersion theory (Mohan and Siddiqui, 1997)

3) Combination of theory and empirical results: Draxler’s method (Draxler, 1976)

**A.1.1 Empirical Methods**

Values of $\sigma_y$ and $\sigma_z$ may be obtained from empirical results. Perhaps the most well-known diffusion study is the 1956 Project Prairie Grass in O’Neill, Nebraska. During Project Prairie Grass, sulfur dioxide (SO$_2$) was used as the tracer gas and released at a height of 0.5 m above ground over a large flat field (Arya, 1999). Concentrations at 1.5 m above ground were measured at distances between 50 and 800 m from the source and were based on 10 min sampling periods (Arya, 1992). This experiment resulted in the Pasquill and Gifford (P–G) dispersion curves. To estimate $\sigma_y$ and $\sigma_z$ from the P-G curves, the stability class must first be determined based on simple measurements such as sunshine, cloud cover, and wind speed
measured at 10 meters above ground, as shown in Table 3.2. “Strong” incoming solar radiation corresponds to a solar angle greater than 60° with clear skies, while “slight” incoming solar radiation corresponds to a solar angle less than 35° with clear skies (Arya, 1999). Along with solar angle, cloud cover is also considered when determining the strength of incoming solar radiation. The use of computers for solving the Gaussian dispersion equations make it desirable to convert the P-G curves into analytical equations. The equations of Martin (1976) represent a reasonable fit to the Pasquill and Gifford (P–G) dispersion curves.

Table A.1: Pasquill Stability Classes

<table>
<thead>
<tr>
<th>Surface Wind Speed (m s⁻¹)</th>
<th>Daytime Incoming Solar Radiation</th>
<th>Nighttime Cloud Cover</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Strong</td>
<td>Moderate</td>
</tr>
<tr>
<td>&lt; 2</td>
<td>A</td>
<td>A-B</td>
</tr>
<tr>
<td>2-3</td>
<td>A-B</td>
<td>B</td>
</tr>
<tr>
<td>3-5</td>
<td>B</td>
<td>B-C</td>
</tr>
<tr>
<td>5-6</td>
<td>C</td>
<td>C-D</td>
</tr>
<tr>
<td>&gt; 6</td>
<td>C</td>
<td>D</td>
</tr>
</tbody>
</table>

Note: Class D applies to heavily overcast skies, at any wind speed day or night

The following equations were developed by Martin for calculating the dispersion coefficients:

\[
\sigma_y = ax^{0.894} \quad \text{A.1}
\]

\[
\sigma_z = cx^d + f \quad \text{A.2}
\]

where the constants \(a, c, d, \text{ and } f\) are given in Table A.2. The values of \(\sigma_y\) and \(\sigma_z\) are in meters when the value of the downwind distance \(x\) of the receptor from the source is given in kilometers.
In addition to Project Prairie Grass, a number of other diffusion experiments have been conducted to empirically estimate the dispersion coefficients. For rural situations, the three main diffusion experiments are the 1956 Project Prairie Grass, the Brookhaven National Laboratory (BNL) experiments, and the Tennessee Valley Authority (TVA) experiment. Results from Project Prairie Grass are for non-buoyant, near ground level releases at downwind distances less than 1 km. The BNL experiment results reflect effects of elevated releases (~108 m above ground) over a rougher surface at downwind distances extending to tens of kilometers (Arya, 1999). The TVA results reflect effects of highly buoyant releases from higher elevations over gently rolling terrain (Arya, 1999). In 1973, Briggs synthesized the results from Project Prairie Grass, BNL, and TVA to produce a set of formulas for open country situations, which are shown in Table A.3 (Arya, 1999). Briggs’ formulas use the Pasquill stability classes.

<table>
<thead>
<tr>
<th>Stability</th>
<th>$a$</th>
<th>$x &lt; 1$ km</th>
<th>$x &gt; 1$ km</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$c$</td>
<td>$d$</td>
<td>$f$</td>
</tr>
<tr>
<td>A</td>
<td>213</td>
<td>440.8</td>
<td>1.041</td>
</tr>
<tr>
<td>B</td>
<td>156</td>
<td>106.6</td>
<td>1.149</td>
</tr>
<tr>
<td>C</td>
<td>104</td>
<td>61.0</td>
<td>0.911</td>
</tr>
<tr>
<td>D</td>
<td>68</td>
<td>33.2</td>
<td>0.725</td>
</tr>
<tr>
<td>E</td>
<td>50.5</td>
<td>22.8</td>
<td>0.675</td>
</tr>
<tr>
<td>F</td>
<td>34</td>
<td>14.35</td>
<td>0.740</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Stability</th>
<th>$\sigma_y$ (m)</th>
<th>$\sigma_z$ (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>$0.22x(1 + 0.0001x)^{-5}$</td>
<td>$.20x$</td>
</tr>
<tr>
<td>B</td>
<td>$0.16x(1 + 0.0001x)^{-5}$</td>
<td>$.12x$</td>
</tr>
<tr>
<td>C</td>
<td>$0.11x(1 + 0.0001x)^{-5}$</td>
<td>$.08x(1 + 0.0002x)^{-5}$</td>
</tr>
<tr>
<td>D</td>
<td>$0.08x(1 + 0.0001x)^{-5}$</td>
<td>$.06x(1 + 0.0015x)^{-5}$</td>
</tr>
<tr>
<td>E</td>
<td>$0.06x(1 + 0.0001x)^{-5}$</td>
<td>$.03x(1 + 0.0003x)^{-1}$</td>
</tr>
<tr>
<td>F</td>
<td>$0.04x(1 + 0.0001x)^{-5}$</td>
<td>$.016x(1 + 0.0003x)^{-1}$</td>
</tr>
</tbody>
</table>
A.1.2 Theoretical Methods

Another way of determining the dispersion coefficients is through Taylor’s (1921) theory of diffusion. This theory considers one-dimensional diffusion of a cloud of particles released from the same point in a stationary and homogeneous turbulent flow using a frame of reference moving with the mean flow (Arya, 1999). Taylor’s theory is difficult to use since it requires exact knowledge of the behavior of the Lagrangian autocorrelation $R(\xi)$ at different times which is often very difficult to obtain (Mohan et al., 1997). Several methods have been suggested for determining $\sigma_y$ and $\sigma_z$ which do not require the autocorrelation function but still retain the essential features of Taylor’s theory. Pasquill suggested the following relationships, which are more applicable for use with readily measured Eulerian quantities (Mohan et al., 1997):

\begin{align*}
\sigma_y &= \sigma_v t F_y \quad \text{(A.3)} \\
\sigma_z &= \sigma_w t F_z \quad \text{(A.4)}
\end{align*}

where $\sigma_v$ and $\sigma_w$ are the standard deviation of velocity fluctuations in the y and z directions, $t$ is the travel time, and $F_y$ and $F_z$ are universal functions of a set of parameters that specify the characteristics of the atmospheric boundary layer. For this application, measurements of wind components $u$, $v$, and $w$ were measured at 10 Hertz frequency using a sonic anemometer. A coordinate rotation was applied to the $u$ and $v$ components so that the $u$ component is along the mean wind direction and the mean $v$ component of the wind is zero. The $w$ component was not used in the coordinate conversion since its magnitude is small compared to the other two components. The standard deviation of the $v$ and $w$ wind components were calculated for every 30-min period to get the $\sigma_v$ and $\sigma_w$ respectively.
For receptors close to the emission source, the suggested relationships above can be reduced to:

\[ \sigma_{y,z} = \sigma_{v,w} t, \quad t = \frac{x}{u} \]

where \( x \) is the downwind distance of receptor from source in meters and \( u \) is the wind speed in meters per second. Since the field data used for this project utilized receptors that were relatively close to the source (~400 m downwind), use of the relationship above is appropriate.

### A.1.3 Combination of Theory and Empirical Results

In 1975, Draxler combined Taylor’s theory with empirical data from various field experiments including Project Prairie Grass and others (Draxler, 1976). Draxler’s equations are:

\[ \sigma_{y,z} = \sigma_{\theta,\phi} x f_{1,2} \left( \frac{T}{T_i} \right) \]

\[ f_{1,2} (T / T_i) = \frac{1}{1 + 0.9 \left( \frac{T}{T_i} \right)^{0.5}} \]

where \( \sigma_{\theta,\phi} \) are the standard deviations of vertical and horizontal wind direction fluctuations, \( x \) is the downwind distance of receptor from source in meters, and \( T_i \) is a time scale proportional to the Langrangian scale (Draxler, 1976). Values of \( T_i \) for different stabilities and source heights for use in \( f_1 \) and \( f_2 \) are given in Table A.4 (Draxler, 1976). Values of \( T_i \) are mean values from each category of experiments.

<table>
<thead>
<tr>
<th></th>
<th>Surface Sources</th>
<th>Elevated Sources</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Stable</td>
<td>Unstable</td>
</tr>
<tr>
<td><strong>Std Dev</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Horizontal</td>
<td>300</td>
<td>300</td>
</tr>
<tr>
<td>Vertical</td>
<td>50</td>
<td>100</td>
</tr>
</tbody>
</table>
A.2 Results

The various methods described were used to simulate eight 30-minute periods of sulfur hexafluoride (SF$_6$) tracer diffusion data collected from the potato fields (see Table A.5). This section presents the results from the various $\sigma_y$ and $\sigma_z$ schemes for the eight 30-minute periods of SF$_6$ tracer diffusion data. Predicted and observed SF$_6$ concentrations are compared graphically and statistically.

Table A.5: Periods of diffusion data used for comparing various diffusion schemes

<table>
<thead>
<tr>
<th>Date</th>
<th>Time (PST)</th>
</tr>
</thead>
<tbody>
<tr>
<td>7/19/07</td>
<td>11:30 am to 12:00 pm</td>
</tr>
<tr>
<td>7/19/07</td>
<td>12:30 pm to 1:00 pm</td>
</tr>
<tr>
<td>7/19/07</td>
<td>1:00 pm to 1:30 pm</td>
</tr>
<tr>
<td>7/19/07</td>
<td>1:30 pm to 2:00 pm</td>
</tr>
<tr>
<td>7/19/07</td>
<td>2:30 pm to 3:00 pm</td>
</tr>
<tr>
<td>7/29/08</td>
<td>9:20 am to 9:50 am</td>
</tr>
<tr>
<td>7/29/08</td>
<td>11:20 am to 11:50 am</td>
</tr>
<tr>
<td>7/29/08</td>
<td>11:50 am to 12:20 pm</td>
</tr>
</tbody>
</table>

A.2.1 Comparison graphs

Concentration distributions obtained from using various $\sigma_y$ and $\sigma_z$ schemes for 7/19/07 (12:30 pm to 1:00 pm PST) and 7/29/08 (11:50 am to 12:20 pm PST) are shown in Figures A.1-A.2. Results for other periods are given in figures and tables at the end of this appendix. The receptor position represents the location of each syringe sampler employed during the SF$_6$ tracer studies. For all eight sampling periods, Martin’s, Briggs’, and Draxler’s methods performed well in terms of predicting the shape of the measured concentration distributions while Taylor’s method did not. In addition, Taylor’s method severely underestimated the maximum observed values and overestimates the outside concentrations.

One to one ratio comparisons (Figures A.3 and A.4) provide another way of evaluating the accuracy of each method. For four out of the eight sampling periods, Martin’s method
yielded the best results in terms of accurately predicting observed concentration values at every receptor. For three out of the eight sampling periods, Brigg’s method closely predicted measured concentrations for every receptor.

Figure A.1: Predicted and observed SF₆ concentration distribution from 12:30 pm to 1:00 pm on 7/19/07 (PST).

Figure A.2: Predicted and observed SF₆ concentration distribution from 11:50 am to 12:20 pm on 7/29/08 (PST).
Figure A.3: A 1:1 comparison of observed vs. predicted SF$_6$ concentrations from 12:30 pm to 1:00 pm on 7/19/07 (PST).

Figure A.4: A 1:1 comparison of observed vs. predicted SF$_6$ concentrations from 11:50 am to 12:20 pm on 7/29/08 (PST).
A.2.2 Statistical Analysis

Various statistical measures were calculated for further analysis of each method. The statistical measures were: Fractional Bias (FB), Unpaired Peak Prediction Error (UPPE), and Mean Normalized Gross Error (MNGE). These statistical measures were performed for all measured SF\textsubscript{6} concentrations greater than 50 pptv. The FB, UPPE, and MNGE are defined as:

\[
\text{Fractional Bias (FB)} = \frac{1}{N} \sum_{i=1}^{N} \frac{C_p(x,t_i) - C_o(x,t_i)}{C_p(x,t_i) + C_o(x,t_i)}
\]

\[
\text{Unpaired Peak Prediction Error (UPPE)} = \frac{C_p(x,t')_{\text{max}} - C_o(x,t')_{\text{max}}}{C_o(x,t')_{\text{max}}}
\]

\[
\text{Mean Normalized Gross Error (MNGE)} = \frac{1}{N} \sum_{i=1}^{N} \frac{|C_p(x,t_i) - C_o(x,t_i)|}{C_o(x,t_i)} \times 100\%
\]

where \(N\) is the number of concentrations at a given receptor location, \(C_o(x,t)\) and \(C_p(x,t)\) are the observed and predicted concentrations at the receptor location at \(x\) for hour \(t\), respectively, and \(C_o(x,t')_{\text{max}}\) and \(C_p(x,t')_{\text{max}}\) are the observed and predicted maximum concentrations at a specific receptor location, respectively. The statistical measures for all sampling periods were averaged and are given in Table A.6. The lowest value in each category is highlighted in yellow.

Table A.6: Average Values of Statistical Measures for Various Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>FB</th>
<th>UPPE</th>
<th>MNGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martin</td>
<td>-0.06</td>
<td>0</td>
<td>41 %</td>
</tr>
<tr>
<td>Briggs</td>
<td>-0.11</td>
<td>-0.08</td>
<td>43 %</td>
</tr>
<tr>
<td>Draxler</td>
<td>0.04</td>
<td>-0.06</td>
<td>60 %</td>
</tr>
<tr>
<td>Taylor</td>
<td>-0.17</td>
<td>-0.7</td>
<td>53 %</td>
</tr>
</tbody>
</table>

Bias statistics measure the tendency of a model to over or under predict observed values. FB equally weighs positive and negative bias estimates. An FB of zero is ideal. Draxler’s
method resulted in the lowest average FB values while Taylor’s method yielded the highest. Martin’s method resulted in the second lowest average FB.

Error is a measure of the extent that the model deviates from the observations. The UPPE compares the maximum observed and maximum predicted concentrations at a single sampling period. For the MNGE, a value of zero would indicate that the model exactly matches the observed values at all points in space/time. Martin’s method resulted in the lowest average UPPE and MNGE values.

A.2.3 Conclusions

Several methods (Martin, Briggs, Taylor, and Draxler) for determining \( \sigma_y \) and \( \sigma_z \) have been compared with real diffusion data. Through analysis of comparison plots between predicted and observed SF6 values as well as statistical measures including FB, FGE, and UPPE, Martin’s method yielded the best results. The average UPPE was zero for Martin’s method, showing that maximum observed and maximum predicted concentrations closely matched at every period. The average FB and MNGE for all sampling periods were -0.06 and 41 % for Martin’s method. Thus, Martin’s method was used for estimating the diffusion coefficients for SIMFLUX to simulate NH\(_3\) and N\(_2\)O emission rates from potato fields.
A.3 Additional Results

Figure A.5: Predicted and observed SF$_6$ concentration distribution for Tracer 7/19/07 (Syringe #2, 11:30 am to 12:00 pm PST)

Figure A.6: A 1:1 comparison of observed vs. predicted SF$_6$ concentrations for Tracer 7/19/07 (Syringe #2, 11:30 am to 12:00 pm PST)
Figure A.7: Predicted and observed SF$_6$ concentration distribution for Tracer 7/19/07 (Syringe #5, 1:00 pm to 1:30 pm PST)

Figure A.8: A 1:1 comparison of observed vs. predicted SF$_6$ concentrations for Tracer 7/19/07 (Syringe #5, 1:00 pm to 1:30 pm PST)
Figure A.9: Predicted and observed SF₆ concentration distribution for Tracer 7/19/07 (Syringe #6, 1:30 pm to 2:00 pm PST)

Figure A.10: A 1:1 comparison of observed vs. predicted SF₆ concentrations for Tracer 7/19/07 (Syringe #6, 1:30 pm to 2:00 pm PST)
Figure A.11: Predicted and observed SF₆ concentration distribution for Tracer 7/19/07 (Syringe #8, 2:30 pm to 3:00 pm PST)

Figure A.12: A 1:1 comparison of observed vs. predicted SF₆ concentrations for Tracer 7/19/07 (Syringe #8, 2:30 pm to 3:00 pm PST)
Figure A.13: Predicted and observed SF$_6$ concentration distribution for Tracer 7/29/08 (Syringe #4, 9:20 am to 9:50 am PST)

Figure A.14: A 1:1 comparison of observed vs. predicted SF$_6$ concentrations for Tracer 7/29/08 (Syringe #4, 9:20 am to 9:50 am PST)
Figure A.15: Predicted and observed SF₆ concentration distribution for Tracer 7/29/08 (Syringe #8, 11:20 am to 11:50 am PST)

Figure A.16: A 1:1 comparison of observed vs. predicted SF₆ concentrations for Tracer 7/29/08 (Syringe #8, 11:20 am to 11:50 am PST)
### Table A.7: Values of Statistical Measures for Various Sigma Methods for Tracer 7/19/07 (Syringe #2, 11:30 am to 12:00 pm PST)

<table>
<thead>
<tr>
<th>Method</th>
<th>FB</th>
<th>MNB</th>
<th>MAE</th>
<th>MNGE</th>
<th>UPPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martin</td>
<td>-0.12</td>
<td>-25%</td>
<td>74.24</td>
<td>45%</td>
<td>-0.22</td>
</tr>
<tr>
<td>Briggs</td>
<td>-0.17</td>
<td>-40%</td>
<td>85.44</td>
<td>48%</td>
<td>-0.28</td>
</tr>
<tr>
<td>Draxler</td>
<td>0.02</td>
<td>25%</td>
<td>70.70</td>
<td>54%</td>
<td>-0.12</td>
</tr>
<tr>
<td>Taylor</td>
<td>-0.12</td>
<td>-32%</td>
<td>130.88</td>
<td>46%</td>
<td>-0.72</td>
</tr>
</tbody>
</table>

### Table A.8: Values of Statistical Measures for Various Sigma Methods for Tracer 7/19/07 (Syringe #5, 1:00 pm to 1:30 pm PST)

<table>
<thead>
<tr>
<th>Method</th>
<th>FB</th>
<th>MNB</th>
<th>MAE</th>
<th>MNGE</th>
<th>UPPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martin</td>
<td>0.02</td>
<td>18%</td>
<td>79.25</td>
<td>44%</td>
<td>0.15</td>
</tr>
<tr>
<td>Briggs</td>
<td>-0.04</td>
<td>-1%</td>
<td>63.44</td>
<td>38%</td>
<td>0.06</td>
</tr>
<tr>
<td>Draxler</td>
<td>0.11</td>
<td>59%</td>
<td>100.59</td>
<td>59%</td>
<td>0.09</td>
</tr>
<tr>
<td>Taylor</td>
<td>-0.12</td>
<td>-38%</td>
<td>85.71</td>
<td>38%</td>
<td>-0.66</td>
</tr>
</tbody>
</table>

### Table A.9: Values of Statistical Measures for Various Sigma Methods for Tracer 7/19/07 (Syringe #6, 1:30 pm to 2:00 pm PST)

<table>
<thead>
<tr>
<th>Method</th>
<th>FB</th>
<th>MNB</th>
<th>MAE</th>
<th>MNGE</th>
<th>UPPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martin</td>
<td>0.03</td>
<td>16%</td>
<td>58.05</td>
<td>23%</td>
<td>0.28</td>
</tr>
<tr>
<td>Briggs</td>
<td>-0.03</td>
<td>-5%</td>
<td>65.36</td>
<td>32%</td>
<td>0.17</td>
</tr>
<tr>
<td>Draxler</td>
<td>0.13</td>
<td>78%</td>
<td>128.32</td>
<td>78%</td>
<td>0.36</td>
</tr>
<tr>
<td>Taylor</td>
<td>-0.10</td>
<td>-25%</td>
<td>115.45</td>
<td>50%</td>
<td>-0.59</td>
</tr>
</tbody>
</table>

### Table A.10: Values of Statistical Measures for Various Sigma Methods for Tracer 7/19/07 (Syringe #8, 2:30 pm to 3:00 pm PST)

<table>
<thead>
<tr>
<th>Method</th>
<th>FB</th>
<th>MNB</th>
<th>MAE</th>
<th>MNGE</th>
<th>UPPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martin</td>
<td>-0.05</td>
<td>14%</td>
<td>115.18</td>
<td>70%</td>
<td>0.58</td>
</tr>
<tr>
<td>Briggs</td>
<td>-0.10</td>
<td>-7%</td>
<td>101.55</td>
<td>66%</td>
<td>0.45</td>
</tr>
<tr>
<td>Draxler</td>
<td>0.14</td>
<td>90%</td>
<td>137.90</td>
<td>93%</td>
<td>0.40</td>
</tr>
<tr>
<td>Taylor</td>
<td>-0.07</td>
<td>-22%</td>
<td>55.70</td>
<td>27%</td>
<td>-0.33</td>
</tr>
</tbody>
</table>

### Table A.11: Values of Statistical Measures for Various Sigma Methods for Tracer 7/29/08 (Syringe #4, 9:20 am to 9:50 am PST)

<table>
<thead>
<tr>
<th>Method</th>
<th>FB</th>
<th>MNB</th>
<th>MAE</th>
<th>MNGE</th>
<th>UPPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martin</td>
<td>-0.05</td>
<td>-15%</td>
<td>69.36</td>
<td>23%</td>
<td>-0.24</td>
</tr>
<tr>
<td>Briggs</td>
<td>-0.09</td>
<td>-29%</td>
<td>98.61</td>
<td>29%</td>
<td>-0.24</td>
</tr>
<tr>
<td>Draxler</td>
<td>-0.05</td>
<td>-13%</td>
<td>118.44</td>
<td>31%</td>
<td>-0.51</td>
</tr>
<tr>
<td>Taylor</td>
<td>-0.28</td>
<td>-71%</td>
<td>264.82</td>
<td>71%</td>
<td>-0.87</td>
</tr>
</tbody>
</table>
Table A.12: Values of Statistical Measures for Various Sigma Methods for Tracer 7/29/08
(Syringe #8, 11:20 am to 11:50 am PST)

<table>
<thead>
<tr>
<th>Method</th>
<th>FB</th>
<th>MNB</th>
<th>MAE</th>
<th>MNGE</th>
<th>UPPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martin</td>
<td>-0.18</td>
<td>-48%</td>
<td>178.40</td>
<td>48%</td>
<td>-0.26</td>
</tr>
<tr>
<td>Briggs</td>
<td>-0.22</td>
<td>-57%</td>
<td>214.62</td>
<td>57%</td>
<td>-0.33</td>
</tr>
<tr>
<td>Draxler</td>
<td>-0.11</td>
<td>-28%</td>
<td>184.98</td>
<td>50%</td>
<td>-0.27</td>
</tr>
<tr>
<td>Taylor</td>
<td>-0.25</td>
<td>-57%</td>
<td>350.85</td>
<td>68%</td>
<td>-0.84</td>
</tr>
</tbody>
</table>
APPENDIX B:
ADDITIONAL N₂O RESULTS
Figure B.1: N₂O flux rate, soil temperature, and soil volumetric water content (0-10 inches below the soil surface) as a function of time after fertigation from 9:15 am to 2:45 pm on 07/17/07 (PDT).

Figure B.2: N₂O flux rate, soil temperature, and soil volumetric water content (0-10 inches below the soil surface) as a function of time after fertigation from 12:10 pm to 5:10 pm on 06/20/08 (PDT).
Figure B.3: N₂O flux rate, soil temperature, and soil volumetric water content (0-10 inches below the soil surface) as a function of time after fertigation from 9:45 pm to 3:15 pm on 06/21/08 (PDT).

Figure B.4: N₂O flux rate, soil temperature, and soil volumetric water content (0-10 inches below the soil surface) as a function of time after fertigation from 9:45 am to 2:30 pm on 07/22/08 (PDT).
APPENDIX C:
ADDITIONAL NH₃ RESULTS
C.1 Application Rates Calculations:

N application rate (12 deg hr\(^{-1}\) sprinkler arm rotation rate):
\[
\frac{\text{lbsN}}{\text{acre} \times \text{rotation}} \times \frac{1 \text{ rotation}}{360 \text{ deg}} \times \frac{6 \text{ deg}}{1 \text{ min}} \times \frac{1 \text{ min}}{60 \text{ s}} \times \frac{453.6 \text{ g}}{1 \text{ lb}} \times \frac{1}{10 \text{ m} \times 400 \text{ m}} = 7.8 \times 10^{-2} \text{ gN m}^{-2} \text{s}^{-1}
\]

N application rate (18 deg hr\(^{-1}\) sprinkler arm rotation rate):
\[
\frac{\text{lbsN}}{\text{acre} \times \text{rotation}} \times \frac{1 \text{ rotation}}{360 \text{ deg}} \times \frac{9 \text{ deg}}{1 \text{ min}} \times \frac{1 \text{ min}}{60 \text{ s}} \times \frac{453.6 \text{ g}}{1 \text{ lb}} \times \frac{1}{10 \text{ m} \times 400 \text{ m}} = 1.17 \times 10^{-1} \text{ gN m}^{-2} \text{s}^{-1}
\]

NH\(_3\) application rate by sprinkler arm (400 m by 10 m) (12 deg hr\(^{-1}\) sprinkler arm rotation rate):
\[
\frac{\text{lbsN}}{\text{acre} \times \text{rotation}} \times \frac{1 \text{ rotation}}{360 \text{ deg}} \times \frac{6 \text{ deg}}{1 \text{ min}} \times \frac{1 \text{ min}}{60 \text{ s}} \times \frac{453.6 \text{ g}}{1 \text{ lb}} \times \frac{1}{10 \text{ m} \times 400 \text{ m}} \times \frac{17 \text{ g/molNH}_3}{14 \text{ g/molN}} = 9.5 \times 10^{-2} \text{ gNH}_3 \text{ m}^{-2} \text{s}^{-1}
\]

NH\(_3\) application rate by sprinkler arm (400 m by 10 m) (18 deg hr\(^{-1}\) sprinkler arm rotation rate):
\[
\frac{\text{lbsN}}{\text{acre} \times \text{rotation}} \times \frac{1 \text{ rotation}}{360 \text{ deg}} \times \frac{9 \text{ deg}}{1 \text{ min}} \times \frac{1 \text{ min}}{60 \text{ s}} \times \frac{453.6 \text{ g}}{1 \text{ lb}} \times \frac{1}{10 \text{ m} \times 400 \text{ m}} \times \frac{17 \text{ g/molNH}_3}{14 \text{ g/molN}} = 1.425 \times 10^{-1} \text{ gNH}_3 \text{ m}^{-2} \text{s}^{-1}
\]
C.2 Additional NH₃ Concentrations Results

Figure C.1: Time series plot of measured 30-minute average NH₃ concentration, wind speed, wind direction, air temperature, and soil surface temperature from 5 pm on 07/08/07 to 6 pm on 07/09/07.

4:30 pm to 6 pm, 07/09/07

Figure C.2: Polar plot of NH₃ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 4:30 pm to 6 pm 07/09/07 (PDT).
Figure C.3: Time series plot of measured 30-minute average NH₃ concentration, wind speed, wind direction, air temperature, and soil surface temperature from 11 am on 07/11/07 to 3 pm 07/12/07.

Figure C.4: Polar plot of NH₃ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 11 am to 7:30 pm 07/11/07 (PDT).
Figure C.5: Polar plot of NH$_3$ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 9 am to 3 pm on 07/12/07 (PDT).

Figure C.6: Time series plot of measured 30-minute average NH$_3$ concentration, wind speed, wind direction, air temperature, and soil surface temperature from 12 pm on 07/18/07 to 4 pm on 07/19/07.
Figure C.7: Polar plot of NH$_3$ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 12 pm to 5:30 pm on 07/18/07 (PDT).

Figure C.8: Polar plot of NH$_3$ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 12 pm to 4 pm on 07/19/07 (PDT).
Figure C.9: Time series plot of measured 30-minute average NH₃ concentration, wind speed, wind direction, air temperature, and soil surface temperature from 11 am on 07/23/08 to 1 pm on 07/24/08.

Figure C.10: Polar plot of NH₃ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 11 am to 4 pm on 07/23/08 (PDT).
Figure C.11: Polar plot of NH$_3$ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 4:30 pm to 11:30 pm on 07/23/08 (PDT).

Figure C.12: Polar plot of NH$_3$ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 11:30 pm on 7/23/08 to 2 am on 07/24/08 (PDT).

Figure C.12:  Polar plot of NH$_3$ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 4:30 pm to 11:30 pm on 07/23/08 (PDT).
Figure C.13: Polar plot of NH₃ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 2:30 am to 5 am on 07/24/08 (PDT).

Figure C.14: Polar plot of NH₃ concentration versus sprinkler arm rotation angle (red circles) and wind direction (blue squares) for 7 am to 1 pm on 07/24/08 (PDT).
Figure C.15: Time series plot of measured 30-minute average NH$_3$ concentration, wind speed, wind direction, air temperature, and soil surface temperature from 10 am on 07/28/08 to 2:30 pm on 07/29/08.
C.3 Additional NH₃ Flux Rates Results

Figure C.16: Wind speed vs. wind direction sigma (given in 30-min averages) for all days of NH₃ measurement. Low wind speeds are associated with large variation in wind direction. Periods with large variation in wind direction (greater than 30 deg) and low wind speeds (less than 1 m s⁻¹) should not be included in determining NH₃ flux rates since the theoretical basis of the Gaussian plume dispersion model becomes invalid.

Figure C.17: NH₃ concentrations (ppbv) normalized by wind speed (m s⁻¹) as a function of wind speed (m s⁻¹).
Figure C.18: NH$_3$ concentrations normalized by wind speed (m s$^{-1}$) as a function of wind direction (deg) for fertigation periods when the DOAS measurement path was in the SE quadrant of the potato fields.

Figure C.19: NH$_3$ concentrations normalized by wind speed (m s$^{-1}$) as a function of wind direction (deg) for fertigation periods when the DOAS measurement path was in the NE quadrant of the potato fields.
Figure C.20: NH₃ concentrations normalized by wind speed (m s⁻¹) as a function of wind direction (deg) for periods of no irrigation or fertigation.

Figure C.21: NH₃ concentrations normalized by wind speed (m s⁻¹) as a function of soil surface and air temperature (°C) for fertigation periods.
Figure C.22: NH₃ flux rates (µg m⁻² s⁻¹) as a function of soil surface and air temperature (°C) for fertigation periods.

Figure C.23: NH₃ concentrations normalized by wind speed (m s⁻¹) as a function of soil surface and air temperature (°C) for periods of no irrigation or fertigation.
Figure C.24: NH$_3$ flux rates (µg m$^{-2}$ s$^{-1}$) as a function of soil surface and air temperature (°C) for periods of no irrigation or fertigation.

Figure C.25: NH$_3$ flux rates (µg m$^{-2}$ s$^{-1}$) as a function of time of day for fertigation periods.
Figure C.26: Temperature (°C) as a function of time of day (PDT) for periods of no irrigation or fertigation

Figure C.27: Adjusted and non-adjusted NH₃ flux rates (µg m⁻² s⁻¹) as a function of the angle between the sprinkler arm and wind direction (deg).
Figure C.28: Angle between sprinkler arm and wind direction (deg) as a function of wind speed (m s\(^{-1}\)).