EVALUATING FROZEN BEEF AND MEAT PACKAGING MATERIAL EXPOSED TO 
LOW LEVELS OF AMMONIA GAS

by

FARIS A. KARIM HUSSAIN

B.S., Al-Mustansiriya University, Iraq, 1991

A THESIS
Submitted in partial fulfillment of the requirements for the degree

MASTER OF SCIENCE

Food Science

KANSAS STATE UNIVERSITY
Manhattan, Kansas

2009

Approved by
Major Professor
J. Scott Smith
Animal Sci & Ind
ABSTRACT

Ammonia leaks in meat chilling/frozen storage or processing facilities are not uncommon. Often the meat products are packaged in polymer films that theoretically protect the product from contamination. Unfortunately, there is almost no data on whether ammonia can permeate packaging films. The objectives of this study were to evaluated meat contaminated by low levels of ammonia under frozen storage conditions (-17 ± 3 °C), and further evaluated the permeability of common meat packaging films including: low density polyethylene (LDPE), 3 mil Cryovac (E-2300), and 3 mil vacuum (V-PA/PE) at freezing (-17 ± 3 °C) and room temperatures (21 ± 3 °C).

Fresh beef Semitendinosus muscles were fabricated to form 10 x 5 x 2.3 cm steaks. The packaging films were fabricated into 10 x 5 cm pouches and filled with 50 mL deionized water. The meat and the pouches were exposed in a plexiglass enclosure, contained in a freezer, to 50, 100, 250, and 500 ppm ammonia gas (85 mL/min) for exposure times of 6, 12, 24, and 48 hr. The ammonia levels in the meat samples were 34.2 (50 ppm exposure), 51.5 (100 ppm exp.), 81.1 (250 ppm exp.), and 116 ppm (500 ppm exp.), and the pH values ranged from 5.56 to 5.75 (control ranged from 5.31 to 5.43) at 48 hr. At freezing temperatures, ammonia residues remained undetected, and no differences in pH were found in the pouches. At room temperature, all pouches were slightly permeable to ammonia; the levels observed in the pouches were, 7.77 ppm (pH, 8.64) for E-2300, 5.94 ppm (pH, 8.38) for LDPE, and 0.89 ppm (pH, 7.23) for V-PA/PE at 500 ppm exposure for 48 hr (unexposed samples pH ranged from 5.49 to 6.44).

The results showed that meat packaging materials have low ammonia permeability and thus protect meat products exposed to ammonia exposure during frozen storage. Moreover, meat content is low even with ammonia exposures as high as 500 ppm for up to 48 hr.
# Table of Contents

LIST OF FIGURES ........................................................................................................................................ vi

LIST OF TABLES ......................................................................................................................................... viii

ACKNOWLEDGMENTS ................................................................................................................................ ix

PART I: REVIEW OF LITERATURE ........................................................................................................ 1

INTRODUCTION ........................................................................................................................................ 1

CHEMICAL AND PHYSICAL PROPERTIES OF AMMONIA ................................................................. 5

AMMONIA PRODUCTION, USE, AND DISPOSAL ............................................................................... 9

AMMONIA TOXICITY ............................................................................................................................ 11
  Oral Exposure ....................................................................................................................................... 13
  Toxic Compounds Resulting from Ammoniation .............................................................................. 14

AMMONIA FOOD CONTAMINATION ............................................................................................... 15
  Wisconsin milk contamination 1985 .............................................................................................. 15
  Illinois 2002 ................................................................................................................................... 16

ECONOMIC EFFECTS OF AMMONIA FOOD CONTAMINATION .................................................... 16

AMMONIA REGULATIONS FOR FOOD AND FOOD PRODUCTS ...................................................... 18

EFFECTS OF AMMONIA ON MEAT QUALITY AND PROPERTIES ................................................ 20
  Meat Color .......................................................................................................................................... 20
  Meat Flavor and Odor ......................................................................................................................... 23
  Meat pH ............................................................................................................................................ 24
  Meat Tenderness ............................................................................................................................... 26
  Meat Water Holding Capacity ........................................................................................................... 27

MEAT PACKAGING FUNCTIONS ......................................................................................................... 28

ENVIRONMENTAL EFFECTS ON PACKAGING PERMEABILITY .................................................... 28
  Temperature ......................................................................................................................................... 28
LIST OF FIGURES

FIGURE PAGE
1. Chemical structure of 4-methylimidazole and subsequent interchangeable rearrangement ................................................................. 15
2. The relationship among the three pigments color in the meat .................. 22
3. The change in total pH of Semitendinosus steaks over time when exposed to different concentrations of ammonia ........................................... 26
4. Chemical structure of Nylon 6 and Nylon 11 ........................................ 31
5. Branched polymer polyethylene ............................................................... 32
6. Polypropylene types: (a) Ilostactic, (b) Syndiotactic, and (c) Atactic .......... 34
7. A general flow chart for the experiment process of the meat packaging materials ...... 49
8. Fabricated plastic films Cryovac, low density polyethylene, and vacuum with dimensions of 10 × 5 cm and filled with 50 mL deionized water .............................................................. 50
9. Exposure chamber used in the experiment with two L-shape fittings mounted on the back and 15 cm round front door ................................................................. 51
10. Water filled fabricated pouches inside the exposure chamber inside the freezer suspended on two plexiglass bars ............................................................... 52
11. The exposure system components .......................................................... 54
12. Tubing connection between the system parts ........................................... 55
13. Chamber flushing process and ammonia gas directions in the system .......... 56
14. Movement directions of ammonia gas during exposure .......................... 56
15. Water pH levels for Cryovac E-2300, LDPE, and vacuum (V-PA/PE) bags at 500 ppm exposure for 6, 12, 24, and 48 hr at room temperature ................................. 60
16. Water ammonia level means for Cryovac E-2300, LDPE, and vacuum (V-PA/PE) bags at 500 ppm for 6, 12, 24, and 48 hr exposure at room temperature ......................... 64
17. A general flow chart for processing and exposure of the steaks to ammonia .......... 75
18. Beef Semitendinosus muscle trimmed of external fat and fabricated to dimensions of 10 × 5 × 2.3 cm .......................................................................................................................... 76
19. Meat samples attached to stainless steel hooks and suspended on two plexiglass bars inside the freezer ................................................................. 77
20. Meat pH mean levels as a different from control at 50, 100, 250, and 500 ppm exposure and 6, 12, 24, and 48 hr at -17 ± 3 °C ................................................................. 83

21. Levels of meat ammonia as a different from control at 50, 100, 250, and 500 ppm exposure for 6, 12, 24, and 48 hr at -17 ± 3 °C ................................................................. 85
# LIST OF TABLES

<table>
<thead>
<tr>
<th>TABLE</th>
<th>PAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. List of some chemical and physical properties of ammonia</td>
<td>8</td>
</tr>
<tr>
<td>2. Human physiological response to ammonia</td>
<td>12</td>
</tr>
<tr>
<td>3. Ammonia and its salt regulations in food, processed food, and animal feed</td>
<td>19</td>
</tr>
<tr>
<td>4. Basic properties of various polyethylene films</td>
<td>33</td>
</tr>
<tr>
<td>5. Pouches exposure levels of ammonia measured with the MinRae 2000</td>
<td>57</td>
</tr>
<tr>
<td>6. Cryovac E-2300 bag’s water pH levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature</td>
<td>61</td>
</tr>
<tr>
<td>7. Low density polyethylene bag’s water pH levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48 hr exposure at room temperature</td>
<td>62</td>
</tr>
<tr>
<td>8. Vacuum bag’s water ammonia levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature</td>
<td>62</td>
</tr>
<tr>
<td>9. Vacuum bag’s water pH levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature</td>
<td>63</td>
</tr>
<tr>
<td>10. Cryovac E-2300 bag’s water ammonia levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature</td>
<td>65</td>
</tr>
<tr>
<td>11. Ammonia levels in low density polyethylene bag’s water at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature</td>
<td>65</td>
</tr>
<tr>
<td>12. Physical properties of selected packaging materials used to evaluate ammonia permeability provided by suppliers</td>
<td>67</td>
</tr>
<tr>
<td>13. Meat pH levels at 50, 100, 250, and 500 ppm exposure for 6, 12, 24, and 48 hr at -17 ± 3 °C</td>
<td>81</td>
</tr>
<tr>
<td>14. Meat ammonia levels exposed at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48 hr at -17 ± 3 °C</td>
<td>85</td>
</tr>
</tbody>
</table>
ACKNOWLEDGMENTS

I would like to express my heartfelt gratitude to my advisor Dr. J. Scott Smith for his friendship, patience, and guidance through my degree program. I would also like to thank his family for their support and friendship. I also extend thanks to my committee members, Dr. Curtis L. Kastner and Dr. Tomas Herald for their suggestions, recommendations, encouragement, and advice.

I wish to acknowledge my special thank and gratefulness to my father Abdul-Karim for encouraging my curiosity and instilling the love of science, reading, and discovery.

My deepest thanks and appreciations to my mother Sahira who covered me with love and care during my life. I deeply appreciate my brother Yasser who supported me during my studies and to my brother Ali and my sister Zahraa for their love and care. I am indebted to my cousin Sinna Mahdi and her husband Dr. Imad Khamis and their family for their support.

Finally, I would like to thank Kansas State University and its Department of Animal Sciences and Industries for giving me the opportunity to pursue my master’s degree.
INTRODUCTION

Food safety and quality are goals of food science, food industry, and regulatory agencies, and are a concern for consumer and public health organizations. Because food safety is important to human health and food quality is important to consumers, the relationship between food safety and food quality must be established and clarified. This would benefit not only the food industry but also would facilitate decisions about food-related issues such as food contamination, and help in evaluating critical control points and corrective actions in the hazard analysis and critical control program (HACCP). Moreover, this information could be used in insurance/customer claims involving food contamination (Al-Sahal 2003).

Food can have good quality but may still be unsafe. Many factors affect food quality and safety, including temperature, oxygen, light, and chemicals. Temperature is the most important in food value and acceptance. Temperature directly affects microbial growth, microbial contamination, and chemical reactions, especially enzymes, as well as affecting the physical properties of food. One of the oldest methods of preserving perishable food is freezing. Most perishable foods are stored in a cold storage warehouses using a refrigerant like ammonia for larger cold storage units or halocarbons for smaller or mid-size ones. Halocarbons like carbon dioxide, sulfur dioxide, and methyl chloride have many advantages that make them good refrigerants; they are non-toxic, non-flammable, and non-reactive. Conversely, their low evaporation enthalpy, poor heating transfer, excessive pressure drop, and most importantly, depleting the ozone layer, have made the refrigeration industry consider ammonia as an alterative refrigerant. Ammonia has important advantages as a refrigerant: high efficiency in
providing cooling and freezing for food products (Holmstrom 1994), its relative low environmental impact (Ross 1994), and its low production cost (Arnold 1993), with no impact on the ozone layer. Ammonia is the refrigerant of choice for many food products.

Today, refrigeration with ammonia is the backbone of the food industry. It cools fruits, vegetable, poultry, fish, beverages, dairy products, milk, and meat (Lorentzen 1988). Meat production and processing require a large area for fabrication, handling, and storage. Most fabrication facilities, slaughterhouses, and cold storage warehouses use ammonia as their refrigerant (Arnold 1993; Ross 1994; Ross 1995; Sun 1998).

The disadvantages of using ammonia as a refrigerant include its toxicity and its flammability. Ammonia is very explosive when mixed approximately one volume of ammonia to two volumes of air, and much more so when mixed with oxygen (EPA 1995). Moreover, ammonia itself can contaminate food products (Arnold 1993). Refrigerated warehouses using ammonia as a refrigerant are subject to ammonia leaks due to typical equipment failures and operator error; ammonia leaks are usually just a matter of time in a refrigerated/frozen food storage facility. Leaks may be due to carelessness, but more often, they are due to equipment issues (Ostner 1986).

High concentrations of ammonia affect many quality properties of meat, such as color (Curda and Hruby 1987; Bonne et al. 1993), increased pH (Kassem and Johnston 1965; Anil 1971; Ireland 1988; Al-sahal 1995), increased water holding capacity (Anil 1971; Al-sahal 1995), persistent pink color after cooking (Smolskiy et al. 1985; Shaw et al. 1992), and altered flavor (Ireland 1998; Hagyard et al. 1993; Guerrero and Arnau 1995). Ting and Henrickson (1986) studied the effect of ammonia on meat tenderness, flavor, taste, water holding capacity, and pH. The pH value is not a reliable indicator of the amount of the ammonia absorbed because
of the different buffering capacities of different foods (Kassem 1965). Furthermore, the odor of ammonia from contaminated meat is not a good indicator of concentration level in contaminated meat, because ammonia reacts with water and ice to form ammonium ions, which are odorless (Dworkin et al. 2004).

Regulations and published information on food contaminated with ammonia is limited, so contaminated products are held for an indeterminate period or condemned. The scientific literature offers little guidance on evaluating product quality after exposure to ammonia (Goodfellow et al. 1978). Some research has established the effects of high levels of ammonia concentrations on meat quality, but the effects of low levels of ammonia on contaminated food has not been covered, and the effect of time on contamination remains undetermined (Al-Sahal 2003).

In addition to refrigeration, most perishable foods currently are packed in plastic films that protect food against external/internal influences that cause deterioration. In the 19th century, pioneers like Nicholas Appert, Louis Pasteur, Samuel C. Prescott, and William L. Underwood discovered the basic food packaging and preservation concepts that remain relevant today (Wilson 2007). In the 20th century, inventions like glass bottles, cellophane wrap, aluminum foil, and plastics gave packagers more flexibility (Lord 2008). Plastic packaging material is functional, recyclable, lightweight, and widely used in food industry, so plastic packaging has been increased markedly over the past 40 years. In 1991, plastics account for up to 50% of the primary food packaging in the food industry, and polymers food wraps represent $211 million in sales (Brewer and Harbers 1991).

Packaging encloses product using bags, pouches, cans, tubes, trays, or wraps. Packaging follows the steps needed to contain and assemble food into product units for storage,
distribution, sale, and cooking. The main goal of packaging is to protect the product, increase the shelf life, and maintain the product’s original conditions.

Packaging protects contents from contamination and spoilage, makes it easier to transport food, store goods, in addition to providing a uniform measure of the contents (Hine 1995). Protecting and preserving food is the principal role of food packaging (Robertson 2006).

Beardsell (1961) noted that packages should withstand the following conditions: substantial change in temperature; humidity; vapor pressure; vibration through transportation; low temperature and high air velocities in blast freezers; condensation from changes in temperature and humidity; and weight of stacking under unfavorable conditions.

Package integrity and barrier characteristics are of concern in many areas of food industry. Their importance in preserving commercial sterilized food in sealed packages is well understood (Philips 1985; Denny 1989; Harper et al. 1995); this protection can be achieved with single layer of polymer or multilayered films including different polymers.

Barrier materials are materials that possess low permeability to gases, vapors, and liquid. Under this category, different types of barrier materials can be included: paper, plastic film sheeting, fabrics and metallic foils, particle board, and wood laminates. In food packaging, plastic materials are the most important, especially polyethylene, ethylene-vinyl alcohol copolymers, polyvinylidene chloride, nylons, acrylic latex, high nitrile resins.

Gas permeability (transmission rate) and water vapor transmission rate (moisture transmission rate) are the most important characteristics of barrier materials in food packaging, but meat packaging requires special packaging materials because of special concerns during storage.

Packaging films permeability depends on, among other things, molecular size and shape,
wetability, and the texture of the fabricated materials. Moreover, because permeability is a molecular transport phenomenon, it is affected by the molecular orientation, degree of crystallinity, and temperature. Kassem (1965) concluded that for ammonia vapor, the most permeable packaging material was waxed paper, while Cryovac vacuum packaging was the least permeable.

Package protection depends mainly on the package’s permeability to gases and vapors that are harmful to the quality of the food products, and this permeability could be affected by polymer properties, gas barrier properties of these polymers, and other properties that affect permeability, which includes the size and the shape of the permeant molecule, polarity, and the permeant temperature and pressure (Jasse et al. 1994). Even increasing the oxygen permeability because of flex-cracking of the polymer used in all kinds of the packaging during transportation may adversely affect food shelf-life (Mannheim and Miltz 1987).

Although meat contamination with high levels of ammonia has been investigated extensively, there is almost no data available on meat contamination or whether ammonia can permeate packaging films at levels below 500 ppm. Our research objectives were to measure the levels of ammonia in exposed meat at freezing temperatures and evaluate the permeability of three types of meat packaging materials at freezing and at room temperature after low levels of ammonia exposure and different exposure times.

CHEMICAL AND PHYSICAL PROPERTIES OF AMMONIA

The ammonia molecule has a trigonal pyramid configuration with bond angles of 107°. The pyramid structure consists of tri-hydrogen atoms and a nitrogen atom resonating between equal stable positions at a distance of 0.360 Å above and below the three hydrogen atoms’ plane.
This H-N-H bond angle of $107^\circ$ is close to the tetrahedral angle of $109.5^\circ$. This electronic arrangement of valence electrons in nitrogen is sp$^3$ hybridized atomic orbitals (Jones 1997).

Ammonia is a colorless gas under standard conditions with a mass of 17.031 g/mol, a boiling point of $-33.3 \, ^\circ C$, a freezing point of $-77.7 \, ^\circ C$, and a liquid density of 681.9 kg/m$^3$ at $-33.3 \, ^\circ C$ and one atmosphere. Ammonia vapor density is 0.5967 (air = 1) (Table 1). Ammonia gas is easily detectable by smell; most people can distinguish ammonia at 35 ppm and higher. Trained people can detect ammonia at 5 ppm or even less (Raj 1982). Ammonia boils readily if it is spilled and vaporizes, cooling its surroundings.

Approximately 1 ppm of ammonia (1 mg/liter) is equal to 0.70 mg/m$^3$; however, this value can vary according to the ambient atmosphere pressure and temperature. In liquid form, ammonia is 60% of the weight of the water and as gas, is lighter than air at 25 $^\circ C$ and 1 bar (Ostner 1986). Most commercially available ammonia forms are aqueous solutions. The most common commercial formulation is 28-30% NH$_3$ (Weast et al. 1988); at this concentration, ammonia forms a nearly saturated solution in water.

Ammonia possesses a high affinity to water and dissolves readily in water or any product containing water, forming an alkaline solution (aqua ammonia or ammonium hydroxide). Ammonia in solution is usually called ammonium hydroxide and has been historically called “spirit of hawthorn” (Windholz et al. 2001). Ammonium hydroxide acts as a weak base and is partially ionized to ammonium (NH$_4^+$) and hydroxyl (HO$^-$). This behavior refers to the polarity of ammonia molecules and their ability to form hydrogen bonds. Furthermore, in aqueous solution, ammonia acts as a base, because the nitrogen atom has a pair of unshared electrons that can react with the H$_2$O hydrogen ions, producing ammonium and hydroxide ions; the alkalinity for the aqueous solution of ammonia relates to the hydroxide ions. The alkalinity of ammonia is
related to the availability of this unshared pair of electrons (Anil 1971; Mcmurray and Fay 1995)

\[
\text{NH}_3 (aq) + \text{H}_2\text{O} (l) \rightleftharpoons \text{NH}_4^+ (aq) + \text{OH}^- (aq)
\]

\[
\downarrow
\]

\[
\text{H}^+ + :\text{NH}_3 \rightleftharpoons \text{H}:\text{NH}_3^+
\]

The acid disassociation constant, \(K_b\), is \(1.774 \times 10^5\) at 25 °C (\(pK_b\) is 4.751) and increases slightly with increasing temperature (Weast et al. 1988). The equilibrium between the un-ionized form of ammonia (\(\text{NH}_3\)) and the ionized (\(\text{NH}_4^+\)) is directly affected by pH. At pH 7.25, 99% of the ammonia will be ionized in ammonium form (\(\text{NH}_4^+\)), while at pH 8.25, 90% of the ammonia will be ionized; at pH 9.25, 50% of the ammonia will be ionized, with 50% un-ionized. At pH 11.2, 99% of the ammonia will be un-ionized; therefore, at most environmentally substantial pHs, most ammonia will be ionized, and as the pH rises from pH 7, the proportion of un-ionized ammonia increases. Not all dissolved ammonia reacts with water to form ammonium; some ammonia remains in molecular form in solution (Mcmurray and Fay 1995).

Temperature and pH affect ammonia solubility and volatility. Ammonia solubility in water increases with decreasing the pH, while volatility increases with increasing pH and temperature (ATSDR 2004). Therefore at high pH solutions, the ammonia volatilizes freely. Ammonium salt such as chloride and sulfate are strongly dissociated and very soluble in water (Weast et al. 1988). As a result, changes in pH will not necessarily result in ammonium precipitates.
Table 1: List of some chemical and physical properties of ammonia\(^1\)

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical formula</td>
<td>(\text{NH}_3)</td>
<td></td>
</tr>
<tr>
<td>Molecular weight</td>
<td>17.03</td>
<td>LeBlanc et al. 1978</td>
</tr>
<tr>
<td>Color</td>
<td>Colorless</td>
<td>LeBlanc et al. 1978</td>
</tr>
<tr>
<td>Physical state</td>
<td>Gas at room temp.</td>
<td>LeBlanc et al. 1978</td>
</tr>
<tr>
<td>Melting point</td>
<td>-77.71 °C</td>
<td>LeBlanc et al. 1978</td>
</tr>
<tr>
<td>Boiling point</td>
<td>-33.35 °C</td>
<td>LeBlanc et al. 1978</td>
</tr>
<tr>
<td>Density:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gas</td>
<td>0.7710 g/L</td>
<td>Weast et al. 1988</td>
</tr>
<tr>
<td>Liquid</td>
<td>0.6818 g/L (-33.35 °C; 1 atm)</td>
<td>Windholz 1983</td>
</tr>
<tr>
<td>Vapor density</td>
<td>0.5967 (air=1)</td>
<td>Windholz 1983</td>
</tr>
<tr>
<td>Specific gravity (25 °C)</td>
<td>0.747 g/L</td>
<td>Lide 1998</td>
</tr>
<tr>
<td>Odor</td>
<td>Sharp, Intensely irritating</td>
<td>Sax &amp; Lewis 1987</td>
</tr>
<tr>
<td>Odor threshold in air</td>
<td>48 ppm (34 mg/m(^3))</td>
<td>Leonardsos et al. 1969</td>
</tr>
<tr>
<td>Odor threshold in water</td>
<td>1.5 ppm</td>
<td>Amoore 1983</td>
</tr>
<tr>
<td>(pK_a)</td>
<td>9.26 (25 °C)</td>
<td>Lide 1998</td>
</tr>
<tr>
<td>Solubility in water at 0 °C</td>
<td>47.8% (w/w)</td>
<td>Budavari et al. 1996</td>
</tr>
<tr>
<td>Solubility in water at 25 °C</td>
<td>34% (w/w)</td>
<td>LeBlanc et al. 1978</td>
</tr>
<tr>
<td>Vapor pressure:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anhydrous (\text{NH}_3)</td>
<td>8.5 atm (20 °C)</td>
<td>Sax &amp; Lewis 1987</td>
</tr>
<tr>
<td>Autoignition Temperature</td>
<td>650 °C</td>
<td>LeBlanc et al. 1978</td>
</tr>
<tr>
<td>Flammability limits in air</td>
<td>16-25%</td>
<td>LeBlanc et al. 1978</td>
</tr>
<tr>
<td>(pH) in water</td>
<td>11.6 (1 N)</td>
<td>Windholz 2001</td>
</tr>
</tbody>
</table>

\(^1\)Adapted from: ATSDR (2004)
AMMONIA PRODUCTION, USE, AND DISPOSAL

Ammonia is both a natural and manufactured chemical compound. Ammonia is an essential product in the nitrogen cycle. In nature, the major source of the ammonia is microbial production. Most nitrogen fixation in nature to usable forms (e.g., NH₃) occurs because of lightning strikes; microbial nitrogen fixation at the beginning of the 20th century has been estimated at 90-130 teragrams (Tg; 1 teragram = one million metric tons) per year. Human production also fixes nitrogen in useful forms. In 1999, human production of fixed nitrogen (NH₃) is estimated to be 140 Tg N per year, an amount that is similar to non-anthropogenic source (NSF 1999; Socolow 1999).

Ammonia from decomposing animal excreta probably accounts for most of the ammonia produced, with decaying organic materials from plants, dead animals, and the like contributing significant amount (Dawson 1977; Crutzen 1983; Dawson and Farmer 1984; Galbally 1985; Irwin and Williams 1988).

However, production of ammonia in the United States has declined over the past several years. Increasing natural gas prices and weather related decreases in demands and closure of several production plants have caused this decline. In 1999, the U.S. annual commercial production capacity for ammonia was 16.6 million metric tons (CMR 1999), 15.7 million metric tons in 2000 (SRI 2000), and 9.5 million metric tons in 2001 (Kramer 2002). In 2002, ammonia production levels increased slightly to reach 10.8 million metric tons (Kramer 2003) because natural gas prices went down.

Ammonia can be produced commercially in several ways. The modified Harber-Bosch process is the usual commercial method for producing ammonia. This process was first demonstrated in 1909 (Kramer 2000) and was commercially developed in Germany. The first
U.S. plant to use this process was built in Syracuse, New York, in 1921 (DOI 1985). In 1979, 98% of the produced ammonia in the United State was produced by Harber-Bosch process (EPA 1980; HSDB 2003). In this process, hydrogen (from the natural gas) and nitrogen (from the atmosphere) are mixed at a ratio of 1 to 3 over a catalyst at high temperature and high pressure. Any remaining unreacted gases are recirculated through the reactor and the ammonia is collected. Ammonia may be produced as a by-product of coking coal, but the amount produced is small compared to the amount produced by the modified Harber-Bosch process. Because natural gas is used to produce ammonia, most industrial ammonia production occurs in areas where natural gas is cheap and accessible.

The main commercial use of ammonia and ammonium compounds is in agricultural fertilizers, representing 89-90% of commercially produced ammonia. Plastics, synthetic fibers and resins, explosives, and other uses account for the rest (Kramer 2002, 2004). Kramer (2003) categorized the direct use of ammonia as fertilizer based on the mass of nitrogen percentage in each compound as follows: anhydrous ammonia, 26%, urea/ammonium nitrate solutions, 23%; urea, 20%; ammonium nitrate, 4.5%; ammonium sulfate, 2%; other forms, 3%; and multiple nutrient forms, 21%. Most ammonium compounds and nitric acid are used directly in the fertilizer production, usually to produce anhydrous ammonia.

Ammonia has a wide application in other industries as well. Ammonia is used as a household cleaner, to inhibit corrosion, as a refrigerant, to purify water supplies, to make pulp and paper, in metallurgy, in the food and beverage industry, and to make rubber. Ammonia is used to manufacture pharmaceuticals and explosives and to produce various chemical intermediates (LeBlanc et al. 1978; Sax and Lewis 1987). Ammonia may be released into air or water as a gas or liquid, as regulated by federal, state, local governments. Because ammonia has
a high affinity to water, disposed ammonia may be diluted with water and neutralized with HCl before discharging the chemical into surface waters. However, this generates a lot of heat, so this method of disposing of ammonia is not desirable because of the related danger to personnel involved in the process. Recovering ammonia from aqueous waste solutions is, however, a viable option for many industries (HSDB 2003).

**AMMONIA TOXICITY**

Ammonia is strong local irritant and has a pungent odor. Ammonia is a corrosive because of the alkalinity of ammonium hydroxide; when the chemical forms on mucous membranes, it dissolves the cellular protein and causes corrosion. Ammonia primarily affects the respiratory system and may cause severe damage, possibly leading to death. In general, the following symptoms can be observed: pharyngitis, laryngitis, tracheobronchitis, nausea, vomiting, increases salivation, and reflectoric bardycardia; edema of the glottis, laryngospasm, bronchospasm, and interstitial lung edema are life-threatening symptoms (Coplin 1949). The degree of ammonia toxicity and hazard to the human life depends on the concentration of ammonia, length of exposure, and mechanism of exposure (Lessenger 1985). Table 2 illustrates human physiological response to different levels of ammonia exposure. The general cause for ammonia irritation in humans comes from the alkalinity when ammonia dissolves in body fluids. The respiratory system, eyes, and skin are more susceptible to ammonia than other parts of the body. Death by suffocation can be caused by breathing air containing 5000 ppm of ammonia. Exposure to ammonia at 2000 ppm for a few seconds may result in skin burns and blistering and may lead to acute lung edema. Moreover, exposure to ammonia concentrations higher than 700 ppm will cause eye injure that may cause loss of sight (Slack and James 1973; WHO 1986). For
most people, a moderate degree of eye, throat, and nose irritation can result from a short exposure to 50 to 100 ppm (Swotinsky and Chase 1990).

Table 2: Human physiological response to ammonia (Adopted from Al-Sahal 2003)

<table>
<thead>
<tr>
<th>Physiological response</th>
<th>Ammonia levels (ppm)</th>
<th>Observations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Odor detectable</td>
<td>1-5</td>
<td>Trained or most sensitive people</td>
</tr>
<tr>
<td>Mildly-distinct but not annoying</td>
<td>10-20</td>
<td>Average people</td>
</tr>
<tr>
<td>Recognizable for all human</td>
<td>35</td>
<td>Least sensitive people</td>
</tr>
<tr>
<td>OSHA¹ &amp; NIOSH² limits</td>
<td>50</td>
<td>Exposure on time-weight average (TWA)</td>
</tr>
<tr>
<td>Very noticeable</td>
<td>100-200</td>
<td>Can be tolerated for several hours</td>
</tr>
<tr>
<td>Eye and throat irritation</td>
<td>400-500</td>
<td>Very annoying to throat, nose, and eyes</td>
</tr>
<tr>
<td>Irritation and eye injury</td>
<td>700-1000</td>
<td>Toxic, accessible with cartridge type gas mask but for shot time</td>
</tr>
<tr>
<td>Permanent injury, coughing</td>
<td>1500-2000</td>
<td>Difficulties in breathing laryngospasm</td>
</tr>
<tr>
<td>Could be fatal</td>
<td>2500-4500</td>
<td>Threat to life after 30 min exposure</td>
</tr>
<tr>
<td>Rapidly fatal for short time</td>
<td>≥5000</td>
<td>Death rapidly result from respiratory arrest</td>
</tr>
</tbody>
</table>

¹OSHA = The U.S. Occupational Safety and Health Administration
²NIOSH = The U.S. National Institute for Safety and Health.
Oral Exposure

Humans suffer ammonia toxicity by inhalation, dermal contact, and ingestion. In the food industry, oral ingestion of ammonia is a major concern. One ammonia contamination occurred in Wisconsin in 1985, the second incident occurred in Illinois in 2002. In the first incident, children suffered from mouth/throat burn and nausea when children consumed contaminated milk with ammonia (530 to 1,524 ppm; normal 1.4 ppm), while vomiting, nausea, headache, and stomachache were the symptoms among the children who consumed contaminated chicken tenders (138 to 2,468 ppm; normal 75 to 95 ppm) in the second incident. The symptoms in the Illinois incident were more severe because of the high levels of ammonia in the tenders. The literature supplies little information about ammonia oral toxicity; more research is needed in this topic.

The scientific literature provides a few animal studies that document the effects of ammonia contamination. Karplyuk et al. (1989) fed three generations of rats with meat that was contaminated with ammonia levels of 0.1% (1000 ppm) and 0.3% (3000 ppm). At 3000 ppm, each generation suffered from system damage during the first six months of the experiment: liver damage, central nervous system damage, and reduced cholinesterase activity in the blood. At 1000 ppm, the rats still suffered from reduced lactate dehydrogenase and alanineaminotransferase activity.

In another study, ammonium chloride was given to mice, guinea pigs, rats, rabbits, and dogs to study the effects of metabolic acidosis, defined as formation of hydrogen ions from the metabolism of ammonium ions to urea. The ingestion of ammonium chloride in doses of 500 to 1,000 mg/kg/day for 1-8 days induced metabolic acidosis in mice, guinea pigs, rats, rabbits, and dogs (WHO 1986). Metabolic acidosis can cause a variety of changes in neurological,
pulmonary, cardiovascular, gastrointestinal, musculoskeletal function, as well as hematological and clinical chemistry parameters.

**Toxic Compounds Resulting from Ammoniation**

Ammonia can be added to a cattle feed to increase food digestibility, dry matter intake, and the nutritive value of poor quality roughage (Birkelo et al. 1986). Ammoniation is also used to reduce the spoilage in high quality forages baled or stored with high moisture content. Ammoniation prevents heating and subsequent microbial proliferation, fermentation, and mold (Thorlacius and Robertson 1984; Atwal et al. 1986). However, feeding cattle molasses treated with ammonia can cause “bovine bonkers” or “crazy cow syndrome”. Many cases of ammonia intoxication in cattle were reported after cattle were fed treated grass straw or grain straw (Bergstrom 1991). Winggins and Wise (1955) reported pyrazine and 4-methylimidazole (4-MeI) in molasses treated with ammonia while Morgan and Edward (1986) referred to 4-methylimidazole as a cause of the intoxication. Cattle fed ammoniated feed exhibited toxic effects, especially in the central nervous system (Perdok and Leng 1987). Symptom included ear twitching, salivation, stampeding, convulsions, trembling, and death (Morgan and Edward 1986). Nishie et al. (1969, 1970) reported similar effects in mice fed different types of imidazole, 1-methylimidazole, 2-methylimidazole, 4-methylimidazole, and pyrazine derivatives. The toxicity of the pyrazine derivatives was low; most probably, the stronger effects were caused by 4-MeI from the ammoniated feed.

4-methylimidazole is a colorless heterocyclic compound formed from the reaction between ammonia and a reducing sugar with heat (Simeone 1992; Yuan and Burka 1995). Bergstrom (1991) found heating, moisture content, amount of reducing sugar, and amount of the
ammonia were all important in forming 4-MeI (Figure 1). Theoretically, 4-MeI formed in ammoniated grass through the reaction between ammonia and the sugar of the hays or forage during the preservation and heating process. Currently, hay is treated with anhydrous ammonia (35 g/kg dry matter) and heated it to 90 °C for 15 hr during curing (Manson et al. 1989).

\[
\begin{align*}
\text{H}_3\text{C} & \quad \text{N} \\
\text{HC} & \quad \text{N} \\
\text{CH} & \quad \text{CH}
\end{align*}
\]

\[
\begin{align*}
\text{H}_3\text{C} & \quad \text{N} \\
\text{HC} & \quad \text{N} \\
\text{CH} & \quad \text{CH}
\end{align*}
\]

Figure 1: Chemical structure of 4-methylimidazole and subsequent interchangeable rearrangement

AMMONIA FOOD CONTAMINATION

Ammonia food contamination was reported as a food outbreak twice in the United States. The first incident occurred in Wisconsin in 1985, and the second incident occurred in Illinois in 2002.

Wisconsin milk contamination 1985

Two elementary schools in Wisconsin suffered ammonia contamination of milk on October 30, 1985 (CDC 1986). The school children reported burning in the mouth and throat within 1 hr of drinking contaminated milk. The incident occurred when the milk was packed in half-pint containers that were contaminated by a high-pressure ammonia leak in the product cooler. Analysis of the remaining milk in the containers showed an ammonia level of 530 ppm to 1,524 ppm (normal is 1.4 ppm; Hiroshi 2000). The contaminated milk had elevated pH of 9.1
to 10.0 (normal is 6.7-6.9) due to ammonia contamination.

**Illinois 2002**

Several dozen schoolchildren in Illinois suffered from ammonia exposure from chicken tenders on November 25, 2002 (Dworkin et al. 2004). The schoolchildren reported nausea, headache, and stomach ache. The USDA-FSIS laboratory analyzed the remaining contaminated chicken tenders and found a level of 138 ppm to 2,468 ppm of ammonia. It should be noted that typical background levels for uncontaminated chicken tenders (nuggets) for the test used in this evaluation ranged from 75-95 ppm (Hijaz et al. 2007).

**ECONOMIC EFFECTS OF AMMONIA FOOD CONTAMINATION**

Even though ammonia has many advantages as refrigerant, its toxicity remains the major disadvantage. In many cases, ammonia leaks have resulted in multi-million dollars losses (EPA 2001). Arnold (1993) noted what affects the degree of ammonia food contamination: exposure time, temperature, ammonia concentration, product types, and packaging. According to Goodfellow et al (1978), three different methods to determine if ammonia contaminates food is suitable for human consumption: pH values, sensory analysis, and ammonia concentration levels. The food is safe for humans if the pH does not exceed the normal (uncontaminated) pH by more than 1.0, its sensory analysis panel score is 5.0 or higher, and its ammonia level is not more than 1000 ppm (Goodfellow et al. 1978).

Ammonia leaks are not uncommon. In January 1964, a leak occurred in a commercial plant in Knoxville, Tennessee, involving several food products. The amount of ammonia released was estimated at 100 pounds (Kassem 1995). The largest ammonia leak incident
occurred in Florida in 1974 in a commercial cold storage warehouse (Goodfellow et al. 1978). The total value of damaged product was $45,000,000, including 121 types of products in 150,281 cases and 939 lots. In February 1994, another ammonia leak occurred in Olympic Cold Storage in Washington. Fifty eight tons of frozen seafood and vegetables worth around several hundred thousand dollars were exposed in an ammonia spill. All the contaminated foods were condemned because of high levels of ammonia in the analyzed samples and a strong ammonia odor.

Another ammonia leak occurred in Virginia on September 30, 1999, in the S. Wallace Edwards & Sons cold storage facility. The contaminated food product was 70,000 pounds of ham. Because of the public safety concerns, the product was destroyed (USCOA 2003).

Two recalls were issued in 2001 for ammonia contamination. In December 19, 2001, Stewart’s Beverages, Inc., in New York recalled several beverage products because of possible contamination with ammonia. The recalled products involved different beverage flavors in different sizes. All the contaminated products were produced by a third party company, Hillside Bottling in New Jersey (FDA 2008h). Also in December 19, 2001, another product, sparkling soda in 16 oz glass bottles also produced by Hillside Bottling in New Jersey was recalled by Mistic Brands, Inc., because of potential ammonia contamination (FDA 2008i). Obviously, in these cases, the possible ammonia contamination occurred at Hillside Bottling.

Recently on November 17, 2006, Birds Eye Foods voluntarily recalled frozen cooked winter squash due to possible ammonia contamination. In all, Birds Eye recalled approximately three million packages. The cooked winter squash was produced by a third party, Chase Farms, in Walkersville, Michigan (FDA 2006).
AMMONIA REGULATIONS FOR FOOD AND FOOD PRODUCTS

The FDA (1973) has determined that normal concentrations of ammonia and ammonium compounds present in food are GRAS (generally recognized as safe) food ingredients presented no risk. Ammonia and ammonium ions are essential components of normal metabolic processes. However, some restrictions have been established by FDA for allowable levels of ammonium salts in food and food processing: 0.04–3.2% ammonium bicarbonate in baked goods, grain, snack foods, and reconstituted vegetables; 2.0% ammonium carbonate in baked goods, gelatins, and puddings; 0.001% ammonium chloride in baked goods; 0.8% ammonium chloride in condiments and relishes; 0.6–0.8% ammonium hydroxide in baked goods, cheeses, gelatins, and puddings; 0.01% monobasic ammonium phosphate in baked goods; 1.1% dibasic ammonium phosphate in baked goods; 0.003% dibasic ammonium phosphate in nonalcoholic beverages; and 0.012% dibasic ammonium phosphate for condiments and relishes (ATSDR 2004).

Anhydrous ammonia is consider GRAS and approved for use as an antimicrobial agent in lean finely textured beef according to good manufacture practice (USDA-FSIS 2009). In addition, ammonium ion is regulated by the Clean Water Effluent Guidelines for the meat products (EPA 2002j). Beef Products, Inc. (BPI), produces lean trim treated with anhydrous ammonia to increase its pH and to eliminate pathogenic microorganisms (Niebur and Dickson 2003). Table 3 lists ammonia and ammonium salt regulations in food, processed food, and animal feed.
<table>
<thead>
<tr>
<th>Agency</th>
<th>Ammonium salt/ion</th>
<th>In</th>
<th>Regulations</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>EPA</td>
<td>Ammonium ion</td>
<td>Meat products</td>
<td>Clean water effluent guidelines</td>
<td>EPA 2002j</td>
</tr>
<tr>
<td>EPA</td>
<td>Ammonium chloride</td>
<td>Pesticide for raw agricultural commodities</td>
<td>Tolerance exempt when used with GAP</td>
<td>EPA 2002e</td>
</tr>
<tr>
<td>EPA</td>
<td>Ammonium hydroxide</td>
<td>Corn grain feed oranges, etc.</td>
<td>Tolerance exempt if used with GAP</td>
<td>EPA 2002f</td>
</tr>
<tr>
<td>FDA</td>
<td>Ammonium chloride</td>
<td>Direct food substances</td>
<td>GRAS</td>
<td>FDA 2008a, 21CFR 184.1138</td>
</tr>
<tr>
<td>FDA</td>
<td>Ammonium hydroxide</td>
<td>Direct food substances</td>
<td>GRAS</td>
<td>FDA 2008b, 21CFR 184.1139</td>
</tr>
<tr>
<td>FDA</td>
<td>Ammonium sulfate</td>
<td>Direct food substances</td>
<td>GRAS</td>
<td>FDA 2008c, 21CFR 184.1143</td>
</tr>
<tr>
<td>FDA</td>
<td>Anhydrous ammonia</td>
<td>Animal feeding and drinking water</td>
<td>Permitted</td>
<td>FDA 2008e, 21CFR 57.180</td>
</tr>
<tr>
<td>FDA</td>
<td>Ammonium hydroxide</td>
<td>Animal feeding</td>
<td>GRAS when use with GMP or GFP</td>
<td>FDA 2008f, 1CFR 582.1139</td>
</tr>
<tr>
<td>USDA</td>
<td>Ammonium hydroxide</td>
<td>Meat brine</td>
<td>Less than 11.6 pH</td>
<td>USDA-FSIS 2009. 7120.1, 18</td>
</tr>
</tbody>
</table>

EPA = Environmental Protection Agency; FDA = Food and Drug Administration; GFP = Good Feeding Practice; GMP = Good Manufacture Practice; GRAS = Generally Recognized as Safe; USDA = United State Department of Agriculture
Ammonia contamination of meat is obvious if the change in the meat pH rises by more than 1.0 unit or a smell of ammonia presents when contaminated meat is cooked. For meat to develop signs of contamination, the ammonia contamination must be relatively high (>15 ppm) and/or the exposure time relatively long (> 120 min) (CSIRO 2002). Not only does contaminated meat have elevated pH, but water holding capacity, color, tenderness, flavor, juiciness, and texture also change (Al-Sahal 2003; Ting and Henrickson 1986). The degree of change in the meat quality depends on ammonia exposure concentration, exposure time, type of meat, type of cut, temperature, humidity, meat surface condition, and the distance between the meat and the leakage source (Tuengerthal 1979). Several researchers have pointed out the color darkening effect (Curda and Hruby 1987; Bonne et al. 1993; Al-Sahal 1995), while the presence of pink color after cooking was noticed by Smolskiy et al. (1985) and Shaw et al. (1992). Ammonia’s effect on meat flavor was studied by Ireland (1988), Hagyard et al. (1993), and Guerrero and Arnau (1995). Increased meat pH was characterized by Kassem and Johnston (1965), Herrmann and Johnston (1966), Anil (1971), Ireland (1988) and, Al-Sahal (1995). Meat tenderness was investigated by Anil (1971) and Herman (1965). Anil (1971) and Al-Sahal (1995) indicated that meat contaminated by ammonia had higher water holding capacity than control. The type of food packing materials also affects the level of contamination. Wax paper was most permeable to ammonia, while Cryovac vacuum packaging was less permeable (Kassem 1965).

Meat Color

Meat color and meat appearance are most important to consumers purchasing meat. The
meat myoglobin molecule is responsible for the fresh meat color, and any change in this pigment will change meat appearance and color. Moreover, the level of the myoglobin not only affects the color of meat but the type of myoglobin and its chemical form, along with characteristics of other meat components, also affect the color (Lawrie 1998).

To prevent meat color from deteriorating, most fresh meat producers have recently begun to introduce carbon monoxide gas into packages. Carbon monoxide binds to myoglobin to produce a bright color (carboximyoglobin) identical to the bright color produced by the oxygen-myoglobin reaction (oxymyglobin). Figure 2 shows the reaction between meat pigments with oxygen and the developing color resulting from this interaction.

Treating meat with ammonia can also prevent meat from losing its color during storage or cooking. Kassem (1965) exposed ground beef to 10 mL of ammonium hydroxide (NH₃OH) for 60 hr at -23 °C, and after aerating for 30 min at room temperature, the ground beef was wrapped in three different types on packaging materials (waxed paper, regular, polyethylene, and Cryovac wraps). Kassem (1965) concluded that a grayish color formed on the surface of the ground beef (most noticeable in the regular wrap and the waxed paper wrap) with no color change inside the ground beef. His results showed a significant difference between the treated ground meat samples and the control in color, with no significant differences among the treated ground beef samples. Smolskiy et al. (1985) concluded that sausage produced from meat treated with 0.1 - 0.15% (1000-1500 ppm) ammonia had a more intensive pinkish color than the control. Bonne et al. (1993) investigated beef and sheep carcasses in a refrigeration unit leaking ammonia. They noticed meat contaminated with ammonia had a bright red color that did not change even after 24 hr. Ammonia preserves meat color even after cooking. Shaw et al. (1992) studied the effects of ammonia exposure and the pH on the pink color in cooked pork. Their
conclusion was ammonia treatment enhanced the pink color in the cooked pork meat rather than changing in the cooked meat pH. Al-Sahal (2003) studied the effect of high ammonia exposure on cooked beef muscles. He found that contaminated cooked beef muscle was more pink in color than the uncontaminated muscle. Ammonia increases meat pH, and high pH protects the myoglobin molecule from denaturation, which could explain the bright red color of the contaminated meat. Kropf and Hunt (1998) indicated that high pH is important to produce the bright red color in cooked meat.

In general, meat color depends on the availability of myoglobin and the fiber content of the meat. However, meat type affects the formation of the bright red color. For example, chicken has less myoglobin, so color in chicken meat will not change as color in beef or pork meat does. Al-Sahal (1995) showed chicken meat contaminated with ammonia changes color less than contaminated beef or pork.

**Figure 2:** The relationship among the three pigments color in the meat. Adapted from Robertson (2006)
Meat Flavor and Odor

The higher ammonia concentration and the longer the exposure time the more effect has on meat flavor and odor (Al-Sahal 2003). Bonne et al. (1993) investigated an ammonia leak in a slaughterhouse where it was easy to detect ammonia odor on the contaminated carcasses surface. However, aeration the carcasses for several hours in air clean cold room was effective to eliminate ammonia odor. Kassem (1965) investigate an ammonia leak incident in a Knoxville company in 1964 where different foods were packed in different packaging material (regular, polyethylene, waxed paper, and Cryovac), concluding that ammonia altered the flavor and the odor of the packed food. The sliced beef in a barbecue sauce had an ammonia odor and of flavor. In addition, whole chicken in Cryovac packaging was not affected due to the impermeability of the package. He also concluded that the packaging type changed how ammonia affected the food and wax paper was the most permeable, while Cryovac was the least permeable. Another study he performed. It was an organoleptic evaluation to a contaminated ground beef packaged in the same packaging materials. The testers were unable to notice any difference between the contaminated samples and the control, but they were able to detect ammonia odor.

The effect of ammonia on meat was studied by Ireland (1988), who exposed lamb loins to 2 M of ammonia gas at 4, 6, 16, and 32 min. Even with low ammonia concentrations, meat flavor and odor changed from the control, as did the texture (more sloppy) and the color (much paler).

Guerrero and Arnau (1995) noticed a change in ham flavor when they treated an exposed ham with mites and their eggs with air saturated with ammonia for 24 hr to study the effect of ammonia on eliminating the mites and their eggs from the ham. They concluded that, after one
day of treatment, ammonia was effective to eliminate all mites stages but also altered the ham flavor and it was noticeable even after 7 days. In a study conducted by Hagyard et al. (1993), levels of ammonium hydroxide solution (2 M), caused lamb loins to develop a rancid flavor after 3 months of storage at freezing temperature of -20 °C (twenty four month maximum storage time for lamb loins at -18 °C; USDA-FSIS 2008); they concluded that ammonia contamination reduced meat shelf life and freezing temperatures did not prevent damage to meat flavor.

In general, people can smell ammonia at a level as low as 10 to 20 ppm (Raj 1982). In the Wisconsin incident (1985), no one detected any ammonia odor in the contaminated milk level (CDC 1986). In a previous research in our own laboratory, we could not detect ammonia odor in beef contaminated with ammonia at 1000 ppm. The ammonium ion has no odor and thus when ammonia reacts with water in the food, it forms ammonium ions, which are odorless (Massachusetts Department of Public Health 1998). Also the longer contaminated food is stored the more chance that the ammonia odor will not be present due to ammonia volatility.

**Meat pH**

Meat pH is important to meat quality. pH can affect meat color, texture, enzymatic reactions, and the water holding capacity of the meat. Exposing meat to ammonia will mostly affect the outer layer of the meat surface (Anil 1971). Herrmann (1965) studied the effect of high ammonia concentration on pork and beef meat surfaces. He noticed an increase in the exposed meat pH compared to unexposed meat. Herrmann and Johnston (1966) studied the change in the pH in four layers of a meat contaminated with ammonia. They found a significant increase in pH for the first outer layer, with less increase in pH in the second layer, the third layer, and the fourth layer. They concluded storage temperature (-28, -18, and -9.4 °C) had little
effect on pH level.

Anil (1971) studied the effect of temperature and ammonia absorption on the pH in beef muscle. Anil (1971) exposed beef muscle to 10 milliliters of liquid ammonia for 72 hr at -4 °C. The pH and the ammonia levels increased significantly in the exposed muscle, the ammonia concentration in the first outer layer (0.6 cm) was higher than in the second layer (0.6 cm). The third layer did not differ significantly from the control. Anil (1971) concluded that storage temperature did not influence the distribution of ammonia within the meat layers.

Al-Sahal (2003) studied pH in beef muscle after it was exposed to high levels of ammonia (500, 1000, 2500, and 5000 ppm and 0, 5, 10, and 20 min of exposure times). Exposing meat to 5000 ppm for 20 minutes elevated the pH by 1 unit (Figure 3). The pH of the beef steak surface was higher than the inner portion. Moreover, the total pH for a ground sample from the same steak was lower than surface pH; the sample’s pH was diluted by the grinding and by mixing the outer and the inner portions of the sample. In other research, adding 0.1 to 1.5% (1000 to 15000 ppm) ammonia to meat processed into sausage elevated the sausage pH by one pH unit (Smolskiy et al. 1985). However, pH is not a reliable indicator for ammonia contamination in meat (Kassem 1965; Ireland 1988; Hagyard et al. 1993; Hijaz et al. 2007). The buffering capacity of different meat types may affect the ammonia sufficiently to influence the pH (Al-Sahal 1995).
Figure 3: The change in total pH of *Semitendinosus* steaks over time when exposed to different concentrations of ammonia. Adapted from Al-Sahal (2003)

**Meat Tenderness**

Muscle fibers, connective tissues, and adipose tissues each influence meat tenderness (Judge et al. 1989). Cover et al. (1962) described meat tenderness as softness to tongue and cheek, ease of fragmentation, resistant to tooth pressure, resistance after chewing, adhesion, and mealiness. Increasing meat’s water holding capacity by increasing the pH of the meat increases tenderness. Herrmann (1965) studied the effect of ammonia on the tenderness of meat. He exposed beef samples to ammonia for 24 hr at -28 °C, -18 °C, and -9.4 °C. Tenderness in exposed meat improved at -28 °C and -9.4 °C compared with the unexposed control. However, a sampling error may have occurred because the tenderness from the control did not differ significantly from the ammonia treated sample at -18 °C.

Anil (1971) also noted that exposing beef samples to 10 mL ammonium hydroxide solution for 72 hr effectively increased tenderness of meat. Anil pointed out that the meat surface was more tenderer than the inner portion.
Meat Water Holding Capacity

During meat processing, meat loses water as well as any added water. Meat water holding capacity (WHC), is the ability of meat to retain water under external forces like grinding, cutting, centrifuging, heating, or pressing (Jauregui et al. 1981; Judge et al. 1989). Hamm (1986) studied WHC using freezing, pH, thawing, heating, and postmortem changes. The WHC of meat is mainly controlled by the protein net charge in the meat muscle, if the meat pH is normal (around 7) at slaughter. Postmortem glycolysis in muscle causes an accumulation of lactic acid, which results in a rapid decline in muscle pH to 5.5-5.6 (Greaser 1986). The iso-electric point (equal charges of positive and negative in the protein net) for the meat is near pH 5 (Aberle et al. 2001). Equality in the net charge means WHC is minimal, so adding ammonia to meat either intentionally or unintentionally will elevate meat pH and increase the net charge, resulting in increased WHC and thus tenderness.

Al-Sahal (2003) assigned meat to different ammonia exposures (500, 1000, 2500, and 5000 ppm) and different exposure times (0, 5, 10, and 20 min) to test WHC. His results confirmed a significant increase in WHC for meat exposed to concentrations of 2,500 ppm and 5,000 ppm at 5, 10, and 20 min. At 10,000 ppm ammonia exposure and 0, 3, 6, 12, 24, and 48 hr exposure time, Al-Sahal (1995) noticed an increase in WHC of beef, chicken, and pork; at 5,000 ppm, WHC was significantly increased for the beef. He concluded that the different buffering capacities of beef and the chicken may affect WHC. Smolskiy et al. (1985) concluded that ammonia contaminated ground beef used to make sausage showed increased pH and WHC. In another study, WHC increased in ammonia contaminated uncooked beef muscles and ammonia contaminated cooked muscle (Anil 1971).
MEAT PACKAGING FUNCTIONS

Protecting meat from external hazards is the primary function of meat packaging. At the same time, the packaging materials themselves should have no effect on meat. The packaging materials chosen should meet minimum technical, legislative, environmental, and commercial requirements (George 2000). Moreover, the food package should function as preservation containment, protection, communication, identification, convenience, and distribution (Robertson 1993; Soroka 1999).

ENVIRONMENTAL EFFECTS ON PACKAGING PERMEABILITY

Temperature

Both the diffusion coefficient \( (D) \) and the solubility coefficient \( (S) \) of a penetrant determine the permeability coefficient \( (P) \). Both the diffusion coefficient and the solubility coefficient depend on the temperature. Measuring the permeability coefficient or the vapor transmission rate is not valid without reference to the test temperature (Plastic Design Library 1995). The effect of the temperature on the solubility coefficient can differ according to type of penetrans. Permanent gases like oxygen and sparingly soluble gases and vapors increase with increasing temperature, in such gases increasing the temperature increases the solubility coefficient resulting in increasing the permeability, while increasing the temperature in condensable gases such as ammonia and sulfur dioxide decreases the solubility coefficient resulting in decreasing the permeability in general (Plastic Design Library 1995). Water vapor permeation usually increases with increasing temperature, depending on the barrier and the moisture content of the barrier. Increasing the temperature of organic vapors generally increases the permeability but interferes with the swelling of the barrier materials.
**Relative Humidity and Pressure**

Diffusion of water through certain packaging materials depends on water concentration due to the plasticizing effects of water on those materials that can lead to increase the permeability. Therefore, water vapor transmission rate is affected by differences in relative humidity. Thus, measuring relative humidity is important in correctly interpreting the permeability measurement. Moreover, some packaging materials absorb water, which can increase their permeability because of the elasticizing effect of the water. At normal pressure, the solubility coefficient for permanent gases is proportional to their partial pressure, because at normal pressure, the permanent gases follow Henry’s Law “At a constant temperature, the amount of a given gas dissolved in a given type and volume of liquid is directly proportional to the partial pressure of that gas in equilibrium with that liquid”. Thus, their permeation rates are generally independent of pressure. For readily soluble gases, which do not obey Henry’s Law, the permeation rates relate complexly to pressure. For these gases, measuring the pressure is very important in correcting the permeation rate. For sparingly soluble vapors such as water vapor in polyolefin, the permeation rate is proportional to the vapor pressure different across the barrier wall.

**FACTORS AFFECTING PACKAGE PERMEABILITY**

The permeability of the polymeric packaging materials depends on two factors: the diffusion and the sorption of gases and vapors. Both chemical structure and methods of fabrication, including processing conditions of the packaging materials, influence permeability coefficients. Moreover, the permeability of the packaging materials depends on polymer structural parameters, including crystallinity, polarity, cross-linking, grafting, tacticity, and free
volume (Jasse 1994). In general, gas permeation rates through packaging plastic depends on type of plastic, thickness, surface area, methods of processing, concentration or partial pressure of the permeant molecule, and storage temperature (Kirwan et al. 2003). In addition, the permeability of packaging materials to water vapor is depending upon material permeability, surface area, and temperature (George 2000).

**PROPERTIES OF SOME PLASTIC POLYMERS**

**Polyamide**

Polyamide, or nylon, is a linear condensation thermoplastic made from monomers with amine and carboxylic acid functional groups resulting in amide (-CONH-) linkage in the main polymer chain that provide mechanical strength and barrier properties (Robertson 2006). Because of the polarity of the polyamide (PA), the PAs are highly permeable to water vapor, although their permeability to $O_2$ and other gases is fairly low when dry. For most applications, PAs are combined with low density polyethylene (LDPE) to add heat stability and a moisture barrier to the packaging materials. Multilayer films containing a PA layer are used principally in vacuum packages for processed meat and cheese (Robertson 2006). Identification for PA refers to the number of carbon atoms in the diamine followed by the diacid. Nylon 6 and nylon 11 are the most used forms of PA (Figure 4). Nylon 11 has twice the distance between amide groups than nylon 6 and consequently is intermediate in properties between nylon 6 and polyethylene (Brydson 1999a). Biaxially oriented PA film is highly heat resistant and resistant to stress, cracking, and puncture. It has good clarity and is easily thermoformed with good barrier property (Coles et al. 2003).
Polyolefin

Polyolefin refers to group of plastics based on propylene and ethylene, an important class of thermoplastic packaging materials. Olefin means oil-forming and was originally the name of ethylene (Robertson 2006). Polyolefin is widely used for meat and meat products especially for poultry products for it tight, hermetically sealed package. However, Polyolefin are known to absorb oil-based flavors (Jenkins and Harrington 1991).

Under Polyolefin, two major classes can be classified including polyethylene (PE) and polypropylene (PP)

Low density polyethylene (LDPE)

LDPE is the largest volume single polymer used in the food packaging in both the film and blow-molded form (Robertson 2006). LDPE is commonly used for packaging individually quick frozen foods although it provides relatively poor protection from oxygen and good protection from water vapor (George 2000). Figure 5 shows the polyethylene structure which is basically a polymerization of -CH₂-. A wide range of pressures and temperatures can be applied in polymerization of polyethylene. In commercial production, pressure between 1,000 to 3,000
atmospheres and temperatures of 100 to 350 °C are usually applied. Because of this, the simple structure of polyethylene (unbranched -CH₂-) is disturbed and long branches and short branches form (Robertson 2006).

![Branched polymer polyethylene](C:\example.png)

**Figure 5**: Branched polymer polyethylene

Polyethylene is classified into three major types: low density (LDPE), medium density (MDPE), and high density (HDPE). LDPE is a good water vapor and poor oxygen barrier and can form heat seals (Ozen and Mauer 2004). The number of CH₃ groups in 1,000 atoms of carbon identifies each type of polyethylene. Table 4 shows the major types of polyethylene and some of their physical properties.

The different of properties of the many grades of LDPE follow (Brydson 1999):

1- Variation in the degree of short chain branching in the polymer.

2- Variation in the degree of the long chain in the polymer.

3- Variation in the average molecular weight.

4- Variation in the molecular weight distribution (which may in part depend on the long chain branching).

5- The presence of small of co-monomer residue.
6- The presence of impurities or polymerization residues, some of which may be combined with the polymer.

Table 4: Basic properties of various polyethylene films\(^1\)

<table>
<thead>
<tr>
<th>Type of Polyethylene</th>
<th>Moisture Vapor Transmission</th>
<th>Gas Transmission</th>
<th>CH(_3) Group per 1000 Cs</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDPE (920 kg m(^{-3}))</td>
<td>1.4</td>
<td>500</td>
<td>1350</td>
</tr>
<tr>
<td>MDPE (940 kg m(^{-3}))</td>
<td>0.6</td>
<td>225</td>
<td>500</td>
</tr>
<tr>
<td>HDPE (960 kg m(^{-3}))</td>
<td>0.3</td>
<td>125</td>
<td>350</td>
</tr>
</tbody>
</table>

\(^{1}\)Transmission rate unite (g mm m\(^{-2}\) day\(^{-1}\)). Adapted and modified from (Robertson 2006)

Polypropylene

Polypropylene has similar molecular structure to PE in some respect but it is more complicated and has lower density, low water vapor transmission, and medium gas permeability (Robertson 2006). It is a linear polymer containing little or no unsaturation and can be subcategorized into three different stereo types: Isotactic; Syndiotactic; and Atactic depending on the distribution of the methyl group on either side of the polymer chain. Types of the catalyst and polymerization conditions determine the stereo type of the PP. The three types of polypropylene are displayed in Figure 6. Isotactic PP, is the most common commercial form of polypropylene, it has good chemical and heat resistance but poor transparency. The relatively high temperature resistant of PP permits its use as a sealing layer in retortable pouches (Robertson 2006). In addition, PP films have limited food packaging application because of their low cold temperature resistant. Copolymer mixtures with ethylene are used to improve cold
resistant and heat stability as well as materials strength and seal strength (Johannes and Otto 2000)

Figure 6: Polypropylene types: (a) Ilostactic, (b) Syndiotactic, and (c) Atactic

GENERAL MATERIALS REQUIREMENTS FOR MEAT PACKAGING

Jenkins and Harrington (1991) summarized the general requirements for meat packaging materials that can apply for all products including: clarity; resistance to cracking, puncture, and pinhole formation; good oxygen barrier properties; good sealability; and easy opening/closing. Moreover, the FDA and USDA regulations are usually concerned about migration of unapproved chemical substances through the contact layer of the package and reach the food. U.S. legal requirements concerning food packaging are published in the Code of Federal Regulation (CFR) title 21 volume 2 subpart 110.80. The section stated that "Packaging process and material shall not transmit contaminant or objectable substances to the product". Extending meat shelf life, prevent color discoloration, and fat rancidity are main requirements for packages that intended to use for processed meat products, these adverse reactions are due to oxygen and accelerated by light. In today’s applications, low oxygen permeability materials (0.2
cc/100 in\(^2\).atm/24 hr at 50% R.H.) are used to structure meat processed packaging materials (Jenkins and Harrington 1991). In addition, preventing freezing effects including discoloration and frost formation inside the package are the most important features in this regard (Robertson 2006).
REFERENCES


Anil N. 1971. Effect of ammonia on water-holding capacity and tenderness of frozen beef muscle. MS. Theses, University of Tennessee, Knoxville.

Arnold M. 1993. The big chill ammonia finds its niche as a refrigerant. Food Processing 54(12):45,47.


Guerrero L, Arnau J. 1995. Dry cured ham, chemical methods to control mites. Fleischwirtschaft Int. 2:84-86.


Soroka W. 1999. Fundamentals of food packaging technology, 2nd ed. Institute of packaging professionals


PART II: PERMEABILITY OF MEAT PACKAGING MATERIALS TO LOW LEVELS OF AMMONIA GAS

ABSTRACT

Meat products are packaged in polymer films that theoretically should protect the product from exterior contamination. Unfortunately, there is almost no data on ammonia permeability of packaging films. Thus, we chose to investigate common meat packaging films 2.2 mil low density polyethylene (LDPE), 3 mil Cryovac multilayer polyolefin (E-2300), and 3 mil vacuum 0.6 mil polyamide/2.4 mil polyethylene (V-PA/PE) and their permeability to ammonia. The films were fabricated into 10 x 5 cm pouches and filled with 50 mL deionized water. Pouches were placed in a plexiglass enclosure in a 1.3 CUF freezer and exposed to 50, 100, 250, and 500 ppm ammonia gas at 6, 12, 24, and 48 hr in -17 ± 3 °C and 21 ± 3 °C. At freezing temperatures, no ammonia residues were detected and no differences in pH were found. At room temperature, the ammonia concentrations and the pH increased significantly ($P < 0.05$) in the water with increasing exposure time and ammonia concentrations. The maximum average levels observed in the water were 7.77 ppm for Cryovac (E-2300), 5.94 ppm for LDPE, and 0.89 ppm for V-PA/PE at 500 ppm exposure for 48 hr at 21 ± 3 °C. The maximum average pH values observed were 8.64 for E-2300, 8.38 for LDPE, and 7.23 for V-PA/PE (unexposed ranged from 5.49 to 6.44) with exposure of 500 ppm at 48 hr. The results showed that meat packaging materials have low ammonia permeability and thus protect meat products exposed to ammonia exposure during frozen storage.

*Presented in part at IFT Annual Meeting, June 7, 2009, # 060-12 (http://www.am-fe.ift.org/cms)
INTRODUCTION

Ammonia is the refrigerant of choice for food products and the refrigeration backbone in the food industry (IIAR 2006). Ammonia has important advantages as a refrigerant: high efficiency in providing cooling and freezing for food products (Holmstrom 1994), relatively low environmental impact (Ross 1994), and low production costs (Arnold 1993) with no impact on the ozone layer.

The food industry uses refrigeration with ammonia to cool fruits, vegetables, poultry, fish, beverages, dairy products, milk, and meat (Lorentzen 1988). Most fabrication facilities, slaughterhouses, and cold storage warehouses use ammonia as their refrigerant (Ross 1995; Sun 1998). With typical equipment failures and operator error, ammonia leaks will occur in refrigerated/frozen food storage facilities. Carelessness may be part of the problem, but more often equipment failures are the issue (Ostner 1986).

Beside refrigeration and freezing, most perishable foods currently are packed in plastic films to protect food against external/internal deterioration. Package integrity and barrier characteristics are important in many areas of the food industry. Their importance in preserving and protecting commercial sterilized food in sealed packages has been well investigated (Denny 1989; Harper et al. 1995) A single layer or multilayered polymer films of different polymer materials can sufficiently protect food (Jasse et al. 1994).

Package protection means the material is not permeability to gases and vapors harmful to the quality/safety of the food products. Permeability through any plastic film is measured by the permeability coefficient \( P = D \times S \) that describes the transfer of a total mass at a steady state through plastic materials (Crank 1975). In this equation, \( P \) is the permeability coefficient; \( D \) is the diffusion coefficient (a measure of the transportation speed of the molecule through a plastic
film); and S is the solubility coefficient that measures the number of the permeant molecules moving through a plastic film. The properties of polymers that affect the permeability are crystallinity; polarity; chain to chain packing ability; glass transition temperature; size, shape, and polarity of the permeant; temperature; and pressure (Pascat 1986; Sperling 1992; Robertson 1993). In addition, the permeant temperature and pressure are important in permeability to gases and vapors (Jasse et al. 1994). The stages through which a permeant molecules permeates polymers are the following (Ashley 1985): (1) absorption of the permeant onto the polymer surface; (2) solubilization of the permeant in the polymer texture; (3) diffusion of the permeant through the polymer along a concentration gradient; and (4) desorption of the permeant from the other polymer surface. Mannheim and Miltz (1987) concluded that an increase in oxygen permeability due to flex-cracking of the polymer during transportation may adversely affect food shelf life.

The literatures offer almost no data regarding permeability of food packaging to ammonia gas. In addition, there is a lack of information related to the level of protection that food packaging materials can provide during an exposure to ammonia gas. The objectives of this research were to evaluate the permeability of three different packaging materials commonly used in the meat industry to low levels of ammonia exposure at different times in freezing and room temperature.

**MATERIALS AND METHODS**

**Experimental Design**

Three different meat packaging materials, 2.2 mil low density polyethylene (LDPE) (Manchester Packaging Co., Saint James, Missouri, USA); 3 mil Cryovac multilayer polyolefin
Prefabricated (10 × 5 cm) 50 mL water filled Cryovac E-2300, LDPE, and V-PA/PE pouches

↓

Placed in freezer for 4-6 hr

↓

Inserted four pouches of each type (total of 12 samples) into freezer 1.3 CUF (-17 ± 3 °C)

↓

Applied flow rate of 85 mL/min of ammonia gas (50, 100, 250, and 500 ppm)

↓

Withdrew one pouch of each type only at certain time of 6, 12, 24, and 48 hr

↓

Thawed overnight at 4-8 °C in a refrigerator

↓

Measured pH values and ammonia levels for each pouches water

↓

Repeated the process, applying the same ammonia concentrations and exposure times at room temperature (21 ± 3 °C) without freezing the samples.
Sample Preparation

Plastic films were sealed to 10 cm × 5 cm pouches using a Midwest Pacific impulse heat sealer (J. J. Elemer Corp., St. Louis, Missouri, USA), and filled with 50 mL of deionized water (Figure 8).

Figure 8: Fabricated plastic films Cryovac, low density polyethylene, and vacuum with dimensions of 10 × 5 cm and filled with 50 mL deionized water

A pinhole was made in one corner of the pouch to remove any trapped air. Each pouch was approximately 2.3 cm thick. In the experiment performed under freezing conditions, the pouches were placed inside an air tight 1.7 L plastic container (Lock and Lock, Heritage Mint, Ltd. Scottsdale, Arizona, USA) and inserted into a freezer for 4-6 hr until the pouches were frozen. Frozen pouches were assigned to different ammonia exposures for different times. In the room temperature experiment, pouches were fabricated and filled with deionized water and assigned directly to the experiment. Pouches were coded and placed randomly in a treatment chamber for a total of twelve samples for each treatment replicate combination. One pouch of each kind was removed at 6, 12, 24, and 48 hr for freezing temperature treatment and room temperature treatment.
Exposure Chamber

A clear plexiglass box measuring 38W × 13.5D × 36H cm (18.5 L) was used as a treatment chamber (see Figure 9). The exposure chamber was tested for leaks and proper seal by inserting a ToxiRae PGM-30 Photo-Ionization Detector (PID) (accuracy ± 2 ppm or ± 10% of the reading; RAE Systems, California, USA) into the chamber and locking the chamber. The detector was operated and calibrated according to the ToxiRae PGM-30 instruction manual. The ToxiRae PGM-30 can detect ammonia levels and store the data. The detector is also portable and rechargeable and has a screen that displays readings. A concentration of 50 ppm of ammonia gas was introduced into the chamber from a calibrated 50 ppm (± 2%) ammonia gas cylinder.

Figure 9: Exposure chamber used in the experiment with two L-shape fittings mounted on the back and 15 cm round front door

When the concentration of 50 ppm was reached and detected, the ammonia gas supply was stopped, and the chamber was locked. After 7 hr, the detector was removed, and the readings were recorded, after analyzing the detector’s data no depleting in the ammonia
concentration was found inside the chamber, the box had no leaks. In addition, the chamber was pressurized with air, locked, and immersed in a water bath. No bubbles were observed coming from the chamber during this test.

The pouches were inserted into the chamber through a round front door (15 cm dia) and were attached to stainless steel hooks, suspended on two plexiglass bars attached horizontally between the left and the right side of the box (Figure 10). On the back side of the chamber, two L-shape threaded fittings were mounted to serve as gas inlet and outlet, one in the upper right and the other in the lower left; these allow the ammonia gas to be homogeneously distributed throughout the chamber.

**Figure 10:** Water filled fabricated pouches inside the exposure chamber inside the freezer suspended on two plexiglass bars
Exposure System Design

The exposure system (See Figure 11) was designed to simulate the true conditions of low level of ammonia exposure and to manually control the ammonia flow rate entering the chamber while maintaining the desired level of ammonia concentration. The ammonia cylinder was connected to a 0-65 mm gas flow regulator inlet (Cole-Parmer Instrument Company, Vernon, Illinois, USA), and the gas regulator outlet was fitted to a T-shaped plastic connector. The exposure chamber and the detector were attached to the connector. The chamber outlet was assembled with a T-shaped tube connector hocked to a one-way plastic valve (TETRA Technologies, Inc., The Woodlands, TX, USA) to prevent the ambient air from entering into the chamber and diluting the ammonia concentration, especially under freezing conditions. The valve served as a pressure release device to prevent breakage in the chamber if pressure levels became too high during flushing. A second gas flow regulator identical to the first one connected to the other end of the one-way valve to monitor the ammonia exhaust flow rate.

The other end of the T-shaped connector was attached to a glass heat exchanger in order to warm the ammonia gas before reaching the detector, especially in the experiments that were performed under freezing temperatures. Plastic tubing (TYGON S-50-HL, class VI, size 0.46 × 0.16 cm) was used to combine the system parts together. Figure 12 shows the tubing connections between the freezer and the inlet gas flow regulator; MiniRae 2000 PID; heat exchanger; and the outlet gas regulator.

Procedure

The ammonia gas cylinders were obtained from Linweld, Inc., (Manhattan, Kansas, USA) and were manufactured by Matheson Tri-Gas, Inc., (Joliet, Illinois, USA) with concentrations of 50, 100, 250, 500, and 1000 ppm (calibrated concentration were 48.8, 90.0,
238.7, 474.0, and 992.0 ppm; ±2% in air). Two cylinders were used in each treatment. The 1000 ppm ammonia concentration cylinder introduced ammonia at the desired level to flush the chamber (Figure 13) while maintaining a continuous exposure of the required ammonia concentration for the required time with 50, 100, 250, and 500 ppm concentration cylinders (Figure 14).

Figure 11: The exposure system components: low ammonia concentration tank, inlet gas flow regulator, MiniRae 2000 detector, glass heat exchanger, freezer, and outlet flow regulator
Figure 12: Tubing connection between the system parts

A stainless steel, single stage regulator (Air Products, Brown Welding, Salina, Kansas, USA) was mounted on the low concentration cylinder to control the flow rate pressure of ammonia gas into the gas flow regulator. The temperature inside the chamber was monitored with a Smart button data logger (Accuracy: ±1.0 °C from -30 °C to 45 °C) (ACR Systems, Inc., Surrey, British Columbia, Canada).
Figure 13: Chamber flushing process and ammonia gas directions in the system. The chamber was flushed using the higher ammonia concentration cylinder (1000 ppm), and the reading was recorded using MiniRae 2000 photo ionization detector.

Figure 14: Movement directions of ammonia gas during exposure.
The chamber was flushed using ammonia gas from the higher ammonia concentration cylinder, and the reading was taken using MiniRae 2000 self sampling photo ionization detector (PID; accuracy 2 ppm or 10% of the reading from 0-2000 ppm; RAE Systems, California, USA). The detector inlet port was connected to the glass heat exchanger end, and the detector outlet port was connected to a tube going into the chamber inlet. To avoid losing ammonia from the chamber, the detector takes an ammonia gas sample from the chamber, detects the gas, and circulates the gas back into the chamber. The detector was operated and calibrated according to the MiniRae 2000 instructions manual. After the desired concentration inside the chamber was reached, the ammonia gas from the lower concentration cylinder was applied by adjusting the gas cylinder regulator to a pressure of 2.5 psi, and the flow rate was adjusted to 85 mL/min. Ammonia exhaust flow rate was monitored by the second gas flow regulator, with the dial opened to the maximum to freely flow exhaust the ammonia. At the end of each experiment, the ammonia cylinder supply was turned off, and the ammonia concentration inside the chamber was measured using the MiniRae 2000 detector (see Table 5).

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>Ammonia concentration (ppm)(^{1,2})</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>50</td>
</tr>
<tr>
<td>6</td>
<td>30 ± 2</td>
</tr>
<tr>
<td>12</td>
<td>33 ± 2</td>
</tr>
<tr>
<td>24</td>
<td>29 ± 2</td>
</tr>
<tr>
<td>48</td>
<td>29 ± 2</td>
</tr>
</tbody>
</table>

\(^{1}\text{Values are means ± standard deviation of 3 replicates.}\)

\(^{2}\text{The measurements were taken at the end of each run.}\)
Indophenol Assay Ammonia Measurement

Sodium hypochlorite solution (10-15% available chlorine) and sodium nitroferricyanide (III) dihydrate were purchased from Sigma Chemical Co. (St. Louis, Missouri, USA.). Phenol, sodium hydroxide, sodium phosphate dibasic dodecahydrate, and a GenesysTm 10 visible spectrophotometer were purchased from Fisher Scientific Co. (Pittsburgh, Pennsylvania, USA). The reagents were prepared and stored according to Broderick and Kang (1980).

Ammonia in the water pouches exposed at freezing temperatures was determined by defrosting the pouches overnight at 4-8 °C in a refrigerator and transferring 200 µL of the water to a glass test tube. The water was mixed with 2.5 mL phenol reagent and 2 mL hypochlorite reagent. The test tube was placed in a 95 °C water bath for 5 min and allowed to stand at room temperature until ~25 °C. To construct the standard curve, solutions of 0.05, 0.1, 0.25, 0.5, 0.75, 1.0, 2.5, 5.0, 7.5, and 10 ppm ammonia as nitrogen (N) were prepared from the commercial standard 1000 ppm ammonium chloride (Thermo Electron Corporation, Miami, Florida, USA). The blank was prepared using 200 µL of distilled water. Finally, the 630 nm spectrum was used to scan the samples and the standards using a GenesysTm 10 visible spectrophotometer. To measure the ammonia levels in water pouches at room temperature, 200 µL of the water was transferred directly to a glass test tube and the procedure continued from there.

pH Determination

Water pH was measured directly by inserting the pH meter probe in the pouches (Denver Instruments UP-5 pH meter, single electrode, Denver Instrument, Denver, Colorado, USA). pH of each sample was measured twice after temperature stabilized between 20-25 °C. Frozen samples were defrosted at room temperature before pH was determined.
**Statistical analysis**

The statistical analysis system (Version 9.1, SAS Institute, Inc., Cary, N.Y., USA; 2002) was used for data analysis. The experiment design was a three way analysis of variance 4×4×3 for ammonia levels and 4×4×3 and the control time (0 hr) for pH values using PROC GLM/TUKEY to determine significant differences among the treatment means at $P < 0.05$.

**RESULTS AND DISCUSSION**

**pH**

No significant change in the pouches water pH was observed when the pouches exposed to ammonia gas at freezing temperature (-17 ± 3 °C). In the experiment conducted at room temperature (21 ± 3 °C), the overall model (time and concentration) showed significant increases in pH ($P < 0.05$) for all packaging types as exposure time increased and as ammonia concentration increased (Table 6, 7, and 9). Figure 15 shows the pH levels for E-2300, LDPE, and V-PA/PE at 500 ppm ammonia exposure and 6, 12, 24, and 48 hr of exposure time. Post hoc analysis showed that the main effect of exposure time and concentration was significant ($P < 0.05$). Although, time and concentration showed significant interaction ($P < 0.05$). There was no significant interaction between bag type and ammonia concentration ($P < 0.0523$). That lack of significant interaction indicates that pH increased independently of increasing the ammonia concentrations and the bag type. In addition, there was no significant interaction between bag type, ammonia concentration, and time ($P < 0.9997$). However, the interaction between time and bag type was significant ($P < 0.05$). The maximum differences values in the pH from the control mean were 2.71 ± 0.8 for the E-2300, 2.45 ± 0.97 for LDPE, and 1.30 ± 1.04 for V-PA/PE when the pouches were exposed to 500 ppm for 48 hr.

Generally, the alkalinity of ammonia increases water pH because ammonium hydroxide
forms. Ammonium hydroxide is a weak base that is partly ionized in water to form ammonium
(NH$_4^+$) and hydroxyl (OH$^-$) ions (the alkalinity property for the aqueous solution of ammonia
relates to hydroxide ions) according to the equilibrium:

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

Table 6 lists the Cryovac E-2300 pH values. The water in E-2300 exhibited the
maximum pH level. Statistical analysis showed significant increases in pH ($P < 0.05$) as
exposure time increased and as ammonia concentration increased. However, the interaction
between exposure time and ammonia concentration showed no significant relationship ($P <
0.0893$).

![Graph showing pH levels for Cryovac E-2300, LDPE, and vacuum (V-PA/PE) bags at 500
ppm exposure for 6, 12, 24, and 48 hr at room temperature.]

**Figure 15:** Water pH levels for Cryovac E-2300, LDPE, and vacuum (V-PA/PE) bags at 500
ppm exposure for 6, 12, 24, and 48 hr at room temperature

The LDPE showed pH profile similar to Cryovac E-2300 (Table 7). The statistical
analysis for pH related to exposure time and ammonia exposure concentration showed significant results ($P < 0.05$), whereas no significant interaction was found between them ($P < 0.4591$). V-PA/PE showed significant results ($P < 0.05$) in the overall model as exposure time and ammonia concentration increased. Post hoc analysis showed no significant interaction between the time and the ammonia concentration ($P < 0.9764$). V-PA/PE was less permeable to ammonia (Table 8).

Table 6: Cryovac E-2300 bag’s water pH levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>Ammonia concentration (ppm)$^1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>100</td>
</tr>
<tr>
<td>Control (0)</td>
<td>5.52 ± 0.02$^{aC}$</td>
</tr>
<tr>
<td>6</td>
<td>5.99 ± 0.01$^{bBC}$</td>
</tr>
<tr>
<td>12</td>
<td>6.17 ± 0.05$^{cBC}$</td>
</tr>
<tr>
<td>24</td>
<td>6.40 ± 0.07$^{dAB}$</td>
</tr>
<tr>
<td>48</td>
<td>7.12 ± 0.66$^{eA}$</td>
</tr>
</tbody>
</table>

$^1$Values are means ± standard deviation of 3 replicates.

$^a$-$c$Means with different superscript letters within the row are significant different at $P < 0.05$.

$^A$-$D$Means with different superscript letters within the column are significant different at $P < 0.05$.

This increasing pH over time ($P < 0.05$) may be attributed to the effect of V-PA/PE materials on pH (see Table 9). Test where conducted to determine the effect of the pouches on pH, and water in the V-PA/PE was elevated by 0.2 pH unit; no significant effects were found in E-2300 or LDPE.
Table 7: Low density polyethylene bag’s water pH levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48 hr exposure at room temperature

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>Ammonia concentration (ppm)¹</th>
<th>50</th>
<th>100</th>
<th>250</th>
<th>500</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (0)</td>
<td></td>
<td>5.52 ± 0.02aC</td>
<td>5.52 ± 0.11aD</td>
<td>5.77 ± 0.26aC</td>
<td>5.93 ± 0.48aB</td>
</tr>
<tr>
<td>6</td>
<td></td>
<td>5.93 ± 0.04bBC</td>
<td>5.99 ± 0.12bC</td>
<td>6.15 ± 0.30bBC</td>
<td>6.99 ± 0.50aAB</td>
</tr>
<tr>
<td>12</td>
<td></td>
<td>6.17 ± 0.03cBC</td>
<td>6.28 ± 0.05cBC</td>
<td>6.66 ± 0.04cBC</td>
<td>7.22 ± 0.24aAB</td>
</tr>
<tr>
<td>24</td>
<td></td>
<td>6.42 ± 0.03bAB</td>
<td>6.62 ± 0.07bB</td>
<td>7.11 ± 0.26bAB</td>
<td>7.85 ± 0.62aA</td>
</tr>
<tr>
<td>48</td>
<td></td>
<td>7.09 ± 0.58A</td>
<td>7.30 ± 0.32aA</td>
<td>8.24 ± 0.87A</td>
<td>8.39 ± 0.75A</td>
</tr>
</tbody>
</table>

¹Values are means ± standard deviation of 3 replicates.

a–cMeans with different superscript letters within the row are significant different at P < 0.05.

A–DMeans with different superscript letters within the column are significant different at P < 0.05.

Table 8: Vacuum bag’s water ammonia levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature

<table>
<thead>
<tr>
<th>Time (hr)³</th>
<th>Ammonia concentration (ppm)¹,²</th>
<th>50</th>
<th>100</th>
<th>250</th>
<th>500</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td></td>
<td>ND³</td>
<td>ND³</td>
<td>ND³</td>
<td>ND³</td>
</tr>
<tr>
<td>12</td>
<td></td>
<td>ND³</td>
<td>ND³</td>
<td>ND³</td>
<td>0.23 ± 0.07abc</td>
</tr>
<tr>
<td>24</td>
<td></td>
<td>ND³</td>
<td>ND³</td>
<td>0.14 ± 0.13ac</td>
<td>0.47 ± 0.15b</td>
</tr>
<tr>
<td>48</td>
<td></td>
<td>ND³</td>
<td>ND³</td>
<td>0.34 ± 0.18bc</td>
<td>0.89 ± 0.20d</td>
</tr>
</tbody>
</table>

¹Values are means ± standard deviation of 3 replicates.

²Means with different superscript letters are significant different at P < 0.05.

³At time 0 (control), ammonia level was 0.00 ppm for all concentrations.
Table 9: Vacuum bag’s water pH levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>50</th>
<th>100</th>
<th>250</th>
<th>500</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (0)</td>
<td>5.52 ± 0.02&lt;sup&gt;aC&lt;/sup&gt;</td>
<td>5.52 ± 0.11&lt;sup&gt;aD&lt;/sup&gt;</td>
<td>5.77 ± 0.26&lt;sup&gt;aC&lt;/sup&gt;</td>
<td>5.93 ± 0.48&lt;sup&gt;aB&lt;/sup&gt;</td>
</tr>
<tr>
<td>6</td>
<td>5.84 ± 0.12&lt;sup&gt;bBC&lt;/sup&gt;</td>
<td>5.76 ± 0.09&lt;sup&gt;bC&lt;/sup&gt;</td>
<td>5.94 ± 0.07&lt;sup&gt;bBC&lt;/sup&gt;</td>
<td>6.20 ± 0.62&lt;sup&gt;aAB&lt;/sup&gt;</td>
</tr>
<tr>
<td>12</td>
<td>5.92 ± 0.06&lt;sup&gt;cBC&lt;/sup&gt;</td>
<td>5.92 ± 0.13&lt;sup&gt;cBC&lt;/sup&gt;</td>
<td>6.15 ± 0.04&lt;sup&gt;bBC&lt;/sup&gt;</td>
<td>6.33 ± 0.54&lt;sup&gt;aAB&lt;/sup&gt;</td>
</tr>
<tr>
<td>24</td>
<td>6.04 ± 0.09&lt;sup&gt;AB&lt;/sup&gt;</td>
<td>6.22 ± 0.22&lt;sup&gt;BC&lt;/sup&gt;</td>
<td>6.28 ± 0.21&lt;sup&gt;AB&lt;/sup&gt;</td>
<td>6.65 ± 0.74&lt;sup&gt;aA&lt;/sup&gt;</td>
</tr>
<tr>
<td>48</td>
<td>6.24 ± 0.09&lt;sup&gt;A&lt;/sup&gt;</td>
<td>6.32 ± 0.17&lt;sup&gt;A&lt;/sup&gt;</td>
<td>6.96 ± 0.47&lt;sup&gt;A&lt;/sup&gt;</td>
<td>7.23 ± 1.27&lt;sup&gt;A&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Values are means ± standard deviation of 3 replicates.

<sup>a-c</sup>Means with different superscript letters within the row are significant different at \(P < 0.05\).

<sup>A-D</sup>Means with different superscript letters within the column are significant different at \(P < 0.05\).

Ammonia levels

Non significant change in the pouches water ammonia concentration was observed when the pouches exposed to ammonia at freezing temperature (-17 ± 3 °C). The overall model showed significant increases in ammonia levels in all pouches water \((P < 0.05)\) as exposure time and ammonia concentration increased at 21 ± 3 °C. Post hoc analysis showed that the main effect of exposure time and concentration were statically significant \((P < 0.05)\). Exposure time and concentration showed significant interaction \((P < 0.05)\), as did exposure time and bag type \((P < 0.05)\), concentration and bag type \((P < 0.05)\), and exposure time and bag type and concentration \((P < 0.05)\). Two methods for determining ammonia levels less than 10 ppm in the pouches: the ion selective electrode (ISE) and the indophenol method were evaluated. Hijaz et al. (2007) concluded that the ion selective electrode (ISE) and the indophenol methods are precise enough to
detect ammonia concentrations in meat. The ISE method was unstable and time consuming if used to measure low levels of ammonia. Although, the indophenol method offers many advantages: 1) high sensitivity (up to 0.03 ppm), 2) smaller sample size, 3) easily prepared reagent, and 4) short measuring time. Therefore, we concluded that the indophenol method was more suitable for our research needs. The research does show that the indophenol method gives more precise results when used to measure low levels of ammonia in ruminal fluid (Broderick and Kang 1980). The ammonia levels in water increased significantly ($P < 0.05$) with increasing exposure time (6, 12, 24, and 48 hr) and ammonia concentration (50, 100, 250, and 500 ppm). Figure 16 shows the average ammonia values from the control (0 ppm) for the exposed pouches at 500 ppm ammonia exposure for 48 hr. The maximum ammonia absorptions as they differed from the control were 77.7 ± 0.67 for E-2300, 5.94 ± 0.16 for LDPE, and 0.89 ± 0.20 for V-PA/PE. Table 10 shows the ammonia content of the exposed Cryovac E-2300, Table 11 shows LDPE, and Table 8 shows V-PA/PE. Ammonia levels increased significantly ($P < 0.05$) with increasing ammonia concentrations and exposure times. In addition, the interaction between exposure time and ammonia levels was significant ($P < 0.05$) for all types of packaging materials.

**Figure 16:** Water ammonia level means for Cryovac E-2300, LDPE, and vacuum (V-PA/PE) bags at 500 ppm for 6, 12, 24, and 48hr exposure at room temperature
**Table 10:** Cryovac E-2300 bag’s water ammonia levels at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature

<table>
<thead>
<tr>
<th>Time (hr)³</th>
<th>Ammonia concentration (ppm)₁,²</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>50</td>
<td>100</td>
<td>250</td>
<td>500</td>
</tr>
<tr>
<td>6</td>
<td>NDᵃ</td>
<td>NDᵃ</td>
<td>0.41 ± 0.15ᵇᵇ</td>
<td>0.80 ± 0.24ᵇᶜ</td>
</tr>
<tr>
<td>12</td>
<td>NDᵃ</td>
<td>0.45 ± 0.06ᵇᵇ</td>
<td>0.93 ± 0.17ᵇᶜ</td>
<td>1.78 ± 0.18ᵈ⁻</td>
</tr>
<tr>
<td>24</td>
<td>0.37 ± 0.01ᵇᵇ</td>
<td>0.87 ± 0.05ᵇᶜ</td>
<td>1.95 ± 0.48ᵈ⁻</td>
<td>4.20 ± 0.44ᵉ⁻</td>
</tr>
<tr>
<td>48</td>
<td>0.73 ± 0.05ᵇᵇ</td>
<td>1.46 ± 0.06ᶜᵈ</td>
<td>3.98 ± 0.66ᵉ⁻</td>
<td>7.77 ± 0.67ᶠ⁻</td>
</tr>
</tbody>
</table>

¹Values are means ± standard deviation of 3 replicates.

²Mean with different superscript letters significantly different at \( P < 0.05 \).

³At time 0 (control), ammonia level was 0.00 ppm for all concentrations.

---

**Table 11:** Ammonia levels in low density polyethylene bag’s water at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48hr exposure at room temperature

<table>
<thead>
<tr>
<th>Time (hr)³</th>
<th>Ammonia concentration (ppm)₁,²</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>50</td>
<td>100</td>
<td>250</td>
<td>500</td>
</tr>
<tr>
<td>6</td>
<td>NDᵃ</td>
<td>NDᵃ</td>
<td>0.27 ± 0.24ᵇᵇ</td>
<td>0.67 ± 0.26ᵇᶜ</td>
</tr>
<tr>
<td>12</td>
<td>NDᵃ</td>
<td>0.38 ± 0.04ᵇᵇ</td>
<td>0.73 ± 0.21ᵇᶜ</td>
<td>1.62 ± 0.30ᵈ⁻</td>
</tr>
<tr>
<td>24</td>
<td>0.31 ± 0.03ᵇᵇ</td>
<td>0.70 ± 0.04ᵇᶜ</td>
<td>1.45 ± 0.31ᵈ⁻</td>
<td>3.27 ± 0.12ᵉ⁻</td>
</tr>
<tr>
<td>48</td>
<td>0.58 ± 0.03ᵇᵇ</td>
<td>1.19 ± 0.04ᶜᵈ</td>
<td>2.80 ± 0.49ᵉ⁻</td>
<td>5.94 ± 0.16ᶠ⁻</td>
</tr>
</tbody>
</table>

¹Values are means ± standard deviation of 3 replicates.

²Mean with different superscript letters significantly different at \( P < 0.05 \).

³At time 0 (control), ammonia level was 0.00 ppm for all concentrations.
Table 12 shows the vendors reported transmission rates of water vapor and oxygen for Cryovac E-2300, LDPE, and V-PA/PE used in this research. Cryovac E-2300 is more permeable to oxygen (5000 cc/m²/24hr/atm) and water vapor (9.3 g/m²/24hr/atm) than LDPE and V-PA/PE. It is more permeable to ammonia than LDPE and V-PA/PE. However, we tested the E-2300 under normal conditions without exposing it to a vacuum or to heat, unlike the process described by Teixeira et al. (1986) that required vacuum sealing vacuum packaging machine and inserting of vacuum pack in a hot water (88.7 °C) to shrink.

LDPE had a slightly different permeability profile than E-2300. Oxygen (3743 cc/m²/24hr/atm) and water vapor (8.3 g/m²/24hr/atm) transmission rates were slightly lower than the E-2300. Although E-2300 film is thicker (3 mil) than LDPE (2.2 mil), E-2300 was more permeable to ammonia. The V-PA/PE was the least permeable to ammonia given its oxygen (60-70 cc/m²/24hr/atm) and water vapor (6.3 cc/m²/24hr/atm) transmission rates. This low permeability to ammonia could be due to the film’s two layers properties especially the polyamide one in the V-PA/PE, which is a good barrier to gases and aromatics (Mauer and Ozen 2004). When the pouches were exposed to ammonia gas at freezing temperatures (-17 ± 3 °C), no changes in water ammonia levels or pH values were observed, possibly because of the testing temperature. All pouches were permeable to ammonia at room temperature (21 ± 3 °C). The permeant temperature and pressure make food packaging more permeable to gases and vapors (Jasse et al. 1994). We applied ammonia gas at a specified flow rate, so we should see no change in the permeability transmission rate because of pressure.

Although ammonia has a high affinity to water, dissolving readily in water or any product containing water, the saturation point for ammonia remained unknown in this experiment because the reaction between ammonia and water in the pouches was continuous until the saturation point was reached.
Table 12: Physical properties of selected packaging materials used to evaluate ammonia permeability provided by suppliers

<table>
<thead>
<tr>
<th>Packaging Materials</th>
<th>Thickness (mil)*</th>
<th>Oxygen Transmission Rate (cc/m²/24 hr/atm)</th>
<th>Water Vapor Transmission Rate (g/m²/24hr/atm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cryovac E-2300a</td>
<td>3.0</td>
<td>5000 (23 °C/0% r.h.)</td>
<td>9.30 (37.7 °C/100% r.h.)</td>
</tr>
<tr>
<td>LDPEb</td>
<td>2.2</td>
<td>3743 (23 °C/0% r.h.)</td>
<td>8.32 (37.5 °C/100% r.h.)</td>
</tr>
<tr>
<td>V-PA/PEc</td>
<td>3.0</td>
<td>60-70 (23 °C/75% r.h.)</td>
<td>6.30-10.2 (23 °C/85% r.h.)</td>
</tr>
</tbody>
</table>

*One mil = 0.001 inch or 0.025 mm.

aCryovac Sealed Air Corporation; bManchester Packaging Co.; cUltravac Solutions LLC

CONCLUSION

Ammonia content and the pH levels in the exposed pouches were highest in Cryovac E-2300 and LDPE. Both showed similar permeability profiles when exposed to ammonia at same concentrations and exposure times, although E-2300 was slightly more permeable than LDPE. V-PA/PE was least permeable. The oxygen permeability for the packaging materials affected the ammonia permeability rather than the water vapor property. Temperature affected the rate of permeability. All pouches were permeable to ammonia at room temperature but not at freezing temperatures. Our results showed that the three types of meat packaging material used in this study protect meat from low levels of ammonia exposure during frozen storage.
FUTURE STUDIES

Although the three types of packaging materials tested here protected meat from ammonia contamination, the reaction between those materials and ammonia has not yet been clarified and more investigation is needed. In addition, more researches are needed to investigate the safety of chemical compounds that result from the reaction between ammonia and the packaging materials. Moreover, the literature provides little information about ammonia contamination at high concentrations (6,000 ppm to 10,000 ppm) for less than 8 hr under freezing conditions (-18 °C) that occur when ammonia leak happen during the time needed to relocate the contaminated product to an contaminated area. Because ammonia leaks often occur in frozen/refrigerated warehouses, information regarding the level of protection that the packaging materials can provide at 4 °C is essential and should be researched.
REFERENCES
Arnold M. 1993. The big chill ammonia finds its niche as a refrigerant. Food Processing 54(12):45,47.


PART III. FROZEN BEEF CONTAMINATION AFTER EXPOSURE TO LOW LEVELS OF AMMONIA GAS

ABSTRACT

Regulations and publications about food contaminated during ammonia refrigerant leaks provide limited information, which means that products are often held for an indeterminate period or condemned. Moreover, the scientific literature and regulatory agencies offer little guidance on disposing of products exposed to low levels of ammonia refrigerant gas. In this study, we evaluated meat contaminated with low levels of ammonia under frozen storage conditions. Fresh beef *Semitendinosus* muscles were trimmed of external fat and fabricated to 10H x 5W x 2.3D cm steaks. The steaks were exposed to 50, 100, 250, and 500 ppm ammonia gas (85 mL/min) in a plexiglass enclosure contained in a 1.3 CUF freezer, for 6, 12, 24, and 48 hr at -17 ± 3 °C. The ammonia in the contaminated meat was analyzed using the indophenol method, and the meat pH was measured according to the AOAC Int. # 981.12 method. Both ammonia levels and pH values increased significantly (*P* < 0.05) in the exposed meat with increasing exposure times and ammonia concentrations. The ammonia levels were 34.2, 51.5, 81.1, and 116 ppm, and the pH values ranged from 5.56 to 5.75 (control range 5.31-5.43) when the meat was exposed to 50, 100, 250, 500 ppm at 48hr. However, to declare meat contaminated with ammonia, its ammonia content should be more than 1000 ppm and its pH should be more than 1.0 unit than the background. Our results showed that meat content is low even with ammonia exposures as high as 500 ppm at freezing temperatures.

*Presented in part at IFT Annual Meeting, June 7 2009, # 060-12 (http://www.am-fe.ift.org/cms)
INTRODUCTION

Food safety and quality are objectives for food science, food industry, regulatory agencies, consumer and public health organizations. Research is intended to establish and clarify issues that would benefit the food industry, regulators, and members of the food industry to make an accurate decisions. Establishing critical control points and discovering corrective actions in a hazard analysis and critical control program (HACCP) may be used in insurance/customer claims involving food contamination (Al-Sahal 2003).

Food refrigeration is one of the oldest methods to preserve and keep food safe. Currently, the food industry uses refrigeration with ammonia to cool fruits, vegetables, poultry, fish, beverages, dairy products, milk, and meat (Lorentzen 1988). Meat production and processing requires large areas for fabrication, handling, and storage. Most fabrication facilities, slaughterhouses, and cold storage warehouses use ammonia as their refrigerant (Ross 1995; Sun 1998). With typical equipment failures and operator error, ammonia leaks will occur in refrigerated/frozen food storage facilities. Carelessness may be the problem, but more often equipment failures are the issue (Ostner 1986). Because published information and data on assessing meat contamination with ammonia are limited, especially after low levels of ammonia exposure, what to do in cases of meat contamination with low level of ammonia exposure remains uncertain and complex (Goodfellow et al.1978).

Several factors affect the degree of ammonia contamination in meat, including ammonia concentration, exposure time, temperature, humidity, type of meat, type of cut, condition of the meat surface, and the distance of the meat from the ammonia leakage point (Tuengerthal 1979). Frozen or refrigerated meat contaminated by ammonia is unsuitable for human use and can poison the consumer (Dworkin et al. 2004).
Two food borne illness outbreaks caused by ammonia were reported in the United States. The first outbreak was reported in two elementary school children in Wisconsin in 1985 within one hour of drinking contaminated milk (CDC 1986). The second outbreak occurred in several dozen school children in Illinois in 2002 within one hour of eating contaminated chicken tenders (Dworkin et al. 2004). Symptoms among children in both incidents included stomachache, headache, nausea, vomiting, diarrhea, and mouth burning. However, meat contaminated by low levels of ammonia (less than 1000 ppm) with no change in pH, taste, or smell is safe for human consumption (Goodfellow et al. 1978). No direct regulations specify how much ammonia is allowed in food products. Most forms of ammonia, such as bicarbonate, chloride, sulfate, alginate, citrate, phosphate, and ammonium hydroxide are generally recognized as safe (GRAS) when used under good manufacturing practice (GMP). Ammonium hydroxide is listed as GRAS for leavening, pH control, surface-finishing agent, and as a boiler water additive (21 CFR 2003). According to the Food Safety Inspection Services (FSIS) Directives # 7120.1, anhydrous ammonia is approved for use as an antimicrobial agent in lean finely textured beef according to GMP (USDA-FSIS 2009). In fact, Beef Product Inc. (BPI) treats meat trim with anhydrous ammonia to enhance the pH to eliminate pathogenic microorganisms like \textit{E. coli} O157:H7, \textit{Salmonella}, and \textit{Listeria Monocytogenes} (Niebuhr and Dickson 2003).

High concentrations of ammonia affect many quality properties of meat: darkened color (Curda and Hruby 1987; Bonne et al. 1993); increased pH (Anil 1971; Ireland 1988); increased water holding capacity (Anil 1971; Al-Sahal 1995); persistent pink color after cooking (Smolskiy et al. 1985; Shaw et al. 1992); and meat flavor (Guerrero and Arnau 1995; Ireland 1998). Ting and Henrickson (1986) studied the effect of ammonia on meat tenderness, flavor, taste, water holding capacity, and pH.
pH is not a reliable indicator of the amount of the ammonia absorbed because of the different buffering capacities of different foods (Kassem 1965). Hagyard et al. (1993) exposed lamb loins to 2 M ammonium hydroxide for 16 or 32 min and concluded that pH cannot be used as an indicator of ammonia contamination because 0.5 or 1.0 pH unit is within the normal pH range in lamb. Moreover, ammonia odor in contaminated meat is not a good indicator of the ammonia concentration in contaminated meat because ammonia reacts with water and ice to form ammonium ions, which do not have an odor (Dworkin et al. 2004).

Research has established the effects of high levels of ammonia concentrations on meat quality (Al-Sahal 2003), but for levels of ammonia concentrations below 500 ppm, no research was conducted and little information exists. Our research objectives were to measure the absorbency levels and changes in the pH in meat exposed to low levels of ammonia for different times under freezing conditions ranging from -17 ± 3 °C.

**MATERIALS AND METHODS**

**Experimental Design**

Fresh beef *Semitendinosus* muscle was randomly assigned to a various ammonia concentrations at different exposure times at -17 ± 3 °C. The average ammonia concentrations used were 0 (control), 50, 100, 250, and 500 ppm with exposure times of 0 (control), 6, 12, 24, and 48 hr. This procedure was replicated three times in a plexiglass box inside a freezer 1.3 CUF. The sequenced diagram below (Figure 17) shows the steps of the experiment.
Sample Preparation

Beef *Semitendinosus* muscle meat was obtained from a supermarket, the samples ranged from 2-3 wk postmortem *Semitendinosus* muscles (USDA choice) were trimmed of external fat and sliced into steaks (2.3 cm; Cabela’s meat slicer, Sidney, Nebraska, USA).

Beef *Semitendinosus* fabricated steak (10H x 5W x 2.3D cm)

↓

Freeze for 4-6 hr in a freezer (-18 °C)

↓

Inserted four steaks into exposure chamber into a freezer 1.3 CFT (-17 ± 3 °C)

↓

Applied flow rate of 85 mL/min of ammonia gas with concentrations of 50, 100, 250, and 500 ppm

↓

Withdrawn one steak each at 6, 12, 24, and 48 hr

↓

Thawed overnight at 4-8 °C in a refrigerator

↓

Measured pH value and ammonia level for each sample

Figure 17: A general flow chart for processing and exposure of the steaks to ammonia

The steaks were trimmed with a knife to dimensions of 10 cm × 5 cm (Figure 18). The steaks were placed in a freezer for 4-6 hr inside an air tight 1.7 L plastic container (Lock and Lock, Heritage Mint, Ltd. Scottsdale, Arizona, USA) until frozen. The meat samples were weighed after preparation two times, once when they were fresh and later after removing the ice layer from the top of the frozen samples. All samples were assigned randomly to a treatment with
four samples in each treatment replicate. At the end of each experiment, one meat sample was withdrawn at 6, 12, 24, and 48 hr of exposure to ammonia, the iced layer removed from the sample because the ammonia gas is approximable three times more soluble in ice than water at -30 °C (CSIRO 2002).

Figure 18: Beef *Semitendinosus* muscle trimmed of external fat and fabricated to dimensions of 10 × 5 × 2.3 cm

**Exposure Chamber**

A clear plexiglass box measuring 38W × 13.5D × 36H cm (18.5 L) was constructed as a treatment chamber. The chamber was tested for leaks and proper seal by inserting a ToxiRae PGM-30 Photo-Ionization Detector (PID) (RAE Systems, California, USA) into the chamber and locking the chamber. The detector was calibrated and operated according to the ToxiRae PGM-30 instruction manual. The ToxiRae PGM-30, which is portable, rechargeable, and small with a screen that displays readings, can detect ammonia levels during a specified time period and store the data. A concentration of 50 ppm of ammonia gas was introduced into the chamber from a calibrated 50 ppm (± 2%) ammonia gas cylinder. After a concentration of 50 ppm was reached,
the ammonia gas supply was stopped, and the chamber was locked. After 7 hr, the detector was pulled out of the chamber, and the readings were recorded. Another simple test checked the seal on the chamber: it was pressurized with air, locked, and immersed in a water bath. This ensured that the chamber sealed perfectly, with no observed bubbles.

The meat samples were inserted into the chamber through a round front door (15 cm in dia) and were attached to stainless steel hooks (Figure 19), suspended on two plexiglass bars attached horizontally between the left and the right sides of the box. On the back side of the chamber, two L-shape threaded fittings were mounted to serve as gas inlet and outlet, one on the upper right part and the other on the lower left part; in these positions, the ammonia gas could be homogeneously distributed throughout the chamber to ensure each sample had the same exposure rate.

**Figure 19:** Meat samples attached to stainless steel hooks and suspended on two plexiglass bars inside the freezer
Exposure System Design

The system (Figure 11) was used in this experiment to test meat packaging. The ammonia cylinder was connected to a 0-65 mm gas flow regulator inlet, and the gas regulator outlet was fitted to a T-shaped plastic connector. The exposure chamber and the detector were attached to the connector. The chamber outlet was assembled with a T-shaped tube connector hocked to a one-way plastic valve to prevent the ambient air from entering into the chamber and diluting the ammonia concentration, especially under freezing conditions. The valve also serves as pressure release device to prevent breakage in the chamber if pressure levels became too high during flushing. A second gas flow regulator connected to the other end of the one-way valve to monitor the ammonia exhaust flow rate.

The other end of the T-shaped connector was attached to a glass heat exchanger in order warm the ammonia gas before it reaches the detector, especially in the experiments which was performed under freezing temperatures. Combining together the system parts were assembled using plastic tubing.

Operation Procedure

The ammonia gas cylinders with concentrations of 50, 100, 250, 500, and 1000 ppm (calibrated concentrations were 48.8, 90.0, 238.7, 474.0, and 992 ppm; ± 2% in air) were used in the experiment. Two ammonia gas cylinders were used in each treatment. The 1000 ppm ammonia cylinder was used to flush the chamber, while maintaining a continuous exposure of the required ammonia concentration for the time needed using 50, 100, 250, and 500 ppm cylinders. A stainless steel single stage regulator was mounted on the low concentration cylinder to control the flow rate pressure of the ammonia gas into the gas flow regulator. The temperature inside the
chamber was monitored with a Smartbutton data logger. The chamber was flushed using ammonia gas from the higher ammonia concentration cylinder, and a reading was taken using MiniRae 2000 self sampling photo ionization detector. The detector inlet port was connected to the glass heat exchanger end, and the detector outlet port was connected to a tube going into the chamber inlet. To avoid losing ammonia from the chamber, the detector removed an ammonia gas sample from the chamber, detects it, and circulates it back into the chamber. The detector was operated and calibrated according to the MiniRae 2000 instructions manual. After the desired concentration inside the chamber was reached, the ammonia gas from the lower concentration cylinder was applied by adjusting the gas cylinder regulator to a pressure of 2.5 psi, and the flow rate was adjusted by adjusting the gas flow regulator to 85 mL/min. Ammonia exhaust flow rate was monitored by the second gas flow regulator, with the dial opened to the maximum to give free flow to the ammonia exhaust. At the end of each experiment, the ammonia flow was stopped, and the concentration inside the chamber was measured using the MiniRae 2000 detector (see Table 5).

**Indophenol Ammonia Measurement**

The ion selective electrode and the indophenol methods are precise enough to detect ammonia concentrations in meat (Hijaz et al. 2007). The indophenol method offers many advantages: high sensitivity (up to 0.03 ppm), small sample size (1.0 gram with recovery of 94 to 113%; Hijaz et al. 2007), and easily prepared reagent. Moreover, this method may be used with different protein precipitation acids. Sodium hypochlorite solution (10-15% available chlorine), sodium nitroferricyanide (III) dihydrate, and trichloroacetic acid were purchased from Sigma Chemical Co. (St. Louis, Missouri, USA). Phenol, sodium hydroxide, sodium phosphate dibasic
dodecahydrate, and a Genesys™ 10 visible spectrophotometer were purchased from Fisher Scientific Co. (Pittsburgh, Pennsylvania, USA). The reagents were prepared and stored according to Broderick and Kang (1980). While still frozen and after removing the glazed ice layer from the surface, the steak from each treatment was ground two times through a fine blade (0.4 cm) using an electric grinder (Rival, Fontana, California, USA).

One gram of the ground eye of round beef (*Semitendinosus*) was transferred to a 15 mL screw top disposable centrifuge tube to determine ammonia concentration using the indophenol method. Trichloroacetic acid TCA (10 mL; 2%) was added to precipitate protein from the sample, and the mixture was vortexed for 2 min and then allowed to stand for 10 min. The mixture was centrifuged at 2500 rpm for 5 min and then filtered through No.4 Whatman filter paper. The volume was adjusted to 14 mL by rinsing the filter paper with 2% TCA. Of the filtrate, 200 µL was transferred to a glass test tube and mixed with 2.5 mL phenol reagent and 2 mL hypochlorite reagent. The test tube was placed in a 95 °C water bath for 5 min and allowed to stand at room temperature to cool to 20-25 °C. Finally, the ammonia levels were measured at 630 nm using GenesysTm 10 visible spectrophotometer. To construct the standard curve, solutions of 1.0, 2.5, 5.0, 7.5, 10.0, 25.0, and 35.0 ppm ammonia as nitrogen (N) were prepared from the commercial standard 1000 ppm ammonium chloride (Thermo Electron Corporation, Beverly, Miami, USA); the blank was prepared using 200 µL of distilled water.

**pH Determination**

Two aliquots were used from each ground samples for pH analysis. The meat pH was measured according to AOAC Int. (2002) method #981.12 using a Denver Instruments UP-5 pH meter (single electrode, Denver Instrument, Denver, Colorado, USA). The meat samples were
measured in duplicate (Table 13). From each ground sample, 5 g were diluted with 45 mL of distilled water and blended for 30 sec at medium speed using a Waring blender (Waring Laboratory, Torrington, USA).

**Statistical Analysis**

SAS (Version 9.1, SAS Institute Inc., Cary, NY, USA, 2002) was used for the data analysis. The experimental design was a two way complete factorial design (4 × 4) analysis and the control time (0 hr) using PROC GLM/ TUKEY to determine significant differences among treatments at $P < 0.05$.

**Table 13:** Meat pH levels at 50, 100, 250, and 500 ppm exposure for 6, 12, 24, and 48 hr at $-17 \pm 3 \degree C$

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>Ammonia concentration (ppm)$^{1,2}$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>50</td>
</tr>
<tr>
<td>0 (control)</td>
<td>5.43 ± 0.00$^{bc}$</td>
</tr>
<tr>
<td>6</td>
<td>5.51 ± 0.06$^{cde}$</td>
</tr>
<tr>
<td>12</td>
<td>5.53 ± 0.03$^{cde}$</td>
</tr>
<tr>
<td>24</td>
<td>5.54 ± 0.07$^{def}$</td>
</tr>
<tr>
<td>48</td>
<td>5.62 ± 0.08$^{ef}$</td>
</tr>
</tbody>
</table>

$^1$Values are means ± standard deviation of 3 replicates.

$^2$Mean with different superscript letters significantly different at $P < 0.05$.  

81
RESULTS AND DISCUSSION

Meat pH levels

Overall, pH increased significantly ($P < 0.05$) as exposure time and ammonia concentration increased. Post hoc analysis showed that the main effect of time and concentration were statically significant ($P < 0.05$) for both factors although there was significant interaction between time and concentration ($P < 0.0142$). That significant interaction indicates that the pH increased with increasing time and concentration. The difference values in the pH from the control were 0.19, 0.17, 0.25, and 0.32 when the meat was exposed to 500 ppm at 6, 12, 24, and 48 hr. Generally, the alkalinity of ammonia increases the pH of meat (Anil 1971; Al-Sahal 2003). Figure 20 shows average pH values as a different from the control for the exposed meat for all treatments combinations. Al-Sahal (2003) had similar results after exposing meat to different ammonia concentrations for different times. His results showed that to raise the pH by 1.0 pH unit required 683 ppm of ammonia in the meat, which he achieved by exposing beef to 5000 ppm for 20 min. Moreover, similar results occurred when Al-Sahal (1995) exposed beef to even higher concentrations of ammonia.

Al-Sahal (1995) found 25,000 ppm exposure for 3 hr could increase the pH by 1.0 unit. In our study, the maximum increase in the pH for the control was 0.32 pH unit after exposing the meat to 500 ppm for 48 hr in freezing conditions. Moreover, Al-Sahal (2003) used a 0.9 cm thick steak, while Al-Sahal (1995) used a 1.27 cm thick steak. In this study, the beef steak was 2.3 cm thick. Differences in absorbed ammonia levels were likely due to the thickness of the meat, which is what Al-Sahal (2003) also stated.

Smolskiy et al. (1985) concluded that the pH of ground beef increased by 1 pH unit or more when exposed to 1000-1500 ppm (0.1 - 0.15%) ammonia. Other studies found that
treatment temperature had no effect on the pH of contaminated meat (Anil 1971; Herrmann and Johnston 1966).

Other differences in results may be due to how meat was exposed. This study exposed meat to ammonia continuously, whereas all previous studies exposed meat to ammonia using fixed volumes of ammonia concentrations.

![Figure 20: Meat pH mean levels as a different from control at 50, 100, 250, and 500 ppm exposure and 6, 12, 24, and 48hr at -17 ± 3 °C](image)

**Figure 20:** Meat pH mean levels as a different from control at 50, 100, 250, and 500 ppm exposure and 6, 12, 24, and 48hr at -17 ± 3 °C

**Meat Ammonia levels**

The ammonia content of exposed meat is presented in Table 14. Ammonia levels of samples increased significantly ($P < 0.05$) with increasing exposure time (6, 12, 24, 48 hr) and ammonia concentration (50, 100, 250, and 500 ppm). Figure 21 shows average ammonia values as a different from the control for exposed meat for all treatments combinations. The ammonia absorptions, as they differed from the control, were 34.2, 51.4, 81.1, and 116.3 ppm when meat was exposed to 50, 100, 250, and 500 ppm for 48 hr. The overall model showed significant
increases ($P < 0.05$) in ammonia levels for all samples as both exposure time and ammonia concentration increased. Post hoc analysis showed that the main effect of time and concentration are both statically significant ($P < 0.05$). There was a significant interaction between time and concentration ($P < 0.05$), which indicates that ammonia level increases depended on different exposure times and concentrations levels. The high affinity of the ammonia to water means that ammonia is mainly absorbed by meat water (75%). In this research, the saturation point for ammonia remained unknown because the reaction between ammonia and water in the meat was continuous until the saturation point was reached.

Al-Sahal (2003) exposed meat in an enclosure to a fixed volume of ammonia gas with different ammonia concentrations (500, 1000, 2500, and 5000 ppm) and different exposure times (5, 10, and 20 min) at room temperature (20-25 °C). His results indicate that ammonia concentrations increased 1-10 times in meat 0.9 cm thick. In our research, ammonia concentration increased by 1-5 times when the meat was exposed to a flow rate of 85 mL/min for continuous exposure at -17 ± 3 °C. The differences in the absorption levels between our results and Al-Sahal’s (2003) report may be due to differences in exposure temperature, sample size, and ammonia exposure concentrations and times.
**Table 14:** Meat ammonia levels exposed at 50, 100, 250, and 500 ppm for 6, 12, 24, and 48 hr at -17 ± 3 °C

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>50</th>
<th>100</th>
<th>250</th>
<th>500</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 (control)</td>
<td>51.5 ± 0.00^a</td>
<td>97.8 ± 0.00^{cde}</td>
<td>110 ± 0.00^{efgh}</td>
<td>141 ± 0.00^{gh}</td>
</tr>
<tr>
<td>6</td>
<td>57.1 ± 2.10^{ab}</td>
<td>108 ± 4.64^{def}</td>
<td>125 ± 3.21^{efgh}</td>
<td>175 ± 10.9^{ij}</td>
</tr>
<tr>
<td>12</td>
<td>68.5 ± 16.5^{abc}</td>
<td>124 ± 10.7^{efgh}</td>
<td>146 ± 8.71^{hi}</td>
<td>191 ± 7.11^{i}</td>
</tr>
<tr>
<td>24</td>
<td>74.9 ± 7.67^{abc}</td>
<td>130 ± 7.61^{fgh}</td>
<td>153 ± 10.9^{hi}</td>
<td>199 ± 8.46^{i}</td>
</tr>
<tr>
<td>48</td>
<td>85.8 ± 13.4^{bcd}</td>
<td>149 ± 10.5^{hi}</td>
<td>191 ± 12.2^{i}</td>
<td>257 ± 24.6^{k}</td>
</tr>
</tbody>
</table>

1Values are means ± standard deviation of 3 replicates.

2Meat protein was precipitated with 2% trichloroacetic acid.

3Mean with different superscript letters significantly different at $P < 0.05$.

**Figure 21:** Levels of meat ammonia as a different from control at 50, 100, 250, and 500 ppm exposure for 6, 12, 24, and 48 hr at -17 ± 3 °C
CONCLUSION

The maximum increase of ammonia in exposed meat was 116.3 ppm. While pH raised 0.32 units when meat was exposed to 500 ppm ammonia for 48 hr. According to Goodfellow (1978) recommendation to consider meat is contaminated with ammonia including: meat ammonia content should be more than 1000 ppm (0.1%), its pH should be more than 1.0 unit than the background, and its sensory panel score is less than 5.0 on a hedonic scale. Our research result showed that, even with a concentration of 500 ppm of ammonia and 48 hr of exposure time the meant ammonia content was less than 1000 ppm and the pH did not exceed by 1.0 unit than the normal back ground. However, the reaction between meat water and the ammonia is continuous until the saturation is reached, which remains unknown in this study. Furthermore, pH is not a good indicator of the level of ammonia contamination in meat; sensory analysis would be better to evaluate contaminated meat safety and acceptability.

FUTURE STUDIES

The literature provides little information about contamination with ammonia at high concentrations (6,000 ppm to 10,000 ppm) for less than 8 hr in frozen conditions (-18 °C). More studies are also needed to investigate the effects of low ammonia contamination (< 500 ppm) on meat quality and acceptability with exposure for 72 hr or more. Moreover, ammonia is often added to meat to retain its color during storage or to prolong the shelf life by reducing microbial count. Studies of food safety should evaluate levels of 4-methylimidazole due to its carcinogenicity to human, which forms in the presence of heat, reducing sugar, and ammonia, in marinated meat cooked on a grill. The literature provides little information regarding the mechanism of how ammonia affects the meat color during storage or cooking. In addition, more
research in needed concerning ways that the meat industry can treat/recycle the contaminated meat and make it safe and acceptable for human consumption. Ammonia is added to ground beef and steaks to preserve the color and to prolong the shelf life and considered by FDA as GRAS. However, there is no regulation about what levels of ammonia differentiate between contaminated and non contaminated meat. Extensive research is needed to determine the threshold values for contamination by ammonia in meat and meat products, such information would help the meat industry and the regulatory to facilitate a decision when ever a contamination with ammonia occurs. In addition, it would inform the costumer about the safety of a product if become contaminated with ammonia.
REFERENCES


SUMMARY AND CONCLUSIONS

The temperature affected the permeability of the tested packaging materials when exposed to ammonia. In freezing conditions experiment (-17 ± 3 °C), all packaging films showed no permeability to ammonia or changing in their water pH. While, in experiment performed in room temperature conditions (21 ± 3 °C), Cryovac E2300, LDPE, and V-PA/PE encountered permeability to ammonia and changed in their water pH. Each packaging material used in this research showed low permeability to ammonia. Due to the vacuum (V-PA/PE) multilayer film structure, it was least permeable to ammonia gas followed by LDPE and Cryovac E-2300. In addition, high permeability of Cryovac E2300 to oxygen affected the overall transmission rate through the film materials and thus resulted in greater transmission of ammonia compared to LDPE and V-PA/PE. However, the ammonia levels passing through the packaging materials were low even with 500 ppm ammonia exposure for 48 hr. The ammonia levels in the water in the pouches, affected the pH values with a ratio that corresponded to ammonia concentrations. The highest pH level, expressed as a difference from the control, resulted in Cryovac E-2300 and was 2.71 pH units with exposure of 500 ppm for 48 hr. However, our results showed that the three types of meat packaging material used in this study protect meat from low levels of ammonia exposure during frozen storage.

Meat ammonia and pH increased with increasing ammonia concentrations and exposure time. The maximum ammonia and pH values were 116.3 ppm and 0.32 pH unit with exposure of 500 ppm and 48 hr. The requirements for food to be considered safe for humans consumption, its ammonia content not more 1000 ppm and its pH is not more than 1.0 above the normal background. Our results showed that, the ammonia level and pH were within the acceptable levels. Frozen meat contaminated with 500 ppm ammonia gas for up to 48 hr can be considered safe for humans consumption.
APPENDIX

Ammonia gas flow rate calculations ................................................................. 94

FIGURES PAGE

Figure A.1. Specifications sheet for vacuum film ................................................... 95
Figure A.2. Specifications sheet for low density polyethylene film ......................... 96
Figure A.3. Specifications sheet for Cryovac E-2300 film ...................................... 97
Figure A.4. Temperature reading using Smartbutton data logger during the experiment ...... 98
Figure A.5. Standard curve used to calculate the ammonia concentrations in the pouches water
(first replicate) with 50 ppm ammonia exposure at 6, 12, 24, and 48 hr in room
temperature exposure .......................................................................................... 99
Figure A.6. Standard curve used to calculate the ammonia concentrations in the pouches water
(first replicate) with 100 ppm ammonia exposure at 6, 12, 24, and 48 hr in room
temperature exposure .......................................................................................... 99
Figure A.7. Standard curve used to calculate the ammonia concentrations in the pouches water
(first replicate) with 250 ppm ammonia exposure at 6, 12, 24, and 48 hr in room
temperature exposure ......................................................................................... 100
Figure A.8. Standard curve used to calculate the ammonia concentrations in the pouches water
(first replicate) with 500 ppm ammonia exposure at 6, 12, 24, and 48 hr in room
temperature exposure ......................................................................................... 100
Figure A.9. Standard curve used to calculate the ammonia concentrations in steaks with 50
ppm ammonia exposure at 6, 12, 24, and 48 hr at freezing temperature
exposure .............................................................................................................. 101
Figure A.10. Standard curve used to calculate the ammonia concentrations in steaks with 100 ppm ammonia exposure at 6, 12, 24, and 48 hr at freezing temperature exposure ................................................................. 101

Figure A.11. Standard curve used to calculate the ammonia concentrations in steaks with 250 ppm ammonia exposure at 6, 12, 24, and 48 hr at freezing temperature exposure (-17 ± 3 °C) ........................................................................................................... 102

Figure A.12. Standard curve used to calculate the ammonia concentrations in steaks with 500 ppm ammonia exposure at 6, 12, 24, and 48 hr at freezing temperature exposure (-17 ± 3 °C) ........................................................................................................... 102

Figure A.13. Pouches water ammonia levels. (a) at 50 ppm, (b) at 100 ppm, (c) at 250 ppm, and (d) 500 ppm ammonia exposure and 6, 12, 24, and 48 hr at 21 ± 3 °C ...... 103

Figure A.14. Pouches water pH levels as different form control for (a) Cryovac E-2300, (b) LDPE, and (c) vacuum V-PA/PE with 50, 100, 250, and 500 ppm ammonia exposure at 6, 12, 24, and 48 hr at 21 ± 3 °C ..................................................... 104

Figure A.15. Ammonia concentration over time measured with the ToxiRae PGM-30 in the sealed exposure chamber to test the chamber integrity ................................. 105

Table A.1. pH and ammonia levels inside the pouches water as different from the control without exposure at 6, 12, 24, and 48 hr at 21 ± 3 °C ..................................................... 105
Ammonia gas flow rate calculations

The flowing equations were used to calculate the ammonia equivalent air flow rate:

\[ Q_{\text{air}} = K_{\text{gas}} \times Q_{\text{gas}} \]

\[ K_{\text{gas}} = \sqrt{\frac{G \times T_{\text{act}}}{T_{\text{o}}} \times \frac{P_{\text{act}}}{P_{\text{o}}}} \]

Where

\( Q_{\text{air}} \) = equivalent air flow capacity at Standard Conditions (SPT).

\( Q_{\text{gas}} \) = maximum flow of metered gas.

\( G \) = specific gravity of metered gas which is 0.5963 for ammonia gas (\( G = 1 \) for air).

\( T_{\text{act}} \) = absolute temperature at flow condition, deg R or deg K.

\( T_{\text{o}} \) = absolute temperature at STP, deg R (530) or deg K (294)

\( P_{\text{act}} \) = pressure at flow conditions (psia)

\( P_{\text{o}} \) = pressure at STP (14.7 psia)

In our research the testing conditions were:

\( Q_{\text{gas}} = 45 \text{ mil/min}; T_{\text{act}} = 25 \text{ °C (298.2 °K)}; P_{\text{act}} = 2.5 \text{ psi} \)

\( G = 0.5963 \) for ammonia gas (\( G = 1 \) for air)

After Calculations

\( Q_{\text{air}} = 84.9 \text{ mil/min} \)
### Figure A.1. Specifications sheet for vacuum film

<table>
<thead>
<tr>
<th>Thickness: 3 mil 3-sided Seal Vacuum Pouch (PA 0.60 mil / PE 2.4 mil)</th>
<th>Test Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>μ</td>
<td>DIN 53122 23°C / 85% r.F.</td>
</tr>
<tr>
<td>μ</td>
<td>DIN 53380 23°C / 75% r.F.</td>
</tr>
<tr>
<td>μ</td>
<td>DIN 53380 23°C / 75% r.F.</td>
</tr>
<tr>
<td>μ</td>
<td>DIN 53455 23°C / 50% r.F.</td>
</tr>
<tr>
<td>μ</td>
<td>DIN 53445 23°C / 50% r.F.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Test Method</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>μ</td>
<td>75</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>μ</th>
<th>Water Vapor</th>
<th>Oxygen (O₂)</th>
<th>Carbon Dioxide (CO₂)</th>
<th>Nitrogen (N₂)</th>
<th>Mechanical Strength: MD</th>
<th>Tensile Strength: TD</th>
</tr>
</thead>
<tbody>
<tr>
<td>μ</td>
<td>6.3 – 10.2</td>
<td>60 – 70</td>
<td>180 – 200</td>
<td>41</td>
<td>41</td>
<td>31</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>μ</th>
<th>Sealing Temperature:</th>
<th>Temperature Consistency:</th>
</tr>
</thead>
<tbody>
<tr>
<td>μ</td>
<td>100 – 180</td>
<td>-50/+90</td>
</tr>
</tbody>
</table>

Information is based on our general experience and is given in good faith but no warranty is given with respect to such information.
CLEAR FILM (L D P E) low density polyethylene.

Water Vapor Transmission Rate = 0.5365 grams/100 in day  Conditions 100% RH, 37.5 C.

Oxygen Transmission Rate = 241.5 cc's / 100 in2 - day - atm  Conditions 0% RH 23 C

Film Gauge = 2.2 mils

Figure A.2. Specifications sheet for low density polyethylene film
### PROPERTIES

<table>
<thead>
<tr>
<th>Property</th>
<th>ASTM Test Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Appearance</td>
<td>Transparent</td>
<td></td>
</tr>
<tr>
<td>Forms</td>
<td>Bags, taped &amp; loose</td>
<td></td>
</tr>
<tr>
<td>Width</td>
<td>8-14</td>
<td></td>
</tr>
<tr>
<td>Length</td>
<td>12-30</td>
<td></td>
</tr>
<tr>
<td>Tensile</td>
<td>I-12, 760</td>
<td></td>
</tr>
<tr>
<td>Elong. (%)</td>
<td>1-172</td>
<td></td>
</tr>
<tr>
<td>Peak Load (%)</td>
<td>297</td>
<td></td>
</tr>
<tr>
<td>Energy to Break (J)</td>
<td>3.5</td>
<td></td>
</tr>
<tr>
<td>Tear Prop. (grams)</td>
<td>20-50</td>
<td></td>
</tr>
<tr>
<td>Sealing Tech</td>
<td>Heat Seal or Clip</td>
<td></td>
</tr>
<tr>
<td>Shrink Temp. Water °F</td>
<td>185-190</td>
<td></td>
</tr>
<tr>
<td>Unrestrained Shrink (%)</td>
<td>L-36 T-43</td>
<td></td>
</tr>
<tr>
<td>Low Temp. Prop (J)</td>
<td>Excellent</td>
<td></td>
</tr>
<tr>
<td>Min. Use temp °F</td>
<td>20°</td>
<td></td>
</tr>
<tr>
<td>Oxygen Transmission Rate (cc/m²/24 hrs., 1 atm) (%)</td>
<td>5000</td>
<td></td>
</tr>
</tbody>
</table>

This information represents our best judgment based on the work done, but the Company assumes no liability whatsoever in connection with the use of information or findings contained herein.

---

**Figure A.3.** Specifications sheet for Cryovac E-2300 film

---

To find out more about Cryovac's total systems approach to packaging, please contact Cryovac specialist at the nearest regional office.

**CRYOVAC**
Sealed Air Corporation
Cryovac, Duncan, SC 29334

1010 Commerce Way
Greensboro, CA 95953
(800) 620-0481
(916) 328-1206
(916) 328-1214

1055 Johnson Dr
Buffalo Dr, E 50000
(716) 520-6727

Room 250
150 Lenox Rd, Suite A-270
Denver, CO 80215
(303) 225-6658

157 North Concourse Way
Bethlehem, PA 18017-8955
(610) 494-5000

100 Rogers Bridge Rd., Build. B
Duncan, SC 29334
(864) 415-3000
(864) 415-3001

529 520-313229
Figure A.4. Temperature reading using Smartbutton data logger during the experiment
Figure A.5. Standard curve used to calculate the ammonia concentrations in the pouches water with 50 ppm ammonia exposure at 6, 12, 24, and 48 hr at 21 ± 3 °C

Figure A.6. Standard curve used to calculate the ammonia concentrations in the pouches water with 100 ppm ammonia exposure at 6, 12, 24, and 48 hr at 21 ± 3 °C
Figure A.7. Standard curve used to calculate the ammonia concentrations in the pouches water with 250 ppm ammonia exposure at 6, 12, 24, and 48 hr at 21 ± 3 °C

Figure A.8. Standard curve used to calculate the ammonia concentrations in the pouches water with 500 ppm ammonia exposure at 6, 12, 24, and 48 hr at 21 ± 3 °C
Figure A.9. Standard curve used to calculate the ammonia concentrations in steaks with 50 ppm ammonia exposure at 6, 12, 24, and 48 hr at -17 ± 3 °C

Figure A.10. Standard curve used to calculate the ammonia concentrations in steaks with 100 ppm ammonia exposure at 6, 12, 24, and 48 hr at -17 ± 3 °C
Figure A.11. Standard curve used to calculate the ammonia concentrations in steaks with 250 ppm ammonia exposure at 6, 12, 24, and 48 hr at -17 ± 3 °C

\[
y = 0.0561x + 0.0118
\]
\[
R^2 = 0.9999
\]

Figure A.12. Standard curve used to calculate the ammonia concentrations in steaks with 500 ppm ammonia exposure at 6, 12, 24, and 48 hr at -17 ± 3 °C

\[
y = 0.0684x - 0.0053
\]
\[
R^2 = 0.9998
\]
Figure A.13. Pouches water ammonia levels. (a) at 50 ppm, (b) at 100 ppm, (c) at 250 ppm, and (d) at 500 ppm ammonia exposure and 6, 12, 24, and 48 hr at 21 ± 3 °C
Figure A.14. Pouches water pH levels as different form control for (a) Cryovac E-2300, (b) LDPE, and (c) vacuum V-PA/PE with 50, 100, 250, and 500 ppm ammonia exposure at 6, 12, 24, and 48 hr at 21 ± 3 °C.
Figure A.15. Ammonia concentration over time measured with the ToxiRae PGM-30 in the sealed exposure chamber to test the chamber integrity.

The depletion in ammonia concentration over time is correlated to the detector performance not due to losing ammonia concentration from the chamber (personal communication).

Table A.1. pH and ammonia levels inside the pouches water as a different from the control without exposure at 6, 12, 24, and 48 hr at 21 ± 3 °C

<table>
<thead>
<tr>
<th>Time</th>
<th>Cryovac E-2300</th>
<th>LDPE</th>
<th>V-PA/PE</th>
<th>Cryovac E-2300</th>
<th>LDPE</th>
<th>V-PA/PE</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>0.03</td>
<td>0.00</td>
<td>0.20</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>12</td>
<td>0.09</td>
<td>0.00</td>
<td>0.18</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>24</td>
<td>0.06</td>
<td>0.00</td>
<td>0.18</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>48</td>
<td>0.05</td>
<td>0.00</td>
<td>0.17</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
</tbody>
</table>