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Energy Use and Energy Intensity of the U.S. Chemical Industry

Ernst Worrell, Dian Phylipsen, Dan Einstein, and Nathan Martin

Environmental Energy Technologies Division

April 2000



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Abstract

The U.S. chemical industry is the largest in the world, and responsible for about 11% of the U.S. industrial production measured as value added. It consumes approximately 20% of total industrial energy consumption in the U.S. (1994), and contributes in similar proportions to U.S. greenhouse gas emissions. Surprisingly, there is not much information on energy use and energy intensity in the chemical industry available in the public domain. This report provides detailed information on energy use and energy intensity for the major groups of energy-intensive chemical products.

Ethylene production is the major product in terms of production volume of the petrochemical industry. The petrochemical industry (SIC 2869) produces a wide variety of products. However, most energy is used for a small number of intermediate compounds, of which ethylene is the most important one. Based on a detailed assessment we estimate fuel use for ethylene manufacture at 520 PJ (LHV), excluding feedstock use. Energy intensity is estimated at 26 GJ/tonne ethylene (LHV), excluding feedstocks.

The nitrogenous fertilizer production is a very energy intensive industry, producing a variety of fertilizers and other nitrogen-compounds. Ammonia is the most important intermediate chemical compound, used as basis for almost all products. Fuel use is estimated at 268 PJ (excluding feedstocks) while 368 PJ natural gas is used as feedstock. Electricity consumption is estimated at 14 PJ. We estimate the energy intensity of ammonia manufacture at 39.3 GJ/tonne (including feedstocks, HHV) and 140 kWh/tonne, resulting in a specific primary energy consumption of 40.9 GJ/tonne (HHV), equivalent to 37.1 GJ/tonne (LHV). Excluding natural gas use for feedstocks the primary energy consumption is estimated at 16.7 GJ/tonne (LHV).

The third most important product from an energy perspective is the production of chlorine and caustic soda. Chlorine is produced through electrolysis of a salt-solution. Chlorine production is the main electricity consuming process in the chemical industry, next to oxygen and nitrogen production. We estimate final electricity use at 173 PJ (48 TWh) and fuel use of 38 PJ. Total primary energy consumption is estimated at 526 PJ (including credits for hydrogen export). The energy intensity is estimated at an electricity consumption of 4380 kWh/tonne chlorine and fuel consumption of 3.45 GJ/tonne chlorine, where all energy use is allocated to chlorine production. Assuming an average power generation efficiency of 33% the primary energy consumption is estimated at 47.8 GJ/tonne chlorine (allocating all energy use to chlorine).

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1. Introduction

In 1994 the manufacturing sector consumed 26 EJ of primary energy in the United States, almost one-quarter of all energy consumed that year (U.S. DOE, EIA 1997).¹ Within manufacturing, a subset of raw materials transformation industries (primary metals, pulp and paper, cement, chemicals, petroleum refining) require significantly more energy to produce than other manufactured products.

In this report we study the energy consumption and intensity in the U.S. chemical industry. The chemical industry is one of the largest energy consuming industrial sub-sectors in the U.S. The chemical industry is complex, encompassing the production of over 50,000 chemical compounds. The industry is also an important part of the global economy, accounting for approximately 7% of global income and 9% of international trade (WEC, 1995).

The chemical industry produces many intermediate compounds that are used as the basis for many chemical products. For example, ethylene, one of the most important bulk chemicals from an energy point-of-view, is used to produce products varying from solvents to plastics. Also, many processes in the chemical industry produce different co-products. This makes energy analysis of the chemical industry more complicated compared to other industries.

In this report we study the energy consumption of the U.S. chemical industry. We assess current energy consumption and production levels. This information has been used to develop a detailed baseline energy consumption and intensity for three of the most energy intensive and energy consuming chemicals, i.e. ethylene, ammonia and chlorine. We focus on the year 1994, as this is the last year for which the Energy Information Administration has published energy consumption data (the Manufacturing Energy Consumption Survey 1994). No detailed studies on the energy intensity of the U.S. chemical industry have recently been published. The energy intensity baseline will be used for future detailed analyses of the potential of energy efficiency improvement, and carbon dioxide emission reduction in the U.S. chemicals industry.

We start with a discussion of the major energy consuming sub-sectors and products in the chemical industry (Chapter 2). This is followed by detailed energy analyses of energy use and energy intensities of three major bulk chemicals, i.e. ethylene (Chapter 3), ammonia and nitrogenous fertilizers (Chapter 4) and chlorine (chapter 5). The reliability and usefulness of the results are discussed in Chapter 6.

2. Energy Use in the Chemical Industry

Primary energy use by and associated CO_2 emissions from the U.S. chemicals sector (SIC 28) were roughly 20% of total manufacturing primary energy use and CO_2 emissions in 1994. The chemical industry produced 11% of total manufacturing value added in the U.S. (see Table 1). Figure 1 depicts carbon dioxide emissions from energy use and value added for SIC 28 between 1985 and 1994.² Emissions have grown at an annual rate of 2.9% during that period, while value added has increased at a slightly faster rate of 4.6%.

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¹ In this report we will use SI-units. To convert from GJ to MBtu, multiply by 0.95; to convert from metric tons to short tons, multiply by 1.1; to convert from GJ/metric ton to MBtu/short ton, multiply by 0.86.

² Carbon emissions estimates include fuels and electricity for energy consumption and feedstocks used for ammonia manufacture less carbon embedded in urea. For other chemical products most of the carbon is not emitted as CO_2 . Emissions factors are given by EIA (1996).

Table 1. Primary Energy Use (excluding feedstocks), Carbon Dioxide (CO₂) Emissions from Energy Use, and Value Added for U.S. Chemicals and U.S. Manufacturing in 1994. Source: EIA (1997); EIA (1996); BOC (1998).

Sector	Primary Energy (Tbtu)	CO ₂ Emissions (MtC)	Value Added (Billion \$1992)
Chemicals (SIC 28)	5,141	77	172
Manufacturing total	26,047	398	1,570
Share of SIC 28 to Total Manufacturing	20%	19%	11%

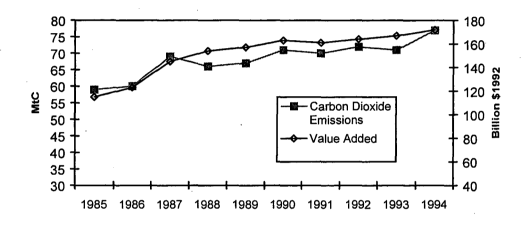


Figure 1. Carbon Dioxide Emissions from Energy Use, and Value Added for U.S. Chemicals, 1985-1994. Source: EIA (1997); EIA (1996); EIA (1994); EIA (1991); EIA (1988) BOC (1998).

The U.S. Manufacturing Energy Consumption Survey provides energy consumption information for selected chemicals subsectors. The subsectors that accounted for the largest share of primary energy (see also Figure 2 and Table 2) in 1994 were: industrial organic chemicals not elsewhere classified (SIC 2869), industrial inorganic chemicals not elswhere classified (SIC 2819), plastic materials and resins (SIC 2821), and nitrogenous fertilizers (SIC 2873), industrial gases (SIC 2813) and alkalies and chlorine (SIC 2812). With the exception of plastics and resins, the primary energy requirements associated the production of one dollar of shipped product are three to nine times greater than for the average of the chemicals sector as a whole. This demonstrates that there are several highly energy-intensive production processes within these subsectors that are often used to produce intermediate chemicals. These subsectors will be described in greater detail below.

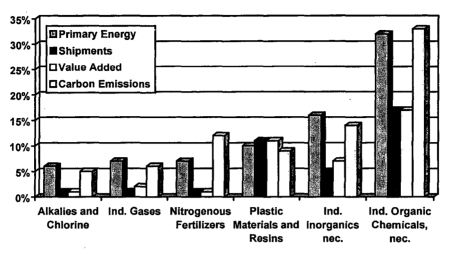


Figure 2. Share of Primary Energy Use, Shipments, Value Added, Carbon Dioxide (CO₂) Emissions, for Selected U.S. Chemicals Subsectors, 1994. Source: EIA (1997); EIA (1996); BEA, (1998); BOC (1998).

Table 2. Primary Energy Use, Shipments, Value Added, Carbon Dioxide (CO₂) Emissions, for Selected U.S. Chemicals Subsectors in 1994. Sources: EIA (1997); EIA (1996); BEA (1998), BOC (1998). Energy use and emission data exclude feedstocks.

Chemical Subsector	SIC Code	Primary Energy (PJ)	Shipments (Millions \$1992)	Value Added (Millions \$1992)	CO ₂ Emissions from Energy Use (MtC)
Industrial Organics, nec	2869	1,653	53,983	22,623	25
Industrial Inorganics, nec	2819	830	15,638	9,491	11
Plastic Materials and Resins	2821	518	35,307	14,307	7
Nitrogenous Fertilizers	2873	344	3,648	1,689	10
Industrial Gases	2813	364	3,128	2,188	4
Alkalies and Chlorine	2812	286	2,034	951	4
Others		1,146	203,316	120,796	16
Total Chemicals	28	5,141	317,054	172,045	77

2.1 Industrial organic chemicals, not elswhere classified (SIC 2869)

As noted above, industrial organic chemicals not elsewhere classified accounted for the largest share of energy use in the chemicals sector as a whole in 1994 (33%). From an energy perspective, some of the key chemical products included within this category are ethylene and other steam cracking derivatives (propylene and butadiene) and methanol.

Ethylene and its derivatives are important petrochemicals in the U.S. economy, and are feedstocks for many plastics and resins products produced in SIC 2821 (see below) as well as fibers and detergents. In 1994, ethylene was the fourth largest chemical produced while propylene was the seventh largest chemical produced (Chemical and Engineering News, 1995). The U.S. is currently the worlds largest ethylene producer accounting for 28% of world installed capacity (Oil and Gas Journal, 1997. Since 1974, ethylene production has grown by 3% annually while propylene has grown by over 4% annually. Propylene has grown more rapidly in the last decade—5% per year. Overall, however, industrial organic chemicals as a group have grown more slowly (2% per year) since 1985, due in part to a drop in output in 1996 (Chemical and Engineering News, 1997).

In ethylene production, hydrocarbon feedstocks (such as ethane or naphtha) are heated in pyrolysis furnaces, separated into gaseous products, and then rapidly cooled, compressed, and purified into final products with the largest energy requirements required in the pyrolysis, refrigeration and rapid cooling (WEC, 1995). Lighter feedstocks such as ethane produce higher ethylene yields. More severe processing conditions (higher temperatures and pressures) used on heavier feedstocks require more energy to crack but also result in a more co-product yields (methane, butadienes, benzene, and toluene) (Phylipsen et al, 1998a). In the US, ethane remains the primary feedstock used in steam cracking, followed by propane, naphtha and gas oil. Chapter 3 describes energy use and energy intensity for ethylene manufacturing in more detail.

Methanol is produced through the reaction of carbon monoxide and hydrogen, with the production of hydrogen being a significant energy use. Methanol demand has been driven up in recent years due to increasing demand for Methyl tertiair-butyl ether (MTBE) as a reformulated gasoline additive. Growth over the last decade has averaged 8.5% annually. However, MTBE-use in the U.S. will be phased out in the foreseeable future due to water pollution problems associated with MTBE-use. Estimated energy intensity for methanol (including feedstocks) is 38 GJ/tonne with most of the energy use being used for hydrogen production (Lipinsky and Ingham, 1994).

2.2 Industrial inorganic chemicals, not elsewhere classified (SIC 2819)

Industrial inorganic chemicals not elsewhere classified accounted for the second largest share of carbon emissions within the US chemicals sector in 1994 (14%). This category includes a wide variety of inorganic chemicals including sulfuric and hydrochloric acid, potassium fertilizers (potash), alumina, and aluminum oxide. Hydrochloric acid and Potash are produced in bulk quantities and were among the top 40 chemicals produced in the US in 1994.

2.3 Plastic Materials and Resins (SIC 2821)

While not as energy-intensive as the production of bulk chemicals, the production of plastic materials in SIC 2821 accounts for a significant share of carbon dioxide emissions (9% of chemical industry emissions in 1994) due to primarily the large volume of production. Some of the main plastic products include polyethylene, (low and high density), polypropylene, polystyrene, and polyvinyl chloride. Ethylene (within SIC 2869) is used as a primary feedstock for polyethylene manufacture. Estimates of per ton energy requirements for polymerization processes are shown in Table 3 below. Estimates from Worrell et al (1994a), except for polystyrene, likely reflect best practice levels for the US.

Product	Estimate 1*	Estimate 2*
Polyethylene (LDPE)	9.3	1.6
Polypropylene	10.5	1.2
Polystyrene	9.3	11.3
Polyvinyl Chloride*	11.6	9.9

Table 3. Energy Requirements for Plastics Production (GJ/tonne)

*Estimate (1) is based on Lipinsky and Wesson (1995); Estimate 2 is based on Worrell et al. (1994a).

Table 4 shows the production growth rates of high and low density polyethylene, polypropylene, polystyrene, and PVC over the last two decades. As table 4 shows, plastics production has grown rapidly at rates of over 3-8% since 1974, with particularly strong growth in PVC, polypropylene, and high-density polyethylene. Given the continued demand for plastics in a variety of end uses, we expect continued growth in this subsector.

Table 4. Average Annual Growth Rate of selected U.S. plastic products. Source: Chemical and Engineering News (1985); Chemical and Engineering News (1997).

Product	1974-1996	1985-1996
Polyethylene – low density	4.0%	4.3%
Polyethylene - high density	6.9%	5.8%
Polypropylene	7.9%	8.0%
Polystyrene	2.7%	3.7%
Polyvinyl Chloride and	4.7%	6.3%
copolymers		

2.4 Industrial Gases (SIC 2813)

The production of industrial gases is a relatively energy intensive process, and this sector accounted for 6% of carbon dioxide emissions from within the chemical sector in 1994. Nitrogen and oxygen production have historically counted for about half the value of shipments in the subsector (Lipinsky and Ingham, 1994) and are typically the second and third largest produced chemicals. These two gases are produced mainly through cryogenic air separation where air is cooled and pressurized until it becomes a liquid with the various gases extracted through fractional distillation. However other technologies are increasingly being used such as pressure swing absorption and membrane separation. Energy consumption for oxygen production has been estimated at 2.0 GJ/tonne (OTA, 1993). Assuming these intensities, energy consumption for these two gases in 1994 would account for over 85% of SIC 2813. Since 1985 the production of nitrogen and oxygen has grown by over 4% annually.

2.5 Nitrogenous Fertilizers (SIC 2873)

The production of ammonia, a key component in the manufacture of nitrogenous fertilizers, is a highly energy intensive process. Roughly 80% of ammonia production is used as fertilizer feedstock in the U.S. (Lipinsky and Ingham, 1994). Like methanol, ammonia is produced through the high-pressure synthesis of gases (carbon dioxide, hydrogen, and nitrogen). Ammonia production accounts for over 85% of the energy consumption in SIC 2873. The production of these products has grown on the order of 1% annually since 1974.

2.6 Alkalies and Chlorine (SIC 2812)

One of the main uses of chlorine (around 30%) is as an intermediate feedstock for polyvinyl chloride (PVC) which has been growing rapidly over the past decade (Lipinsky and Ingham, 1994). Chlorine is also used as a bleaching agent in pulping operations. The production of chlorine is a highly electricity-intensive process requiring between 3065 kWh/tonne and 3960 kWh/tonne depending on the cell type (Pletcher and Walsh, 1989). In the process an electric current is used to separate molecules into their constituents. The products of the process include chlorine and caustic soda. Table 5 shows estimated energy consumption for the various cell types. Steam consumption in diaphragm cells is greater since more energy is required to concentrate the brine. We assume a weighted energy intensity of 12.8 GJ/tonne based on the 1994 shares (see Chapter 5).

Table 5. Energy Consumption for Chlorine production with Mercury, Diaphragm, and Membrane Cells (kWh/tonne and GJ/tonne) and estimated shares in the U.S. in 1994 (percent). See also chapter 5. Electricity is converted to primary energy using a generation efficiency of 33%. All energy use in Table 5 is allocated to chlorine production. In reality the product-mix consists of approximately 55% caustic soda and 45% chlorine (on weight-basis).

Component	Units	Mercury	Diaphragm	Membrane
Steam	GJ/tonne	0.1	2.6	0.5
Electricity	kWh/tonne	3420	3140	2720
Total – Primary Energy	GJ/tonne	37.4	36.8	30.2
Shares in U.S. (1994)	Percent	15%	75%	8%

2.7 Summary of Energy Use for Key Chemicals

Table 6 summarizes the energy consumption and estimated energy intensity of the key chemical compounds produced in the U.S. chemical industry. A few chemical products dominate energy use, i.e. ethylene and co-products, ammonia, chlorine and methanol. In the following sections we will assess energy intensities for the production of ethylene, ammonia and chlorine in more detail.

Table 6. 1994 Estimated U.S. final energy consumption (HHV) for selected key chemicals. (including feedstocks)

Product	Estimated Final	1994	Estimated Total	Percent Share of
	Energy SEC	Production	Energy Use in	SIC 28 Energy Use
			1994	
	(GJ/tonne)	(million tonnes)	(PJ)	(%)
Ethylene and co-products	67.5	26.2	1768	29.3%
Methanol	38.4	4.9	188	3.1%
Polyethylene	9.3	5.7	53	0.9%
Polypropylene	10.5	4.4	. 45	0.7%
Polyvinyl Chloride	11.6	5.4	62	1.0%
Polystyrene	9.3	2.6	24	0.4%
Nitrogen	1.8	28.6	49	0.8%
Oxygen	1.8	22.7	44	0.7%
Ammonia	39.8	16.2	645	10.5%
Urea	2.8	7.6	21	0.3%
Chlorine	19.2	11.1	213	3.5%
Total			3112	51.5%

¹Co-products include propylene, benzene, and butadiene. SEC reflects energy per ton of all high value products from steam cracking.

²The SEC, or Specific Energy Consumption estimates are preliminary. Sources for SEC are as follows: ethylene and co-products (see Chapter 3), methanol and urea (Lipinksy and Ingham, 1994), polyethylene, polypropylene, polyvinyl chloride (Lipinsky and Wesson, 1995), Nitrogen and Oxygen (OTA, 1993), ammonia (see Chapter 4), chlorine (see Chapter 5). Production estimates are from CMA (1996).

3. The Ethylene and Petrochemical Industry

This chapter reflects an in-depth analysis of a part of one of the energy-intensive sub-sectors: the petrochemical. In the petrochemical industry mostly relatively simple organic chemicals are produced such as ethylene, propylene and benzene. These chemicals (some through intermediates, e.g. mono vinyl chloride or styrene) form the building blocks for many products such as plastics, resins, fibers, detergents, etc.

The single most energy-consuming step in the petrochemical industry is the steam cracking of hydrocarbon feedstocks to produce ethylene, propylene, butadiene and aromatics (benzene, toluene and xylenes). Recent estimates of global energy consumption for the production of ethylene and co-products are not available. In 1990 energy consumption was estimated to be about 1 EJ (or 950 TBtu, excluding feedstock energy consumption), with ethylene production amounting to 50 Million tonnes (WEC, 1995). By 1997, global ethylene production had risen to 70Mt/a (O&GJ, 1998). The U.S. is currently the world's largest ethylene producer, accounting for about 28% of world capacity (Rhodes, 1997).

In section 3.1 we first discuss the major process used to produce ethylene and its co-products, followed by a discussion of the U.S. petrochemical industry (section 3.2) and its energy consumption and intensity (section 3.3).

3.2. Process Description

In the cracking process, hydrocarbon feedstocks are preheated in the convection section to 650° C (using fuel gas and waste heat), mixed with steam and cracked in the radiant section at a temperature of about 850° C (Worrell et al., 1994). Subsequently, the gas mixture is rapidly cooled to 400° C (or quenched) to stop the reaction, during which process high pressure steam is produced. Injection of water further decreases the temperature to about $40-50^{\circ}$ C and a condensate, rich in aromatics, is formed. The liquid fraction is extracted, while the gaseous fraction is fed to a series of low temperature, high pressure distillation columns. The fractionation sequence varies from plant to plant (Zeppenfeld et al., 1993). Figure 3 shows a typical arrangement.

Feedstocks used in steam cracking are ethane, LPG, naphtha, gas oils (GOs) and sometimes coalderived feedstocks. Many of the installations used today can handle different (if not all) types of feedstock (Chemfacts, 1991). The choice for a particular feedstock, together with processing conditions (heat, pressure, steam dilution rate) will determine the yield of ethylene, propylene and other co-products in steam cracking. Table 7 shows how product yield varies with feedstock type (Chauvel and Lefevbre, 1989).

Product				Feedstock		
	Ethane	Propane	Butane	Naphtha	Atmospheric gas oil	Vacuum gas oil
Hydrogen (95% purity)	9	2	2	2	1	1
Methane	6	28	22	17	11	9
Ethylene	78	42	40	34	26	21
Propylene	3	17	17	16	16	14
Butadiene	2	3	4	5	5	5
Pyrolysis gasoline Of which:	2	7	7	19	18	19
- Benzene	2	3	3	7	6	4
- toluene	0	1	1	3	3	3
Fuel oil	-	1	2	5	18	25

Table 7. Influence of feedstock on steam cracker yield¹ (weight %) (Chauvel and Lefebvre, 1989)

Note¹ These values are obtained at high severity and with recycling of unconverted ethane/propane streams

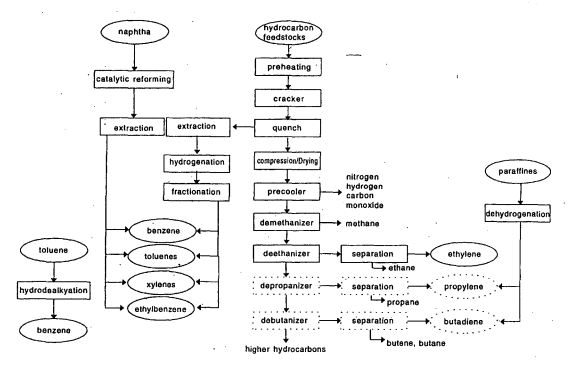


Figure 3. Process routes for the production of ethylene and its co-products (Phylipsen et al., 1998).

If ethane is used as feedstock, almost no propylene, butadiene and aromatics are formed as byproducts. Therefore, other processes are used for producing these chemicals in countries that predominantly use ethane cracking. In the U.S. propylene is also retrieved from refinery fractions (resulting from fluid catalytic cracking or hydro cracking). Another possibility is catalytic dehydrogenation of propane, but the share of this process is negligible at this moment. Aromatics (such as benzene, toluene and xylenes, or BTX) are usually produced by hydrogenation and desulphurisation of pyrolysis gasoline, after which the different components are separated by solvent/solvent extraction. Another route for BTX production is catalytic reforming of naphtha, followed by extraction of individual compounds out of the reformate. Furthermore benzene can be produced by hydro-de-alkylation of toluene. The different routes for petrochemicals production are shown in Figure 3.

3.2 The U.S. Petrochemical Industry

The U.S. is currently the largest ethylene producer in the world, producing about 20 Million tonnes in recent years (1994-1996) (Chemical and Engineering News, 1997). Table 8 compares the U.S. ethylene capacity and feedstock mix with other major producers. The shares of feedstocks used in U.S. ethylene production in 1995 are listed in more detail in Table 9 (Chemfacts, 1991; O&GJ, 1994, 1997, 1998). These tables show the relatively high share of ethane used for ethylene production in the US compared to the global average feedstock input (Appendix A provides a plant-by-plant overview of feedstock input). The difference is caused by the availability of abundant, cheap ethane as a by-product of natural gas and oil production.

Country	try Share of world ethylene capacity	
U.S.	28%	42%
Western Europe ⁴	22%	8% ¹
Japan	8%	0% ²
FSU	5%	
China (including Taiwan)	5%	
South Korea	5%	0%3
Saudi Arabia	4%	76% ³
Canada	4%	72% ³
Brazil	3%	3% ³
Mexico	2%	100% ³
India	2%	
Others	12%	
World	90 Mt	28%

Table 8. The US ethylene capacity and feedstock input compared to other major producers (all countries/regions with a capacity of over 1 Million tonne/year are shown (O&GJ, 1998), unless otherwise noted)

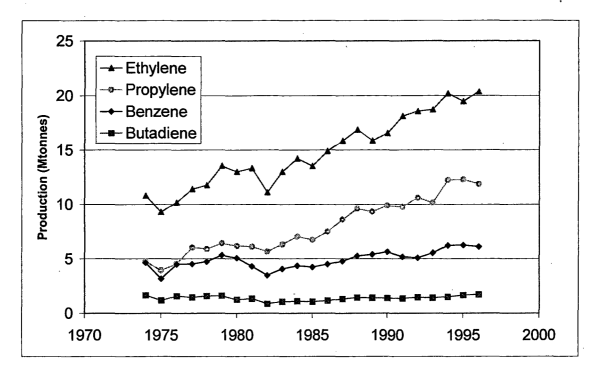
Notes: 1) in 1989 (Joosten, 1998); 2) 90% naphtha, 10% LPG in 1994 based on capacity (HP, 1995); 3) typical feedstock, based on 1996 plants (Rhodes, 1997); 4) includes all ethylene producers in Western Europe.

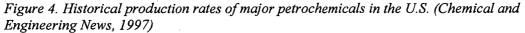
Table 9. Estimated feedstock input for the U.S. and total world ethylene production capacity in	
1995 (as % of ethylene capacity) (Chemfacts, 1991: O&GJ, 1994, 1998; Rhodes, 1997)	

	Ethane	Propane/Butane	Naphtha	Gas oil	Others
U.S.	42%	25%	20%	10%	3%
World	28%	12%	52%	6%	2%

Figure 4 shows historical production rates of ethylene and its co-products for the U.S. The average annual growth rate for ethylene production has been around 3% since 1974. Propylene production increased 4% per year (Chemical and Engineering News, 1997b). In recent years, global capacity for ethylene production has outgrown ethylene demand (O&GJ, 1998; Chemical Market Reporter, 1998). In spite of already existing over-capacity, global ethylene capacity is expected to grow from 89 Million tonnes/year in 1997 to 103 Million tonnes/year in 2000. According to Chemical Markets Associates Inc. (a Houston-based consulting firm) ethylene demand in 2000 is expected to be about 90 Million tonnes/year (O&GJ, 1998). Utilisation rates are expected to drop below 90%, with ethylene margins decreasing 40-50% compared to the March 1998 levels (Chemical Market Reporter, 1998). The expected capacity growth in the U.S. of about 7 Million tonnes/year between 1998 and 2002 is second only to that in East Asia (Rhodes, 1997; O&GJ, 1998).

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Average plant capacity for the U.S. is about 600,000 tonnes ethylene per year (O&GJ, 1998). A typical size for new U.S. plants is somewhat higher: 800-900,000 tonnes per year. Ethane-based plants generally have a higher capacity than liquid (naphtha/gas oil) crackers. Liquid crackers require a higher throughput of feedstock per unit of ethylene produced and get more easily contaminated. Larger plants are, generally, more efficient (Solomon, 1995).

Age of technology

Estimates of the average age and age distribution of ethylene production facilities vary. For example, according to Exxon Chemical Company Basic Chemicals Americas, as of 1998 75% of U.S. ethylene capacity is no older than 10 years (Chemical Market Reporter, 1998b). However, Los Alamos National Laboratory reports that 75% of current plants in the U.S. olefins industry are older than 20 years (LANL, 1996). Often plant sites and the basic design of a plant may be much older (dating back to 1962, see also Appendix A). It is very difficult to determine which part of the installations has been replaced since the time of construction. As might be expected newer plants are, generally, more efficient than older ones. The difference between plants built in the early seventies and the early nineties is on average 40% (worldwide) (Solomon, 1995).

Economics

Manufacturing plants fed with ethane and propane can be constructed at much lower investment costs than naphtha crackers (Chemical Market Reporter, 1998b). For a 450,000 tonnes/year plant, investment costs vary from \$430/tonne ethylene for ethane crackers, \$645/tonne ethylene for naphtha crackers to \$755/tonne ethylene for gas oil crackers (1993-\$) (Gielen et al., 1996)³. For larger plants, investment costs (per tonne of ethylene) will be lower. Production costs vary from

³ These estimates are based on Dutch production plants. However, according to (Gielen et al., 1996) investment costs are similar for U.S. Gulf Coast plants.

\$295-430/tonne ethylene for ethane-based production (with the US on the high side of the range), about \$755/tonne ethylene for naphtha crackers to \$940/tonne ethylene for gas oil-based production (Gielen et al., 1996). The variation in production costs are caused by differences in e.g. investment costs, labor costs, feedstock and energy prices, ethylene yield (per unit of feedstock). Heavier feedstocks will result in by-products, generating additional revenues. Investment costs for flexible crackers (able to use different types of feedstock) are generally higher than dedicated crackers (designed for one particular feedstock). However, flexible crackers have the opportunity to choose a different (cheaper) type of feedstock if feedstock prices change Overall profitability of flexible crackers versus dedicated crackers depends on the price developments in different feedstocks and different products (ethylene versus propylene, butadiene and aromatics). In August 1998 cash margins for liquid-based crackers were higher than for ethane-based crackers (see Table 10) (Bonner & Moore, 1998b).

Table 10. Cash margins for ethylene production for different feedstocks during August 1998 (Bonner & Moore, 1998b).

Feedstock	Cash margin (cents per kg) ¹
Ethane	11.3-11.8
Propane	13.3-14.0
Butane	14.2-14.7
Naphtha	14.2-14.9
Gas oil	16.4-18.2

Note: Includes feedstock costs, co-product credits, variable costs, fixed costs and ethylene price

Purchased energy costs in North America in 1995 for the production of ethylene amounted to \$2.11/GJ (\$2.23/MBtu), compared to \$3.90/GJ (\$4.11/MBtu) for Europe and \$6.02/GJ (\$6.35/Mbtu) for Asia (Solomon, 1995). Between 1993 and 1995 energy costs in North America decreased 20%, while in the same time energy costs in Europe rose by 3%. Differences in energy prices are caused by the indigenous availability of energy resources, tax levels and competition of specific energy carriers for other applications (e.g. the application of fuels for space heating).

Import/export

1

U.S. ethylene net imports amounted to 230,000 tonnes in 1995 (equal to about 1% of U.S. production). Since then net imports have been decreasing to 5,000 tonnes in 1997 and as of 1998 the U.S. is expected to be a net exporter (Bonner & Moore, 1998). Exports, however, are limited (less than 1% of total production). In general, transport costs will be too high to allow large-scale exports over long distances. As an example, the transportation costs for ethylene from the U.S. to India (excluding tariffs and import taxes) have been estimated at 1.5-2.5 times the U.S. production costs per tonne of ethylene (Chemistry & Industry, 1996). This implies that the risk of U.S. companies relocating because of rising energy prices to supply the U.S. market from abroad is limited. In Europe and Asia energy prices are considerably higher. In the Middle East energy prices are lower, but capital costs are higher because of a lack of sufficient infrastructure and skilled labor (Gielen et al., 1996). The same is expected to be valid for Latin America. In 1993, the only countries/regions producing ethylene at significantly lower costs (20-40% lower) than the U.S. were Canada, Venezuela and (ethane-based capacity in) the Middle East (Hydrocarbon Processing, 1994). It seems likely that transportation costs (of both feedstock from the Gulf coast to the production site and of products to the U.S. market) and unstable political and economic climate (in the case of the Middle East and Latin America) will limit the relocation of U.S. production facilities.

3.3 Energy Use and Intensity in the U.S. Petrochemical Industry

Average Energy Efficiency

Because of the emphasis on ethylene as the most important product from steam cracking in the past, the specific energy consumption per tonne of ethylene (SEC_e) is a common measure of energy consumption for cracking. Modern plant values for SEC_e are 14 GJ/tonne of ethylene for ethane cracking (13 MBtu/ton, HHV) and 20-27 GJ/tonne of ethylene for naphtha/gas oil cracking (19-26 MBtu/ton, HHV) (Phylipsen et al., 1998a).

The cracker product mix is influenced by feedstocks and processing conditions (or severity which includes parameters such as temperature, pressure and furnace residence time). Using the specific energy consumption per tonne of ethylene (SEC_e) as a measure of energy efficiency would mean that all energy consumption is allocated to ethylene, and none to the other products. This means that using another feedstock or severity will affect the SEC_e, even if total energy consumption does not change. In order to be able to compare different processes and feedstocks (with different yields for the various products) another allocation has to be used. In order to exclude effects from changing product yields, energy consumption should be allocated over all products formed in a particular process (on a mass basis).

Solomon Associates Ltd., a company performing energy efficiency analyses and comparisons for refineries and petrochemical plants world-wide, has developed an extensive network for this type of analysis. Participating companies provide Solomon with very detailed data on production, throughput, energy consumption, installed technology. In return, Solomon offers companies a comparison of their own plant's performance with that of all the other participating plants. Plant performance is measured by comparing the plant's actual energy consumption to a reference level of energy consumption, based on the most efficient technology available, using the plant's own configuration, including feedstock input, severity, and product mix. Specific energy consumption is defined as the net energy consumption (lower heating value, LHV⁴) per unit of high value chemicals (including hydrogen, ethylene, propylene, a mixed butenes fraction and a BTX fraction⁵). The energy efficiency is presented as the ratio of the actual SEC to the reference SEC; the energy efficiency index, in which 100 equals the efficiency of the state-of-the-art reference system.

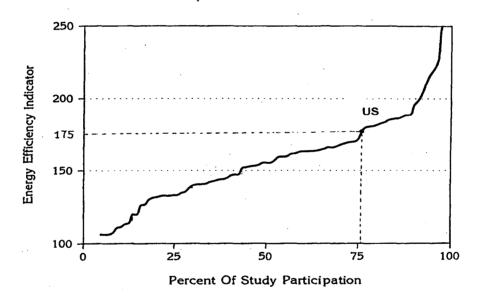
According to Solomon the energy efficiency index (EEI) of the naphtha/gas oil-based crackers in the U.S. is 163 (in 1995). These crackers, located in Texas and Louisiana, account for about 40% of ethylene production (10Mt) (Solomon, 1998). The EEI of all cracking activities in North America (including the U.S., Canada and Mexico) equals 175 (Solomon, 1995). The U.S. accounts for 85% of the capacity in North America (O&GJ, 1998).

⁴ Lower heating value (LHV) and higher heating value (HHV) are both measures of energy consumption. Higher heating values measure the heat that is freed by the combustion of fuels in case the chemically formed water is condensed. Lower heating value measure the heat of combustion in case the water formed remains gaseous. Although the energy difference between HHV and LHV may not always be technically recoverable, from the perspective of energy efficiency analysis HHV is considered preferable, because it is a better measure of the energy inefficiency of processes. In the US HHV is commonly used in energy reporting, in many other countries LHV is more common. Organisations such as Eurostat, the IEA and the UN also use LHV (Phylipsen et al., 1998).

⁵ Measured in tonnes of benzene

In the 1995 analysis, ethane/propane crackers were less efficient than naphtha/gas oil crackers⁶. The global average EEI of ethane crackers was 200, while including all propane crackers improves the EEI to 180 (Solomon, 1995). Assuming 40% of U.S. ethylene production had an EEI of 163 and 60% of ethylene production had an EEI of 180, leads to an estimated overall EEI for the U.S. of 173. This matches closely with the EEI of 175 reported for North America. Considering the US product mix (an average ethylene yield of 51-53%), our best estimate of the specific energy consumption would be about 13 GJ/tonne (LHV) of total product (or 13.3 MBtu/ton, HHV). For the U.S. product mix the SEC per tonne of high value chemicals amounts to about 20 GJ (LHV)/tonne, while SEC per tonne of ethylene amounts to 26 GJ (LHV)/tonne of ethylene (HHVs are 19.1 MBtu/ton high value chemicals and 24.8 MBtu/ton ethylene respectively) (Phylipsen et al., 1998b).

Figure 5 depicts the estimated average U.S. performance compared to all the plants in the Solomon survey (Solomon, 1995). Table 11 shows the EEI for several regions with the U.S. included in 'North America'. The U.S. accounts for about 85% of the ethylene capacity in North America. The Solomon survey covers on average about 70% of the plants worldwide, with somewhat higher coverage in the U.S. and Europe, and somewhat lower in Asia.



ENERGY EFFICIENCY INDEX

Figure 5. The relative performance of the US ethylene plants.

⁶ There is no specific reason why ethane/propane crackers are less efficient than liquid crackers. The observed difference may be caused by the fact that the highest share of liquid crackers are generally located in more efficient countries (such as Japan, South Korea, Germany, The Netherlands). Factors that are likely to play a role are the higher feedstock price for naphtha/gas oil (per tonne of ethylene produced) and the generally higher energy prices in these countries.

Table 11. The energy efficiency index for several regions according to the Solomon survey (stateof-the-art-technology = 100) Source: Solomon (1995).

Region	Energy Efficiency Index
North America	175
Europe	151
Asia	126
Total study	164

With an ethylene production of about 20Mt total U.S. energy consumption for the production of ethylene and its co-products is estimated to be about 520 PJ excluding feedstocks (LHV), and 2225 PJ including feedstocks (LHV) (0.53 Quads excluding, and 2.28 Quads including feedstocks in HHV, respectively). Assuming a predominant use of petroleum products as energy source, carbon emissions resulting from the production of ethylene and its co-products would amount to 10 Mt C⁷ (excluding feedstocks).

Energy consumption by process step

Individual process steps in the production process of ethylene and its co-products are shown in Figure 3. In general, fuels are used to fire the cracking furnace and to generate steam. High pressure steam is used to drive compressors and pumps, medium pressure steam for dilution steam generation and low pressure steam for direct process heating (Di Cintio et al., 1993). A further breakdown of energy consumption in ethane cracking, the predominant feedstock in the U.S., is shown in Table 12 (derived from Brown et al., 1985).

Unit process	Contribution to specific energy consumption			
	Share (%)	Estimated SEC (GJ/tonne)		
Cracker	47%	11.0 GJ/t		
Heat of reaction	23%	5.4 GJ/t		
Dilution steam	6%	1.4 GJ/t		
Heating + losses	18%	4.2 GJ/t		
Compression	22%	5.2 GJ/t		
Separation	31%	7.3 GJ/t		
Chiller	21%	5.0 GJ/t		
Condensor	16%	3.8 GJ/t		
Ethane separator	5%	1.2 GJ/t		
Steam	10%	2.3 GJ/t		
Acetylene removal	3%	0.7 GJ/t		
Heavy separation	7%	1.6 GJ/t		
Specific energy consumption	100%	23.5 GJ/t		

Table 12. Detailed breakdown of energy consumption in ethane cracking (derived from Brown et al., 1985). Specific energy consumption is expressed per tonne of ethylene. Feedstock energy is excluded.

The breakdown of energy consumption in unit processes will vary with feedstock. Because of the lower ultimate ethylene yield per tonne of feed with heavier feeds, higher feed volumes are required to produce the same amount of ethylene than with ethane. This will result in higher compression and separation energy requirements for naphtha and gas oils compared to ethane. Ethane, however, has a lower initial conversion rate. As a result, recycle streams are larger,

⁷ In the calculation we use an emission factor of 20 kg C/GJ

requiring more heating, compression, pumping and separation. These effects will partly outweigh each other. Table 13 shows how energy consumption varies with feedstock (Stratton, 1983).

				Feed	stock			
	Eth	ane	Naphtha Gas Oil		Gas Oil		Mix	
	GJ/t	%	GJ/t	%	GJ/t	%	GJ/t	%
Heat of Reaction	4.8	21%	6.2	20%	12.4	33%	6.0	23%
Compression	3.5	16%	3.5	13%	4.1	11%	3.7	14%
Heating and Separation Losses	14.2	63%	17.7	66%	20.6	55%	16.8	63%
SEC	19.4		26.2		31.9		26.5	

Table 13. Breakdown of primary specific energy consumption (SEC) for ethylene production for different feedstocks. Feedstock energy is excluded. Source: Stratton (1983).

Note: The U.S. mix is assumed to be 60% ethane, 30% naphtha and 10% gas oil.

Although the energy consumption data underlying the shares listed in Table 13 are relatively old, the efficiency level is comparable to the current U.S. situation. Calculating the weighted average, based on the U.S. feedstock mix and the total specific energy consumption figures listed in Table 13, leads to a SEC of 26.5 GJ/tonne ethylene, compared to our estimate of 26 GJ/tonne ethylene for the U.S. The ethane cracker described in Table 12 is slightly less efficient than the one included in Table 13.

Table 13 shows only a limited breakdown In order to estimate energy savings potentials by individual measures a more detailed breakdown is needed, especially for the category of 'heating, separation and losses'. For ethane we use the subdivision listed in Table 12. For other feedstocks a more detailed breakdown of energy consumption is given in Table 14, showing a naphtha cracker and a naphtha/ethane-propane-butane cracker (Di Cintio et al, 1993).

Table 14. Detailed breakdo	n of energy	consumption	for a	naphtha	cracker	and a flexible
cracker. Source: Di Cintio et	ıl (1992).	•				

	Naphtha (Cracker	Flexible C	Cracker
	SEC (GJ/tonne)	Share (%)	SEC (GJ/tonne)	Share (%)
Feedstock Heating	20.7	65%	13.1	55%
Compression	4.8	15%	3.6	15%
Separation	6.3	20%	7.2	30%
Total	31.5		23.9	

Notes: The EEI of a naphtha cracker is about 150 (Phylipsen et al., 1998b) and of a high efficiency cracker, the EEI is approximately 120. In this calculation we assume a feedstock-mix of 70% naphtha, and 30% ethane/propane/butane (Di Cintio et al., 1993).

According to Tables 12, 13 and 14 the energy consumption for separation varies between 6.3-7.3 GJ/tonne ethylene. Based on the U.S. feedstock mix and efficiency, we estimate separation energy consumption for the U.S. to be 7 GJ/tonne ethylene. The compression energy ranges from 3.5-5.2 GJ/tonne ethylene. Taking into account feedstock and efficiency, energy consumption for compression in the U.S. is estimated to be 4 GJ/tonne ethylene. With a total specific energy consumption of 26 GJ/tonne ethylene, cracker/feedstock heating energy consumption results in 15 GJ/tonne. Based on dilution steam rates listed by Di Cintio et al (1993) for various feedstocks and the U.S. feedstock mix, energy consumption for dilution steam can be estimated at 2 GJ/tonne

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ethylene. Table 15 shows the results for the detailed breakdown of energy consumption in U.S. crackers.

Table 15. Estimated energy	consumption by process	step in U.S.	ethylene production.

	U.S. Feedst	tock Mix
	SEC (GJ/tonne ethylene)	Share (%)
Cracker	15.0	58%
Reaction Heat	7.0	26%
Dilution Steam	2.0	8%
Heating and Losses	8.0	31%
Compression	4.0	15%
Separation	7.0	27%
Total	26.0	

4. The Ammonia and Nitrogenous Fertilizer Industry

This chapter reflects an in-depth analysis of the ammonia and nitrogenous fertilizer industry. The nitrogen fertilizer industry is a large energy consumer, with an estimated worldwide annual production capacity of over 100 Mtonnes N and estimated energy consumption equal to 1% of global primary energy use. The production of ammonia is the most energy intensive production step in the manufacture of fertilizers and other nitrogen containing products. In the U.S. ammonia is one of the major chemicals produced, with an estimated production of 16.3 Mtonnes (18.0 Million short tons) (CMA,1996). In the U.S. about 80% of the ammonia is used for fertilizer production, the remainder for a variety of products, mainly explosives and plastics. The most important fertilizers produced in the U.S. are ammonium nitrate (AN), nitric acid (NA), urea, compound fertilizers, and liquid ammonia. Ammonium sulfate (AS) is most commonly produced as a co-product of nylon manufacturing.

The world fertilizer market grows slowly, due to growth especially in developing countries. The world market price of ammonia has been depressed since the late 1980's due to cheap exports from producers in Central and eastern Europe and the former Soviet Union, limiting expansion in the Western World (especially Western-Europe). The U.S. fertilizer market is still slowly growing (IFA,1998), and ammonia prices have been high since 1994 (USGS,1998). The U.S. is a net importer of ammonia, and of some fertilizer types, e.g. urea. The main imports are from countries with cheap natural gas resources, i.e. Trinidad and Tobago, and Canada. Some U.S. firms operate or construct plants abroad, e.g. in Trinidad (e.g. Mississippi Chemical).

In this chapter we first discuss the major process used to make ammonia and nitrogenous fertilizers (section 4.1), followed by a discussion of the U.S. ammonia industry (section 4.2) and energy consumption and intensity (section 4.3).

4.1 Process Description

Ammonia is produced by the reaction of nitrogen and hydrogen, the so-called Haber-Bosch process. The main hydrogen production processes used in the U.S. are steam reforming of natural gas and partial oxidation of oil residues. Hydrogen is produced by reforming the hydrocarbon feedstock, producing synthesis gas, contining a mixture of carbon monoxide and hydrogen. The carbon monoxide is then reacted with steam in the water-gas-shift reaction to produce carbon dioxide and hydrogen. The carbon dioxide is removed from the main gas stream. The carbon dioxide is recovered for urea production or exported as a co-product, or vented. The hydrogen then reacts with nitrogen in the final synthesis loop, to form ammonia. The anhydrous ammonia is sold as product, or used to produce a variety of fertilizers, or other products. For a detailed description of the ammonia production process, see Worrell and Blok (1994). Ammonia production typically requires between 28 and 40 GJ/tonne (LHV)⁸ of ammonia, including feedstocks.⁹ U.S. energy consumption for ammonia manufacture is roughly estimated at 790 PJ (HHV) (Lipinsky and Ingham,1994). The specific energy consumption (SEC) of modern partial oxidation units is 30 GJ/tonne (Lurgi,1987),

⁸ The heating value of a fuel can be expressed in lower (or net) heating value (LHV) and higher (or gross) heating value (HHV). The difference is the condensation heat of the water vapor of the combustion process, which is included in the HHV. LHV is commonly used in international statistics and in Europe, while HHV is used in the U.S. and Canada. Natural gas in the U.S. has a set HHV of 38.4 MJ/Nm3 or 1,030 Btu/scf. The LHV is approximately 11% lower.

⁹ Feedstock consumption is estimated to be 19 to 22 GJ/tonne (LHV) ammonia. Note that although energy use is earmarked as feedstock, the carbon may be emitted to the atmosphere if it is not used for urea manufacture, or recovered for other purposes (e.g. soft drink manufacture). Total energy consumption is equivalent to 24-34 MBtu/short ton ammonia.

and upto 30% higher for older units (Nitrex, 1989). The SEC of modern natural gas steam reforming is 27 to 28 GJ/tonne (Appl, 1994).

Although, ammonia is used directly as a fertilizer in the U.S., most of the ammonia is converted to other compounds to be used as fertilizer. We give a short description of the main processes for fertilizer production.

<u>Urea</u> is produced in two steps by the reaction of NH₃ and CO₂. The CO₂ is produced in the NH₃ synthesis. In the first step, carbamate (NH₂CO₂NH₄) is synthesized. Feedstock is provided in an NH₃/CO₂ ratio varying from 2.5 to 3.5 to achieve a high conversion rate. In the second step, the carbamate is dehydrated to urea. The reaction is not complete. Therefore NH₃ and CO₂ are both stripped from the urea solution and recycled. Commercial processes differ in the dehydration step. The main processes are solution recycle (NH₃ and CO₂ are recycled to the synthesis reactor as an aqueous solution) and stripping process (the non-reacted carbamate is removed by partial pressure reduction, using one of the reactants). Several techniques are used to dry the urea solution. Normally the solution is treated in a prilling tower, to produce granulate. In Europe the urea plants consume between 3.2 and 4.6 GJ/tonne of primary energy (Worrell and Blok,1994). The CO₂ is extracted from the NH₃-process, so no energy is consumed. However in some NH₃-processes there is a CO₂ deficiency (depending on the production volumes of NH₃ and urea) which is met by extra CO₂ production (which consumes energy and emits carbon dioxide) in a separate plant.

<u>Nitric Acid</u> (NA) is used mainly for the production of ammonium nitrate (NH₄NO₃). HNO₃ is also used to produce non-fertilizer products. NH₃ is burned over catalysts to produce nitrous oxides. The NO_x are passed to an absorber column to produce HNO₃ (reaction with water). The total process is highly exothermic, so waste heat boilers are installed to generate superheated high pressure steam. Two main types of processes can be identified: mono-pressure (oxidation and absorption pressures are the same) and dual-pressure (absorption pressure is higher than oxidation pressure). For both types a large number of different processes (with different operating pressures) have been developed. New designs have been developed which have a lower steam consumption (due to the incorporation of expansion turbines) or which have increased heat recovery.

<u>Ammonium nitrate</u> (AN) is produced by the neutralization of HNO_3 with NH_3 , in an exothermic reaction. The released heat can be used in the process internally (to evaporate the water of the HNO_3 solution, or to preheat the HNO_3 , or evaporate the liquid NH_3) and can produce steam, which can be exported. Whereas older plants work at atmospheric pressures (which require the import of steam), most modern processes work at elevated pressures (and export low pressure steam).

<u>Ammonium sulfate</u> (AS) is produced as a byproduct of nylon 6,6 manufacture. In the synthesis of caprolactam ammonia is added to control the reaction. As much as three to five times as much ammonium sulfate may be produced than caprolactam (Lipinsky and Ingham, 1994). After separation of the ammonium sulfate solution, the water is evaporated to produce a marketable product.

4.2 Characterization of the U.S. Ammonia Industry

In the U.S. ammonia is produced in 41 plants (1996) which predominantly use natural gas as a feedstock, as can be seen from energy statistics (EIA,1997). Natural gas is the favorable feedstock for ammonia, as it has the highest hydrogen to carbon ratio, leading to reduced energy consumption, and carbon dioxide emissions. Natural gas based plants also have lower capital costs, although ammonia manufacture remains capital intensive. The capital costs for a new greenfield plant are estimated at roughly \$300 per tonne annual capacity (Worrell and Blok,1994). Although currently available data makes it impossible to calculate the average age of

ammonia production facilities in the U.S, the original production capacity in the U.S. seems relatively old. It is important to note that many plants have undergone various reconstructions and expansions, changing the process lay-out and improving performance, compared to the original plant. Concentration of production in medium to large-scale plants has taken place in the last decade. Future capacity or expansion will probably take place only in areas with low natural gas prices, e.g. Texas, Louisiana, and Oklahoma. Almost 60% of U.S. ammonia capacity is located in these states (USGS,1998).

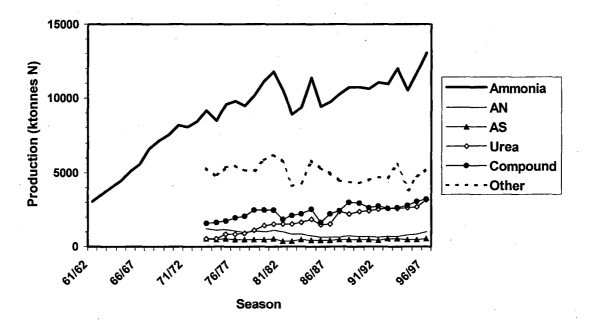


Figure 6. Historic U.S. production of ammonia, and some major fertilizers, expressed in 1000 metric tonnes of Nitrogen. The data are given for the agricultural season, as provided by the International Fertilizer Association (IFA,1998). Data of the Chemical Manufacturers Association (CMA,1996) are given by calendar year, and vary slightly from the IFA data.

4.3 Energy Use and Intensity in the U.S. Ammonia Industry

Methodology

The U.S. Manufacturing Energy Consumption Survey reports only the energy consumption for the nitrogenous fertilizer industry as whole, which includes not only the manufacture of ammonia, but also the production of urea, nitric acid, ammonium nitrate, and compound fertilizers. For part of the fertilizers it also contains the granulation of the fertilizers. The MECS does not report on the ammonia process itself. Hence, we have to estimate the energy intensity of ammonia making in the U.S. based on the total energy consumption in the sector, as well as on other sources, e.g. an annual survey of North-American ammonia producers by The Fertilizer Institute.

Energy Use and Carbon Dioxide Emissions

Energy consumption for ammonia and fertilizer production consists of consumption as energy for fueling the chemical process, as well as the energy used as feedstock. With respect to consumption for fueling the chemical processes, the U.S. Manufacturing Energy Consumption Survey estimated the consumption for the production of nitrogenous fertilizers (SIC 2873) to be

288 PJ in fuels and 13.7 PJ electricity in 1994. Consumption of energy sources for feedstock is estimated at 368 PJ (EIA,1997). The specific feedstock consumption is estimated to be 22.6 GJ/tonne ammonia (HHV).

Carbon emissions can be estimated on the basis of total fuel consumption of the sector, accounting for the carbon dioxide that is used as feedstock in urea manufacture,¹⁰ and carbon dioxide recovered for other purposes. Urea production in 1994 is estimated at 8,067,000 tons (CMA,1996). On a molecular basis, 20% of the urea is carbon. The amount of carbon exported in urea is estimated at 1.46 MtC.¹¹ There are no data available on the amount of carbon dioxide recovered for other purposes in the U.S. ammonia industry. The total carbon emission in the U.S. (1994) is estimated to be minimal 9.07 MtC.¹² This is relatively low due to the high use of natural gas as the main feedstock and energy carrier (99% of fuels used).

Non-CO₂ Greenhouse Gas Emissions

The production of nitric acid is one of the industrial emission sources of nitrous oxide (N₂O), besides the production of adipic acid (for Nylon 6,6).¹³ In the combustion of ammonia to produce nitric acid, some nitrous oxide may be formed as an accidental co-product. Off-gas measurements at DuPont showed a wide range in emissions, varying from 2 to 9 g N₂O/kg HNO₃ (IPCC,1995). It is not known if the DuPont emission factors are representative for all processes used, or for the whole industry. The total emission of the U.S. fertilizer industry is estimated at 16-72 Gg N₂O, equivalent to 1.4-6.1 Mt C (using a GWP of 310 for a 100 year time period (IPCC,1996)). US DOE estimates the 1994 nitrous oxide emission from nitric acid production at 47 ktonnes (EIA,1996), equivalent to 3.9 Mtonne C. At high concentrations (as with adipic acid production) the N₂O emissions can be reduced by oxidation in a flame. It is not clear if this technology can also be used in the production of nitric acid. However, the wide range shows that good process control may influence the N₂O emission. Future emission controls may be based on catalytic processes, but no technologies are commercially available yet.

Energy Intensity and Carbon Dioxide Emissions

U.S. energy statistics do not give direct information on the energy consumption for ammonia manufacture. As 80-90% of the primary energy in SIC 2873 is consumed in ammonia manufacture, the total consumption gives some rough information on the energy consumption (see above). According to a survey of the Fertilizer Institute among 77% of North American ammonia producers (incl. Canada) the energy intensity ammonia making is estimated at 39.3 GJ/tonne (HHV) of natural gas and 140 kWh per tonne of ammonia (Vroomen, 1998). This is equivalent to a primary energy consumption of 40.9 GJ/tonne (HHV) or 37.1 GJ/tonne (LHV)).¹⁴

¹⁰ Carbon dioxide is actually released to the atmosphere as soon as the urea is applied.

¹¹ This assumes that all carbon dioxide is recovered from ammonia manufacture, and that no extra carbon dioxide is produced for urea manufacture. However, back-up carbon dioxide generators are often available to meet peak CO_2 demands. In that case, the actual CO_2 will be higher than the figure mentioned above.

¹² The following emission factors are used for the carbon calculation: coal: 27.0 kg/GJ; natural gas: 15.3 kg/GJ, and for electricity: 50.5 kg/GJe (EIA,1996).

¹³ Note that the industrial emissions of nitrous oxide are relatively small compared to that of other sources, i.e. mobile sources. Also note that the application of fertilizers may lead to the emission of nitrous oxides, depending on the application method, soil type, groundwater level and fertilizer type.

¹⁴ Lipinsky and Ingham (1994) estimated the 1992 total energy consumption for ammonia manufacture at 42.2 Mbtu/ton ammonia (16.1 Mbtu/ton for energy and 26.1 Mbtu/ton for feedstock). This is equivalent to 49.1 GJ/tonne (HHV). This seems to be too high, compared to the information from the industry's survey, and to intensities in other countries (Worrell et al., 1994b).

An analysis of energy intensities of ammonia manufacture in Europe showed that 1989 energy consumption varied between 28 GJ/tonne (Spain, LHV) and 40 GJ/tonne (Belgium, LHV), while the average 1989 SEC in the European Union is estimated at 35.5 GJ/tonne (LHV) (Worrell et al.,1994b).

The theoretical minimum energy consumption for ammonia manufacture through steam reforming is approximately 21.6 GJ/tonne ammonia (HHV). This illustrates the maximum potential for efficiency improvement, which is, however, not practically achievable, not even on the long term.

Energy Balance for Ammonia Making

For a detailed analysis of the impact of energy efficient practices and technologies on energy intensity and carbon dioxide emissions we generated a detailed energy balance of the ammonia manufacturing process. We constructed a detailed balance on the basis of the specific energy consumption of ammonia plants in the U.S. (based on the Fertilizer Institute Survey (Vroomen,1998)), the feedstock consumption given by MECS (EIA,1997), and energy balances given by Nielsen (1995) and Brown et al. (1985). An recent analysis provides an energy and exergy balance for an energy efficient process (Nielsen, 1995) while the study by Brown et al. (1985) reports on older plant, being more energy intensive than the current U.S. situation. We combined the insights of both analyses to estimate an average energy balance for the U.S. In Table 16 we give the energy balance broken down to the major process steps and unit-operations.

Unit Operation	LHV, GJ/tonne				HH	V, Mbtu/	ton			
	Gas	Steam	Losses	Electricity	Heat	Gas	Steam	Losses	Electricity	Heat
Reformer Feed	20.4				20.4	19.4			· · · ·	19.4
Reformer Fuel	9.9				9.9	9.4				9.4
Primary reformer	1	4.8			4.8	1	4.1			4.1
Secondary reformer		0.0			0.0		0.0			0.0
Waste heat Boiler		-5.6			-5.6		-4.8			-4.8
Shift + CO2 removal		1.2		0.2	1.2		1.0		0.2	1.0
Methanator			0.3		0.3		0.0	0.2		0.2
Synthesis loop	[-2.0		0.2	-2.0		-1.7		0.2	-1.7
Aux. Boiler	4.5	-3.9			0.6	4.2	-3.3			0.9
Turbines/Compressor		5.5			5.5		4.7			4.7
Miscellaneous			0.3	0.1	0.3			0.3	0.1	0.3
Flare	0.3				0.3	0.3				0.3
Total	35.0	0.0	0.6	0.5	35.6	33.4	0.0	0.5	0.4	33.8
Summary										
Feedstock					20.4					19.4
Fuel					15.2					14.4
Electricity			139	KWh/t	0.5			119	kWh/ton	0.4
Final			L	·	36.1					34.3
Primary (incl. feedstock)					37.1					35.1
Primary (excl. feedstock)					16.7	*				15.7

Table 16. Estimated energy balance for U.S. ammonia manufacturing (1996). Boiler efficiency is assumed to be 78% (HHV) or 86% (LHV). Power generation efficiency is assumed to be 33%.

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5. The Chlorine Industry

This chapter reflects an in-depth analysis of the chlorine and alkaline industry. The chlorine and alkalines production is closely connected, due to the feedstock used (salt). This sub-sector is one of the most energy and electricity intensive industries. The U.S. is the world's largest chlorine manufacturer.

The major markets for chlorine are PVC (37%), inorganic chemicals (22%), other organic chemicals (17%), propylene oxide (7%), pulp and paper (6%), water treatment (6%), solvents (5%). The major markets for caustic are: pulp and paper (26%), soaps and detergents (9%), propylene oxide (9%), petroleum (8%), water treatment (6%), other organic chemicals (13%), inorganic chemicals (12%) (Florkeiwicz, 1998). These areas are expected to continue to grow, but below the historical rate of growth (CMA, 1996). In part, the slower growth is attributable the collapse of the Asian economies and to environmental concern about the use of chlorine in industrial processes (Hileman et al., 1994).

In this chapter we first discuss the major process used to make chlorine (section 5.1), followed by a discussion of the U.S. chlorine industry (section 5.2) and energy consumption and intensity (section 5.3).

5.1 Chlorine manufacturing Processes

The production of chlorine gas is an energy intensive chemical process requiring between 25-40 GJ (worldwide average) primary energy per tonne chlorine produced (Phylipsen et al., 1998a). In the process a brine solution is converted into two co-products through electrolysis: chlorine gas and sodium hydroxide (caustic soda). The three main electrolysis cell types that are used to separate and produce the chlorine gas and caustic are the mercury flow, diaphragm, or ion-selective membrane. In the diaphragm and membrane cells the caustic soda requires an additional step of concentrating the solution so that it can meet market specifications for most products. Of the three cell types the membrane cell requires the least energy to operate and is currently considered the state-of-the-art technology. Figure 7 shows the main process stages for chlorine production.

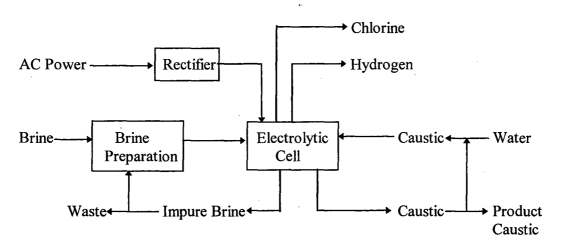


Figure 7. Simplified Chlorine Production Flow Chart

5.2 The U.S. Chlorine Industry

Figure 8 below shows the age distribution of US chlorine plants. As the figure shows, most of the plants are 20-25 years old, and some considerably older. Appendix C provides data on chlorine production by plant and cell type for the US in 1994. As the table indicates there were 24 companies engaged in chlorine production with a total cell capacity of 11,525 Ktonnes. Some of the largest companies include Dow, Occidental, PPG Industries, and Olin with a share of total capacity of 27%, 25%, 13%, and 7% respectively. The vast majority (83%) of production took place in the South, where companies are able to take advantage of low electricity prices and low labor costs. Total capacity utilization in 1994 was 95%. A total capacity of 2.1 million tonne/year is expected to come on line by 2000 in the US (Westervelt, February 25, 1998).

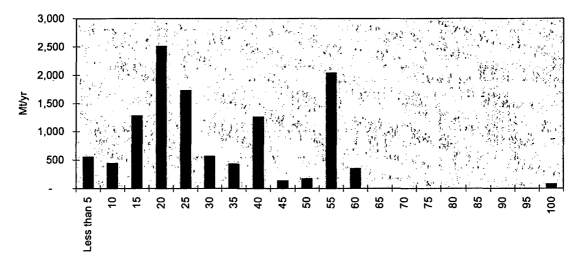


Figure 8. Age of U.S. Chlorine Plants. Source: SRI International, 1997

5.3 Energy Use and Energy Intensity of Chlorine Making

US energy statistics only report energy consumption for the chlor-alkali sector as a whole (which includes several products including chlorine) (SIC 2812) while excluding chlorine produced at an integrated facility where the final product is outside of the chlor-alkali sector (such as PVC). Therefore, in estimating 1994 baseline energy use and carbon emissions for the chlorine sector, we use available data to estimate the age and distribution of cell types in the US and estimate weighted average energy intensity, energy use, and carbon emissions. We also assume that cells which are 20 years old or greater are less efficient than new cells of the same type.

Table 17 identifies estimated final energy¹⁵ consumption and carbon emissions in 1994 for chlorine/caustic production. Energy intensity estimates are calculated using a technological analysis of all three major technologies (mercury, diaphragm, and membrane). These analyses are based on energy consumption in a modern Dutch chlorine plant (Doesburg, 1994). This data is then combined with information on the market share of each of the technologies in the US (Kirk-Othmer, 1994). In addition, we assumed that chlorine plants constructed 20 years ago are 10% less efficient than modern plants. Combining these two sources we calculate a weighted-average US energy intensity. The intensity values are multiplied by 1994 production estimates from the

¹⁵ Final energy simply represents all the energy consumed at an industrial facility. Purchased electricity and steam are accounted by simple energy content as they enter the facility. This is in contrast to primary energy which includes the energy used to make that steam and electricity. All primary energy numbers in this paper assume electricity production is 40% efficient.

Chemical Manufacturers Association to produce estimates of total consumption and carbon emissions. We estimate the specific electricity consumption at 4380 kWh/tonne and specific fuel consumption at 3.45 GJ/tonne. Net primary energy consumption is estimated at 47.9 GJ/tonne, crediting the energy content of hydrogen export at 3.35 GJ/tonne chlorine. For these calculations total energy use has been allocated to chlorine production.

Process Stage	Final	Fuel	Final	Final	Final	Final	Carbon
	Electricity	Intensity ^a	Energy	Electricity	Fuel	Energy	Emissions ^c
	Intensity ^a		Intensity ^a	Use ^b	Use ^b	Use ^b	
	GJ/tonne	GJ/tonne	GJ/tonne	PJ	PJ	PJ	KtC
	Cl ₂	Cl ₂	Cl ₂				
Rectifier	0.28	0.00	0.28	3.1	0.0	3.1	132
Brine Preparation	0	0.02	0.02	0.0	0.2	0.2	10
Cell Use	13.63	0.00	13.63	149.6	0.0	149.6	6365
NaOH Concentration	0	3.42	3.42	0.0	37.6	37.6	435 ^d
NaOH Cooling	0.27	0.00	0.27	2.9	0.0	2.9	125
Hydrogen Cooling/Drying	0.58	0.00	0.58	6.4	0.0	6.4	272
Chlorine Cooling/Drying	0.39	0.00	0.39	4.3	0.0	4.3	181
Chlorine Compression	0.63	0.00	0.63	6.9	0.0	6.9	295
Total	15.78	3.45	19.23	173.20	37.81	211.02	7817
Hydrogen Heating Value			-3.35°			-36.8°	-

Table 17. Chlorine Industry Energy Use and Carbon Dioxide Emissions. Sources: Doesburg (1994); CMA (1996); Kirk-Othmer (1994).

^a1994 Industry Final Energy Intensity based on a calculation of weighted energy use by cell type and plant age. Cells and intensities for a modern 1989 plant were: Membrane (9.8 GJ/tonne electricity, 0.5 GJ/tonne fuel), Diaphragm (11.3 GJ/tonne electricity, 2.6 GJ/tonne fuel), Mercury (12.3 GJ/tonne electricity, 0.1 GJ/tonne fuel) (Doesburg, 1994). Shares of cell type for 1994 are: membrane 8%, diaphragm 75%, mercury 15%. These are based on Kirk-Othmer, 1994 and currently exclude plants that manufacture chlorine from sources other than Sodium chloride (e.g. magnesium chloride). Using the 1989 efficiencies for each cell type, and an assumed 10% efficiency improvement over 20 years, we calculated an efficiency for each plant and took a production weighted average.

^bIndustry total energy use based on a production of 11.05 Mt of chlorine in 1994. Electricity production from cogeneration is currently not included in the calculation due to lack of reliable data.

^cCarbon emissions are based on the following factors from EIA, 1995. Electricity 42.6 KtC/PJ, Oil 20.4 KtC/PJ.

^dThis value assumes that half the hydrogen produced is used to concentrate the caustic. The rest is sold or used in other production processes.

^eThis is the fuel value of hydrogen. It is listed as a negative because it represents energy that can be used to fuel other processes.

The US Manufacturing Energy Consumption survey reports a total final energy consumption in 1994 for the chlor-alkali industry of 136 PJ with a net electricity consumption of 49 PJ. The MECS also reports that if one includes onsite cogeneration, the total electricity consumption for the sector is 62 PJ which is still lower than our total calculated electricity use in Table 17. Since MECS includes products we are not considering, and excludes some chlorine production, we have chosen to currently rely on our technical estimates for energy consumption.

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6. Discussion and Conclusions

The U.S. chemical industry is the largest in the world, and responsible for a large part of U.S. industrial production (11% of U.S. manufacturing value added). It also consumes approximately 20% of total industrial energy consumption in the U.S. (1994), and contributes in similar proportions to U.S. greenhouse gas emissions. Surprisingly, there is not much information on energy use and energy intensity in the chemical industry available in the public domain. This report provides detailed information on energy use and energy intensity for the major groups of energy-intensive chemical products.

Ethylene production is the major product in terms of production volume of the petrochemical industry. The petrochemical industry (SIC 2869) produces an extremely wide variety of products, although most energy is used for a small number of intermediate compounds, of which ethylene is the most important one. Based on a detailed assessment we estimate fuel use for ethylene manufacture at 520 PJ (LHV), excluding feedstock use. Ethylene production is responsible for approximately 34% of energy use in the petrochemical industry. Energy intensity is estimated at 26 GJ/tonne ethylene (LHV), excluding feedstocks.

The nitrogenous fertilizer production is also a very energy intensive industry, producing a variety of fertilizers and other nitrogen-compounds. Ammonia is the most important intermediate chemical compound, used as basis for almost all products. In the U.S. ammonia is produced mainly from natural gas. Fuel use is estimated at 268 PJ (excluding feedstocks), while 368 PJ natural gas is used as feedstock.¹⁶ Electricity consumption is estimated at 14 PJ. We estimate the energy intensity of ammonia manufacture at 39.3 GJ/tonne (including feedstocks, HHV) and 140 kWh/tonne, resulting in an estimated primary energy consumption of 40.9 GJ/tonne (HHV), equivalent to 36.6 GJ/tonne (LHV).

The third most important product from an energy perspective is the production of chlorine and caustic soda. Chlorine is produced through electrolysis of a salt-solution. Chlorine production is main electricity consuming process in the chemical industry, next to oxygen and nitrogen production. We estimate final electricity use at 173 PJ (48 TWh) and fuel use of 38 PJ. Total primary energy consumption is estimated at 526 PJ (including credits for hydrogen export). The energy intensity is estimated at an electricity consumption of 4380 kWh/tonne and fuel consumption of 3.45 GJ/tonne, if all energy use is allocated to chlorine production.

Our estimates are based on surveys of individual plants in the industry as well as engineering estimates on the basis of literature values and surveys. This introduces uncertainties in the results. However, we are not able to estimate the magnitude of the uncertainties. Comparison with other official energy consumption statistics is difficult due to the aggregation levels, as well as different sub-sector divisions used in the statistics and this report.

Compared to earlier studies (e.g. Brown et al., 1985; Lipinsky and Ingham, 1994) the current report provides a detailed baseline of energy use in these processes, based on a detailed assessment of the industry and technologies used. The report also provides breakdowns for energy use in the various production steps and unit-operations for recent estimates of the energy intensity of processes.

¹⁶ Feedstock use in the ammonia industry may result in direct CO_2 emissions, in contrast to the petrochemical industry, where most of the carbon in the feedstock is used in the chemical product.

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Company	Location		Year of start up	Ethylene Capacity	Feedstock Mix (%)						Licensor, Remarks
	1			(ktonne/a)	Ethane	Propane	Butane	Naphtha	Gas oil	Other	
Атосо	Chocolate Bayou	TX		1,451	36%	40%	0%	24%	0%		Braun ?
Chevron	Cedar Bayou	TX		681	25%	40%	5%	30%	0%		Stone & Webster
	Port Arthur	TX		784	80%	20%	0%	0%	0%		
Concea Vista	Lake Charles	LA	1968	430	100%	0%	0%	. 0%	0%		Lummus; Including expansion of 150kt in 1990; Formerly Vista Chemicals
Dow	Freeport	ΤX		1,224	50%	50%	0%	0%	0%		
	Freeport	TX			50%	50%	0%	0%	0%		
	Plaquemine	LA		1,102	50%	50%	0%	0%	0%		
	Plaquemine	LA			25%	25%	25%	25%	0%		
DuPont	Orange	TX		590	50%	50%	0%	0%	0%		Lummus
Eastman	Long-view	TX		675	30%	50%	5%	15%	0%		Lummus/Kellogg ?
Equistar	Channelview	TX		873	5%	5%	5%	40%	45%		Before 1997: Lyondell
	Channelview	TX		873	5%	5%	5%	40%	45%		Before 1997: Lyondell
	Clinton	IA	1968	435	95%	5%	0%	0%	0%		Kellogg; Before 1997 Quantum; Expansion planned of 37kt in 1997
	LaPorte	TX	1991	789	80%	20%	0%	0%	0%		Before 1997: Quantum; Expansion planned of 182kt in 1996
	Morris	IL	1971	512	90%	10%	0%	0%	0%		Lummus; Before 1997: Quantum; Including expansion of 32kt in 1996
Exxon	Baton Rouge	LA	1973	882	0%	0%	0%	0%	0%		Kellogg; Including expansion in 1993
-	Baytown	TX		1,890	0%	0%	0%	0%	0%		Kellogg; Including expansion of 700kt in 1997 by Lummus/Exxon
Formosa Plastics	Point Comfort	TX	1994	714	45%	25%	0%	15%	15%		Kellogg Millisecond; Expansion planned of 204kt in 1997
Huntsman	Port Arthur	TX	1978	551	0%	0%	0%	60%	0%	40% LPG	Stone & Webster; Before 1994: Texaco
	Port Neches	TX		136	50%	50%	0%	0%	0%	RG	Scientific Design
Javelina Co.	Corpus Christi	TX		108	0%	0%	0%	0%	0%	100% RG	· · · · · · · · · · · · · · · · · · ·
Mobil	Beaumont	TX	1975	566	70%	24%	6%	0%	0%		Stone & Webster; Expansion planned of 250kt in 1995
	Houston	TX		342	70%	30%	0%	0%	0%		Kellog

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Appendix A: Overview of U.S. Ethylene Plants (Situation End of 1997).

Occidental	Chocolate Bayou	TX	1980	500	0%	0%	0%	75%	25%	Brown & Root, Lummus	4
	Corpus Christi	TX	1980	773	15%	30%	0%	35%	20%	Stone & Webster	
	Lake Charles	LA	1986	364	50%	50%	0%	0%	0%	Lummus Crest	
Philips	Sweeny	TX	1978	2,040	70%	30%	0%	0%	0%	Selas, debottlenecking 1990	
Philips	Sweeny	TX		Ī	92%	8%	0%	0%	0%		
	Sweeny	TX			75%	25%	0%	0%	0%		
	Sweeny	TX	1990	Ţ	30%	60%	10%	0%	0%	1,000kt Braun, 1990; Including ex 227kt in 1996	pansion of
Rexene Prod.	Odessa	TX		230	50%	50%	0%	0%	0%	Expansion planned of 188kt in 199	98
Shell	Deer Park	TX	T	952	10%	0%	0%	60%	30%	Kellog	
	Norco	LA	1976	793	0%	0%	0%	30%	70%	Kellog	
	Norco	LA	1976	535	50%	0%	0%	50%	0%	Kellog: Including expansion of 18	2kt in 1996
Sun	Brandenburg	KY		45	100%	0%	0%	0%	0%	Lummus	
	Marcus Hook	PA		102	0%	0%	0%	0%	0%	· · · · · · · · · · · · · · · · · · ·	
Union Carbide	Seadrift	TX	1962	415	80%	20%	0%	0%	0%		
	Taft	LA	1978	680	30%	30%	0%	40%	0%	Wulff/Lummus, restarted in 1989; planned of 318kt in 1997	expansion
	Texas City	TX		680	0%	60%	10%	30%	0%		
Union Texas	Geismar	LA	1968	545	82%	18%	0%	0%	0%	Lummus	
Westlake Pol.	Calvert City	KY	1964	170	0%	100%	0%	0%	0%	Braun; Before 1994: BF Goodrich	
	Lake Charles	LA	1992	1,043	80%	20%	0%	0%	0%	Kellog Millisecond; Including 590 1997	kt expansion in
Total				25,475						· · · · · · · · · · · · · · · · · · ·	

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RG = refinery gas

Company	Ammonia Plan	State	Capacity	Feed-	Licensor/Construction
	Location		(tonnes/year)	stock	
Agrium Inc.	Borger	TX	439,000	NG	Kellogg, modernized in 1989
Air Products	Pensacola	FL	46,000	NG	Chemico
Allied Signal	Hopewell	WV	409,000	NG	Kellogg, revamp in 1986
Ampro	Donaldsonville	LA	386,000	NG	
Arcadian	Geismar	LA	501,000	NG/RG	Kellogg .
	Clinton	IA	237,000	NG	Haldor Topsoe. Original plant 1963, expanded 1982
	Augusta	GA	576,000	NG	Haldor Topsoe (or Braun)
	La Platte	NE	182,000	NG	
	Lima	ОН	523,000	NG	Revamped 1971 and 1982
	Memphis	TN	340,000	NG	
	Woodstock	TN	356,000	NG	Ammonia Casale
Avondale Ammonia	Fortier	LA	399,000	NG	Kellogg
Borden Chemical	Geismar	LA	363,000	NG	Chemico, revamp 1982
CF Industries	Donaldsonville	LA	1,740,000	NG	Kellogg 1966/68, revamp 1988
Coastal St. Helens Chemical	St. Helens	OR	85,000	NG	
Coastal Chem	Cheyenne	WY	172,000	NG	1965 (31,000 tpy), revamp 1985/87
Cytec Industries	Avondale	LA	385,000	NG	
Dakota Gasification Co.	Beulah	ND	91,000	n.a.	
Du Pont	Beaumont	TX	363,600	NG	Kellogg, revamp 1986
Farmland Industries	Beatrice	NE	255,000	NG	
	Dodge City	KS	255,000	NG	Kellogg
	Enid	ОК	919,000	NG	Kellogg
	Fort Dodge	10	241,000	NG	Kellogg
	Lawrence	KS	409,000	NG	Kellogg
	Pollock	LA	459,000	NG	Kellogg
Greenvalley Chemical	Creston	IA	33,000	NG	
IMC Agrico	Donaldsonville	LA	482,000	NG	Kellogg
IMC Nitrogen Co.	East Dubuque	IL	269,000	NG	N-Ren
J.R. Simplot Co.	Pocatello	ID	93,000	NG	Haldor Topsoe 1964, expansion 1977
Koch Industries	Sterlington	LA	1,110,000	NG	Kellogg, two units, second 1977 (379,500 tpy)
LaRoche Ind.	Cherokee	AL	159,000	NG	Kellogg, modernized 1986
Missippi Chemical	Yazoo City	MS	363,600	NG	Kellogg, revamp 1986
Co.	Donaldsonville	LA	409,000	NG	Kellogg
	Donaldsonville	LA	500,000	NG	· · · · · · · · · · · · · · · · · · ·
Monsanto	Luling	LA	446,000	NG	Kellogg
Nitromite Fertilizer	Dumas	TX	128,000	NG	
Shoreline Chem.	Gordon	GA	31,000	H2	
Terra International	Blytheville	AR	364,000	NG/FO	Bechtel/Haldor Topsoe
	Sergeant Bluff	IA	319,000	NG	Kellogg
	Verdigris	OK	955,000	NG	Kellogg 1975, revamp 1986
· · · · · · · · · · · · · · · · · · ·	Woodward City	ОК	446,000	NG	Haldor Topsoe
Unocal	Finley	WA	150,000	NG	Restarted 1995
	Kenai	AK	1,180,000	NG	1968, expanded 1977 (two units)
Wil-Grow Fertilizer Co.	Pryor	OK	86,000	NG	
Total			17,461,000		<u> </u>

Appendix B: Ammonia Plants in the United States (1998 Situation)

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Notes: Feedstock: Natural Gas (NG), Refinery Gas (RG), Fuel Oil (FO), Hydrogen (H2)

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BF Goodrich C Dow F DuPont N Elf Atochem F Formosa Plastics E Fort Howard C GE F Georgia Gulf Corp. F	Ashtabula Calvert City Freeport Plaquemine Niagara Falls Portland Baton Rouge Point Comfort Green Bay Muskogee Rincon Burkville Mount Vernon	OH KY TX LA NY OR LA TX WI OK GA	Capacity (Kt/year)* 36 109 2,041 1,075 77 169 180 505 8 5	Age of Process in 1994* ^b 31 28 54 36 96 47 13 0 26	KCl electrolysis; mercury cell; KOH by-product Salt; mercury cell; diaphragm MgCl containing brines; diaphragm and by-product of magnesium metal production Brine; diaphragm Downs; By-product of metallic sodium production Salt; diaphragm Brine; diaphragm (upgraded in 1981)
BF Goodrich C Dow F DuPont N Elf Atochem F Formosa Plastics E Fort Howard C GE F Georgia Gulf Corp. F	Calvert City Freeport Plaquemine Niagara Falls Portland Baton Rouge Point Comfort Green Bay Muskogee Rincon Burkville	KY TX LA NY OR LA TX WI OK GA	36 109 2,041 1,075 77 169 180 505 8	31 28 54 36 96 47 13 0	Salt; mercury cell; diaphragm MgCl containing brines; diaphragm and by-product of magnesium metal production Brine; diaphragm Downs; By-product of metallic sodium production Salt; diaphragm Brine; diaphragm (upgraded in 1981)
BF Goodrich C Dow F DuPont N Elf Atochem F Formosa Plastics E Fort Howard C B GE E Georgia Gulf Corp. P	Calvert City Freeport Plaquemine Niagara Falls Portland Baton Rouge Point Comfort Green Bay Muskogee Rincon Burkville	KY TX LA NY OR LA TX WI OK GA	109 2,041 1,075 77 169 180 505 8	28 54 36 96 47 13 0	Salt; mercury cell; diaphragm MgCl containing brines; diaphragm and by-product of magnesium metal production Brine; diaphragm Downs; By-product of metallic sodium production Salt; diaphragm Brine; diaphragm (upgraded in 1981)
Dow F DuPont P Elf Atochem F Formosa Plastics E Fort Howard C GE GE Georgia Gulf Corp. P	Freeport Plaquemine Niagara Falls Portland Baton Rouge Point Comfort Green Bay Muskogee Rincon Burkville	TX LA NY OR LA TX WI OK GA	2,041 1,075 77 169 180 505 8	54 36 96 47 13 0	MgCl containing brines; diaphragm and by-product of magnesium metal production Brine; diaphragm Downs; By-product of metallic sodium production Salt; diaphragm Brine; diaphragm (upgraded in 1981)
P DuPont N Elf Atochem P Formosa Plastics E P Fort Howard C M Georgia Gulf Corp. P	Plaquemine Niagara Falls Portland Baton Rouge Point Comfort Green Bay Muskogee Rincon Burkville	LA NY OR LA TX WI OK GA	1,075 77 169 180 505 8	36 96 47 13 0	metal production Brine; diaphragm Downs; By-product of metallic sodium production Salt; diaphragm Brine; diaphragm (upgraded in 1981)
DuPont N Elf Atochem P Formosa Plastics E P Fort Howard C GE E Georgia Gulf Corp. P	Niagara Falls Portland Baton Rouge Point Comfort Green Bay Muskogee Rincon Burkville	NY OR LA TX WI OK GA	77 169 180 505 8	96 47 13 0	Downs; By-product of metallic sodium production Salt, diaphragm Brine; diaphragm (upgraded in 1981)
Elf Atochem P Formosa Plastics E P Fort Howard C GE E Georgia Gulf Corp. P	Portland Baton Rouge Point Comfort Green Bay Muskogee Rincon Burkville	OR LA TX WI OK GA	169 180 505 8	47 13 0	Salt; diaphragm Brine; diaphragm (upgraded in 1981)
Formosa Plastics E P Fort Howard C GE Georgia Gulf Corp. P	Baton Rouge Point Comfort Green Bay Muskogee Rincon Burkville	LA TX WI OK GA	180 505 8	13 0	Brine; diaphragm (upgraded in 1981)
Fort Howard C M GE E Georgia Gulf Corp. P	Point Comfort Green Bay Muskogee Rincon Burkville	TX WI OK GA	505 8	0	
Fort Howard C M GE Georgia Gulf Corp. P	Green Bay Muskogee Rincon Burkville	WI OK GA	8		
GE E Georgia Gulf Corp. P	Muskogee Rincon Burkville	OK GA		26	Membrane
GE E Georgia Gulf Corp. P	Rincon Burkville	GA	5	20	Rocksalt; diaphragm
GE E M Georgia Gulf Corp. P	Burkville		-	9	Rocksalt; membrane (upgraded 1985)
Georgia Gulf Corp. P			6	4	
Georgia Gulf Corp. P	Mount Vernon	AL	24	7	Membrane
		IN	50	18	Captive brine; diaphragm
E	Plaquemine		410	19	Captive brine; diaphragm
	Bellingham	WA	82	29	Salt; mercury cell
HoltraChem A	Acme	NC	48	31.	Salt; mercury cell
	Orrington	ME	73	27	Salt; mercury cell
La Roche Holdings Inc. C	Gramercy	LA	181	36	Brine; diaphragm
Miles Inc. E	Baytown	TX	82	22	By-Product HCl; HCl electrolysis
Niachlor Inc.	Niagara Falls	NY	218	7	Brine; membrane (upgraded 1987)
Occidental C	Convent	LA	279	13	Captive salt dome; diaphragm
	Corpus Christi	TX	417	20	Brine; diaphragm
lr.	Deer Park	TX	347	56	Captive salt dome; mercury cell; diaphragm
I	Delaware City	DE	126	29	Salt mercury cell
L	La Porte	ТΧ	480	20	Captive salt dome; diaphragm
N	Mobile	AL	41	3	Mercury cell, KOH is produced; membrane (upgraded 1991)
IN IN	Muscle Shoals	AL	132	42	Mercury cell; KOH is produced
N	Niagara Falls		293	20	Captive salt dome; diaphragm (upgraded 1974)
-	Tacoma	WA	195	6	Rock salt; diaphragm; membrane (upgraded 1988)
T	Taft	LA	581	19/8	Captive salt dome; diaphragm (upgraded 1975); membrane
					(upgraded in 1986)
	Augusta	GA	102		Salt; mercury cell
1-	Charleston		230	32 .	Rock salt; mercury cell
	McIntosh	_	365	17	Brine diaphragm (upgraded 1977)
		_	82	34	Rock salt, mercury cell (upgraded 1960)
	Albany	OR	2	23	Magnesium chloride; by-product of metallic magnesium
	Henderson		104	18	Salt; diaphragm (upgraded 1976)
	St. Gabriel	LA	160	24	Salt; mercury cell
	Lake Charles	LA	1,126	25/17	Brine; mercury cell (upgraded 1969); diaphragm (upgraded 1977)
	Natrium		356	36/10	Brine; mercury cell (upgraded 1958); diaphragm (upgraded 1984)
Renco Group (Magnesium R Corp. of America)	Rowley	UT	14	17	Brine; by-product of magnesium metal production
	Vicksburg	MS	33	32	By-product of production of potassium nitrate from KCl
	Geismar		243	18	Brine; diaphragm
	Port Edwards	-	65	27	Salt; mercury cell; KOH is also produced
	Wichita		239		Brine; diaphragm (upgraded 1975); membrane (upgraded 1983)
the second s	Longview		136	19	Brine; diaphragm (upgraded 1975)
TOTAL		_	11,525	29 ^d	
Total Production	· · · · ·		10,973°	<u></u>	<u> </u>

Appendix C. Chlorine Plants in the United States (situation end of 1997)

a. Capacity numbers taken from Directory of Chemical Producers (DCP), SRI International b. Cell type data and year of plant upgrade taken from Chlorine Institute Pamphlet 10 (1994). c. Process data taken from DCP

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d. Capacity weighted average age
e. Chemical Manufacturers Association
*Time since last major upgrade

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