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Linear alpha-Olefins (681.5030)

by Elvira O. Camara Greiner with Yoshio Inoguchi

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SUMMARY

This report covers supply and demand for linear alpha-olefins (LAOs) with carbon chain-lengths of four (butene-1) and higher that are produced in facilities that generate an entire range of LAOs. Most of the current commercial LAO plants produce even-numbered alpha-olefins based on ethylene oligomerization. The exceptions are the coal-based synthetic fuel plant of Sasol Chemical Industries in South Africa, which currently produces hexene-1, octene-1 and small quantities of pentene-1; Q-Chem and Sinopec Beijing Yanhua Petrochemical, which produce hexene-1 from ethylene; and Dow, which produces octene-1 from butadiene in Tarragona, Spain. Butene-1 that is produced from refinery and steam cracker C_4 hydrocarbon streams is not included in this discussion.

The following table presents world supply/demand for linear alpha-olefins:

World Supply/Demand for Linear alpha-Olefins by LAO-Producing Region-2009 ^a (thousands of metric tons)						e		
	Annual Capacity (year-end)		Annual Capacity (year-end)			Apparent Consumption		Average Annual Consumption Growth Rate,
	2009	2014	Production	Imports	Exports	2009	2014	(percent)
North America	1,889	2,189	1,575	200	269	1,506	1,649	1.8
Central and South America				61	<u></u>	61	74	3.9
Western Europe	730	730	551	229	135	645	711	2.0
Central and Eastern	90	120	15	5	7	13	16	4.2
Europe								
South Africa	420	460	340	<u> </u>	270	70	90	5.2
Middle East	197 ^b	860	55	25	10	70	113	10.1
Japan	58 ^c	88	90	28	30	88	90	0.5
Other Asia	50	85	40	168		208	335	10.0
Oceania				5		5	6	3.7
Total	3,434	4,532	2,666	721	721	2,666	3,084	3.0%

a. Data include C₄ produced in alpha-olefin plants for all regions, except Western Europe.

b. Jubail United Petrochemicals' 150 thousand metric ton-per-year plant came on stream in December 2009.

c. Mitsubishi Chemical discontinued LAO production in Mizishima, Japan (annual capacity of 60 thousand metric tons) in May 2009.

SOURCE: CEH estimates.

In 2009, world alpha-olefins production was 2.7 million metric tons (reflecting a global capacity utilization rate of 82%—based on an effective capacity of approximately 3,250 thousand metric tons) valued at approximately \$3.7 billion. North America accounted for 59% of world production during 2009, followed by Western Europe (21%), South Africa (13%) and Japan (3%), with Russia, the Middle East and Other Asia accounting for the remainder.

World linear alpha-olefins capacity (excluding butene-1 from refinery streams) amounted to approximately 3.4 million metric tons by year-end 2009. There are currently (as of mid-2010) ten producers of alpha-olefins operating twelve plants internationally. The following table presents world producers of LAOs:

World Producers of Linear alpha-Olefins

Chevron Phillips Chemical Company LP CPChem) Cedar Bayou, TX, United States Now Chemical Tarragona, Spain	703		
CPChem) Cedar Bayou, TX, United States Oow Chemical Tarragona, Spain	703		
Oow Chemical Tarragona, Spain	705	CDCL	
Dow Chemical Tarragona, Spain		CPChem	
Tarragona, Spain			
	100	Telomerization of butadiene	Plant came on stream in 2007. Produces octene from butadiene.
demitsu Petrochemical Company, Ltd.			
Ichihara, Chiba Prefecture, Japan	58	Idemitsu	Plant came on stream in 1989.
NEOS			
Joffre, Alberta, Canada	265	Ethvl	Plant came on stream in late 2001.
Feluy, Belgium	300	Ethyl	Plant came on stream in July 1992.
ubail United Petrochemical Company (IUPC)			
Al Jubail, Saudi Arabia	150	alpha-SABLIN™	Plant came on stream in December 2009.
Jizhnekamskneftekhim			1
Nizhnekamsk, Russia	90	Ethyl	Original plant capacity was 180 thousand metric tons.
Datar Chemical Company (O-Chem)			
Mesaieed, Qatar	47	CPChem	Single-fraction 1-hexene plant (hexene- 1 is produced from ethylene). Operational since February 2003.
asol Chemical Industries Secunda, South Africa	420	Separation and distillation of the Fischer Tropsch stream from the Secunda coal-to- fuels process	High purity hexene-1, octene-1 and pentene-1 are isolated, primarily for co- monomers applications. C_{11} and higher LAOs are extracted for the local production of detergent alcohols. The first hexene unit came on stream in 1994. Hexene-1 capacity is 188 thousand metric tons. Octene-1 capacity is 186 thousand metric tons, including the 90 thousand metric ton unit that became operational in June 2008.
hell Chemical Company			
Geismar, LA, United States	921	Shell (SHOP)	
Ellesmere Port, United Kingdom	330	Shell (SHOP)	
td.			
Beijing, China	50	Ethylene trimerization	Came on stream in May 2007. Produces hexene-1 from ethylene.
Total	3,434		

The following table presents the largest world producers of LAOs by capacity:

	Major V	World Producers of Linear	alpha-Olefins	
		Annual Capacity as of September 1, 2010 (thousands of metric tons)	Percent of World Capacity	
	Shell	1.251	36.4	
	CPChem	703	20.5	
	INEOS	565	16.5	
	Sasol	420	12.2	
	Other	495	14.4	
	Total	3,434	100.0%	
	SOURCE:	CEH estimates.		
The following table present	ts planned w	vorld capacity additions	for LAOs:	
1	Planned Worl	d Capacity Additions for L	inear alpha-Olefi	ns
Company and Plant Location		Annual Capacity Addition (thousands of metric tons)	Es	stimated On-Strea Date/Remarks
CPCC Jubail Jubail Industrial City, Saudi Ara	abia	100	2012. Hexene-1.	
CPChem Cedar Bayou, Texas		200	First-quarter 2014	4. Hexene-1.
Dow Thailand		na	2012. Octene-1. I in Thailand in 20	Dow is starting a J 10.
Gurevesky Karabatan, Kazakhstan		20-40	2011. Butene-1.	
Jam Petrochemical Company Asulayeh, Iran		168	2011. LAOs. It i built. Butene-1 (1	is not likely that t 00 thousand metric
Mitsui Chiba, Japan	5	30	2010. Hexene-1.	
ONGC Petroadditions Ltd. (OPaL Dahej, Gujarat, India)	35	Fourth-quarter 20	12. Butene-1.
PetroRabigh Rabigh, Saudi Arabia		50	Butene-1. On stre	eam date for C_4 is u
Qatar Chemical Ćompany II Ltd. (Ras Laffan, Qatar	(Q-Chem II)	345	Fourth-quarter C_{4} is for C ₆ is 60 thous	2010. Broad ran s 58 thousand metri sand metric tons.
Sasol Lake Charles, Louisiana		100	2013. Hexene-1 technology. Cons	and octene-1. Sasol struction to commen
Total		1.059		
Total		1,058		

SOURCE: CEH estimates.

The largest uses for linear alpha-olefins are as comonomers for polyethylene production, for the production of alcohols (used largely in detergents and plasticizers) and for making polyalphaolefins (used in synthetic lubricants). The following table presents a comparison of end-use patterns for the major producing regions during 2009:

	North America	Western Europe	Japan
Polyethylene Comonomer	51	35	59
Oxo Alcohols	15	25	7
Polyalphaolefins	13	30	a
Other	21	10	34
Total	100%	100%	100%

Forecasts of growth rates vary significantly by region. Annual growth during 2009-2014 is expected to average 2% in North America, 2% in Western Europe and 0.5% in Japan. However, growth for comonomer-grade LAOs is expected to be considerably higher, or 5-6% globally. On a global scale, total LAO consumption should average 3% annually through 2014.

Key findings in the overall LAO market include the following:

- Mitsubishi Chemical discontinued production of linear alpha-olefins in May 2009.
- Jubail United Petrochemicals started its 150 thousand metric ton-per-year LAO plant in Al Jubail, Saudi Arabia in December 2009.
- The LAO market was relatively balanced in most of 2008. However, by the end of the year weak demand brought about by the global economic crisis saw LAO consumption decrease significantly, especially for polyethylene.
- Continued weak demand in the major LAO-consuming markets in 2009 resulted in lower LAO consumption and therefore production. By late 2009 and early 2010, when demand started to pick back up for polyethylene and polyalphaolefins, supplies were tight, especially for hexene-1 and decene-1.
- As of September 2010, the LAO market was relatively balanced as a result of additional capacity that came on stream in late 2009 and the overall ramp-up in production due to stronger demand from most sectors.
- Most of the announced capacity additions (for 2011-2013 commencement) are for C₄, C₆ and C₈ for the polyethylene market.
- Supplies of C_{10+} cuts will become tighter with capacity expansions focused on C_4-C_8 . However, some producers in markets that have historically consumed some of the higher cuts (e.g. detergent alcohols and alkyldimethylamines) have switched to using "natural" feedstocks.

World Average Annual Growth Rate.

2009-2014

(percent)

5-6%

The following table presents key findings and future concerns and implications for the LAO market by major consuming market:



Major Linear alpha-Olefin Consuming Market

Polyethylene (PE) Comonomer

- As of mid-2010, hexene-1 supply was relatively balanced after being tight in the beginning of the year.
- The market for octene-1, long in early to mid-2010, is being driven by high demand for elastomers (Dow announced plans to increase its specialty elastomers capacity in Texas and Spain in August 2010).
- The short-term outlook for PE indicates that from the middle of 2010, HDPE has been in larger supply. LLDPE has been tight and shows the best supply/demand balance of the two.
- In the long term it is expected that the new volumes from the Middle East will refocus trade flows in Asia and Western Europe. North America may become a more regional net importing market. However, developments leading to low/competitively priced U.S. ethane and ethylene feedstocks (namely natural gas in shale) are likely to be very positive for U.S.-based PE producers.
- Oversupply of PE in the market in the future will drive differentiation toward the high performance PE (from C₄ to C₆, C₈).

Polyalphaolefins

- Global PAO capacity utilization in 2009 was approximately 85% due to weak demand worldwide brought about by the global economic crisis (compared to typical operating rates exceeding 90% in recent years).
- PAO demand, and thus production, has rebounded strongly in 2010, with most producers running at or near effective capacity. If the global economy continues strong in the latter part of 2010, utilization rates could reach higher than 90% for the year.
- Growth for PAOs will not necessarily be limited by the expansion of Group III lubricants or gas-toliquid base stocks, as the high performance of PAOs versus Group III base oils has allowed them to retain market share in those sectors that require this performance.
- PAO demand will be driven by tightening emissions standards in the transportation industry (passenger vehicles and heavy-duty diesel).
- It is difficult to predict the growth of PAO for gear boxes in wind turbines. Although an important market, there are regulatory issues, among others, that come into play, making it a challenge to estimate consumption in the next five years.

Oxo Alcohols

Detergent Alcohols

- LAOs compete with natural fats and oils as raw materials in the production of detergent alcohols.
- LAO consumption for detergent alcohols saw a decrease from 2006 to 2009 with increasing imports of detergent alcohols from natural alcohol plants that came on line in Southeast Asia and Brazil.

Natural alcohol plants planned in Southeast Asia will increase detergent alcohol supplies and Asian exports will likely continue to increase.

Oxo Alcohols

Plasticizer Alcohols

- Competition from branched plasticizers contributed to decreased LAO consumption in this application.
- The global average annual growth rate for LAOs used in the production of plasticizer alcohols during 2009-2014 is estimated to be approximately 0.5%.

SOURCE: CEH estimates.

5%

2%

0-1%

The use of linear alpha-olefins in the production of polyethylene, for polyalphaolefins and for use in oil field drilling applications will be the fastest growing segments in North America. Globally, comonomergrade LAO consumption will also be the fastest growing application, particularly LAO consumption for the production of linear low-density polyethylene resins.

INTRODUCTION



alpha-Olefins are straight-chain hydrocarbons characterized by a double bond in the terminal or alpha position. Production of linear alpha-olefins (LAOs) based on ethylene oligomerization yields an entire range of even-numbered carbon chain-length LAOs rather than a single product. Chain lengths vary from four carbons (butene-1) to more than thirty carbons (expressed as C_{30+}). Since 1994, Sasol Chemical Industries has recovered C₅ and C₆ alpha-olefins contained in product streams for its coal-based synthetic fuel plant in Secunda, South Africa. It is the only producer of pentene-1 (C₅) in commercial quantities and has the potential to recover other LAOs. The following table lists the chemical names and chemical formulas for C₄ through C₃₀ LAOs:

	Linear alpha-Olefin N	omenclature
Carbon Chain Length	Chemical Name	Chemical Formula
4	Butene-1	CH ₂ =CH–(CH ₂)–CH ₃
5	Pentene-1	$CH_2 = CH - (CH_2)_2 - CH_3$
6	Hexene-1	CH ₂ =CH–(CH ₂) ₃ –CH ₃
8	Octene-1	$CH_2=CH-(CH_2)_5-CH_3$
10	Decene-1	CH ₂ =CH-(CH ₂) ₇ -CH ₃
12	Dodecene-1	CH ₂ =CH-(CH ₂) ₉ -CH ₃
14	Tetradecene-1	CH ₂ =CH–(CH ₂) ₁₁ –CH ₃
16	Hexadecene-1	CH ₂ =CH–(CH ₂) ₁₃ –CH ₃
18	Octadecene-1	CH ₂ =CH-(CH ₂) ₁₅ -CH ₃
20	Eicosene-1	CH ₂ =CH–(CH ₂) ₁₇ –CH ₃
22	Docosene-1	CH ₂ =CH-(CH ₂) ₁₉ -CH ₃
24	Tetracosene-1	CH ₂ =CH–(CH ₂) ₂₁ –CH ₃
26	Hexacosene-1	CH ₂ =CH–(CH ₂) ₂₃ –CH ₃
28	Octacosene-1	CH ₂ =CH–(CH ₂) ₂₅ –CH ₃
30	Triacontene-1	CH ₂ =CH-(CH ₂) ₂₇ -CH ₃
SOURCE:	G. R. Lappin and J	D. Sauer, alpha-Olefins
)	Applications Handbook,	Marcel Dekker, Inc., New
	York, 1989.	

Physical properties, and, therefore, commercial applications, vary depending on the chain length. Butene-1 is a gas at room temperature and C_6 - C_{18} LAOs are clear, colorless liquids, whereas C_{20+} LAOs are waxy solids. The utility of LAOs is usually because of the reactivity of the double bond. The most common commercial reactions involving LAOs include oxo reactions (hydroformylations), oligomerization/polymerization, simple addition reactions, alkylation reactions, sulfations/sulfonations and oxidations. Even-numbered alpha-olefins below C_{20} are available individually or in narrow carbon ranges. C_{20+} alpha-olefins are generally consumed in a broad range of chain lengths and some of these products contain high levels of other impurities. The following table summarizes major applications for LAOs by carbon chain length:

Linear alpha-Olefin Applications by Chain Length

C_4-C_8	Polymers and polyethylene comonomer
C_6-C_8	Low-molecular-weight fatty acids and mercaptans
C ₆ -C ₁₀	Plasticizer alcohols
$C_{10}-C_{12}$	Polyalphaolefins and other additives for lubricants,
	amine oxides and amines
C ₁₀ -C ₁₆	Detergent alcohols, nonionics and oil field chemicals
$C_{16}-C_{18}^{a}$	Oil field chemicals, lube oil additives and surfactants
$C_{20}-C_{30+}$	Oil field chemicals and wax replacement

a. Includes some C_{14} for oil field chemicals.

- SOURCES: (A) D. L. Burdick and W. L. Leffler, *Petrochemicals in Nontechnical Language*, Pennwell Publishing Company, Tulsa, Oklahoma, 1990.
 - (B) CEH estimates.

MANUFACTURING PROCESSES

Most linear alpha-olefins are currently commercially produced by ethylene oligomerization. The products of the oligomerization consist of even-numbered carbon chains ranging from C_4 to C_{30+} .* They have relatively small amounts of branched and internal olefin isomers and paraffin impurities. Sasol operates a purification facility in South Africa that recovers alpha-olefins contained in streams from coal-based synthetic fuel plants. This plant can recover odd-numbered alpha-olefins in addition to even-numbered products.

ETHYLENE OLIGOMERIZATION

The first commercial production of linear alpha-olefins by ethylene oligomerization was in 1966. All but one LAO producer worldwide currently use an oligomerization process, and the catalyzed chain-growth chemistry is similar in all of the methods. However, modifications by some producers at different stages of the reaction sequence lead to different product and chain-length distributions.

There are three main processes currently in use—Chevron Phillips Chemical (CPChem), Ethyl and Shell. The CPChem route uses a single stage for chain growth and displacement reactions, but is the least flexible of these processes in product distribution. The Ethyl process, which uses two separate steps for chain growth and displacement, gives greater flexibility in product distribution at the expense of increased process complexity and increased branching of heavier fractions. Shell's process is the most complex process of these three routes. It feeds selected alpha-olefins to an isomerization-disproportionation unit that can produce internal olefins for subsequent internal consumption or merchant sale at generally higher market values than the feed materials. The SHOP (Shell Higher Olefins Process) process is the most flexible with regard to product distribution. The isomerization-disproportionation part of the SHOP process converts alpha-olefins to internal olefins.

^{*} Butene-1 is also commercially produced by separation from various refinery and steam cracker C_4 hydrocarbon streams. In fact, butene-1 is most commonly classified as belonging to the C_4 hydrocarbon group of products, regardless of its source. For additional information on butene-1 see the CEH *Butylenes* marketing research report.

LINEAR ALPHA-OLEFINS

The chemical composition and quality of the alpha-olefin products vary somewhat from process to process. The CPChem and Shell processes result in a higher alpha-olefin content in C_{12+} cuts compared with the Ethyl process. Higher linearity in alpha-olefins is desired in some end-use applications, while the higher branching in C_{14} - C_{18} olefins produced by the Ethyl process is preferred in some other applications.

The different processes also yield different chain-length distributions. The Ethyl process (owned by INEOS) results in a Poisson distribution. The CPChem and Shell alpha-olefin processes produce a Schulz-Flory distribution. The following table presents typical product distributions for each of these processes, although each has some flexibility in chain-length distributions:

	Linear alpha-Olefins by Process ^a (percent by weight)			
	CPChem	Ethyl ^b (Canada/ Western Europe)	Shell	
\mathbb{C}_{4}	14	0	7-14	
$C_{6}-C_{10}$	41	70-77	25-41	
$C_{12} - C_{14}$	19	21-28	15-18)
$C_{16} - C_{18}$	12	0	11-15	
20+	14	2	14-42	
. See the	e following section	ons for more information	on product	

b. The Ethyl process (owned by INEOS) in Canada and Western Europe recycles C_4 to produce higher olefins.

SOURCES: (A) G. R. Lappin and J. D. Sauer, *alpha-Olefins Applications Handbook*, Marcel Dekker, Inc., New York, 1989 (data for ETHYL [now INEOS]).

(B) CEH estimates (all other data).

The processes used by INEOS and CPChem are based on Ziegler chemistry using ethylene and a triethylaluminum catalyst to promote chain growth of even-carbon-number alpha-olefin molecules. In both of these methods, the chain-length distribution of the linear alpha-olefin product can be changed to a limited extent by altering operating conditions. Notable differences in the two routes include the following: (1) olefins can be recycled for further growth in the Ethyl process, thus maximizing yields of the C_6-C_{14} olefins, and (2) INEOS' product contains a slightly higher percentage of branched material, particularly in the detergent range, than the product obtained via the CPChem process.

Shell's SHOP ethylene oligomerization route to alpha-olefins differs from the others in that it uses a proprietary nickel complex catalyst in the initial chain-growth step and it can convert alpha-olefins to internal olefins of a desired carbon number by isomerization and disproportionation.

The major commercial processes, in addition to other processes available for commercialization, are described in the following sections. For more detailed information on manufacturing processes for the production of linear alpha-olefins, see the Process Economics Program *Linear alpha-Olefins* report.

CHEVRON PHILLIPS CHEMICALS' ETHYLENE OLIGOMERIZATION (ZIEGLER PROCESS)

Chevron Phillips' process uses a technology developed by Gulf Oil Chemicals Company, which Chevron (now Chevron Phillips Chemical Company LP [CPChem]) acquired in 1985. This one-step conventional Ziegler process combines high-temperature addition of ethylene and displacement of the generated aluminum alkyls to produce linear alpha-olefins. Ethylene reacts with the triethylaluminum growth products to yield even-numbered carbon olefins by displacement. Typical reaction conditions are 175-290°C at pressures of 2,000-4,000 psi. The resulting carbon number range for this process is C_4 - C_{30+} , with distribution skewed toward the lower-carbon-number olefins in the C_4 - C_{10} range. The basic reactions occurring in this process can be represented as follows:



The linear alpha-olefin reaction product is a mixture of olefins that are subsequently purified and can also be fractionated into specific carbon chain-length fractions. The metal alkyl catalyst is deactivated and removed before the products are separated to avoid further isomerization.

The distribution of chain lengths may be varied to some extent by altering process conditions in order to meet merchant market demands. The curve typically is a geometric distribution with ranges as follows:

	Percent by Weight
C_4	14
$C_{6}-C_{10}$	41
$C_{12} - C_{14}$	19
$C_{16} - C_{18}$	12
C ₂₀₊	14

Typical linear alpha-olefin content ranges from 90.5% to 98.3% for C_4 - C_{18} , with the butene-1 fraction being the most pure. Impurities include internal olefins, paraffins and branched olefins. Typical alpha-olefin content for C_{20} - C_{24} is 88.3% and 33.0-48.0% for C_{24} - C_{30+} .

CPChem currently uses this technology in the United States at its plant in Cedar Bayou, Texas. Mitsubishi Chemical Corporation licenses the CPChem technology for its plant in Mizushima, Okayama Prefecture, Japan, but Mitsubishi Chemical discontinued its production in May 2009. Another plant using this technology started up in September 1992 at Chemopetrol's SPOLANA chemical complex in Neratovice, the Czech Republic (but this plant closed in 2003). CPChem has expanded its U.S. production site several times. CPChem (formed from the chemical businesses of Chevron and Phillips) acquired the Phillips hexene-1 technology. Q-Chem, a joint venture between CPChem and Qatar Petroleum, has a hexene-1 plant in Qatar based on Phillips' technology. However, Q-Chem's hexene-1 is produced from ethylene.

INEOS' ETHYLENE OLIGOMERIZATION (MODIFIED ZIEGLER PROCESS)

A second oligomerization process was developed and first used by Ethyl Corporation. Ethyl's LAO business became part of Albemarle Corporation in 1994, when the former split out its chemical activities that were not related to lubricants and fuel oil additives. Subsequently, Albemarle sold its LAO, polyalphaolefin and detergent alcohol businesses to Amoco Corp. in 1996. The latter then merged with British Petroleum plc (BP) to form BP Amoco Corp. within North America. The company later changed its name to BP. BP's refining, olefins and derivatives business (Innovene) was fully acquired by INEOS in 2005.

In the first stage, ethylene is oligomerized by trialkylaluminum growth promotion in a process similar to the one-step process. Typical reaction conditions are temperatures of 116-132°C and pressures of 2,700-3,000 psi.

In the displacement step, the trialkylaluminum compounds formed in the first stage are displaced with ethylene to yield linear alpha-olefins in the C₄-C₁₈ range and to regenerate the triethylaluminum for recycle. Reaction conditions range from temperatures of 260-316°C and pressures of 230-250 psi.

The product separation step processes unconverted ethylene from the growth and displacement reactor stages for recycle and separates the individual C_4 - C_{18} olefins. The linear alpha-olefins are distilled at 90-125°C.

Typical linear alpha-olefin content ranges from 93.7% to 99.7% for C_4 - C_{12} , with the butene-1 fraction being the most pure. Impurities include paraffins, internal olefins and branched olefins. This process yields higher amounts of the branched oligomers than the one-step reaction because of the continued reaction of alpha-olefin product in the recycle step. These branched products are concentrated in the higher-carbon-number C_{14} - C_{18} ranges, whereas the shorter-chain-length linear alpha-olefins have lower concentrations of the branched impurities. Typical alpha-olefin content for C_{14} is 81.0% and 62.7% for C_{16-18} .

To satisfy changing market demands, distribution of chain lengths can be altered by changing reaction conditions. Automated computer control allows for shifting the production of chain-length distributions. Current production emphasis is on the lower-carbon-number olefins, to supply the comonomer, plasticizer and polyalphaolefin markets. The typical product distribution by chain-length range exhibits a Poisson distribution, as follows:

	Percent by Weight
C_4	12
$C_{6}-C_{10}$	54-60
$C_{12}-C_{14}$	20-25
$C_{16}-C_{18}$	5-7
C ₂₀₊	2

BP used this process in its plant at Pasadena, Texas (which closed in 2005). INEOS uses similar technology with the addition of a butene recycle loop in a plant at Feluy, Belgium that began operations in mid-1992 and at Joffre, Alberta (Canada) that became operational in May 2002. The resulting product distribution is as follows:

Percent by Weight

C_4	0
$C_{6}-C_{10}$	70-77
C ₁₂ -C ₁₄	21-28
C ₂₀₊	2

Due to INEOS' product distribution, the company is the second-largest producer of C_6 - C_{10} LAOs. A linear alpha-olefin plant using this ethylene oligomerization process started up in 1990 at Nizhnekamsk in the Tatar Republic of Russia. Capacity for the plant is currently 90 thousand metric tons. Only one of two lines is operational since it restarted in 1998 after a period of inactivity.

SHELL'S ETHYLENE OLIGOMERIZATION (SHELL HIGHER OLEFINS PROCESS-SHOP)

Shell first commercially used the SHOP technology at its Geismar, Louisiana plant in 1977. Unlike the Ethyl and CPChem processes, which use a triethylaluminum catalyst, the ethylene oligomerization step in SHOP is based on a non-Ziegler catalyst system. SHOP's initial products are even-numbered carbon linear alpha-olefins from C_4 to C_{20+} . Shell distills selected individual carbon-number products in the C_4 - C_{20} range and also supplies blends. The alpha-olefins not separated for sale are converted to C_8 - C_{18+} internal olefins via a series of isomerization and disproportionation steps. The internal olefins are subsequently converted via a modified oxo process to detergent-range primary alcohols or sold.

This process selectively yields high-purity linear alpha-olefins and detergent-range internal olefins. The SHOP process is very flexible and alpha-olefins can be withdrawn for sale or converted into internal olefins as desired. Detergent-range internal olefins can be used for manufacture of alcohols, sold or recycled. This flexibility allows Shell to meet a wide range of product demands. Shell developed this process to fit its own special situation as a major supplier of detergent surfactants and intermediates, in particular detergent-range alcohols and/or their ethoxylates.

The initial step of the process is conducted at moderate temperatures (80-120°C) and pressures (1,000-2,000 psi). The ethylene is oligomerized to C_4 - C_{20+} linear alpha-olefins using a proprietary liganded nickel catalyst system. The chain-length distribution of alpha-olefins from the oligomerization step can be varied over the following range:



The C_4 - C_{20} alpha-olefins are distilled from the mixture and separated into high-purity individual components and blends to be marketed as Shell's Neodene[®] linear alpha-olefins. Typical linear alpha-olefin content ranges from 94% to 99% for C_4 - C_{20} , with the hexene-1 fraction being the most pure. Impurities include internal olefins, paraffins and branched olefins. Shell has the highest linearity of alpha-olefin producers.

In the next step of the process, Shell converts C_4 and C_{20+} alpha-olefin fractions and a selected portion of the alpha-olefin products to C_{11} - C_{14} detergent-range internal olefins by isomerization and disproportionation reactions. The olefin fractions are first purified using absorbent beds and then isomerized at moderate temperatures (80-140°C) and pressures (50-250 psi) using a proprietary catalyst. The internal olefins are then disproportionated, using a catalyst and reaction conditions similar to the isomerization step, to new pairs of internal C_4 - C_{20+} odd- and even-numbered carbon chain lengths. This disproportionation of the internal olefins yields 10-15% by weight of detergent-range C_{11} - C_{14} product per pass. These olefins are then recovered by distillation. The remaining olefins are recycled through the disproportionation or isomerization and disproportionation steps.

After distillation, the C_{11} - C_{14} internal linear olefins are converted via a modified oxo process to Neodol[®] detergent-range alcohols; 80% is linear alcohol and the remainder is the 2-methyl isomers. In addition, some quantities of C_9 - C_{11} alcohols are made from processing C_8 - C_{10} internal olefins. (alpha-Olefins can also be processed directly through the oxo alcohol unit.) Internal olefins in the C_8 - C_{18} range are also offered on the merchant market.

In addition to the Geismar, Louisiana facilities, SHOP is also used in the Shell Chemicals UK Ltd. plant in Ellesmere Port (Stanlow), United Kingdom, which has been operating since the fall of 1982.

IDEMITSU'S ETHYLENE OLIGOMERIZATION PROCESS

Idemitsu commercialized its own process at a plant that started up in Japan in mid-1989. This process, also based on ethylene oligomerization, utilizes a proprietary zirconium-based catalyst system. The reported advantage of this process is that it produces fewer branched and internal olefins and fewer waxes are formed. Recommended reaction conditions are 100-150°C at pressures of more than 355 psi. Product distribution can be controlled within the constraints of the geometric pattern by appropriately selecting the reaction conditions and catalyst concentration and composition.

VISTA'S ALFENE® PROCESS

CONDEA Vista Company (now Sasol since the company acquired CONDEA Vista in 2001), developed its Alfene[®] process for the production of linear alpha-olefins, but has not commercialized it. This process parallels its Alfol[®] ethylene oligomerization process for the production of linear alcohols and uses a Ziegler-type low-temperature ethylene growth stage with triethyl-aluminum similar to the Amoco process. The triethylaluminum is separated from the displacement product by formation of a complex and a proportion of the lower olefins is recycled to the growth reaction step. This process also yields a Poisson distribution of LAOs.

EXXON'S ETHYLENE OLIGOMERIZATION PROCESS

Exxon Research and Engineering Company has a low-pressure, low-temperature variation of the ethylene oligomerization process that uses a soluble catalyst derived from alkyl aluminum halides and transition metal compounds. Products are high-purity (greater than 98%) linear alpha-olefins with an even number of carbon atoms. Product distribution, which follows a geometric pattern, can be controlled by altering the process conditions. This process has not been commercialized.

DUPONT'S VERSIPOLTM PROCESS

DuPont has developed a one-step ethylene oligomerization process using its Versipol[™] family of catalysts (such as a mixture of an iron tridentate catalyst and an alkyl aluminum cocatalyst) in the production of LAOs with low capital and operating costs and high product purity (98%). This system offers greater flexibility in the selection of the LAO product distribution. DuPont is offering this technology for license.

SABIC/LINDE ALPHA-SABLINTM PROCESS

Linde and SABIC (Saudi Arab Basic Industries Corporation) developed their linear alpha-olefins technology in partnership under the commercial name of alpha-SABLINTM. alpha-SABLINTM utilizes a two-component catalyst system for homogenous, liquid-phase ethylene oligomerization. The catalyst system consists of a proprietary zirconium (Zr) compound used in conjunction with an aluminum alkyl cocatalyst, providing narrow carbon-number distributions. The product has more branching in the higher-range alpha-olefins.

The LAOs are fed into a series of conventional separation columns to obtain the desired product cuts. The high-selectivity, one-stage reactor is claimed to produce high-purity products that do not require secondary stages, such as superfractionation distillation towers for removal of side products. Technical features of the process include reduced reaction temperature and pressure in the oligomerization reactor, improved control of molecular weight distribution of reactor products, and high product selectivity and purity because of a nonnoble metals catalyst. The relative distribution of the different olefins produced by alpha-SABLINTM technology is shown in the table below.



According to Linde, alpha-SABLINTM offers more moderate reaction conditions and the highest flexibility to produce alpha-olefins in different product ranges as compared to other LAO technologies. Thus, the technology has an advantage that it can be changed to match the products demand.

OTHER ALPHA-OLEFIN SYNTHESIS ROUTES

SASOL'S PURIFICATION PROCESS

Sasol Chemical Industries of South Africa has built a plant for recovering C_5 - C_8 alpha-olefins. The Sasol process is unique in that the alpha-olefins are produced from a gasification plant and Fischer-Tropsch conversion. Coal is converted to crude gas under pressure, high temperature and in the presence of steam

and oxygen. After purification, the synthesis feed gas is sent to reactors for Fischer-Tropsch conversion, in which hydrogen and carbon monoxide react in the presence of an iron-based catalyst and pressure to yield a broad spectrum of hydrocarbons in the C_1 - C_{20} range. Prefractionation splits out the alpha-olefins and a second (etherification) stage converts branched-chain olefins into heavy ethers that are distilled out from the primary linear olefin stream. The final stage consists of superfractionation and distillation to separate out the impurities (mainly paraffins and cyclic olefins) and to produce pure hexene-1, pentene-1 and octene-1.

The purification facility came on stream in 1994. Higher olefins can also be recovered and the company is planning to do so.

PARAFFIN WAX CRACKING

Until the introduction of the oligomerization route, linear alpha-olefins were produced by thermally cracking waxy paraffins found in crude oils. The LAOs produced by cracking contained both odd- and even-numbered carbon chains with lengths ranging from C_5 to C_{20+} . Paraffin wax cracking was less efficient than ethylene oligomerization processes since it was energy-intensive, it did not yield as pure a product and it contained higher levels of internal olefins, dienes and paraffin impurities. Limited feedstock supplies also contributed to its demise.

The process was conducted in the vapor phase at relatively low cracking temperatures followed by rapid quenching to prevent side reactions such as isomerization or cyclization. Using a purified paraffin feedstock resulted in linear alpha-olefins of 90% purity by weight.

This technology is no longer in use commercially. Until 1985 Chevron Chemical Company produced linear alpha-olefins by thermal cracking of paraffin waxes at Richmond, California. The Royal Dutch/Shell Group also produced linear alpha-olefins by wax cracking at Pernis, Netherlands and at the Berre, France facility, but these plants have been permanently shut down since 1986.

DEHYDRATION OF ALCOHOLS

The dehydration of primary alcohols was the first commercial process for linear alpha-olefins. Linear primary alcohols, when vaporized and passed over a heated catalyst, lose a water molecule to yield an alpha-olefin of the same carbon number.

 $RCH_2CH_2OH \xrightarrow{catalyst} RCH=CH_2 + H_2O$

This process has not been used since 1966 when Archer Daniels Midland Company discontinued production of alpha-olefins.

ISOMERIZATION OF INTERNAL OLEFINS

Although internal olefins can be isomerized to yield linear alpha-olefins, they are themselves valuable intermediates in processes such as the synthesis of oxo alcohols and linear alkylbenzenes. They are not considered to be an economical feedstock for the manufacture of alpha-olefins.

SUPPLY AND DEMAND BY REGION

NORTH AMERICA

PRODUCING COMPANIES

The following table presents North American producers of LAOs (all three can produce their own ethylene feedstock):

North A	merican H	Producers	of Linear	alpha-Olefins
	Ann as (tl n	nual Capa of Mid-20 housands netric tons	city)10 of s)	
Company and Plant Location	C ₁ a	C6+	Total	Remarks
	-4	- 0+		
Chevron Phillips Chemical Company LP				
Olefins and Polyolefins Business Unit	100	601	500	
Cedar Bayou, TX	102	601	703	Listed capacity is for LAOs in the C_4-C_{30+} range. The company markets individual linear alpha-olefins in the C_4-C_{18} range and $C_{20}-C_{24}$, $C_{24}-C_{28}$ and C_{30+} fractions. Captive uses include polyalphaolefins, polyethylene comonomers and lube additives. alpha-Olefins are sold under the trademark AlphaPlus [®] NAO.
NFOS				
Joffre, Alberta, Canada		265	265	Current capacity represents the normal alpha- olefins production in the C_6 - C_{18} range, but is flexible. C_4 is recycled to produce higher olefins. The company markets individual linear alpha-olefins in the C_6 - C_{14} range and C_{12} - C_{14} , C_{14} - C_{16} and C_{16} - C_{18} fractions. Captive uses in- clude polyalphaolefins. In 2005, Innovene (formerly BP's olefins and derivatives business) was fully acquired by INEOS. The company started production at its 250 thousand metric ton-per-year plant at Joffre, Alberta, Canada in late 2001. Capacity was increased by 15 thousand metric tons since 2007.
Geismar I A	34-41 ^b	880°	921	Existing capacity is for C ₁ -C ₂₂ alpha-olefing
			/21	Capacity includes alpha-olefins that are converted to internal olefins that, in turn, are consumed to make detergent and other alcohols or sold. The company markets individual linear alpha-olefins in the C ₄ -C ₂₀ range and C ₁₂ -C ₁₄ , C ₁₄ -C ₁₆ , C ₁₆ -C ₁₈ and C ₁₄ -C ₁₈ fractions. Most butene-1 is captively consumed in the SHOP process and for polybutene-1 production. Internal olefins and alpha-olefins are sold under the trademark Neodene [®] .
Total	136-143	1,746	1,889	

- a. Includes only C_4 produced by ethylene oligomerization. There are other producers of C_4 (butene-1) from other sources not included in this report.
- b. Shell consumes some of its 1-butene internally in its I/D unit for conversion to longer-chain internal olefins. The assignment of the volume of C_4 and C_{6+} is arbitrary, but may approximate current practices.
- c. The volume assigned to C_{6+} olefins includes alpha- and internal olefins. This volume is estimated based on the difference between Shell's total capacity and its C_4 capacity of 41 thousand metric tons.

SOURCE: CEH estimates.

The preceding table includes capacity to produce butene-1 by means of ethylene oligomerization. Chevron Phillips Chemical and Shell isolate butene-1 from their ethylene oligomerization plants. Both producers use captive ethylene for making LAO. For more information on butene-1 from other sources, see the CEH *Butylenes* marketing research report.

All three LAO producers use their own process to manufacture alpha-olefins, each process yielding different chain-length distributions. See the **MANUFACTURING PROCESSES** section for a description of the distribution patterns for each process. An ongoing concern of producers is the need to adjust process conditions to maximize the value of their production by increasing production of chain lengths in high demand and minimizing production of chain lengths with low demand. However, limitations of each process, laws of stoichiometry and practical considerations (e.g., the high costs of repeated recycling) place limits on the flexibility of the chain-length distribution pattern. Producers inevitably manufacture some chain lengths in excess of existing demand, although they have often been successful in finding new consumers for these cuts. Producers try to minimize butene-1 production since they compete on the merchant market with lower-cost raffinate-derived butene-1 produced by ExxonMobil and Texas Petrochemicals. On some occasions, a very small proportion of fractions that cannot be sold at prices above fuel value can be used as a fuel in the plant or sold for this purpose.

Shell's higher olefin process (SHOP) includes the capability to convert excess alpha-olefins to C_8-C_{18+} internal olefins via a series of isomerization and disproportionation steps. Its annual capacity for internal olefins is approximately 250 thousand metric tons. Most of the internal olefins are subsequently converted via a modified oxo process to detergent-range primary alcohols or plasticizer-range alcohols, but some are sold. See the **MANUFACTURING PROCESSES** section for information on this process.

Sasol plans on constructing a hexene-1 and octene-1 plant in the United States with a combined annual capacity of 100 thousand metric tons. The process will use Sasol tetramerization technology and construction is slated to commence in 2011. The plant will be located at Sasol's Lake Charles, Louisiana chemical complex.

CPChem plans to build a hexene-1 plant with over 200 thousand metric tons of capacity at its Cedar Bayou, Texas, complex. The plant is expected to start up in first-quarter 2014, using CPChem's proprietary selective hexene-1 process.

PRODUCTION

The following table presents North American production of LAOs:

North American Production of Linear alpha-Olefin	lsa
(thousands of metric tons)	

1995		1,202	
1996		na	
1997		1,313	
1998		1,366	
1999		1,409	
2000		1,415	
2001		1,508	
2002		1,558	
2003		1,732	
2004		1,815	
2005		1,760	
2006		1,791	
2009		1,575	
2010 ^b		1,650	
a. Includes bute		zation.	
	···· · · ·····························		
b. Projected est	imate.		
SOURCES: (A)) NPRA Quarterly Report,	National	
	Petrochemical Refiners As	ssociation	
	(data for 1997-2001).		
(B)) CEH estimates (data for 1	1995 and	
	2002-2010).		

North American production of LAOs (including butene-1) from ethylene oligomerization reached 1,575 thousand metric tons, valued at \$1.8 billion in 2009. North American capacity utilization during 2009 was approximately 84%.

The following table presents North American production of butene-1 and C₆ and higher alpha-olefins:

	North An	nerican Produ (thousands	ction of Linear alpha- s of metric tons)	Olefins	
	Butene-1 ^a	C ₆ -C ₁₀	C ₁₁ and Higher	Total	Total
2003	147	787	798	1,585	1,732
2006	118	745	928	1,673	1,791
2009	115	680	780	1,460	1,575

a. Data include only butene-1 produced by ethylene oligomerization.

SOURCE: CEH estimates.

For historical production and sales of linear alpha-olefins, please see APPENDIX.

CONSUMPTION

The following table presents North American consumption of linear alpha-olefins:

	North Ameri	can Consumptio	n of Linear a	lpha-Olefins		
	Main	200	9	2014	4	Average Annual
	Carbon Number Range	Thousands of Metric Tons	Percent of Total	Thousands of Metric Tons	Percent of Total	Growth Rate, 2009-2014 (percent)
C_6 and Above (alpha and internal)						
Polvethylene Comonomers	6-8	673	45	762	46	2.5
Oxo Alcohols						
Detergent Alcohols	12-20+ ^a	200	13	221	13	2.0
Plasticizer Alcohols	6-10 ^a	25	2	22	1	-2.5
Polyalphaolefins ^b	10.8-12	201	13	256	6	5.0
Oil Field Drilling Fluids	16-18	90°	6	100	6	2.1
Other Lubricant Additives	16-24	38	3	39	2	0.5
Alkyldimethylamines	10 2 .	20	U			010
and Dialkylmethylamines	8-18	66	4	15		-25 6 ^d
Alkenylsuccinic Anhydrides	16-20	25	2	29	2	3.0
Fatty Acids (C ₇ -C ₀)	6-8	19	1	22	-	3.0
alpha-Olefin Sulfonates	14-16	14	1	15	1	14
Linear Alkylbenzene	10-14	9	1	9	1	-1.0
Linear Mercaptans	6-16	8		8	1	2.0
Aluminum Alkyls	6-8	7	0	8	1	2.7
Chlorinated alpha-Olefins	10-16	5		5	0	0
Alkyldinhenylether Disulfonates	12-14	4	0	4	Ő	2.0
Other ^e	6+	22	ů 1	24	1	1.8
omer	01					1.0
Total		1,406	93%	1,539	93%	1.8%
				,		
C ₄ (Butene-1)						
Polyethylene Comonomers ^f	4	90	6	g		
Polybutene-1 ^f	4	g		g		
Aluminum Alkyls	4	g		g		
Butylene Oxide ^f	4	g		g		
C_{ϵ} Aldehydes ^f	4	g		g		
Linear Mercantans ^f	Å	g		g		
Other ^f	4	10	1	110		2.5 ^h
						2.5
Total ⁱ		100	7%	110	7%	1.9%
		100	, , , ,	-		2.270
Total		1.506	100%	1.649	100%	1.8%
		1,000	100/0	1,012	10070	1.070

- The carbon number range is for olefins in the form consumed for these end uses. Data include internal olefins that are made a. from C₄ and higher alpha-olefins; estimates, therefore, include C₄ in addition to higher alpha-olefins.
 - Most PAOs are based on C_{10} ; C_8 and C_{12} are also used to make PAOs.

b.

с.

- Although relatively small compared with C16-C18 usage, beginning in 2003, includes 9-11 thousand metric tons of C14 for offshore and onshore drilling as well as C_{14} ester (from LAO).
- d. The consumption of LAOs for the production of ADMA will decline at a high average annual rate as a result of P&G's feedstock switch from LAOs to natural fatty alcohols in 2010.
- Other uses include those described later in the consumption section, as well as inventory changes and any products burned for e. fuel value.

LINEAR ALPHA-OLEFINS

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- f. Most C_4 (butene-1) markets are also supplied from refinery sources. Estimates in the table include only butene-1 supplied by LAO producers. See the CEH *Butylenes* marketing research report for estimates of total butene-1 consumed. Because producers have a choice of butene-1 suppliers, the amount supplied by LAO producers can vary from year to year.
- g. Since butene-1 from both LAO production and refinery sources is sold into these applications, the total volume of butene-1 from LAO production consumed is not easily estimated for these applications. Thus, any such use of LAO-derived butene-1 is included in the value shown in the Other category.
- h. The volume and growth rate shown for the butene-1 derived from LAOs does not represent a composite of growth in the individual market segments as shown in the table. Instead, it reflects the assumption that all butene-1 produced from ethylene oligomerization will find a home in the overall butene-1 market and that growth will be driven by the higher fractions of LAOs.
- i. Total butene-1 excludes butene-1 consumed by Shell to make internal olefins for oxo alcohol production or for export. The volume of butene-1 consumed for Shell's oxo alcohol production is included in the consumption shown under plasticizer and detergent alcohols in the table.

SOURCE: CEH estimates.

North American alpha-olefin consumption, which reached 1,506 thousand metric tons in 2009, decreased at an average annual rate of approximately 2% during 2006-2009 compared to an average annual increase of 3.6% during 2003-2006. Most of this decrease took place in 2008 and throughout most of 2009. The economic crisis negatively affected consumption of LAOs in most applications, particularly comonomers used for the production of polyethylene (LLDPE and HDPE), the largest consuming market. By the end of 2009, production of HDPE and LLDPE was up significantly over 2008, but not quite to 2006 levels. As of August 2010, consumption of LAOs in most major-consuming markets was up as the economy improved and markets tried to regain volume lost during the economic recession.

The fastest-growing markets are those that use LAOs as raw materials (largely C_{10}) for polyalphaolefins (for synthetic lubricants), as comonomers (C_4 - C_8) in polyethylene, in drilling fluids (C_{16} - C_{18}), and for alkenylsuccinic anhydrides (ASA). Although growing more slowly, detergent-range alcohols absorb large volumes of LAOs as raw materials, mainly in the mid-range (C_8 - C_{14}). These variations in current and projected growth of end-use markets that require different chain lengths present special problems to producers. As already described, alterations in the process conditions permit producers to make adjustments to meet market demand, but since some imbalance persists, wide variations in prices for the various chain lengths often are employed to further balance demand.

Total North American alpha-olefin consumption is expected to increase at an average annual rate of approximately 2% during 2009-2014, reaching 1.65 million metric tons in 2014.

The following table presents historical consumption of North American linear alpha-olefins:

Historical North American Consumption of Linear alpha-Olefins (thousands of metric tons)

	Main Carbon Number Range	1995	1999	2003	2006
C_6 and Above (alpha and internal)					
Polyethylene Comonomers	6-8	277	413	483	656
Alcohols					
Detergent Alcohols	12-20+ ^a	231	277	256	242
Plasticizer Alcohols	6-10 ^a	125	125	116	70
Polyalphaolefins ^b	10, 8-12	131	147	178	219
Oil Field Drilling Fluids	16-18	23	73	79 ^c	84 ^c
Other Lubricant Additives	16-24	27	39	40	42
Alkyldimethylamines					
and Dialkylmethylamines	8-18	29	31	58	64
Fatty Acids (C ₇ -C ₉)	6-8	20	29	29	25
Alkenylsuccinic Anhydrides	16-20	16	21	23	25
alpha-Olefin Sulfonates	14-16	11	13	14	13
Linear Alkylbenzene	10-14	11	23	11	10
Linear Mercaptans	6-16	6	8	9	8
Aluminum Alkyls	6-8	5	6	7	8
Chlorinated alpha-Olefins	10-16	7	7	6	6
Alkyldiphenylether Disulfonates	12-14	3	4	4	4
Other ^d	6+	20	21	22	23
Total		941	1,235	1,336	1,499
C ₄ (Butene-1)					
Polyethylene Comonomers ^e	4	55	95	91	95
Polybutene-1 ^e	4	23	f	f	f
Aluminum Alkyls	4	4	5	f	f
Butylene Oxide ^e	4	14	f	f	f
C ₅ Aldehydes ^e	4	5	f	f	f
Linear Mercaptans ^e	4	3	f	f	f
Other ^e	4	3	19	16	12
Total ^g		107	119	107	107
Total		1,048	1,354	1,443	1,606

a. The carbon number range is for olefins in the form consumed for these end uses. Data include internal olefins that are made from C_4 and higher alpha-olefins; estimates, therefore, include C_4 in addition to higher alpha-olefins.

Most PAOs are based on C_{10} ; C_8 and C_{12} are also used to make PAOs. Beginning in 2006, includes C_8 - C_{12} .

- c. Although relatively small compared with C_{16} - C_{18} usage, beginning in 2003, includes 9-11 thousand metric tons of C_{14} for offshore and onshore drilling as well as C_{14} ester (from LAO).
- d. Other uses include those described later in the consumption section, as well as inventory changes and any products burned for fuel value.

- e. Most C₄ (butene-1) markets are also supplied from refinery sources. Estimates in the table include only butene-1 supplied by LAO producers. See the CEH *Butylenes* marketing research report for estimates of total butene-1 consumed. Because producers have a choice of butene-1 suppliers, the amount supplied by LAO producers can vary from year to year.
- f. Since butene-1 from both LAO production and refinery sources is sold into these applications, the total volume of butene-1 from LAO production consumed is not easily estimated for these applications. Thus, any such use of LAO-derived butene-1 is included in the value shown in the Other category.
- g. Total butene-1 excludes butene-1 consumed by Shell to make internal olefins for oxo alcohol production or for export. The volume of butene-1 consumed for Shell's oxo alcohol production is included in the consumption shown under plasticizer and detergent alcohols in the table.

SOURCE: CEH estimates.

The estimates for C_4 (butene-1) demand include only the material supplied from alpha-olefin plants. See the discussion sections for estimates of total butene-1 consumed from all sources. Butene-1 from LAO production is disposed of as a relatively low-value coproduct (often below the price of ethylene) in a larger butene-1 market served by other sources. Consequently, its distribution within market segments is difficult to quantify and can vary significantly from year to year. Furthermore, growth for butene-1 from LAOs is driven by its supply, which depends upon demand for the other LAO fractions, rather than demand within the actual markets for butene-1. Thus, LAO producers will dispose of butene-1 within its markets at whatever price they can obtain, so long as it is above fuel value. For this reason, the markets for butene-1 are shown separately in the table, following those that consume the higher fractions. In addition, some butene-1 is converted to higher carbon-numbered internal olefins for conversion to alcohols by Shell in its isomerization/disproportionation unit. Most of the resulting olefins are internally consumed to produce plasticizer and detergent alcohols and this volume is accounted for within these two categories in the consumption table. Some internal olefins are exported to other countries.

For historical U.S. consumption (excluding Canada) of linear alpha-olefins, please see APPENDIX.

The end-use market discussion that follows presents each segment in order of market size (largest first), regardless of chain length (i.e., whether from butene-1 or higher LAOs).

Polyethylene Comonomers

Linear alpha-olefins, specifically butene-1 (C₄), hexene-1 (C₆) and octene-1 (C₈), are used as comonomers in the production of high-density polyethylene (HDPE) and linear low-density polyethylene (LLDPE) resins. This category also includes the even lower-density products (i.e., polyolefin plastomers and elastomers) that are made by the same processes. HDPE and LLDPE are both linear polyethylenes produced in low-pressure processes; LAOs are not consumed for production of conventional branched low-density polyethylene (LDPE) resins made in high-pressure processes. The LAO comonomers create short-chain branching in the polymer that prevents polymer chains from packing together tightly, thus lowering product density. The higher the comonomer concentration, the lower the density of the resin. In addition to controlling density, alpha-olefin comonomer also influences these properties. Hexene-1 and butene-1 are comonomers used in HDPE (slurry phase), and in gas phase and solution phase LLDPE. Octene-1, demand for which has remained strong in recent years, is used in solution-phase LLDPE plants.

The following table presents North American consumption of comonomers in PE production in 2009:



193

129

322

235

116

351

619

116

129

864

The following table presents North American comonomer use in PE by type

Plastomers/Elastomers

LLDPE

HDPE

Total

SOURCE:

Overall use of LAO comonomer from LAO producers is expected to grow at a more modest average annual rate of 2.1% during 2009-2014. This growth primarily reflects increasing production of plastomers and elastomers in North America during this time period and increased production of LLDPE in 2010. In terms of LAO component consumption for the combined production of LLDPE and HDPE, growth will be relatively flat. As of mid-2010, North American production of HDPE and LLDPE was higher than in 2009, driven by exports to regions with high demand and improved domestic demand. Although there is a shift in production and consumption of PE to regions such as China where demand is considerably higher or to regions where feedstocks are cheaper (Middle East) and demand is strong, North American shale gas developments should positively affect U.S.-based producers of ethane and ethylene in terms of low/competitively priced feedstocks. This will in turn have a positive effect on North American production of PE.

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CEH estimates.

New PE process and catalyst technologies have been commercialized during the last decade that have provided a better balance of processability and physical properties. Conventional Ziegler-Natta catalysts contain many reactive sites with varying levels of reactivity, resulting in variations in the polymers produced. The newer metallocene catalysts are single-site catalysts that provide identical reactive sites that allow the resin producer to maximize desired physical properties. The distinguishing characteristic of resins produced using metallocenes is a narrow molecular weight distribution. Metallocenes also provide

far better control of comonomer incorporation, molecular weight and crystalline structure. This overall capability to control polymer properties results in improvements in tailoring polyethylenes to meet processor and user needs. In some cases, these improvements will also permit the use of lower levels of comonomer to achieve the same properties as existing products containing higher comonomer levels.

Further developments of this technology have led to the development of low-modulus copolymers designed to compete with elastomers and flexible polymers in the 0.860-0.910 density range. These resins have very high comonomer levels, ranging from 10% to 30%; resins with 10-20% comonomer content are being called polyolefin "plastomers," while resins with 20-30% comonomer content are considered polyolefin elastomers. The new resins are opening new markets for PE, competing with materials such as plasticized PVC, EPDM elastomers and EVA copolymers.

Linear low-density polyethylene (LLDPE) resins

All linear low-density polyethylene (LLDPE) resins are copolymers of ethylene with higher alpha-olefin comonomers such as butene-1, hexene-1 and octene-1. Polyethylene resins are generally characterized by density and melt index. In general, LLDPE resins have densities ranging from 0.916 to 0.940.

Generally, LLDPE resins are produced by both liquid-phase and gas-phase processes. Almost all plants that can make LLDPE as a primary product are actually linear polyethylene plants that are also capable of producing HDPE; however, these "swing plants" are usually operated on a dedicated basis to make predominantly one or the other product for extended periods. Frequent switches between LLDPE and HDPE are costly and not practiced routinely. The most commonly used commercial process for making LLDPE is Dow's (formerly Union Carbide's) gas-phase Unipol® process. In this process, butene-1 or hexene-1 is copolymerized with ethylene in a fluidized-bed reactor. Another widely used process is INEOS' gas-phase fluidized-bed process, which can use C_4 or C_6 (hexene-1 or 4-methylpentene-1) feed; no 4-methylpentene-1 is currently used in the United States. In 2009, INEOS Licensing announced that their Innovene process can use C_8 comonomer instead of the traditional C_4 or C_6 . Significant quantities of LLDPE are also produced by Dow's solution-phase process, which copolymerizes octene-1 with ethylene. Several other producers can also use octene-1 to make LLDPE.

Desired properties, economic considerations and process determine the choice of comonomer. Butene-1 is less expensive than hexene-1 and octene-1. However, LLDPE resins based on hexene-1 and octene-1 are increasingly attractive, despite higher costs, because of the excellent properties imparted to the copolymer product, including greater tear strength and stress crack resistance.

Comonomers are used either alone (copolymer) or in combinations (terpolymer). Although levels of comonomers incorporated in LLDPE can be as low as 3%, an average level is now between 10% and 14%. Until 1984, most LLDPE resins were based on butene-1 comonomer, which provided sufficiently high performance properties to compete with LDPE. During the mid-1980s, Union Carbide began to increase its production of resins based on hexene-1. Currently, butene-1 is still the largest-volume comonomer consumed for LLDPE resins. However, hexene-1 and octene-1 are expected to continue to take market share from butene-1 during the next few years.

North American production of LLDPE resins amounted to approximately 6.0 million metric tons in 2009. Approximately 65% of this amount was in the United States and the balance in Canada. Consumption in film is by far the largest market. Other significant end uses for LLDPE in North America include injection molding, rotomolding, sheet, and wire and cable. The rapid growth of LLDPE following its introduction in the early 1980s was due primarily to its success in capturing a major portion of existing polyethylene

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film markets. LLDPE quickly penetrated established LDPE film markets where its lower prices and higher strength properties could offer improved performance and cost savings through product downgauging. LLDPE has been most successful in consumer, institutional and industrial garbage bags and liners.

Octene-based LLDPE and further improvements in hexene-1-based resins have further expanded the market for LLDPE resins and fueled a rapid growth rate that has exceeded GDP. Improvements in manufacturing and fabrication technologies continue to open up new markets and applications. North American consumption of LAOs for the production of LLDPE is expected to increase at an average annual rate of 1.0% during 2009-2014. The largest-volume growth will be in film applications, followed by injection molding and sheet.

For detailed information on LLDPE resins, see the CEH *Linear Low-Density Polyethylene (LLDPE) Resins* marketing research report.

High-density polyethylene (HDPE) resins

Approximately 13% of comonomer demand is for the production of high-density polyethylene (HDPE) resins. Linear polyethylene resins with densities of over 0.940 are classified as HDPE resins. They are produced in a variety of low-pressure processes based on different catalyst systems. The processes can be classified into three basic groups according to their reaction conditions—gas phase, slurry phase and solution phase. The most widely used process for producing HDPE in the United States is the Phillips process (slurry phase), followed by Dow's Unipol® process (gas phase). The Phillips process uses hexene-1 comonomer, while the Unipol® process can use either hexene-1 or butene-1 comonomer. Octene-1 is used only in facilities that use the solution process, the largest of which is Dow's. See the CEH *High-Density Polyethylene Resins* marketing research report for information on other processes to make HDPE.

High-density polyethylene homopolymers (i.e., those made solely from ethylene) have high stiffness properties; however, they are susceptible to environmental stress cracking in the presence of many liquids, such as household detergents. The incorporation of C_4 - C_8 alpha-olefins at a rate of 0.5-3% by weight greatly increases environmental stress crack resistance (ESCR). Comonomers also increase the impact strength and provide some increase in chemical resistance. The addition of comonomer also decreases the density of the polymer, resulting in a softer, more flexible plastic. The resulting reduction in stiffness can be compensated for in the product design by molding a container with slightly thicker walls. HDPE copolymers have densities of 0.94-0.96.

In 2009, the average comonomer content for all HDPE was 1.5%. Hexene-1 accounted for the majority of comonomer demand, followed by butene-1; octene-1 accounted for a negligible amount of comonomer demand. The relative comonomer share is not expected to change significantly during the next few years.

In 2009, 7.7 million metric tons of HDPE resins were produced in North America. HDPE resins are converted to end products by blow molding or injection molding and by variations of the extrusion process. Blow-molded applications represent the largest end use for HDPE resins in North America, accounting for 38% of domestic consumption in 2009. The largest application in the blow-molding area is for milk bottles; this application and the smaller-volume market for water bottles do not require copolymer resin. This application has surpassed that of the bottles for household industrial chemicals (e.g., bleaches, dishwashing and laundry detergents, pesticides, and swimming pool chemicals), now the second-largest use in the blow-molding category. The resins used in this market are generally copolymers

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with good ESCR. The use of virgin HDPE resin in household industrial chemicals containers has declined because of the increasing use of recycled resin and source reduction, including lightweighting of bottles and the development of reusable rigid containers that can be refilled using flexible pouches. These trends will continue for the next few years, resulting in a further decline in the use of virgin HDPE in this market. Also included in the blow-molded category are other consumer-oriented bottles, pails and drums, motor oil bottles, and automotive fuel tanks.

Injection-molding applications account for approximately 19% of HDPE demand. A wide variety of items are made by injection molding, including shipping pails; crates, trays and cases; food containers; and housewares.

Extrusion processes convert HDPE into film and sheet, pipe and conduit, wire and cable insulation, and profiles. Film and sheet applications are expected to grow strongly during the next few years. Film applications have been growing well because of the continued displacement of paper products by plastics, as well as substitution of high-molecular-weight HDPE for LLDPE in high-strength film applications. Uses include production of retail bags, including T-shirt bags, trash can liners, and food and nonfood packaging. The fastest-growing use for HDPE is for sheet applications, especially for geomembranes used as liners for landfills, ponds, reservoirs and canals.

North American consumption of LAOs for production of HDPE resins is expected to remain relatively flat during 2009-2014.

For additional information on HDPE and its markets, see the CEH *High-Density Polyethylene Resins* marketing research report.

Elastomers and plastomers

Plastomers and elastomers have densities in the 0.860-0.915 range and are produced by the same processes as LLDPE, but the level of comonomer use is in the 20-30% range. Growth in the consumption of plastomers and elastomers has been dramatic in recent years, and growth is expected to continue at a rate of approximately 10% per year during 2009-2014. Approximately 129 thousand metric tons of comonomer were consumed in this segment in 2009 and growth in its usage should continue at the same rate as for the elastomers and plastomers. Octene-1 is the largest-volume comonomer used in this market segment. Dow Chemical and ExxonMobil are the major North American producers of these products. In August 2010, Dow announced plans to increase its specialty elastomers capacity in Texas and Spain.

Oxo Alcohols

Oxo technology utilizing a linear alpha-olefin feed is currently used by Shell in both plasticizer-range and detergent-range alcohol production. As of late 2006, Sterling Chemicals (which also manufactured plasticizer-range alcohols via oxo technology utilizing a LAO feed) ceased production of C_7 , C_9 and C_{11} linear alcohols. For the purposes of this report, plasticizer-range alcohols are defined as monohydric aliphatic primary alcohols containing four to eleven carbon atoms, and detergent-range alcohols are similar products that contain twelve or more carbon atoms. Oxo processes, formally termed *hydroformylation*, involve the addition of synthesis gas (carbon monoxide and hydrogen) to either end of an olefinic bond. Most processes follow this step with hydrogenation of the resultant aldehyde to the respective alcohol in a second reactor. Hydroformylation of even-carbon LAOs yields odd-numbered linear alcohols.

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Shell has developed a major modification to the oxo process. The alpha-olefins not separated for sale are converted to C_8 - C_{18+} internal olefins via a series of isomerization and disproportionation steps in its SHOP plant. (See the **MANUFACTURING PROCESSES** section for more information on this process.) Some internal olefins are subsequently converted via a modified oxo process to detergent-range primary alcohols. Shell's process performs both hydroformylation and dehydrogenation of the intermediate aldehyde in the same reactor.

Detergent alcohols

Approximately 200 thousand metric tons of alpha-olefins were consumed by Shell to produce similar amounts of detergent alcohols in 2009. The remaining detergent alcohol production is based on either ethylene, using Ziegler chemistry, or conversion of natural fats and oils, especially coconut oil, tallow and palm kernel oil.

Shell Chemical is the only company that produces detergent alcohols (C_{12} and higher) from LAOs in the United States, with a nameplate capacity of 286 thousand metric tons at Geismar, Louisiana. In 2002, Shell completed an expansion of 102 thousand metric tons of capacity, for a specific range of monomethyl branched alcohol. In 2003, an equivalent capacity was placed on standby and as of mid-2010 this had not changed.

Most of Shell's plant is devoted to the production of detergent-range alcohols; however, it also produces some C_9 - C_{11} linear alcohols for plasticizer and detergent applications, as well as other oxo alcohols derived from branched olefins. Shell captively consumes most of its alcohol production to make alcohol ethoxylates. The balance is accounted for by merchant sales (including exports) of detergent alcohols, sold under the trademark Neodol[®].

Demand for detergent alcohols from all sources is expected to grow at an average annual rate of approximately 2.5% during 2009-2014. Since 2003, the detergent alcohols industry had experienced some tightness. The idling of Shell's 102 thousand metric tons of capacity greatly reduced the industry's operating capacity. This combined with increasing demand for alcohols resulted in tight supplies of mid-cut (C_{12} - C_{14}) alcohols during intervals of 2006-2007. However this tightness eased with the increase of net imports of detergent alcohols in 2008, as new natural alcohol plants come on line in Southeast Asia and Brazil. It is likely that imports of alcohols from Asia will continue to increase. Consumption of LAOs for detergent alcohols will grow at an average annual rate of 2.0% per year over 2009-2014.

Approximately 84% of 2009 detergent alcohol consumption was accounted for by three surfactants: alcohol ethoxylates (AE), alcohol sulfates (AS) and alcohol ether sulfates (AES). All three are used in household laundry detergents, which is the largest end-use category for these products. Since liquid laundry detergents contain on average higher surfactant levels than laundry powders, the consumer shift to liquids has led to an increasing use of the alcohol-based surfactants in this application. Large amounts of AES, which is derived from sulfating AE, are also used in hand dishwashing liquids. All three surfactants are also used in significant volumes in personal care products (especially shampoos and soaps). In addition, all three are used in many industrial applications (e.g., emulsion polymerization, agricultural emulsifiers, industrial and institutional cleaners, and textile processing).

AS and AES compete with another large-volume anionic surfactant—linear alkylbenzene sulfonate (LAS)—and with nonionic surfactants. The relative costs of the competing surfactants depend largely on the prevailing prices for their raw materials (e.g., coconut oil, ethylene, benzene and n-paraffins). Since

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these prices vary considerably over time, so do the prices for the surfactants derived from them, affecting their relative cost/performance advantages. As a result, reformulations are frequent and this market is internally dynamic.

The slight decrease in consumption of LAOs for the production of detergent alcohols during 2003-2006 can be attributed to several things. The use of AES grew more slowly during recent years, as detergent manufacturers partly offset the growth in the use of AES-containing laundry liquids by reducing its concentration in these detergents—a development that was mainly a consequence of higher alcohol and other raw material costs. The consumption of AS (and alcohol consumed to make it) has fallen since 2003 because of the declining use of laundry powders containing AS, as consumers switch to liquid laundry detergents that do not use AS. A significant reduction in LAO-derived detergent alcohol exports during 2007-2009 contributed to a decrease in consumption of LAOs for the production of detergent alcohols.

Other markets for detergent alcohols include polymethacrylate esters, fatty nitrogen derivatives, thiodipropionate ester and alkyl glyceryl ether sulfonates (AGES). Detergent alcohols are also consumed directly in cosmetics and toiletries as emollients, in formulated lubricants used in the rolling of metallic foil or sheet stock, and in pharmaceutical preparations. For more details on markets for detergent alcohols, see the CEH *Detergent Alcohols* marketing research report.

Plasticizer alcohols

Approximately 25 thousand metric tons of alpha-olefins were consumed to produce similar amounts of C_7 - C_{11} oxo alcohols in 2009. This includes small amounts that were consumed for the production of surfactants.

As of August 2010, Shell is the only company that produces linear plasticizer alcohols (C_7-C_{11}) from LAOs in the United States. Shell produces C_7-C_{11} linear alcohols for plasticizer and detergent applications as an adjunct to its Neodol® detergent-range alcohols. As previously mentioned, Shell converts C_6-C_{10} internal olefins, which are obtained in its SHOP plant via isomerization and disproportionation of linear alpha-olefins, to a mixture of C_7-C_{11} alcohols via a modified oxo reaction. These alcohols, sold as Linevol® 11, Linevol® 911 and Linevol® 79, are approximately 80% linear and 20% branched alcohols. Shell's flexible SHOP process can produce in any desired range; however, most of Shell's production is detergent alcohols and only a relatively small part of Shell's U.S. output is in the plasticizer range.

ExxonMobil purchases linear alcohols for the production of linear plasticizers. Sterling Chemicals ceased production of linear alcohols when it permanently shut down its 102 thousand metric ton-per-year oxo alcohols production unit in late July 2006—citing high raw material costs, raw material supply shortages and competing branched plasticizers as the reasons. BASF had marketing rights to Sterling's alcohol output, which was converted primarily to phthalate esters for BASF's Palatinol[®] line. Sterling will continue to produce phthalate esters for BASF with oxo alcohols supplied by BASF through 2013.

Linear alcohols were introduced into plasticizer markets because of the lower volatility, low-temperature stability and low water extraction that they impart to esters. C_7 - C_{11} alcohols are reacted most commonly with phthalic anhydride or, to a much lesser extent, with other acids, such as adipic acid, to produce the corresponding esters, which are used as plasticizers for flexible polyvinyl chloride (PVC) resins, especially those used in calendering (e.g., coated fabrics and sheet goods) and when outdoor weathering resistance is required. They compete to a limited extent with the largest-volume plasticizers—di(2-ethylhexyl) phthalate (DEHP) and diisononyl phthalate (DINP)—which are branched plasticizers. The

linear C_7 - C_{11} plasticizers (e.g., ExxonMobil's Jayflex[®] 711) have distinct advantages in vinyl plasticizers for automotive upholstery and compartment interiors. While maintaining low-temperature flexibility, the linear phthalates reduce window fogging. Other uses for 711 phthalates include calendered sheet and film for swimming pool liners, roofing membranes, automotive body side molding, tarpaulins, and wire and cable jacketing.

Although their electrical resistivity is poorer than that of branched esters, high-molecular-weight linear phthalates are used in power transmission cable where temperature properties are important. The principal phthalate consumed for this use is diundecyl phthalate (DUP) made from C_{11} alcohol.

Consumption of linear phthalates is also impacted by raw material price relationships (i.e., the ethylene/propylene price ratio). Fluctuations in consumption will continue whenever abrupt changes in typical feedstock price ratios occur.

The future production and consumption of linear plasticizers in the U.S. market will decrease over the next five years. Higher feedstock costs and the termination of feedstock sourcing of LAOs have dampened demand for linear plasticizers. Despite some advantages over branched plasticizers, some industry sources feel that linear plasticizers will be replaced by branched product over the long term. The consumption of LAOs for the production of C_7 - C_{11} linear alcohols is expected to decline at an average annual rate of 2.5% per year during 2009-2014.

For more information on plasticizers, including other smaller-volume plasticizers derived from C_7-C_{11} alcohols and plasticizer alcohols, see the CEH *Plasticizers* and *Plasticizer Alcohols* (C_4 - C_{13}) marketing research reports.

Polyalphaolefins

Polyalphaolefins (PAOs), also known as linear alpha-olefin oligomers, are base stock materials for synthetic lubricants for automotive, industrial and aviation applications. These synthetic base fluids have a uniform isoparaffinic structure similar to paraffinic mineral base stocks. Smaller but significant volumes of PAOs are used in other applications, including dielectric fluids and drilling muds, especially those used in offshore drilling applications. PAOs are produced by oligomerizing short-chain LAOs (largely C_{10} , but also C_8 , C_{12} , and C_{14} ; some companies may mix C_8 , C_{10} or C_{12} , C_{14}) using a promoted boron trifluoride catalyst in a one-step synthesis. Depending on reaction conditions, the typical oligomer mixture is 50-65% trimer and 10-15% dimer, with tetramer and higher oligomers making up the balance. The oligomers are converted to stable, high-purity isoparaffins through hydrogenation.

The following table presents North American producers of polyalphaolefins:

Company and Plant Location	Annual Capacity as of Mid-2010 (thousands of metric tons)	Trademark/ Remarks
Albemarle Corporation		
Pasadena, TX	15.0	ExxonMobil's SpectraSyn [®] high viscosity PAOs are being manufactured under contract.
Chemtura Corporation		
Elmira, Ontario	15.8	Synton [®] . Specialty high viscosity PAOs. Capacity was increased by nearly 2 thousand metric tons in 2006.
Chevron Phillips Chemical Company LP		
Baytown, TX	48.0	Synfluid [®] . Captive LAOs (pure C_{10} or C_{12} for low viscosity PAOs).
ExxonMobil Chemical Company		
Beaumont, TX	84.5	SpectraSyn [®] . Includes capacity for both low and high viscosity PAOs.
INEOS Americas, LLC		
Deer Park, TX		Durasyn [®] . Captive LAOs.
Total	241.3	Y
SOURCE: CEH estimates.		7

North American Producers of Polyalphaolefins

ExxonMobil also produces higher-viscosity PAOs, which are used as base fluids in compounding higherviscosity gear oils, greases, industrial oils, turbine oils and motor oils in blends with other synthetics and mineral oils.

In 2005 ExxonMobil increased capacity for high viscosity PAOs and upgraded low viscosity PAO production. In 2007, ExxonMobil underwent a debottlenecking to boost PAO capacity to help meet increasing market demand for the advanced high viscosity PAO. In December 2009, ExxonMobil announced a custom manufacturing agreement with Albemarle for the production of 15 thousand metric tons of high viscosity SpectraSyn®PAOs in Pasadena, Texas.

A high percentage of ExxonMobil's PAO production is consumed captively by its parent company for the manufacture of automotive and industrial oils and greases. The PAO made by Chevron Phillips, INEOS and Chemtura is sold to the merchant market.

North American production of PAOs in 2009 amounted to 195 thousand metric tons and required 201 thousand metric tons of LAOs, 65-70% of which was decene-1. Except for limited production by Chemtura in Canada, nearly all of the North American production of PAOs was accounted for by the three large U.S. producers. The following table presents U.S. consumption of LAOs for the production of PAOs:

	(thousands of metric tons)	
	Polyalphaolefin Production	Linear alpha-Olefin Consumption ^a
1976	10	11
1979	25	28
1980	25	29
1985	34	38
1990	48	49
1995	125	129
1999	142	147
2003	172	178
2006	213	219
2009	195	201
2010 ^b	210	216

U.S. Consumption of Linear alpha-Olefins for Polyalphaolefins (thousands of metric tons)

a. The amount of decene-1 consumed for PAO varies depending on the process and the viscosity of the oligomer products. On average, 1.09 pounds of decene-1 were consumed per pound of PAO produced during 1976-1988; 1.03 pounds of decene-1 were consumed per pound of PAO beginning in 1989.

b. Projected estimate assuming strong second half of 2010.

SOURCE: CEH estimates.

North American polyalphaolefin production increased steadily from the mid-1980s through 2007 before it decreased in 2008-2009. North American production of PAOs grew at an average annual rate of 5.5% during 2002-2006. During the period of 2004 through most of 2005, many PAO producers reported double-digit growth and operating rates over 90%. Strong demand for PAOs continued into 2006 and 2007 and operating rates remained high. However, by 2008 production (and hence LAO consumption) decreased with Hurricane Ike striking the U.S. Gulf Coast in the third quarter of 2008, followed by weak demand due to the economic recession. Weak economic conditions and a tight decene-1 supply kept production of PAOs from increasing throughout most of 2009. By the fourth quarter of 2009 things had started to improve and production and demand in 2010 (through August) have rebounded strongly compared to 2009 figures. Also, supply of decene-1 should improve with additional oversea LAO capacity.

Consumption of LAOs for the production of PAOs should average at least 5% per year during 2009-2014, assuming the impact of Group III lubricants or gas-to-liquid base (GTL) stocks on the PAO market will be minimal during this period. This also assumes an ample decene-1 supply during 2010-2014. The high performance of PAOs versus Group III base oils has allowed them to retain market share in those sectors that require this performance.

PAOs are usually considered the most versatile of all synthetic lubricants for automotive crankcase applications, which is their largest market. Compared with conventional mineral oil base stocks, they

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have good lubricating properties over a wide temperature range; a high inherent viscosity index; low volatility; high thermal and oxidative stability; excellent response to antioxidants; and higher flash, fire and auto ignition points. The result is reduced engine friction, longer service life and improved efficiency under extreme cold weather conditions and high engine temperatures. PAOs are often used with lower levels of ester-type synthetic lubricants to correct seal-swelling problems that occur when they are used alone in some applications.

The market for PAOs in automotive crankcase lubricants appeared to have great growth potential, since only a very small percentage of U.S. passenger cars currently use synthetic lubricants. However, several refineries have made available increasing volumes of severely hydrocracked mineral oil-based stocks of high-viscosity index that are referred to as Group III lubricants. These new lubricants provide many, if not all, of the superior properties of the PAO-based lubricants and at a far lower cost than PAOs. Furthermore, Castrol won a court ruling in the United States that permitted the company to refer to its Group III-based lubricants as synthetic lubricants. However, PAO consumption in automotive crankcase lubricants may be boosted by the industry's trend toward longer service (drain) intervals—a trend that demands high performance base stocks such as PAOs. The volume of high-performance grade synthetic lubricants such as ExxonMobil's Mobil 1 and other PAO-based products continues to grow.

In addition to their use in automotive crankcase oils, synthetic lubricants are also consumed in automotive gear oils. These products were developed for heavy-duty trucks and other off-road vehicles to provide fuel economy and high-temperature/high-torque performance. The heavy-duty truck market suffered during the economic recession, and this sector primarily uses PAO-based synthetic lubricants for axle, transmission and driveline. Polyalphaolefin consumption in this sector is expected to exhibit above-average growth during 2009-2014, as markets rebound from the weak global economy.

Industrial applications have grown over the years. Because they provide longer drain intervals and may extend the life of equipment, synthetic lubricants can offer cost savings in some industrial applications. Use of synthetic lubricants based on PAO improves the efficiency of industrial gear systems compared with petroleum-based lubricants. A major use of PAO-based gear oils that has been showing rapid growth in recent years is lubrication of the gears of stationary diesel engines used in industry. Consumption of PAOs in industrial gear oils (primarily high-viscosity PAOs) is estimated to grow at an average annual rate of over 3% during 2009-2014.

Other large-volume lubricant applications for PAOs include compressor oils and hydraulic fluids (including automotive, aviation and industrial). Smaller-volume applications for PAO include aviation piston engine oils, greases, bearing oils, gas turbine lubricants, heat-transfer fluids, and transformer dielectric and insulation fluids.

PAOs compete for market share with dibasic esters and polyol esters in some industrial applications, but have the advantage of being less expensive than these other synthetics. For more information on polyalphaolefins and other synthetic lubricants, see the Specialty Chemicals Update Program *Synthetic Lubricants* report.

The consumption of PAOs in nonlubricant applications includes use in heat transfer fluids, transformer dielectric and insulation fluids, and drilling muds. Consumption in nonlubricant applications was 14 thousand metric tons in 2009. PAOs were used in drilling muds to help remove the cuttings, although they also lubricated the drill bit. Consumption of PAO for this purpose amounted to at least 18 thousand metric tons in 1995. However, it is believed they are no longer used in this application in the Gulf of Mexico because they do not meet regulations. PAOs were largely replaced in this application by the less costly C_{16} - C_{18} olefins, either LAOs or isomerized LAOs.

Some industry sources feel that overall U.S. demand for PAOs (both low and high-viscosity) may grow more than 5% per year on average during 2009-2014. Polyalphaolefin demand will be driven by tightening emissions standards in the transportation industry (passenger vehicles and heavy-duty diesel).

Oil Field Drilling Fluids

LAO-based products in the C_{16} - C_{18} range are used in oil field drilling muds as lubricants and to help remove cuttings. This application is identical to that previously described under **Polyalphaolefins**. The C_{16} - C_{18} -range LAOs have been found to provide equivalent performance at a lower price than the C_{10} based PAOs. Consequently, they have been replacing the PAOs as well as the slow-to-biodegrade diesel oils in offshore drilling applications. The LAOs are either used as such or isomerized to internal olefins to lower their melting point. Approximately 90 thousand metric tons of C_{14} - C_{18} -range products and some lower or higher carbon-range products are used in this application. Although C_{16} - C_{18} range is the most widely used for this application, C_{14} usage has increased over the years in offshore drilling applications and it is also being used in onshore drilling, replacing some mineral products. The consumption of oil field drilling chemicals has increased over the years as a result of the shift from oil well drilling to natural gas well drilling. New gas wells are deeper and require more drilling fluids. However, consumption decreased in 2009 from 2008 levels as oil and natural gas exploration and production decreased—a consequence of lower demand for energy during the weak economic times and lower oil and natural gas prices.

It is difficult to predict a growth rate for LAO consumption in oil field drilling muds, especially given the current (August 2010) uncertain regulatory environment in the Gulf of Mexico. Consumption is likely to increase at a reduced rate of approximately 2.0-2.5% per year during 2009-2014. As the economy recovers from the global economic crisis, demand for oil and natural gas will increase, driving drilling activity and hence consumption of oil field drilling fluids. Higher oil and natural gas prices will also contribute to growth in drilling activity. Onshore drilling will help drive this growth. However, this growth rate could be lower if the U.S. EPA places additional environmental constraints on drilling companies in the Gulf of Mexico (where most of the consumption occurs). It does take into account the moratorium placed on drilling activity in the Gulf of Mexico in the second quarter of 2010 (set to expire on November 30, 2010) after the 2010 oil spill in the Gulf of Mexico. Should drilling companies ever be required to recover the cuttings from drilling operations, they might revert to the use of cheaper diesel fuel. Current practice permits dumping the cuttings, provided the material used has acceptable biodegradability and toxicity. Drilling companies have been forced to recover the cuttings off the coast of Norway in the North Sea, however, these regulations are considered unlikely to be imposed in the Gulf of Mexico. Shell supplies C_{15} - C_{18} , while INEOS and CPChem supply C_{16} - C_{18} based fluids for this application. All these fluids are designed for and fully compliant with EPA regulations for use in the Gulf of Mexico.

Other Lubricant Additives

LAOs are incorporated in several types of compounds that have a variety of functions in lube oils. The production of these additives consumed 38 thousand metric tons of alpha-olefins in 2009. This amount can vary from year to year depending on the relative pricing of competing raw materials. Growth in North American consumption of lube oil additives in general is expected to average only 0.5% per year during 2009-2014. Consumption of LAOs for use in this application is estimated to grow at approximately the same rate.

Some of these lube oil additives are described briefly in the following paragraphs.
- Synthetic Heavy Alkylates. alpha-Olefins can be used to alkylate benzene, which is then sulfonated and neutralized to a calcium, magnesium or barium salt. These lube oil sulfonates are used as detergents in lube oils or in metalworking fluids. The main alpha-olefins consumed to produce lube oil additives are C₂₀-C₂₄ olefins, although some products use two alkyl groups containing shorter-chain olefins. The largest North American producers of lube oil additives based on alpha-olefins are The Lubrizol Corporation and Afton Chemical, although these companies may have some of their production toll produced by others (e.g., Pilot Chemical Co.). The two other large lube oil additive manufacturers, Chevron Oronite and Infineum (a Shell and ExxonMobil joint venture), make these products in Europe. Some lube oil sulfonates (probably less than 20%) used in automotive applications are based on natural petroleum sulfonates that do not consume LAO, but their use has been declining for years.
- **Phenates.** Linear alpha-olefins can also be used to alkylate phenols, which are then neutralized to calcium, magnesium or barium salts. Like the synthetic heavy alcohols, the alkyl phenol compounds function as detergent additives in lube oil applications. However, they are generally more expensive than SHA-type compounds and most are based on propylene tetramers and not LAOs. Chevron is believed to make some of these products based on LAO, but none is produced in North America.
- Sulfurized Linear alpha-Olefins. LAOs in the C_{16} - C_{24} range are blended with an oleate ester or a triglyceride and this mixture is then sulfurized to a 5-20% sulfur content. Theoretically, any type of olefin, including internal, can be used for this sulfurized product, but in practice LAOs are the main olefins used. Sulfurized LAOs are used as an extreme-pressure agent in end uses such as metalworking, auto and industrial gear oils and greases, and to some extent hydraulic fluids.
- Alkylnaphthalenes. LAOs in the C₁₆-C₂₄ range are used to alkylate naphthalene, and the resulting product is used as a pour-point depressant. In this application, it competes with polymethacrylate-type pour-point depressants. These products are also used in some crankcase synthetic motor oils (e.g., Mobil 1[®]) as both lubricants and VI improvers. This latter use alone consumed approximately 4 thousand metric tons of LAOs in 2009.
- **Salicylates.** These products compete with sulfonates and phenates in lubricant additives and detergents. They are produced by Infineum (with headquarters based in Britain).

Alkenylsuccinic anhydrides (ASA) based on C_{12} - C_{18} alpha-olefins can also be consumed for lube oil additives as corrosion inhibitors and frictional modifiers for crankcase oils and dispersants. Most, however, are based on propylene tetramers, rather than LAOs. Refer to the section on **Alkenylsuccinic Anhydrides** for information on alpha-olefin demand for ASAs for all other uses for these products. In addition, some dialkylbenzenes, which can be made from alpha-olefins, are also used in lube oil additives.

Alkyldimethylamines and Dialkylmethylamines

Alkyldimethylamines are produced from several different feedstocks in the United States—linear alphaolefins, fatty alcohols or fatty acids. In the United States, two companies produced LAO-based alkyldimethylamines through 2009—Albemarle Corporation and The Procter & Gamble Company (P&G). In 2010, P&G switched feedstock for its tertiary amines from LAOs to natural fatty alcohols to supply its Kansas City plant. Albemarle also manufactures smaller volumes of dialkylmethylamines from LAOs. Lonza makes both types, but from detergent alcohols, while Akzo Nobel makes these amines from fatty nitriles (smaller volumes). It is believed that in 2009 all producers of alkyldimethylamines derived from LAOs or alcohols operated at or near capacity. In 2009, total alkyldimethylamine and dialkylmethylamine production from all raw materials was 93-95 thousand metric tons, of which 83-85 thousand metric tons were derived from alpha-olefins. Approximately 65-67 thousand metric tons of $C_{8-}C_{18}$ alpha-olefins were consumed for their production in 2009. Because of P&G's feedstock switch in 2010, consumption of LAOs for the production of alkyldimethylamines will decrease at an average annual rate of approximately 25% during 2009-2014.

LAO-derived alkyldimethylamines are produced by reacting C_{12} - C_{18} linear alpha-olefins with hydrogen bromide in the presence of peroxide catalysts to form an n-alkyl bromide. Reaction of these bromides with dimethylamine yields the corresponding alkyldimethylamines. Reaction of methylamine with two moles of the alkyl bromide yields a dialkyldimethylamine, but the latter types are based on C_8 - C_{10} LAOs.

P&G manufactures C_{12} - C_{16} alkyldimethylamines at its Kansas City, Kansas plant, with an annual capacity estimated at over 60 thousand metric tons. It consumes most of them captively for the production of fatty amine oxides, but some are sold for use in making alkylbenzyltrimethylammonium salts (quaternaries or quats), used as biocides. P&G would prefer to use worldwide formulations for many of its products and must obtain the necessary raw materials either from local suppliers or by exporting product from its North American plants. For alkyldimethylamines, P&G appears to have found exporting product from its Kansas City plant to be more economical.

Albemarle manufactures ADMA[®] alkyldimethylamines and DAMA[®] dialkylmethylamines in Magnolia, Arkansas. Annual capacity for both the ADMA[®] and DAMA[®] types is 18-20 thousand metric tons. Albemarle's products are marketed as individual cuts of even-numbered carbon chains from C₈ to C₁₈ or as blends. Both ADMA[®] and DAMA[®] amines are sold largely for conversion to quats for biocides. Other applications include use in making fatty amine oxides, as well as quats and amine salts used in fabric softeners, hair conditioners, dye levelers, polyurethane foam catalysts and epoxide resin curing agents.

The largest application for alkyldimethylamines is for the production of fatty amine oxides (FAOs) produced from the reaction of alkyldimethylamines with hydrogen peroxide. The most common amine is C_{12} - C_{14} alkyldimethylamine. Most of the production of fatty amine oxides is accounted for by P&G. Other major producers of fatty amine oxides are Lonza and Stepan Company.

Although mildly cationic in acidic solutions, amine oxides exhibit nonionic properties in neutral and alkaline pH ranges. They are good foam stabilizers, thickeners and emollients, as well as emulsifying and conditioning agents. They are very mild to the skin, an important attribute for dishwashing liquids. When used in combination with alcohol ether sulfates in dishwashing liquids, they provide superior foam stabilization properties in the presence of grease compared with the more commonly used alkanolamides. However, fatty amine oxides are considerably more expensive than alkanolamides and their use in dishwashing liquids is almost exclusively confined to the products of P&G, which has the advantage of a captive source of this surfactant.

Fatty amine oxides are used in some household cleaners and are also used as foam stabilizers and conditioning (i.e., antistatic) agents in shampoos and hair conditioners, as well as in some small-volume industrial applications.

Only slow growth is expected in the U.S. and Canadian markets for fatty amine oxides because of a strong consumer preference for automatic dishwashers, which use detergents that do not consume this surfactant. As already discussed, however, increasing volume may be used in other world areas by P&G, based on exported amines. See the CEH *Surfactants, Household Detergents and Their Raw Materials* marketing research report for additional information on fatty amine oxides.

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The other major application for alkyldimethylamines is for the production of quaternary ammonium compounds. These products are produced by reacting the alkyldimethylamine with benzyl chloride (by far, the most common) or p-ethylbenzyl chloride, to produce alkylbenzyldimethylammonium (ABDM) salts. The alkyl group typically contains twelve to eighteen carbon atoms, depending on the desired properties, but the middle of this range (fourteen and sixteen carbon atoms) normally has the broadest spectrum of biocidal activity.

The other major class of quaternaries that can be derived from LAOs are dialkyldimethylammonium compounds made by quaternizing dialkylmethylamines with methyl chloride. Lonza is the largest producer of these particular compounds. Albemarle also produces DAMA[®] dialkylmethylamines from C_8 - C_{10} alpha-olefins and sells them to the merchant market. Both types of quaternaries are effective bactericides that are used in a variety of disinfectants and preservatives.

 C_{18} alkyldimethylamines are converted to alkyldimethylbenzylammonium chlorides for use as hair conditioning agents in shampoos, conditioners and cream rinses, where they provide softness as well as antistatic properties. For additional information on benzyl quats, see the CEH *Benzyl Chloride* marketing research report.

Lonza is the major producer of biocidal quaternaries derived from alkyldimethylamines, followed by Stepan Company. Lonza's plant for making the amines from alcohols has an estimated capacity of only 7-10 thousand metric tons per year and is believed to have operated at or near capacity in recent years. Stepan purchases its amine requirements. Stepan and Lonza are the major producers of hair-conditioning quats. For further information, see the Specialty Chemicals Update Program *Biocides* and *Cosmetic Chemicals* reports.

Alkyl betaines, which are amphoteric surfactants, are produced by reacting tertiary amines, such as alkyldimethylamine, with sodium chloroacetate. They can be used as surfactants in light-duty dishwashing liquids. P&G used an LAO-derived alkyl betaine in its Ivory[®] dishwashing liquid for several years, until the product was withdrawn in early 1992.

Alkenylsuccinic Anhydrides

Alkenylsuccinic anhydrides (ASAs) are produced by isomerizing an alpha-olefin to an internal olefin, which is then reacted with maleic anhydride to form the corresponding branched alkenylsuccinic anhydride. The most commonly used alpha-olefins are C_{16} - C_{20} , although other chain lengths are also used. Albemarle is the largest U.S. producer of ASAs. Other producers are Dixie Chemical Company, Milliken Chemical and Vertellus Specialties.

ASAs are consumed in a variety of applications, including paper sizes, lube oil additives, detergents, leather treatment and foods. Use in the paper industry for paper sizing is the largest application for ASAs. Sizes are used in and on paper and paperboard, primarily to make the products more resistant to water and other liquids and to control the spread of inks. ASAs are used as internal sizing agents that are added to the paper slurry prior to sheet formation. They are used in alkaline papermaking processes primarily for high-quality fine papers and gypsum wallboard liners. The other principal commercial alkaline internal sizing agents are alkyl ketene dimers (AKDs). AKDs, made from C_{12} - C_{20} natural fatty acids, are used in the production of most paperboard made by alkaline processing; ASA sizes do not harden as well as AKDs, which is important in paperboard grades. Both ASA and AKD have a reactive end that binds to the cellulosic fibers and a hydrophobic tail that contributes to water repellency.

The use of ASAs and AKDs has grown because of the conversion of paper mills to alkaline processes during the late 1980s and early 1990s. Advantages of the alkaline process include reduced wood pulp requirements, increased strength, improved paper formation, improved printing performance, longer-lasting paper and a more cost-effective process overall for many mills.

ASAs and AKDs each have advantages and disadvantages associated with their use in the alkaline process. The main advantage of ASAs is that they cure rapidly on the paper machine, while AKDs require a longer off-machine cure time. A disadvantage associated with ASAs is their high reactivity, which requires that they be made into an emulsion at the mill and used immediately; AKDs are less reactive and can be shipped as an emulsion.

ASAs of various carbon numbers consume small quantities of alpha-olefins for other markets. Octenyl succinic anhydride, made from octene-1, is used in making derivatized starches used as thickening agents in foods, juices and puddings. The potassium or sodium salts of octenyl succinic anhydride are used as surfactants in industrial metal cleaners and institutional floor cleaners. ASA compounds derived from both C_{16} and C_{18} LAOs are used in finishes that soften leather and protect it from water penetration. Epoxy resins cured with C_{16} - C_{20} ASAs have good electrical properties, flexural strength and hardness. Some ASAs based on C_{12} - C_{18} LAOs are sold to the lube oil additives industry. These ASA compounds are processed to make derivatives that are used as corrosion inhibitors and frictional modifiers for crankcase oils and dispersants. Many of the ASA compounds used in lube oil additives, however, are based on propylene tetramers, rather than LAOs.

Approximately 25 thousand metric tons of alpha-olefins or internal olefins derived from LAO were consumed in the production of ASAs in 2009. Paper sizing accounted for 80% of the consumption and this use is expected to grow at an average annual rate of approximately 3% during 2009-2014. Many paper mills have converted from the acidic to the alkaline process. Other applications are not expected to exhibit much growth during this period.

Synthetic Fatty Acids (C7-C9)

OXEA produces heptanoic (C₇) and pelargonic (C₉) acids from hexene-1 and octene-1 olefins, respectively, at a 45 thousand metric ton-per-year plant at Bay City, Texas. OXEA's production of C₇ and C₉ acids was approximately 26 thousand metric tons in 2009, consuming 19 thousand metric tons of LAO. The acids are produced via a rhodium-catalyzed oxo process. Until this plant was commissioned in early 1980, semi-synthetic routes from natural feedstocks were the only sources of C₇ and C₉ fatty acids. Synthetic fatty acids produced by OXEA are more than 95% linear. This degree of purity allows them to be interchangeable in lubricant formulations with fatty acids derived from fats and oils that are available as 100% linear products.

The natural acids are produced as coproducts in the manufacture of other compounds, while the OXEA acids are primary products. Arkema imports natural heptanoic acid from its French operation, which steam-cracks methyl ricinoleate from castor oil to undecylenic acid and C_7 chain-length products, including heptanol and heptaldehyde. Undecylenic acid was previously the driving force for the operation, but increased demand for heptanoic acid has raised the emphasis on the C_7 fraction.

Emery Oleochemicals (formerly Cognis Oleochemicals) coproduces pelargonic acid during the ozonolysis of oleic acid to azelaic acid. The largest use of pelargonic acid is for the production of sodium nonanoyl oxybenzene sulfate (NOBS), which is used by P&G as a bleach activator in some of its laundry detergents. Most of P&G's consumption is accounted for by Tide[®] with bleach, which was introduced in

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the late 1980s. The bleach activator allows sodium perborate to function as a bleaching agent at the washing temperatures commonly used in North America. FutureFuel Chemical Company is the sole U.S. producer of NOBS. This use of pelargonic acid has grown from only 2-3 thousand metric tons in 1988 to approximately 14 thousand metric tons in 2009. However, growth in this application within North America has matured, especially since consumers are increasingly using liquid detergents, such as Liquid Tide[®], which does not contain the bleach activator.

Other outlets for pelargonic acid include its use in the manufacture of synthetic lubricants (discussed later in this section), plasticizers and agrochemicals. For additional information on markets for heptanoic and pelargonic acids, see the CEH *Oxo Chemicals* marketing research report.

The major use of heptanoic acid and the second largest of pelargonic acid is neopolyol esters (NPEs) used for synlube base stock. Approximately 18 thousand metric tons of heptanoic acid and 4 thousand metric tons of pelargonic acid were consumed in NPEs in 2009. NPEs are made by the reaction of one or more organic acids with a polyfunctional alcohol (e.g., neopentyl glycol, trimethylolpropane or pentaerythritol) via a condensation reaction. In addition to heptanoic and pelargonic acids, valeric (C₅), 2-ethylhexanoic (C₈) and C₈ and C₁₀ natural fatty acids are commonly used to produce NPEs; valeric acid can also be derived from alpha-olefins via C₅ aldehydes. The largest market for NPEs is as synthetic lubricant base stocks for both military and commercial aircraft. Other lubricant market segments include automotive crankcases and refrigerators. As with valeric acid, NPEs produced with heptanoic and pelargonic acids are being increasingly used in non-CFC fluorocarbon cooling systems. The C₇ and C₉ acids face competition from C₈ and C₁₀ natural fatty acids derived from coconut oil. For additional information on neopolyol esters, see the Specialty Chemicals Update Program *Synthetic Lubricants* report.

Heptanoic acid was used for the production of tetraethylene glycol diheptanoate (4G7), a plasticizer for PVB resin. U.S. consumption of heptanoic acid in 4G7 declined dramatically during 2002-2005, as a result of DuPont's conversion to triethylene glycol di-2-ethylhexanoate (3GO) as its PVB plasticizer. Both Solutia and DuPont, the U.S. producers of PVB, now consume 3GO for their PVB operations.

Consumption of LAOs in the production of heptanoic and pelargonic acids is forecast to increase at an average annual rate of 2.5-3.0% during 2009-2014.

alpha-Olefin Sulfonates

alpha-Olefin sulfonates (AOS) are produced by the direct sulfonation of C_{14} - C_{16} linear alpha-olefins with dilute sulfur trioxide (SO₃). This is followed by neutralization and hot hydrolysis of the intermediate reaction products with sodium hydroxide. The resulting AOS product, which is produced and sold mainly as a 40% active material, is actually a mixture consisting primarily of alkene-1 sulfonates, hydroxyalkane-1 sulfonates and some disulfonates. The U.S. industry is reportedly using a two-to-one C_{14} : C_{16} blend for most applications. Three U.S. companies currently produce AOS in North America—Stepan Company, Rhodia and Pilot Chemical Company.

The presence of a double bond and a hydroxy group gives good solubility to alpha-olefin sulfonates and they are relatively stable at low pHs compared with the alcohol sulfates (AS) and alcohol ether sulfates (AES), with which they compete in many personal care applications. However, processing costs on a poundage basis are higher for AOS than for most commodity anionics, and various impurities have also made it less attractive for many large household uses, such as for laundry powders. Thus, the use of AOS in North America has been limited largely to some personal care (e.g., liquid hand soaps and shampoos) and industrial applications. The latter include its use in emulsion polymerization processes. AOS has also

been consumed in several oil field applications, including use as a foamer for well clean-out and, in the early 1980s, as a surfactant for steam-driven enhanced oil recovery techniques, but this use nearly disappeared as crude oil prices fell in the late 1980s.

Production of AOS was approximately 19 thousand metric tons in 2009, consuming 14-15 thousand metric tons of LAO. Overall growth for AOS is expected to average 1-1.5% annually during 2009-2014.

Linear Alkylbenzene

Linear alkylbenzene sulfonate (LAS) is a very-large-volume surfactant used in household and industrial detergents. It is produced by sulfonation of linear alkylbenzene (LAB) with oleum or a sulfur trioxide–air mixture. Most LAB is produced by alkylating benzene with normal paraffin–derived C_{10} - C_{14} linear internal olefins or C_{10} - C_{14} monochloroparaffins. However, Huntsman Chemical Corporation supplements its n-paraffin feedstock with purchased LAO. Internal olefins, derived from LAO in Shell's isomerization/disproportionation unit, can also be used for this purpose, but most of these products are exported for use in LAB production in other world areas. The paraffin-derived internal olefins are a far less costly source of raw materials than LAO, but the latter are used when paraffin-derived olefins are not available (as in many plants in Latin America and Southeast Asia) or to increase capacity utilization of plants or to adjust the chain-length distribution of the final product.

Annual U.S. consumption of LAO or internal olefins from LAO for LAB production is usually in the range of 11-14 thousand metric tons. Because of high capacity utilization of LAB plants and tight supplies and high prices for paraffins, consumption of LAOs in North America amounted to 23 thousand metric tons in 1999. However, by 2009, LAO consumption for this application declined to 9-10 thousand metric tons. North American LAB production and consumption decreased during 2002-2009. High raw material costs have caused some detergent manufacturers to reduce their cost by lowering surfactant levels, thus reducing LAB demand. Consumption in the United States is likely to decrease over the 2009-2014 period. (See the CEH *Linear Alkylate Sulfonates, Linear and Branched Alkylbenzenes* and *Surfactants, Household Detergents and Their Raw Materials* marketing research reports for additional information on LAB and LAS.)

Linear Mercaptans

Chevron Phillips and Arkema produce specialty linear, or normal, mercaptans from alpha-olefins. These companies also have facilities in Western Europe and supply virtually all international demand for linear mercaptans. Both U.S. producers have multipurpose equipment that can make a variety of linear, branched and cyclic mercaptans from different feeds. Arkema commercially produces C_4 - C_{12} linear mercaptans from alpha-olefins; it also produces very small amounts of C_{14} and C_{16} linear mercaptans. Chevron Phillips commercially produces normal C_4 , C_6 , C_8 and C_{12} mercaptans from alpha-olefins.

The major markets for linear mercaptans are for agricultural chemicals and polymer modification. They are used primarily as an agricultural intermediate for the production of sodium n-butylmercaptide, which is consumed in the production of the organophosphate herbicides DEF[®] and Folex[®] (tribufos). These products are used as defoliants or desiccants to facilitate machine harvesting of cotton. Approximately 2-3 thousand metric tons of butene-1 were consumed in 2009 for the production of 3-4 thousand metric tons of n-butyl mercaptan; some of the butene-1 may have been supplied by alpha-olefin producers.

Less than 500 metric tons of hexene-1 are consumed annually for the production of n-hexyl mercaptan, also used as an agricultural intermediate.

n-Octyl mercaptan is consumed in fabric coating treatments by the textile industry, as a polymerization modifier in the manufacture of plastics and coatings and in the manufacture of pesticides, special papers and pharmaceuticals.

n-Decyl mercaptan production is very small and is used in only a few specialty proprietary applications. Its production by Arkema is currently alcohol-based and does not consume alpha-olefins.

n-Dodecyl mercaptan is used mainly as a polymerization regulator for polymers, primarily in plastics (e.g., polymethyl methacrylate) but also in rubbers. It is also used for ore flotation in the mining industry, particularly for copper ore; however, this use is more significant internationally than domestically. It is believed to be the largest-volume linear mercaptan.

Consumption of C_6 - C_{12} alpha-olefins for normal mercaptans amounted to approximately 8 thousand metric tons in 2009. In addition, some butene-1 was likely consumed that may have been supplied from alpha-olefin plants. Hexene-1 and dodecene-1 are believed to be consumed in the largest quantities, followed by butene-1, octene-1 and decene-1. Overall growth for linear mercaptans will be modest during the next few years because of the maturity of their markets.

Aluminum Alkyls

Approximately 7 thousand metric tons of LAOs (C_6 - C_8 range) were consumed for the production of aluminum alkyls in 2009. Albemarle produces aluminum alkyls from alpha-olefins. Their largest use is for the production of Ziegler-Natta catalysts, used as polymerization catalysts (e.g., for polyolefins). Other uses include the production of organotins used as stabilizers for PVC or biocides. Growth in these applications was strong in the late 1990s with the increasing use of metallocene catalysts. Future growth is expected to average approximately 3% per year during 2009-2014.

Chlorinated Olefins

Chlorinated paraffin producers consumed approximately 5 thousand metric tons of C_{10} - C_{16} alpha-olefins in 2009 to manufacture chlorinated olefins, which can be used as replacements for chlorinated paraffins. The largest use for chlorinated olefins and paraffins is in the metalworking industry where they are contained in coolants and lubricants. Growth in consumption of LAOs in this category will likely be minimal over 2009-2014. They are also used as secondary plasticizers, especially for polyvinyl chloride and as flame retardants. See the CEH *Normal Paraffins* (C_9 - C_{17}) marketing research report for further information on chlorinated hydrocarbons.

Alkyldiphenylether Disulfonates

Alkyldiphenylether disulfonates were first produced by Dow Chemical U.S.A. and marketed under the trademark DOWFAX[®]. Dow currently supplies these products. Pilot Chemical Company manufactures and supplies alkyldiphenylether disulfonates. Alkyldiphenylether disulfonates can be manufactured by reacting an LAO with diphenyl ether; however, branched-chain olefins are also used as raw materials in their production. Each mole of the resulting alkyldiphenylether is then sulfonated with two moles of sulfur trioxide to produce a double ionic charge upon neutralization. The resulting product has greater dispersing power and higher solubility than most other surfactants.

Dow internally consumes significant quantities of this product for use in emulsion polymerization, which is one of the product's largest uses. Another large use is as an acid dye leveler in nylon dyeing. It is also used in bleach solutions and as a cosurfactant and hydrotrope in various cleaning applications, including machine dishwashing liquids or gels. In the latter application, a specialty version of this surfactant is used that acts as a low-foaming rinse aid to provide a sheeting action that leaves dishes free of water spots. Alkyldiphenylether disulfonates have also been used as crystal growth modifiers and as surfactants in agricultural chemicals. Consumption of LAOs in this application was 4 thousand metric tons in 2009, since most of the product was made from branched olefins.

Other

Other applications for alpha-olefins are small and diverse, accounting for roughly 22 thousand metric tons of C_{6+} and much smaller volumes of butene-1 in 2009. The reactivity of the terminal double bond in LAOs makes them interesting raw materials for the synthesis of a wide variety of chemicals with a linear hydrocarbon chain attached to one or more functional groups.

Other applications for butene-1, which can be supplied by either LAO producers or other butene-1 producers, include the production of o-sec-butylphenol (OSBP) and the smaller-volume para-isomer (PSBP). Producers of OSBP and PSBP are Albemarle and SI Group (formerly known as Schenectady International). OSBP is consumed in the manufacture of a stabilizer for styrene monomer. OSBP was also used in the production of the herbicide dinoseb, which is no longer produced in the United States. PSBP has limited applications in phenolic resins. Butene-1 is also used as a solvent and for miscellaneous uses. See the CEH *Butylenes* marketing research report for information on these small-volume butene-1 derivatives.

Linear alpha-olefins are used in leather- and textile-treating compounds. The alpha-olefins are partially sulfated with 94% or 98% sulfuric acid to the corresponding secondary alkyl sulfates. The sulfurized alpha-olefins are used in combination with natural oils such as vegetable, fish and lard oils with fatty acids, methyl esters or alcohols to yield a formulated product. These sulfurized compounds are used as leather penetrants to soften hides after tanning. They are also consumed by the textile industry as textile lubricants. The proportion of linear alpha-olefins used for these oil mixtures varies in most products from 5% to 30%. The volume of LAOs consumed by this industry varies from year to year, depending on the price differential between the natural oils and LAOs. The leather- and textile-treating industries typically consume between 1-2 thousand metric tons of alpha-olefins annually.

Olefin epoxides can be made by epoxidation of linear alpha-olefins with peracetic acid or t-butyl hydroperoxide. LAO-derived epoxides can be used as modifiers for epoxy resins, as polyether reactants for polyurethanes, as emollient ingredients in cosmetics and lotions and for production of specialty surface-active agents. Arkema produces several specialty epoxides based on C_{10} - C_{30} LAOs at its plant in Blooming Prairie, Minnesota. These epoxides consume more than one thousand metric tons of LAO per year.

 C_{30+} alpha-olefins are waxlike products that are offered only by Chevron Phillips Chemical. They are used as a lubricant that is compounded into polyvinyl chloride to ensure lubrication during extrusion and to control fusion of the PVC compound. They are also used to improve wax blends for candles, crayons and coatings. They are used in the manufacture of wood siding that is pretreated with waterproofing agents based in part on LAOs. An estimated 2-3 thousand metric tons of these higher olefins are used in these applications.

Some surplus LAOs, especially those that are highly contaminated by branched and internal olefins, may be burned for fuel value or sold into low-value end-use markets, including down-hole oil field applications and to the mining industry. These lower-value uses are declining, but may still account for 2-5 thousand metric tons annually.

Polymers of C₄, C₈, C₁₀ and C₁₂ alpha-olefins are used as flow improvers at crude oil pumping stations to reduce pipeline drag, thereby increasing pipeline capacity.

Small amounts of C₁₈ LAO are copolymerized with maleic anhydride by Chevron Phillips in production of PA-18 resin, used primarily in water-resistant sunscreens.

alpha-Olefins are consumed for a variety of small-volume products such as trialkylphosphines and trialkylphosphine oxides, silvlhydrocarbons, silicones and organometallics.

PRICE

The following table presents U.S. prices for the various linear alpha-olefin fractions and blends:

			(c	ents per	pound)				
Chain Length	January 1, 1984	April 1, 1988	October 1, 1991	June 1993	September 1996	June 2000	January 2004	July 2007 ^b	August 2010 ^b
C_4	28	30-33	30-40	40	32	36	42	45-50	45-50
C ₆	42-44	50-55	50.5-70	70	59.5	63.5	69.5	120-130	80-85
C ₈	42-44	50-55	50.5-70	70	59.5	63.5	69.5	75-80	60-70
C ₁₀	42-44	50-55	50.5-70	- 70	70	74	80	80-85	65-70
C ₁₂	42-44	50-55	50.5-70	70	59.5	63.5	69.5	70-75	55-60
C ₁₄	38-44	50-55	50.5-70	70	59.5	63.5	69.5	70-75	50-55
C ₁₆	38-42	50-52	60-70	70	59.5	63.5	69.5	75-80	55-60
C ₁₈	38-44	50-52	60-70	70	59.5	63.5	69.5	75-80	60-65
C ₁₄ -C ₁₆	38-41	50-55	45-60	70	59.5	63.5	69.5	70-75	55-60
C ₁₆ -C ₁₈	38-43	50-55	45-60	70	59.5	63.5	69.5	75-80	60-65
C ₂₀ -C ₂₄	40	40	45	45	25	na	na	50-55	60-65
C ₂₄ -C ₂₈	39	40	X 45	45	25	na	na	50-55	60-65
C ₃₀₊	39	40	45	45	25	na	na	50-55	60-65

U.S. Prices for Linear alpha-Olefins

Bulk quantities, tank trucks or tank cars, fob shipping point. a.

b. Beginning in 2007, prices are estimated transaction prices, and thus are not directly comparable to previous years.

SOURCE CEH estimates based on communication with industry.

Because a variety of alpha-olefins are produced, some of which are in much higher demand than others and each of which has its own markets, the relationship between list and market price varies considerably by olefin. Also, product properties vary by supplier. Producers and consumers are reluctant to divulge typical market pricing because of the small number of consumers for each product.

The majority of merchant butene-1 (C_4) is made from C_4 raffinate and therefore its price tends to be influenced by gasoline prices. Since much of the merchant butene-1 is made from a refinery source, LAO producers must price their butene-1 at similar levels to move their product in the market.

TRADE

Trade data for LAOs are not always complete and probably include some mislabeled material; therefore, trade data are subject to user interpretation. Until 1992, North American imports of LAO were relatively trivial compared with total U.S. consumption of these products. Imports increased over the years, with the completion of the plant in Belgium and the availability of large volumes of hexene-1 from Sasol in South Africa beginning in 1994. Some Canadian imports may be from the United States (which may include those from South Africa, which first enter U.S. ports). North American imports of LAO for selected years based largely on data from the Port Import/Export Reporting Service (PIERS) and the Department of Commerce are presented in the following tables:

	U.S.	Imports of (thousand	Linear alp ls of metric	ha-Olefin c tons)	s ^a		
Country of Origin	1995	1998	1999	2002	2003 ^b	2006 ^b	2009 ^b
North America							
Canada				26	2.9	14.0	30.0
Mexico				<1	3.7		
Total				27	6.6	14.0	30.0
Europe							
Belgium	19	41	43	12	14.9	22.0	0.8
Netherlands	1	7	17	13	1 0.3	14.4	82.0
United Kingdom	3	2		t	2.9	5.1	12.0
Other			_1	2	0.1	1.1	2.7
Total	24	50	61	27	28.2	42.6	97.5
Other Countries							
Japan	10	5.2	5	3	1.9	9.2	3.9
South Africa	16	64	65	91	87.5	91.9	92.4
Miscellaneous ^b	1	_ <u>T</u> _		6	1.9	0.1	6.2
Total	26	69.6	70	100	91.3	101.2	102.5
Total	50	119.6	131	154	126.1	157.8	230.0

a. Data are for C_6 - C_{30} linear alpha-olefins.

b. Includes imports attributed to Sasol through other countries. Includes imports from Saudi Arabia and the Republic of Korea in 2009.

SOURCE: CEH estimates based on data from the U.S. Department of Commerce, Bureau of the Census and the Port Import/Export Reporting Service (PIERS).

Imports from the United Kingdom are attributed to Shell; those from Belgium to INEOS Oligomers; those from the Netherlands to both Shell and INEOS; and those from South Africa to Sasol. Both Shell and INEOS trade LAO with their European affiliates to balance demand for various fractions in each world area. Some material from Russia may be included in imports from the Netherlands.

Large volumes of C_{6+} alpha-olefins are exported, accounting for 20-25% of U.S. production in many years. CEH estimates of U.S. exports of LAO are based on data reported by PIERS and the U.S. Department of Commerce and on communication with industry sources. These estimates include Shell's internal olefins derived from LAO. The 1999, 2002 and 2003 data also include some quantities of butene-1 from producers of LAOs. These data are presented in the following table:

	(thousands of metric tons)						
([Country of Destination	1995	1999	2002	2003	2006	2009
North	America						
Car	nada	79	81.0	51.8	63.0	77.2	71.7
Me	xico	1	0.9	0.9	1.2	1.4	2.0
Т	otal	80	81.9	52.7	64.2	78.6	73.7
South	America						
Arg	entina			19.1	18.6	24.1	24.4
Bra	zil	7	22.1	59	4 1	3.6	3.8
Vo		35	20.0	30.5	15.8	25.8	11.7
Oth	er	1	0.0	0.5	3.7	0.8	0.3
Т	otal	43	43.0	56.1	42.2	54.3	40.2
-		10	1010	0011			
Europ	e						
Bel	gium	18	18.4	43.6	54.3	99.7	76.3
Fra	nce	63	82.8	65.3	52.8	61.5	42.1
Net	herlands	14	3.5	6.0	19.6	7.5	3.8
Uni	ted Kingdom	<1	3.9	6.1	2.8	4.6	2.0
Oth	er			1.3	6.4	0.1	2.4
Т	otal	96	108.6	122.2	135.8	173.4	126.6
Middl	e East/Africa						
Sau	di Arabia	1	0	0	11.9	1.0	
Sou	th Africa	5	5	11.4	8.8	4.0	0.5
Oth	er	0	0	0.2			2.0
7	otal	6	5	11.6	20.7	5.0	2.5
Asia I	Pacific						
Chi	na				b	35.4	29.0
Ind	ia 🔨	6	15.2	43.0	67.4	35.3	30.0
Ind	onesia	20		1.3	2.1	3.3	5.1
Ian	an	1	0.0	2.8	0.1	3.0	74
Jap Ko	ea Republic of	13	0.0 7 7	117	14.2	18.4	20 2
Dhi	linnines	15	20	0.0	0.0	0.4	20.2
	inppines	1	2.0	11.0	10.0	0.0 5 2	0.0
Tai	wafi	10	5.7	11.2	12.7	5.5	2.7
Th	lland			10.6	10.7	0.0	0.0
Oth	er	1	2.8	5.1	19.5	1.9	2.9
	otal	58	33.5	85.6	126.8	102.6	97.3
Other						5.0	2.0
_			272.0	220.2	200 7	410.0	242.2

U.S. Exports of Linear alpha-Olefins by Country^a

b. Included in Other (Asia Pacific).

SOURCE: CEH estimates based on data from the U.S. Department of Commerce, Bureau of the Census and the Port Import/Export Reporting Service (PIERS).

Historical data on exports of alpha-olefins have been reported separately by the U.S. Department of Commerce since 1982. Although the data are not always complete and probably include some mislabeled material, the reported export data are presented in the following table:





Shell Chemical exports both internal olefins and alpha-olefins. Most exports of internal olefins are believed to be used in the production of linear alkylbenzene. North American exports (i.e., exports to areas outside of the United States, Canada and Mexico) of alpha-olefins are estimated to have reached approximately 269 thousand metric tons in 2009.

The North American exports shown in the **SUMMARY** section (269 thousand metric tons) exclude U.S. exports to Canada and Mexico for that year. Likewise, North American imports shown in the **SUMMARY** (200 thousand metric tons) exclude U.S. imports from Canada.

WESTERN EUROPE

PRODUCING COMPANIES

The following table presents Western European producers of linear alpha-olefins:

Wes	tern European Produce	ers of Linear alpha-Olefins
Company and Plant Location	Annual Capacity as of Mid-2010 (thousands of metric tons)	Remarks
Dow Chemical		
Tarragona, Spain	100	Plant came on stream in 2006. Produces octene-1
INEOS Oligomers Feluy, Belgium	300 ^a	from butadiene. Listed capacity is for LAOs in the C_6 - C_{18} range. C_4 is recycled to produce higher olefins.
Shell Chemicals UK Ltd.	220	Mainite and the Band Detak (Shall Cours
Stanlow, Ellesmere Port,	550	Majonity owned by Royal Dulch/Shell Group.
Onneu Kingdoni		C_6 - C_{22} range are produced. Trade name in the United States is Neodene.
Total	730	

a. Part of C_{10} production is used captively as feedstock for INEOS' 126 thousand metric ton-per-year polyalphaolefins (PAO) plant. In July 2009 INEOS announced the start-up of its new LAO technology innovation at Feluy that allows for greater flexibility to produce individual LAO cuts.

SOURCE: CEH estimates.

In the 1990s, there was very strong growth in global demand for linear alpha-olefin products. Driving this growth was the use of LAO as a polyethylene comonomer and the use of polyalphaolefins for synthetic lubricants. After an oversupplied global market and price pressures in 2002-2003, profitability returned in 2004 and by 2005 the market had become balanced. In 2006, the market was tight, due to increased demand for polyethylene and PAOs. Additional capacity that came on stream in mid-2007 (Dow in Tarragona, Spain, and Yanshan in China) and the lifting of Sasol's allocation eased the tightness in the market in 2007. With the global economic crisis, demand and production decreased in 2008 and 2009, but as demand picked back up in the latter part of 2009, the market became quite tight, particularly for C₆.

INEOS plant at Feluy, Belgium produces LAOs in the C_6 - C_{24} range. The main products are C_6 , C_8 and C_{10} cuts; smaller amounts of C_{12} - C_{24} cuts are also produced. The chain length distribution is approximately 28% C_6 , 28% C_8 and 20% C_{10} . This distribution can be altered by a few percent. Part of the plant output is consumed as feedstock for the 126 thousand metric ton-per-year polyalphaolefin plant. In July 2009 INEOS announced the start-up of its new LAO technology innovation at Feluy that allows for greater flexibility to produce individual LAO cuts.

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The Royal Dutch/Shell Group operates a plant based on the Shell Higher Olefins Process (SHOP) for the production of linear alpha- and internal olefins at its Stanlow chemical facility in Ellesmere Port, United Kingdom. As well as producing LAOs, the SHOP process is unique in that it can recycle longer-chain LAOs into internal olefins. Internal olefins derived from alpha-olefins (by the SHOP process) are included in the production data shown below.

The SHOP plant is supplied ethylene via pipeline from the cracker at Mossmorran, Scotland. The process results in even-numbered alpha-olefins ranging from C₆ to C₂₂ and greater, and a range of internal olefins from C_6 to C_{14} . Blends can also be tailored to customers' specific requirements. The linear alpha- and internal olefins produced at Stanlow are destined for both merchant and captive consumption, with Shell a major consumer of both types in the production of detergent alcohols, plasticizer alcohols and lube oil additives (through Infineum, its joint venture with ExxonMobil Chemical Company).

Sasol Italy S.p.A. produces linear internal olefins from n-paraffins in a plant at Augusta in Sicily, with a capacity of 250 thousand metric tons per year; around 90% is used captively for the production of oxo process chemicals, linear alkylbenzene and polyinternal olefins. CEPSA Quimica (formerly Petresa) in Spain also produces linear internal olefins from n-paraffins. These products are not included in this report, although some alpha-olefins are present in the products and they are sometimes used in the same applications as LAOs derived from ethylene (and internal olefins made from LAOs by isomerization using the SHOP process).

Higher olefins (such as those produced by ExxonMobil) that are based on the oligomerization primarily of propylene, but also butylene, are not included in this report. In addition, butene-1 is not covered in the discussion of Western European alpha-olefin markets; for information on this product see the CEH Butylenes marketing research report. All butene-1 generated in alpha-olefin plants in Western Europe is converted to higher alpha-olefins or internal olefins and therefore does not contribute to merchant supply.

SALIENT STATISTICS

The following table presents Western European supply/demand for linear alpha- and internal olefins:

	(thousands of metric tons)				
	Production	Imports	Exports	Net Imports	Consumption
1992	290	125	18	107	397
1995	350	157	30	127	477
1996	375	152	47	105	480
1997	406	156	34	122	528
1998	426	181	31	150	576
1999	461	168	34	134	595
2000	485	195	49	146	631
2001	473	215	45	170	643
2002	428	272	35	237	664
2003	475	248	45	203	678
2006	535	235	120	115	650
2009	551	229	135	94	645

Western European	Supply/Demand for Linear alpha-Olefins ^a
	thousands of metric tons)

a. Data include C_6 and higher (alpha and internal).

SOURCES: (A) Official statistics.

(B) CEH estimates.

The data in the above table include estimates for LAOs, and internal olefins derived from LAOs with a carbon content of C_6 and higher; all butene-1 is converted internally to internal and C_{6+} alpha-olefins. For information on butene-1 generated from other sources, see the CEH *Butylenes* marketing research report.

CONSUMPTION

The following table presents Western European consumption of LAOs and internal olefins derived from them (linear internal olefins are closely related to linear alpha-olefins in use and are interchangeable in certain outlets):

Western European Consumption of Linear alpha-Olefins ^a						
		2009	2009 2014			Average Annual
	Main Carbon Number Range	Thousands of Metric Tons	Percent of Total	Thousands of Metric Tons	Percent of Total	2009-2014 (percent
Polyethylene Comonomers	6-8	225	35	242	34	1.5
Polyalphaolefins ^b	10, 8-12	196	30	235	33	3.7
Oxo Alcohols						
Detergent Alcohols ^c	10-16	118	18	125	18	1.2
Plasticizer Alcohols	6-10	43	7	45	6	0.9
Lube Oil Additives	16-18	36	6	34	5	-1.1
alpha-Olefin Sulfonates	14-16	3	<1	3	<1	0.5
Other ^d		24	4	27	4	2.4
Total		645	100%	711	100%	2.0%

a. Data include C_6 and higher (alpha and internal).

b. Approximately 80% of PAOs are based on C_{10} ; C_8 and C_{12} are also used to make PAOs.

- c. Consumption includes exclusively LAOs and SHOP internal olefins.
- d. Includes consumption for uses such as amines and mercaptans.

SOURCE: CEH estimates.

The following table presents historical Western European consumption of LAOs and internal olefins:

LINEAR ALPHA-OLEFINS

Historical Western European Consumption of Linear alpha-Olefins ^a
(thousands of metric tons)

	(
	Main Carbon Number Range	1992	1995	1999	2002	2003	2006
Polyethylene Comonomers	6-8	60	87	170	213	220	232
Oxo Alcohols							
Detergent Alcohols ^b	10-16	150	160	145	161	163	115
Plasticizer Alcohols	6-10	34	38	40	42	42	42
Polyalphaolefins	10	62	94	186	192	196	191
Lube Oil Additives	16-18	40	42	42	40	40	39
alpha-Olefin Sulfonates	14-16	3	3	3	3	3	3
Linear Alkylbenzene ^c	10-14	40	45				A
Other ^d		8	8	9	13	13	28
Total		397	477	595	664	678	650

a. Data include C_6 and higher (alpha and internal).

b. Starting in 1999, consumption includes exclusively LAOs and SHOP internal olefins

c. Since 1996, LAB is produced nearly exclusively with normal paraffins.

d. Includes consumption for uses such as amines and mercaptans.

SOURCE: CEH estimates.

Linear alpha-olefins are currently consumed primarily in the United Kingdom, Germany, France and Belgium.

Linear alpha- and SHOP internal olefin demand in Western Europe was 645 thousand metric tons in 2009 and is expected to grow at an average annual rate of 2% during 2009-2014, reaching 711 thousand metric tons in 2014. The ability to interchange alpha- and internal olefins in some downstream applications makes it difficult to forecast a specific pattern of future demand for alpha-olefins as a separate entity. Growth in LAO consumption will be driven mainly by the increasing use of alpha-olefins as comonomers in polyethylene production (mainly C_6 and C_8 alpha-olefins) and polyalphaolefin production. Most of the outlets for detergent and plasticizer alcohols are expected to grow at a lower rate as the alpha-olefinderived alcohols compete with alcohols derived from dehydrogenated paraffins and from fats and oils (i.e., natural alcohols).

The following table presents Western European consumption of LAOs by type in 2009:

Western European Linear alpha-O (perce	Consumption of lefins—2009 nt)
$egin{array}{ccc} C_6 & & & \ C_8 & & \ C_{10} & & \ C_{12} & & \ C_{14} & & \ C_{16} & & \ C_{18} & & \ C_{20+} & & \end{array}$	23 16 28 14 9 6 2 2 2
Total	100%
SOURCE: CEH e	stimates.



Polyethylene Comonomers

The largest outlet for linear alpha-olefins and internal olefins from the SHOP process is in the production of comonomers for polyethylene production. Only highly linear alpha-olefins can be used as comonomers; olefins derived from other sources such as paraffin cracking cannot be used for this application, because of their high level of impurities.

 C_4 , C_6 and C_8 LAOs are increasingly being used as comonomers in the production of polyethylene, primarily LLDPE; and some C_4 and C_6 are used for HDPE. Typically, these comonomers are used up to 8-10% by weight of LLDPE, up to 2-3% of HDPE and up to 20-30% by weight in some specialty-grade polyethylenes. Butene-1 is used in the commodity end-use applications of LLDPE. It is still the most-used comonomer but hexene-1 and octene-1 comonomers yield polymers with improved properties. LLDPE with hexene-1 gives a tougher film than with butene-1 that can be produced in thinner thickness as preferred in packaging applications. Polyethylene with octene-1 gives the highest-quality products with good surface finish, good transparency and improved resistance to tearing. It is produced by solution-phase processes. Hexene-1 can be used in gas-phase processes while butene-1 is used in either solution-phase or gas-phase processes.

BP previously used 4-methylpentene-1 as a comonomer but switched to hexene in its Innovene process and closed its 4-methylpentene-1 production plant. Separate fractions of C_6 (hexene-1) and C_8 (octene-1) LAOs have been available for use as comonomers since mid-1983. As a result, there has been a shift from butene-based polymers toward C_6 and C_8 . In 1999, the market share of butene-based polyethylene was approximately 50% of the total polyethylene film market with hexene- and octene-based polymers accounting for 17% and 33%, respectively. Of the total Western European polyethylene film market in 2009, butene-polyethylene had a market share of around 40%, hexene-polyethylene, 27% and octenepolyethylene, 33%. The substitution from C_4 to C_6 is expected to continue at reduced rates as the higher price for C_6 limits its penetration.

The following table presents Western European consumption of C_6 and C_8 LAOs as comonomers:

	Western E Comone P (thou	Curopean Consumptioners for Polyethyle roduction—2009 sands of metric tons	ion of ene
×e	C_6 C_8		150 75
	Total		225
	SOURCE:	CEH estimates.	

Western European consumption of C_6 and C_8 LAOs as polyethylene comonomers was 225 thousand metric tons in 2009. The economic crisis that hit in 2008 and continued through much of 2009 contributed to lower polyethylene demand (and hence lower LAO consumption). LLDPE production in Western Europe is expected to grow at an average rate of 1.5% annually between 2009 and 2014. However, consumption of C_6 and C_8 LAOs for the production of LLDPE is expected to increase at a slightly higher average annual growth rate of 2%. HDPE production is projected to decrease at approximately 2% per year during the forecast period. However, comonomer consumption for the production of HDPE is expected to remain relatively flat compared to HDPE from other raw materials.

LINEAR ALPHA-OLEFINS

The use of new metallocene- and single-site catalyst processes will influence LAO consumption in the production of polyethylene. As such catalysts optimize product performance versus the amount of comonomer added, they yield the same end-use performance by using less comonomer. However, the usage of such catalysts is limited and their penetration in polyethylene production is still slow. Western European consumption of C_6 and C_8 LAOs as comonomers is forecast to grow at approximately 1.5% per year during 2009-2014. Higher growth will be constrained by competing material from the Middle East, where capacity has been recently added and continues to grow. Similar to North America, both the production and consumption of PE is shifting from Western Europe to regions such as China where demand is considerably higher or to regions where feedstocks are cheaper (Middle East) and demand is strong.

Polyalphaolefins

 C_8 linear alpha-olefin, C_{10} linear alpha-olefin (decene-1), C_{12} linear alpha-olefin, C_{14} or in some cases a blend, is converted by oligomerization to produce polyalphaolefins (PAOs), which are used as base stocks for synthetic lubricants. Decene-1 is the mostly widely used LAO. Typically, PAOs contain two to four oligomerized LAOs, with a maximum distribution at three. PAOs offer important advantages over conventional lubricants. They exhibit high thermal stability, low volatility, low pour point and optimum viscosity in automotive crankcases, which accounts for most consumption.

Western European Producers of Polyalphaolefins^a **Annual** Capacity as of Mid-2010 Company and **Plant Location** (thousands of metric tons) Trademark ExxonMobil Chemical Company Gravenchon, France 65 SpectraSyn® INEOS Oligomers^b Feluy, Belgium 126 Durasyn® Neste Oilc Beringen, Belgium **NEXBASE[®]** 60 Total 251 Sasol Italy S.p.A. produced poly-internal-olefins (PIOs) in a plant at Sarroch, Sardinia with an annual capacity of 20 thousand metric tons, but it was mothballed in 2008. b. INEOS increased PAO capacity by 7 thousand metric tons in 2007 and completed a 43 thousand metric ton-per-year expansion in 2009. Neste Oil increased PAO capacity by 10 thousand metric tons in 2008. SOURCE: CEH estimates.

The following table presents Western European producers of polyalphaolefins:

LINEAR ALPHA-OLEFINS

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Western European consumption of alpha-olefins for the production of synthetic lubricant base stock amounted to 94 thousand metric tons in 1995 and increased to 186 thousand metric tons in 1999. Showing high growth rates of above 20% in the past, PAO markets in Western Europe matured during the 1999-2003 period. During 2003-2007 and most of 2008, PAO production plants had high utilization rates. The scarcity of base stocks resulted in the cessation of the trend to reformulate automotive lubricants with Group III. Also, exports to the Middle East, Russia and South America (with the added demand in these regions) increased, attributing to some of the increase in PAO production. However PAO production (and consumption) slowed by year-end 2008 and throughout most of 2009, a result of the global economic crisis.

The following table presents Western European consumption of LAOs for the production of PAOs:

Western I	European Consumption for Polyalphao (thousands of met	of Linear alpha-Olefins lefins aric tons)
	Polyalphaolefin Production	Linear alpha-Olefin Consumption ^a
2006	185	191
2009	190	196
2010 ^b	200	206
a. The a depen- oligon were c	umount of decene-1 co ding on the process a ner products. On average consumed per pound of P	nsumed for PAO varies nd the viscosity of the , 1.03 pounds of decene-1 AO.
b. Projec	ted estimate.	¢
SOURCE:	CEH estimates.	

Consumption of LAOs for PAO production reached approximately 196 thousand metric tons in 2009. Production of PAOs (and therefore consumption of LAOs for this application) is forecast to grow at an average annual rate of 3.7% during 2009-2014, although consumption of PAOs is not expected to grow as quickly in the region as there may be some competition from HVI/VHVI hydrocracked and wax isomer oils. Cost/performance considerations will continue to determine to what extent PAOs will be replaced by HVI/VHVI oils. More decene-1 should become available with recent LAO capacity expansions in the Middle East as well as INEOS' new LAO technology innovation at Feluy that allows for greater flexibility to produce individual LAO cuts.

Historically, consumption of PAO in Western Europe grew more rapidly than in other parts of the world, principally because of the requirements of its automotive industry. Car engines in Western Europe have higher compression ratios than those in North America and thus experience higher temperatures that necessitate motor oils with improved stability. In addition, the specifications that are imposed by the principal European car producers are becoming stricter, in terms of higher thermal and chemical stability, lower VOC emissions and higher viscosity/temperature indexes. Compared with North America, European consumers prefer longer oil-change intervals, on the order of 30,000 kilometers and often use the same oil that was specified by the car manufacturer. In recent years, 10W-40 and 5W-40 motor oil types have been used more frequently, following the trend to lower viscosities and fuel economy as they are multigrade oils. Initially, the 10W-40-grade specifications could be met only by PAOs but now HVI/VHVI oils have improved in quality over the years to the extent that they are used almost

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exclusively in these grades. The 5W-40 grade also started out in 1992 being completely PAO-based; now HVI/VHVI oils account for approximately 80% of the market. The 0W-XX grades are mostly PAO-based because of the extremely low-viscosity and low-volatility requirements (driven by fuel economy), but formulations based on HVI/VHVI oils have been developed and are marketed. The replacement of PAOs by HVI/VHVI oils is driven by cost/performance considerations. Crankcase lubricants for gasoline engines accounted for 75% of total Western European PAO consumption in 2009.

PAO-based oils are also used in diesel engines, although this market is very small as yet, contributing to roughly 8% of Western European PAO consumption in 2009. Legislation that came into force in 2005 requires a reduction of the diesel engine VOC emissions by recycling the exhaust gases. Thus, PAO-based synthetic oils have become an alternative method of meeting this requirement. Therefore, consumption growth through 2014 may be slightly higher than that for other automotive applications for PAOs.

Other applications for PAOs, including automotive gear oils, automatic transmission oils, aviation crankcase lubricants and industrial applications, accounted for 17% of total Western European PAO consumption in 2009.

Sasol Italy produced poly-internal-olefins (PIOs) at its Sarroch, Sardinia plant that was mothballed in 2008. It had an annual capacity of 20 thousand metric tons. PIOs are obtained by oligomerization of internal olefins derived from C_{15} - C_{16} n-paraffins to give a final product that consists of approximately 75% C_{30} , 20% C_{45} and 5% C_{60} . The average molecular weight of a PIO mixture is similar to that obtained with PAOs. Most of the PIOs have kinematic viscosity indexes of 6 and 8 mm²/s (at 100°C). Of the 20 thousand metric tons, 5 thousand metric tons could be treated to give PIOs with a viscosity of 4 (PIO[4]). However, a similar amount (2.5 thousand metric tons) of PIO(8) were also produced as a coproduct, but it was difficult to sell. Oils with a viscosity of 6 are the most commonly used type in car engines, but oils with smaller indexes represent a fast-growing market for high-quality oil. In 2006, Sasol Italy had a market share of 10% in Western Europe. The company produces internal olefins at its own plant in Italy, with a current capacity of 250 thousand metric tons per year. Most of this capacity is devoted to making linear alkylbenzene or detergent alcohols.

For more information on synthetic lubricants see the SCUP Synthetic Lubricants report.

Oxo Alcohols

Commercially, the use of LAOs for linear alcohols is divided into two ranges, C_6 - C_{10} and C_{10} - C_{16} . C_7 - C_{11} linear alcohols derived from C_6 - C_{10} LAOs are consumed primarily for the production of plasticizers while C_{11} - C_{17} linear alcohols, derived from C_{10} - C_{16} , are consumed primarily for surfactants used in detergents. Some consumption in both ranges is for other applications, such as the production of methacrylate esters used as viscosity index improvers (C_8 - C_{15} alcohols) in lube oils and other uses.

Detergent oxo alcohols

The production of detergent-range oxo alcohols in Western Europe based only on LAOs and SHOP olefins is difficult to separate from the production based on other internal olefins. In 2009, approximately 118 thousand metric tons of LAO or SHOP internal olefins were utilized for detergent-range oxo alcohols. The two producers of detergent-range alcohols from LAOs are Shell (which uses its own LAOs from the SHOP plant) and BASF (which purchases linear alpha-olefins). Noroxo, owned by ExxonMobil,

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mothballed its site at Harnes, France in November 2003, closing a 130 thousand metric ton-per-year combined oxo alcohol capacity, mainly for branched alcohols. However, approximately 50 thousand metric tons were dedicated for the production of detergent alcohols. Noroxo used to purchase linear alpha-olefins. Sasol Italy S.p.A., which is back-integrated into internal olefins from paraffins, may also use some LAOs to make detergent alcohols. In Europe, the main use of detergent-range alcohols is for the production of ethoxylates, ether sulfates and alcohol sulfates.

Detergent alcohols produced from linear alpha- and internal olefins compete with natural alcohols that are derived from coconut oil and palm kernel oil. The latter have accounted for about 60% of Western European production in recent years. Annual growth in Western European consumption of LAOs for detergent alcohol production is forecast at 1.2% per year during 2009-2014.

For more information on oxo-derived linear alcohols, refer to the CEH *Oxo Chemicals* marketing research report.

Plasticizer alcohols

As in other parts of the world, linear plasticizer alcohols account for only a minor share (approximately 4% in 2009) of the total plasticizer alcohol market in Western Europe. By far the most commonly used plasticizer alcohols are n-butanol, 2-ethylhexanol (2-EH) and isononyl alcohol (INA). Branched oxo alcohols are also being increasingly used.

A nonplasticizer outlet for C_8/C_{10} olefins is in the manufacture of C_9/C_{11} oxo alcohols that are subsequently ethoxylated to produce nonionic surfactants. These surfactants are used in a number of household and industrial detergent and cleaning formulations.

Western European consumption of C_6 - C_{10} olefins for the production of plasticizer-range alcohols (including those used in nonplasticizer applications) was 43 thousand metric tons in 2009. This includes only plasticizer alcohols produced with alpha- and SHOP internal olefins.

LAOs demand for C_7 - C_{11} oxo alcohol manufacture is expected to increase at a rate of 1.0-1.5% per year during 2009-2014. The more rapid increase in use of nonlinear plasticizers such as diisononyl and diisodecyl phthalates will limit the growth of the linear alcohols.

For additional information on plasticizer alcohols, see the CEH *Plasticizer Alcohols* (C_4 - C_{13}) marketing research report.

Lube Oil Additives

Linear alpha-olefins or polymers in the C_{14} - C_{20} range can be reacted with phenol to produce alkylphenols and with benzene to produce alkylbenzenes. The alkylphenols can subsequently be transformed into calcium phenates and the alkylbenzenes can subsequently be sulfonated and neutralized to produce sodium and calcium sulfonates. Sulfonates and phenates are the two most important detergent additives used in lube oils. Another lube oil additive manufactured from linear alpha-olefins or internal olefins is alkenylsuccinic anhydride (ASA), a dispersant and corrosion inhibitor.

Consumption of detergents used in lube oils is expected to decrease over the next few years in response to their decreasing consumption in automotive engines. Technical improvements in the construction of

engines and in oil formulations have lengthened oil lifetimes and lowered the volumes needed. For further information on lube oil additives see the SCUP *Lubricating Oil Additives* report. Compared with 1999, Western European consumption of linear alpha-olefins for lube oil additive production declined annually by 1.5% to 36 thousand metric tons in 2009. This market is forecast to further decrease at an average yearly rate of 1.1% during 2009-2014.

alpha-Olefin Sulfonates

Western European LAO consumption for alpha-olefin sulfonate (AOS) production was only 3 thousand metric tons in 2009. AOS use in Western Europe is confined mainly to industrial and institutional cleaners, with smaller quantities used in light-duty liquid and personal care products. Future consumption levels in Western Europe are expected to be stable. Akzo Nobel, Clariant, Huntsman and Kao are the main AOS producers in Western Europe and C_{14} - C_{16} olefin sulfonates are their main products. Little growth is expected during 2009-2014.

Linear Alkylbenzene

The use of linear and internal olefins for linear alkylbenzene (LAB) production in Western Europe has essentially ceased since the mid-1980s. In 1991, Shell Chimie S.A. in France and Shell Chemicals in the United Kingdom ceased LAB production based only on linear alpha-olefins. Hüls stopped manufacturing linear alkylbenzene from internal olefins at Marl (Germany) in 1993.

In Western Europe, LAB is produced by alkylating benzene, using either monochloroparaffins or C_{10} - C_{13} internal olefins. LAB production based on LAOs is not significant in Western Europe, although some small LAO quantities may be purchased for that purpose. CEPSA Quimica and Sasol Italy S.p.A. currently produce LAB from internal olefins derived from n-paraffins. In 1996, CONDEA closed its LAB production facility in Mantua, based on internal olefins. Since then, Western European production of LAB has been nearly exclusively from normal paraffins, a less expensive feedstock.

For more information on linear and branched alkylbenzene, please refer to the CEH *Linear and Branched Alkylbenzenes* marketing research report.

Other

Other uses include products such as mercaptans, amines and their derivatives (quaternary ammonium compounds), epoxides and alkenylsuccinic anhydrides.

Linear mercaptans

Long-chain linear and branched mercaptans are produced from linear alpha-olefins, as well as from other raw materials. Arkema at Rotterdam, the Netherlands, and Chevron Phillips Chemicals at Tessenderlo, Belgium produce long-chain mercaptans based on alcohols. The main applications for these products are as rubber chemicals and plastic additives. Demand for LAOs for the manufacture of linear mercaptans was approximately 2 thousand metric tons in 2009. No growth is expected in this application over the next few years.

Alkenylsuccinic anhydride

Alkenylsuccinic anhydrides (ASA) are produced from linear alpha-olefins in the C_{12} - C_{20} range or internal olefins in the C_{15} - C_{18} range by reaction with maleic anhydride. These products are used as paper sizing compounds, surfactants and leather preservatives, as well as in lube oil additives. Consumption of LAO in Western Europe for ASA production more than tripled during the 1999-2009 time period, from 4 thousand metric tons to 20 thousand metric tons, driven mainly by the start-up of several large paper and board machines and by conversion of some other plants from alkyl ketene dimer (AKD) to ASA. The major producer of ASA in Western Europe is Kemira Chemie at Krems, Austria. Other suppliers of ASA in Western Europe include Eka Chemicals, Pentagon and BASF. Growth in the next few years is forecast at 2.0-2.5% per year. For further information on ASA, see the SCUP *Specialty Paper Chemicals* report.

PRICE

LAO contract prices are linked to ethylene prices with a variable add-on depending on customer location and contract timing.

Spot prices for recent years are not given because large quantities of LAOs are not readily available on the open market. The following table presents Western European prices for LAOs:

Western European Prices for Linear alpha-Olefins

	(dollars]	per metric ton)	
	2003 ^a	2007 ^b	2010 ^b
C ₆	700-800	2,500-3,000	1,900-2,100
C_8	1,000-1,100	1,500-1,650	1,400-1,650
C ₁₀	850-950	1,600-1,700	1,500-1,700
C ₁₂	700-800	1,400-1,600	1,200-1,400
C ₁₄	600-700	1,400-1,600	1,200-1,400
C ₁₆ -C ₁₈	850-950	1,500-1,650	1,300-1,500
a. Spot p	rice.		
b. Transa	ction price.		
SOURCE:	CEH estima	tes.	

Spot prices are higher than contract prices and are only indicative, since only small LAO quantities are traded in this way. Prices in 2007 increased primarily because of higher raw material costs, tight supply and increased demand. Weak demand in most sectors brought about by the economic crisis resulted in lower prices in 2008 and 2009.

The following table presents historical midyear prices for several linear alpha-olefins up to 1996. Prices are presented in dollars per metric ton since the LAO market is global.

for Linear alpha-Olefins—1989-1996 ^a (dollars per metric ton)							
	C ₆	C ₈	C ₁₀	C ₁₂			
1989	1,500	1300	1,350	1,200			
1991	1,200	900	900	850			
1992	1,100	800	900	800			
1993	1,050	800	1,000	750			
1994	1,050	900	1,150	750			
1995	800	1,200	1,300	1,200			
1996	800	1,000	1,350	1,250			

a. Prices are for bulk quantities, delivered.

SOURCE: CEH estimates.

TRADE

Western Europe is a net importer of LAOs. Total imports in 2009 were approximately 230 thousand metric tons. The United States was the largest LAO exporter to Western Europe, followed by South Africa and Japan. The following table presents Western European imports of LAO by country of origin in 2009:



Western Europe exported 135 thousand metric tons of LAOs in 2009. The major export destination was the United States, accounting for 97 thousand metric tons in 2009. Other lower-volume destinations are Japan, India, China, the Republic of Korea and Taiwan.

CENTRAL AND EASTERN EUROPE

The following table presents Central and Eastern European producers of linear alpha-olefins:

Central and Eastern	European Producers	of Linear alpha-Olefins
	Bui opean i roudeers	of Emetal alpha Olemb

Company and Plant Location	Annual Capacity as of Mid-2010 (thousands of metric tons)	Remarks
Nizhnekamskneftekhim Inc. (NKNK) Nizhnekamsk, Russia	90	LAOs in the C ₄ -C ₂₆ range are produced. Came on stream in 1990. Plant nameplate capacity was originally 186 thousand metric tons. Operational capacity is 90 thousand metric tons due to demand/feedstock issues, but could actually be lower (75 thousand metric tons).
SOURCE: CEH estimates.		

Nizhnekamskneftekhim's plant is based on technology developed by Ethyl. When it started up in 1990, Nizhnekamskneftekhim produced LAOs under license in two separate production lines with 90 thousand metric tons of annual capacity each. As part of the license, Nizhnekamskneftekhim could sell its products only in countries of the former COMECON and to the United States. Because of serious financial problems in Russia, the plant was inoperative for several years in the 1990s. In 1998, one production line was restarted with an annual capacity of 90 thousand metric tons. Approximately 60% of production is exported. Hexene-1, octene-1 and longer-chain LAO mixtures are sold. Russian production of LAOs was approximately 15 thousand metric tons in 2009, of which 7 thousand metric tons were exported. Because of unresolved quality problems, Nizhnekamskneftekhim experienced difficulties in marketing its LAO products. Therefore, a project for captive consumption of LAOs for the production of LAB and synthetic lubricants was announced. In February 2009, Nizhnekamskneftekhim commissioned a 230 thousand metric ton-per-year LLDPE/HDPE plant.

SOUTH AFRICA

The following table presents South African producers of linear alpha-olefins:



The following table	presents a timeline	of Sasol	production capacity:

Sasol's South African Production Capacity for Linear alpha-Olefins (thousands of metric tons)									
1996 2000 2003 2006 2010									
Pentene-1	10	a	^a	^a	^a				
Hexene-1	45	110	188	188	188 ^b				
Octene-1		48	48	96	186 ^b				
Total	55	158	236	284	420^{c}				

Small quantities of pentene produced. a.

On-purpose capacity. b.

Total capacity, including hexene-1, octene-1, pentene-1 and some c. higher LAO capacity.

SOURCE: CEH estimates.

Sasol installed the first stage of its linear alpha-olefins production in 1994 at its Secunda coal-tosynthetic-fuels plant, with 55 thousand metric tons of combined pentene/hexene annual production capacity. In 1997, pentene/hexene capacity was doubled to 110 thousand metric tons per year by adding a second production line. However, the demand for pentene-1 was smaller than expected; as a result annual pentene-1 capacity has never been fully used and most of it was converted to hexene-1 production. In October 2000, Sasol commissioned a third hexene-1 production line with 78 thousand metric tons of capacity, expanding total annual hexene-1 capacity to 188 thousand metric tons, mainly for the merchant market.

Sasol also started the construction of a new plant for octene-1, sourcing feedstock from its C1-derived synfuel operations. The first plant located in Secunda, South Africa came on stream in the second quarter of 1999 with a capacity of 48 thousand metric tons per year. Its entire production is dedicated to Dow Chemical. The second octene-1 plant, adding another 48 thousand metric tons of annual capacity, was completed in 2005. Sasol built another octene-1 unit with an annual capacity of 90 thousand metric tons that became operational in June 2008.

The following table presents South African production of LAOs:



	South	n African Pi (thousands	rodu of m	ction of LA etric tons)	Os ^a	
199	8					100
200	2					175
200	6					250
200	9					340
a.	Small produce	quantities ed.	of	pentene-1	may	be

SOURCE: CEH estimates.

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The entrance of Sasol into the LAO business in 1994 greatly affected hexene pricing. Sasol produced a good-quality hexene-1 at a price 20% lower than its competitors in Western Europe. In July 2002, Sasol commissioned a 120 thousand metric ton-per-year plant at Secunda to produce detergent alcohols in the C_{12} to C_{13} range, using coal-based longer-chain alpha-olefins as a feedstock. Linear alcohols account for 50% of the output; the balance has varying degrees of branching.

In 2009, South African consumption of LAOs was 70 thousand metric tons, the majority of which was for the production of detergent alcohols. Exports totaled 270 thousand metric tons (34% was exported to the United States, 34% was exported to Western Europe and 32% was exported to other regions). Sasol, South Africa's sole producer of detergent alcohols, is a net exporter, and will continue to be so during 2009-2014. Linear alpha-olefins consumption for the production of detergent alcohols in the region is estimated to grow at an average annual rate of approximately 5% during 2009-2014. However, since most of the LAO production in South Africa is for the export market, it is difficult to predict production growth. It is believed that export growth on an average annual basis will be in the high single digits during 2009-2014.

MIDDLE EAST

The following table presents Middle Eastern producers of linear alpha-olefins:



Middle Eastern Producers of Linear alpha-Olefins (continued)
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Company and Plant Location	Annual Capacity as of Mid-2010 (thousands of metric tons)	Remarks
Qatar Chemical Company II Ltd. (Q-Chem		
I) ^a		
Ras Laffan, Qatar	(345)	Joint venture between Q-Chem II (53%),
		Qatofin (46%) and Qatar Petroleum (1%).
		Planned to be on stream in the fourth quarter
		of 2010. Broad range of LAOs.
Total	197	
Planned	(663)	
Also referred to as Ras Laffan Olefins Co	ompany Limited (RL	DC).
SOURCE: CEH estimates.		

Chevron Phillips Chemical Company (CPCC) and Saudi Industrial Investment Group (SIIG) plan to build a 100 thousand metric ton-per-year hexene-1 plant in Jubail, Saudi Arabia. The planned on-stream date is in 2012.

Jam Petrochemical Company has plans for a 200 thousand metric ton-per-year LAO plant integrated in its olefin cracker No. 10 at Asulayeh, Iran. The plant will most likely use IFP's AlphaSelect LAO technology. However, it is unlikely that this LAO unit will be built.

PetroRabigh plans to have 50 thousand metric tons of butene-1 capacity in Rabigh, Saudi Arabia. The onstream date is not known.

Qatar Chemical Company (Q-Chem), a 51/49 joint venture between Qatar Petroleum and Chevron Phillips Chemicals, uses most of its LAO output as feedstock for the joint venture's polyethylene plant at the same site.

Q-Chem II Project, a joint venture of Q-Chem II (53%), Qatofin (46%) and Qatar Petroleum (QP, 1%) includes a 345 thousand metric ton-per-year LAO plant scheduled to come on stream in the fourth quarter of 2010.

Q-Chem II is a joint venture between QP (51%) and Chevron Phillips Chemicals (49%), while Qatofin is a joint venture of Qapco (63%, which itself is a joint venture between QP [80%], and Total Petrochemicals [20%]), Total Petrochemicals (36%) and QP (1%).

SABIC, through its subsidiary Jubail United Petrochemicals, completed the construction of a LAO plant at Al Jubail, Saudi Arabia with an annual capacity of 150 thousand metric tons in 2006. The plant was commissioned in December 2009. Products in the C_4 - C_8 range can be produced with a focus on hexene-1 and butene-1 monomers for use as polyethylene copolymers. The project uses new technology for LAO production called alpha-SABLINTM, jointly developed by SABIC and Linde, which offers high product selectivity and purity under moderate reaction conditions. See the **MANUFACTURING PROCESSES** section for further information on the SABLINTM process.

 C_4 and some C_6 LAOs are consumed captively. The remainder, 85-90 thousand metric tons, is targeted as exports to Asia and Europe.

JAPAN

PRODUCING COMPANIES

The following table presents the Japanese producer of LAOs:

	Japan	ese Prod	ucer of L	inear alpha-	Olefins
	Annual Capacity as of Mid-2010 (thousands of metric tons)				
Company and Plant Location	C ₄ ^a	C ₆₊	Total	Process	Remarks
Idemitsu Kosan Company, Ltd.					
Ichihara, Chiba	9	49	58	Idemitsu	Plant came on stream in 1989 and was expanded by 8 thousand metric tons in 1999. Annual production capacity was only for C_4 - C_{18} and excluded C_{20+} until 2004, but Idemitsu started producing C_{20} and C_{24} from 2005. Company consumes C_4 and C_8 captively for polyethylene production.
a. Includes only C_4 produced by et	hylene olig	omerizati	ion.		
SOURCE: CEH estimates.				X	

Mitsubishi Chemical discontinued linear alpha-olefins production (annual capacity of 60 thousand metric tons) at its Mizishima plant, in Okayama Prefecture in May 2009. The production was based on ethylene oligomerization. As of 2010, Idemitsu Kosan is the only producer of LAOs in Japan. The preceding table includes capacity to produce butene-1 by means of ethylene oligomerization. Idemitsu isolates butene-1 from their ethylene oligomerization plants; in addition, three other companies refine butene-1 from n-butylene. In 2010, only 15% of Japanese butene-1 capacity was based on ethylene oligomerization. The following table presents Japanese producers of butene-1 from all sources:

	Japanese Producers of Butene-1						
	Company	Annual Capacity as of Mid-2010 (thousands of metric tons)	Source				
	Idemitsu Kosan Company, Ltd.	9	Ethylene oligomerization				
	Mitsui Chemicals, Inc.	22	n-Butylenes				
	Sumitomo Chemical Co., Ltd.	19 ^a	n-Butylenes				
	Tonen Chemical Corporation	10	n-Butylenes				
	Total	60					
5	a. Sumitomo supplies butene-1 to I tract. The plant was expanded to	Dainippon Ink and Chemicals, In 19 thousand metric tons in Septe	nc. under a coproducer con- ember 1993.				
	SOURCE: CEH estimates.						

For more information on butene-1, see the CEH Butylenes marketing research report.

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Idemitsu Kosan began operations at a new linear alpha-olefins plant at Ichihara, Chiba Prefecture in 1989. The annual capacity of this facility was 50 thousand metric tons and the capacity was increased to 58 thousand metric tons in 1999. Idemitsu's capacity did not include C_{20+} fraction until 2005. Since 2005 the company has also produced C_{20} and C_{24} . Idemitsu uses its own process technology incorporating its own zirconium catalyst–based technology. This technology allows for production of narrower fractions with more linear products (less side-chained products) and more terminal (alpha-) double bonds than products made by other technologies. Idemitsu consumes a large amount of its C_4 , C_6 and C_8 production captively as polyethylene comonomer. Idemitsu Kosan offers individual fractions of linear alpha-olefins commercially, although demand for some individual fractions is still very small.

Mitsubishi Chemical Corporation (formerly Mitsubishi Kasei Corporation) had been producing alphaolefins at Kurashiki, Okayama Prefecture using Gulf's (now Chevron Phillips') ethylene oligomerization technology since 1970. The plant's annual capacity was expanded from 36 thousand metric tons to 50 thousand metric tons in 1989 and to 60 thousand metric tons in 2001. However, Mitsubishi Chemical discontinued production of alpha-olefins in May 2009.

Mitsui Chemicals is constructing a new hexene-1 plant at Ichihara Works in Chiba Prefecture. Production capacity of the new plant is 30 thousand metric tons per year and its commercial operation is scheduled to begin in December 2010. The technology is selective ethylene trimerization, utilizing Mitsui Chemicals' new catalyst. The new plant will use the 40 thousand metric tons per year of ethylene surplus at the site, following the termination of EO/EG plant.

SALIENT STATISTICS

The following table presents Japanese supply/demand for linear alpha-olefins:

		(thousands of metric tons)						
		Annual Capacity	Production ^a	Imports	Exports	Apparent Consumption ^b		
	1982	25	25	26	neg	51		
	1986	30	30	18	1	47		
	1990	100	60	10	15	55		
	1995	100	87	7	31	63		
	1996	100	89	5	30	64		
	1997	100	92	6	30	68		
	1998	100	94	9	32	71		
	1999	108	100	10	32	78		
	2000	108	100	12	30	82		
	2001	118	102	15	29	89		
	2002	118	115	15	33	97		
	2003	118	106	16	30	92		
$\overline{}$	2004	118	118	16	31	103		
	2005	118	113	19	30	102		
	2006	118	118	23	42	99		
	2007	118	100	29	32	97		
	2008	118	106	24	40	90		
	2009	118	90	28	30	88		

Japanese Supply/Demand for Linear alpha-Olefins (thousands of metric tons)

- a. Excludes quantities that are burned as fuel.
- b. Equals PRODUCTION plus IMPORTS minus EXPORTS.

SOURCE: CEH estimates.

The following table presents Japanese production of linear alpha-olefins by chain length for selected years:

Japanese Production of Linear alpha-Olefins (thousands of metric tons)						
Carbon Chain Length	1995	1999	2003	2006	2009	Use
C ₄ -C ₈	38	46	49	51	36	Consumed primarily as comonomer for HDPE/LLDPE. Approximately 55-60% of the total was C_4 in 1992, but the percentage of C_6 and C_8 increased to about 70% in 2003.
C ₆ -C ₁₀	17	18	19	22	16	Most of the decene-1 produced is exported as a raw material for PAOs.
C ₁₂ -C ₁₄	13	14	15	18	14	C_{12} is exported and sold domestically as raw material for linear alkylbenzene (LAB).
C ₁₄ , C ₁₆ -C ₁₈	13	16	17	21	18	Sold to Lion Corporation for use in AOS manufacture.
C ₂₀₊	6	6	6		5	Various miscellaneous applications.
Total	87	100	106	.118	90	
SOURCE:	CEH estin	nates.		27		

Production of C_4 - C_8 -range fractions for comonomer has increased, reflecting the growing demand for LLDPE, especially for metallocene LLDPE. In 2009, the production decreased because Mitsubishi Chemical discontinued alpha-olefin production in May of that year.

Production of C_6 - C_{10} -range fraction has gradually increased and the production has been shifted to the lower carbon chain lengths for comonomer applications. Most of the decene-1 produced is exported as a raw material for PAOs.

Most of production of LAOs in the C_{10} - C_{12} range is exported for use in the production of PAOs and LAB, mainly in Asian countries including the Republic of Korea, Taiwan and Indonesia. Idemitsu exports most of its C_{16} and C_{18} linear alpha-olefins to the United States and Europe for drilling mud applications.

CONSUMPTION

The following table presents Japanese consumption of linear alpha-olefins:

		(thousan	ids of metric tons)		
	Dolarothaloro	Oxo A	lcohols	alpha-		
P C	Comonomers	Plasticizers	Detergents	Sulfonates	Other ^b	Total
983	6	17	12	20	1	56
986	8	11	5	20	3	47
.990	18	10	5	17	5	55
995	27	8	5	6	17	63
996	30	6	5	5	18	64
97	35	6	5	4	18	68
998	39	6	5	3	18	71
999	45	5	5	3	20	78
000	48	4	5	3	-22	82
001	50	4	4	3	27	88
002	54	4	3	3	33	97
03	54	4	3	3	29	93
04	58	4	3	4	34	103
005	59	5	3	4	31	102
006	63	5	3	4	24	99
07	56	5	3	3	30	97
08	55	4	3	3	25	90
)9	52	4	2	3	27	88
)14	55	4	2	3	26	90
		Average A	nnual Growth Ra (percent)	ıte ^c		
2009-						
.014	1.3%	0%	0%	0%	-0.8%	0.5%
Dat	a include C ₄ and high	her.				
Poly	valphaolefins, indust	rial synthetic lubric	ants, lube oil addit	ives, mercaptans,	etc.	

Japanese Consumption of Linear alpha-Olefins^a (thousands of metric tons)

c. Data may not equal the calculated growth rate because of rounding.

SOURCE: CEH estimates.

Japanese consumption of linear alpha-olefins has been increasing since 1991, reached a peak in 2004 and then fell to 88 thousand metric tons in 2009. Consumption for plasticizer alcohols and alpha-olefin sulfonates decreased gradually in the 1990s and has been stable since then, while consumption for detergent alcohols has decreased slightly and has been 2-3 thousand metric tons in recent years. Overall demand is expected to be stable during 2009-2014, reaching 90 thousand metric tons in 2014.

During 1979-1983, Japanese consumption of linear alpha-olefins increased slowly but steadily, mainly because of increasing demand from plasticizer alcohols and detergent alcohols production (based on LAOs). After Idemitsu entered the market in 1989, Japan became a net exporter, rather than a net importer, of linear alpha-olefins. Most Japanese imports of LAOs were destined to Shell Japan Ltd. from

Shell Chemicals in the United States for merchant sales to Lion Corporation for AOS production. However, imports declined significantly again in 1990 because Lion began purchasing LAOs from Idemitsu in addition to Mitsubishi Kasei (at that time). In late 1993, Lion began using increasing amounts of natural alcohol-based surfactants in place of AOS in its detergents, further reducing import requirements. However, imports increased again from 1995, because of growing demand for hexene-1 imported from BP (now INEOS) (Belgium) and Sasol (South Africa) for comonomer applications.

Polyethylene Comonomers

The following table presents Japanese consumption of LAOs for use as comonomers in polyethylenes

		(thousands o	i metric tons)		
	Butene-1	_			
	Sources ^a	Butene-1	Hexene-1	Octene-1	Total
1992	59	10	7	4	21
1995	61	12	10	5	27
1999	62	14	25	6	45
2003	49	15	33	6	54
2004	51	17	35	6	58
2005	51		27	6	50
2005	51	16	31	6	59
2006	47	17	40	6	63
2007	51	14	36	6	56
2008	50	15	34	6	55
2009	41	13	33	6	52

Japanese Consumption of Comonomers in Polyethylene Production	
(thousands of metric tons)	

a. Data include butene-1 from ethylene oligomerization and other sources.

b. Data exclude consumption in the production of Tafmer[®], an alpha-olefin copolymer used as a resin modifier.

SOURCE: CEH estimates.

Comonomer demand for linear alpha-olefins for polyethylene production has been increasing gradually in 1990s and has been in the range of 54-63 thousand metric tons since 2003. Consumption of butene-1 from ethylene oligomerization was 17 thousand metric tons in 2009. Consumption of butene-1 from all sources was 41 thousand metric tons in 2009 compared to 50 thousand metric tons in 2008, due to reduced production of HDPE and LLDPE. Consumption of hexene-1 has increased, from 7 thousand metric tons in 1992 to 40 thousand metric tons in 2006, because it is preferred for use in production processes using metallocene catalysts. For example, Mitsui Chemicals Inc., Ltd. has developed a new LLDPE product from a metallocene catalysis process and has licensed this process technology to Ube Industries. Ube is producing LLDPE using hexene-1 and part of its LLDPE production is supplied to Mitsui Chemicals. The consumption of hexane-1 has been in the range of 33-36 thousand metric tons in recent years. Consumption of hexene-1 is expected to recover slightly through 2014.

A Mitsui Chemicals LLDPE plant (operated by Evolue Japan, an affiliate company of Mitsui Chemicals) using metallocene catalysis built at Chiba in 1998 was expanded to 240 thousand metric tons in October 2006 with plans to expand to 300 thousand metric tons in 2011. The plant is now jointly owned by Prime Polymer (75%) and Sumitomo Chemical (25%). Prime Polymer is a joint venture between Mitsui Chemicals (65%) and Idemitsu Kosan (35%), consolidating the polyolefin businesses of both parent companies.

Butene-1 and hexene-1 are also used in the production of Tafmer[®], an alpha-olefin copolymer produced by Mitsui Chemicals, which is used as a resin modifier in the production of thermoplastic olefins (TPOs).

Consumption of octene-1 is limited, since only Idemitsu consumes it captively. Octene-1 consumption has been stable at 6 thousand metric tons per year in recent years. Along with the use of butene-1, consumption for all comonomers in this application is expected to recover slightly during the next five years.

Oxo Alcohols

Mitsubishi Chemical consumed C_6-C_{10} and $C_{12}-C_{14}$ fractions of linear alpha-olefins captively for the production of linear primary alcohols used in the production of plasticizers and household detergents, respectively. However, the company ceased production of linear primary alcohols in 2009.

Plasticizer alcohols

LAO consumption for linear C_7 - C_{11} alcohols for phthalate plasticizers has been 4 thousand metric tons annually for the last two years. Consumption is expected to decrease slightly during the next five years because of the decline of the polyvinyl chloride market. In addition, di(2-ethylhexyl) phthalate (DEHP) and di(2-ethylhexyl) adipate (DEHA), which are derived from 2-EH, are still the preferred products in the Japanese plasticizer market; linear plasticizers are not expected to replace either one.

Detergent alcohols

Japanese detergent producers have been reformulating their products to use "natural" raw materials (i.e., surfactants derived from detergent alcohols from oleochemical sources). Thus, the use of LAOs to make AOS and detergent alcohols has been declining in Japan. Even though the trend toward "natural" product is continuing, some detergent brands still contain synthetic products. Therefore the consumption of alpha-olefins fluctuates largely depending on the detergent brand sold in the market.

Some C_{10} - C_{12} linear alpha-olefins have been used as a raw material for linear alkylbenzene (LAB) production. Mitsubishi Chemical used to consume this fraction captively until 2005, when it ceased production of LAB in March 2005. Idemitsu also exports and sells C_{10} - C_{12} linear alpha-olefins domestically for use in this application. Generally, C_{12} is used in LAB plants that do not have a paraffin dehydrogenation (e.g., Pacol[®]) unit; thus n-paraffins, which are less expensive than LAOs, cannot be used. Some LAB plants that do use n-paraffins also supplement their needs with LAOs to increase reactor throughput or alter the chain-length distribution of the LAB they produce.

Domestic consumption for detergent-range oxo alcohols is expected to be remain at similar levels for the next five years. Although Japanese consumption of detergent alcohols has shifted to natural alcohols

during the past several years, detergent manufacturers are expected to continue to use some synthetic alcohols to ensure an adequate supply of raw materials at reasonable prices. However, natural alcohols will continue to be preferred.

alpha-Olefin Sulfonates

Japanese demand for C_{14} - C_{18} alpha-olefins for AOS has been gradually decreasing since Lion Corporation has been replacing its raw material requirements from synthetic alpha-olefins to natural fatty alcohols and other natural sources. alpha-Sulfonated methyl esters, produced by Lion Corporation, have replaced large amounts of AOS. Indeed, all imports of alpha-olefins for AOS production ceased by the end of 1993. As a consequence, demand for C_{14} - C_{18} alpha-olefins for AOS decreased from 10 thousand metric tons in 1993 to 3 thousand metric tons in 1998. Annual demand has been stable at 3-4 thousand metric tons for the last five years. Demand will remain at 3 thousand metric tons through 2014.

Other

Other applications for LAOs include their use in polyalphaolefins (PAO), synthetic lubricants, lube oil additives, candle manufacturing, wax coating, paper sizing material, resin additives for coating and printing inks and cleaning materials used in combination with solvents. These applications have not increased in recent years.

Mitsui Chemicals operates ethylene/alpha-olefins co-oligomer plants at Ichihara, Chiba Prefecture, with an annual capacity of 75 thousand metric tons and at Kuga-gun, Yamaguchi Prefecture, with an annual capacity of 20 thousand metric tons. However, Mitsui Chemicals primarily uses butene-1 isolated from crude butylenes as the alpha-olefin source for polyalphaolefin production. The co-oligomer is used as a lubricating oil additive (VI improver) and base oil for synthetic lubricants. Although Japanese consumption of LAO-based synthetic lubricants, i.e., polyalphaolefins, was estimated to be 15-16 thousand metric tons in 2009, LAO consumption for this application in Japan is small, since PAOs are mostly produced in and then imported from the United States and Western Europe. Linear alpha-olefin consumption for synthetic lubricants is expected to remain at a similar level through 2014.

PRICE

LAO prices in 2010 were \$200-300 per kilogram (\$2.14-3.21 per kilogram using a conversion rate of \$93.6 per dollar) depending on lot size and fractions. For a large lot size, the price of C₄ was approximately \$120-130 per kilogram (\$1.28-1.39 per kilogram) and C₆ was \$280-330 per kilogram (\$2.99-3.53 per kilogram.) Price fluctuations over the years have been influenced by naphtha prices (the main feedstock for ethylene production).

TRADE

Japan is currently a net exporter of linear alpha-olefins. The following table presents Japanese trade in C_{6+} LAOs:

	(thousands of metric tons)		
	Imports	Exports	Net Trade
1982	26	neg	-26
1986	18	1.5	-16.5
1990	10	15	5
1995	7	31	24
1996	5	30	25
1997	6	30	24
1998	9	32	23
1999	10	32	22
2000	12	30	18
2001	15	29	14
2002	15	33	18
2003	16	30	14
2004	16	31	15
2005	19	30	11
2006	23	42	19
2007	29	32	3
2008	24	40	16
2009	28	30	2
SOURCE:	CEH estimates.		

Japanese Trade in C₆ and Higher Linear alpha-Olefins (thousands of metric tons)

In 2009, Japanese imports of LAOs amounted to 28 thousand metric tons. Historically, until 1986, LAOs were imported from Shell and Ethyl Corporation in the United States. By 1990, most imported LAOs were from Shell (United States) for AOS production. By 1993, imports of LAOs for AOS production had disappeared. However, imports of other ranges, especially C_6 (hexene-1), have gradually increased. Most imports come from Europe (INEOS) and South Africa (Sasol).

Japanese exports of LAOs amounted to 30 thousand metric tons in 2009. Exports of LAOs from 1990 to 1993 were primarily to the Republic of Korea, Taiwan and Singapore. Recently, exports have also gone to other countries, including Western Europe and the United States. Idemitsu has been increasing its exports since 1993. Idemitsu's capacity expansion in 1999 has supported its export activity. The company exports butene-1 to the Republic of Korea, C_{10} for PAO production and C_{12} for LAB production mainly to Western Europe, and C_{16} and C_{18} to the United States and Europe for drilling mud uses.

The following table presents Japanese exports of LAOs by country or region:

	Japanese Exports of Linear alpha-Olefins—2009 (thousands of metric tons)			
	Korea, Republic of	12		
	China	6		
\mathbf{J}	Western Europe	6		
	United States	2		
	Other	4		
	Total	30		
	SOURCE: CEH estimates.			
OTHER ASIA

PRODUCING COMPANIES

The only producer of linear alpha-olefins in Other Asia is in China, as shown in the following table:

Other Asian	Producers of Line	ar alpha-Olefins	
Company and Plant Location	Annual Capacity as of Mid-2010 (thousands of metric tons)	Process	Remarks
Beijing	50	Ethylene oligomerization	Hexene-1. Production started in 2007.
SOURCE: CEH estimates.			

Sinopec Beijing Yanhua Petrochemical Co., Ltd. completed construction of a hexane-1 plant with an annual production capacity of 50 thousand metric tons in May 2007, and started production in 2007. Beijing Yanhua Petrochemical claims to be using its own ethylene oligomerization technology. This is the first LAO plant in Asia outside of Japan.

CONSUMPTION



	Other Asian Consumpt Linear alpha-Olefins – (thousands of metric t	tion of -2009 tons)
	Polyethylene Comonomers alpha-Olefin Sulfonates	85 60-80 40-60
	Total	185-225
R	SOURCE: CEH estimates.	

Other Asian consumption of LAOs for polyethylene was 85 thousand metric tons in 2009. Consumption of LAOs for alpha-olefin sulfonates is mainly in the Republic of Korea, China, Taiwan, Thailand and India. LAB accounts for 40-60 thousand metric tons of alpha-olefin demand in Taiwan, Indonesia and India; the potential demand is believed to be 100 thousand metric tons, shared between LAO and normal paraffins.

APPENDIX

Data for U.S. production in earlier years is largely based on U.S. International Trade Commission (USITC) publications. It reported annual production statistics through 1994 for linear alpha-olefins in two ranges: C_6-C_{10} and C_{11} and higher. Thus, these reported production statistics exclude butene-1. The USITC ended this format with the 1993 report and in 1994 reported the production only of C_6-C_{10} -range LAOs. In 1995, it reported total production of all C_6 and above LAOs only in its preliminary report format (i.e., the *Preliminary Report on U.S. Production of Selected Synthetic Organic Chemicals*, which contained the final revised production numbers; this report was discontinued after 1995). In 1997, the National Petrochemical Refiners Association began reporting production of LAOs. The following table presents reported production of C_6 and higher alpha-olefins and estimated production of butene-1 by alpha-olefin producers for 1988-1995 and 1999*:

	U.S. Production and Sales of alpha-Olefins—1988-1999 (thousands of metric tons)								
		Produ	ction						
			C ₆ and Higher		C	Sales			
	Butene-1 ^a	C ₆ -C ₁₀	C ₁₁ and Higher	Total	C6-C10	C ₁₁ and Higher	Total		
1988	na	395.2	314.8	710.1	na	175.0	na		
1989	na	345.1	346.8	691.8	na	184.4	na		
1990	na	470.9	354.8	825.6	217.8	197.3	415.0		
1991	118	469.0	392.0	861.0	220.4	214.8	435.2		
1992	122	479.0	427.4	906.4	na	212.9	na		
1993	127	439.9	445.6	885.4	na	149.0	na		
1994	134	522.9	na	na	na	153.7	na		
1995	138	na	na	1,063.8	na	na	na		
1999	210	670.0	529.3	1,199.3	na	na	na		

a. Data include only butene-1 produced by ethylene oligomerization.

SOURCES: (A) CEH estimates (data for BUTENE-1 PRODUCTION and all 1999 data).

(B) Preliminary Report on U.S. Production of Selected Synthetic Organic Chemicals, SOC Series C/P 96-2, U.S. International Trade Commission (1995 datum for C₆ AND HIGHER).

(C) *Synthetic Organic Chemicals, U.S. Production and Sales*, U.S. International Trade Commission (all other data).

Data for earlier years are also available; however, data through 1972 cannot be considered an accurate measure of linear alpha-olefin production. Also, in 1973-1974 and 1980-1987, the agency erroneously reported some companies as producers of linear alpha-olefins. Reported data include Shell's total initial production output for LAOs, including those directed to isomerization and disproportionation steps to produce internal olefins. In contrast, CEH estimates of production of C_6 and higher alpha-olefins for these earlier years exclude the LAO that Shell used as feedstock in its SHOP plant. The following table presents historical production and sales data for C_6 and higher alpha-olefins during 1965-1987 from both CEH and the USITC:

^{*} Significant quantities of butene-1 are produced from raffinate sources. See the CEH *Butylenes* marketing research report for information on total butene-1 production.

		Produ	ction					
	Reported Production ^c			ction ^c	Reported Sales			
	CEH Estimates ^b	C ₆ -C ₁₀	C ₁₁ and Higher	Total	C ₆ -C ₁₀	C ₁₁ and Higher	Total	
1965				56.9			52.2	
1966				118.6			73.7	
1967				149.1			106.2	
968	91			153.8			134.0	
1969	113			186.9			154.8	
						T. T		
1970	132			84.0			58.8	
971	145			195.3			153.6	
972	159			234.7		-	181.9	
973	191			188.2			161.1	
974	191			200.1			184.6	
975	170			148.7		<u> </u>	119.0	
976	181			142.3			130.7	
977	191			176.9			160.4	
978	227			223.3			181.3	
979	263			363.1			213.6	
					X			
980	259-263			424.2			246.8	
981	272-277			453.2			150.8	
982	272-277	201.5	262.0	463.4	118.1	117.6	235.7	
983	299	214.0	282.8	496.8	148.5	144.5	293.0	
984	340	283.9			124.0	180.9	304.9	
985	376	234.7	356.2	590.9	152.6	218.9	371.5	
986	433	318.7	317.7	636.4	201.6	182.9	384.5	
1987	469	375.2	273.4	648.6	238.1			

U.S. Production and Sales of C_6 and Higher Linear alpha-Olefins—1965-1987^a (thousands of metric tons)

a. In 1965, statistics include data for alpha-olefins with carbon lengths of C₆-C₂₀ and for an unidentified "all other" range. In 1966-1977, statistics include data for alpha-olefins in the C₆-C₂₀ range and for alpha-olefins in an "all other" category composed of a C₁₆-C₃₀ range. The 1969 sales datum includes alpha-olefins in the C₅-C₇ range. In 1978-1979, the statistics include data for alpha-olefins in the C₆-C₂₀ range. Beginning in 1980, statistics include data for alpha-olefins in the C₆-C₂₀ range and higher.

- b. CEH estimates exclude alpha-olefins that are consumed internally by Shell to produce internal olefins by isomerization/disproportionation.
- c. Data include alpha-olefins that are consumed internally by Shell to produce internal olefins by isomerization/disproportionation.
- SOURCES: (A) CEH estimates (data for PRODUCTION, CEH ESTIMATES).
 - (B) Synthetic Organic Chemicals, U.S. Production and Sales, U.S. International Trade Commission (all other data).

The following table presents historical U.S. consumption of linear alpha-olefins. Total consumption in this table excludes alpha-olefins that were consumed captively by Shell Chemical for the production of internal olefins.

Historical U.S. Consumption of Linear alpha-Olefins—1969-1987
(millions of pounds)

				Oxo Chemicals				
		Resins ^a		Detergent	Synthetic Plasticizer	Fatty A cids	Ca	
	Comonomer	Polybutene-1	Total ^b	Alcohols	Alcohols	(C ₇ -C ₉)	Aldehydes	Total ^b
1969					20			20
1975	14	2	16		61-63		-	61-63
977	19-21	8	27-29	na	77			77
979	41	12	53	na	82		-	82
980	54	11	65	na	68	7		75
981	59	12	70	na	70	5		75
982	85	13	98	na	67	5		71
983	109	15	124	na	70	7	16	93
984	129	16	145	na	73	9)	15	97
985	147	17	163	na	77	11	20	108
986	165	17	182	na	81	13	20	115
007	193	18	211	na	83	15	20	118

	Synt	hetic Lubrican	its		Alkyldimethylam	ines		
	Polyalpha- olefins	Other	Total ^b	Fatty Amine Oxides (FAO)	Quaternary Ammonium Compounds	Other	Total ^b	
1969		na		23			23	
1975	5	na	5	8			8	
1977	14	па	14	8		na	8	
1979	28		28	9-12		2-3	11-15	
1980 1981 1982 1983 1984	29 31 27 31 34	1 	29-30 31-32 27 31 34	9-12 9-12 9-12 9-12 9-11	 3-4 3-4 4-5	3 3-4 1 <1 1	12-15 12-16 13-17 13-17 14-17	
1985 1986 1987	38 43 52		38 43 52	10-11 10-11 10-11	5 5 6	1 2-3 4-5	16-17 17-19 20-22	

Historical U.S. Consumption of Linear alpha-Olefins—1969-1987 (continued) (millions of pounds)

	Surfactants			Surfactants				
	Lube Oil Additives	alpha-Olefin Sulfonates	Other	Total ^b	Alkenylsuccinic Anhydrides	Linear Mercaptans	Other	Total ^b
1969			7	7	7			57
1975		11	1-2	12-13	25			127-131
1977		3	1-2	4-5	34			164-167
1979	18	3-4	11-14	14-18	3	5	18-25	232-243
1980	17	5	2	7	3	6	18-25	234-239
1981	16	8-9	2	10-11	3-4	6	18-25	243-250
1982	15	7	1-2	8-9	4-5	4-5	18-25	258-268
1983	15	5	0-11	5-16	4-5	4-5	18-25	306-329
1984	15	5	0-14	5-19	4-5	5	20-25	339-361
1985	16	5	0-14	5-19	5	5-6	21-25	378-398
1986	17	5	0-14	5-19	5	5-6	23-27	414-435
1987	20	6	0-14	6-20	6	6	23-27	462-482

a. Supplies of butene-1 used as comonomer for HDPE and LLDPE resins, as the monomer for polybutene-1 and for other uses, are not necessarily from linear alpha-olefin plants. High-purity butene-1 may be isolated from ethylene oligomerization processes or purified from refinery and steam cracker C_4 streams (i.e., at an MTBE plant). Therefore, there may be some overcounting, as it is difficult to distinguish the source of the butene-1 used.

b. Totals may not equal the sums of the columns because of conversion and rounding.

SOURCE: CEH estimates.

BIBLIOGRAPHY

Chemical Economics Handbook—The following CEH marketing research reports contain additional information that is related to the subject of this marketing research report:

Benzyl Chloride Butylenes Detergent Alcohols Ethylene High-Density Polyethylene Resins Linear Alkylate Sulfonates Linear and Branched Alkylbenzenes Linear Low-Density Polyethylene (LLDPE) Resins Natural Fatty Acids Normal Paraffins (C_9-C_{17}) Oxo Chemicals Plasticizer Alcohols (C_4-C_{13}) Plasticizers Surfactants, Household Detergents and Their Raw Materials **Process Economics Program**—The following Process Economics Program reports contain more detailed information on the manufacturing processes, process design and process economics of the chemicals discussed in this report. Address inquiries concerning this information to the Process Economics Program, SRI Consulting, Menlo Park, California 94025.

Linear alpha-Olefins, Report Nos. 12, 12A, 12B, 12C, 12D and 12E, June 1966, October 1971, October 1974, July 1990, June 2001 and December 2008.

Specialty Chemicals Update Program—The following SCUP reports contain additional information that is pertinent to the subject of this report. Address inquiries concerning this information to the Specialty Chemicals Update Program, SRI Consulting, Menlo Park, California 94025.

Biocides Cosmetic Chemicals Lubricating Oil Additives Surfactants Synthetic Lubricants

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Polyolefins Multiclient Projects—SRIC has issued the following study of polyolefins technology, markets and competitive strategies and positions:

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